ABSTRACT

Accessory auricle is a rare anomaly with an estimated incidence rate of 0.2-0.5%. The most common form of this malformation is the accessory tragus. It may be a sign of other syndromes, such as oculo-auriculo-vertebral dysplasia (Goldenhar’s syndrome). In this paper, we describe two cases of accessory auricle with a focus on diagnosis and surgical treatment.

Keywords: accessory auricles, surgical treatment, congenital syndromes

INTRODUCTION

Accessory auricle (polyotia) is one of the most common congenital anomalies of the ear. This anomaly is the result of a disruption in the development of branchial arc I or II and is thought to be an autosomal dominantly inherited disorder, with an incidence rate of 0.2-0.5%. Accessory auricles usually present unilaterally and are more frequent in male patients. It can be solitary or multiple with different morphology and localization (1-3). The most frequent localization is on the face in front of the tragus. It could be an isolated malformation or be associated with other anomalies, such as cleft lip or palate, hypoplasia of the lower jaw, and eye and spine abnormalities. It could be a component of another syndrome and is best known as Goldenhar’s syndrome (oculo-auriculo-vertebral dysplasia). The most common and mildest form of this malformation is accessory tragus. Sometimes, larger accessory auricles may be found on the face or neck as a cartilaginous skeleton covered with skin. This anomaly is rarely seen in other regions, such as the middle ear and the pharynx. The definitive diagnosis of an accessory auricle is achieved by post-excisional histological analysis.

CASE REPORT 1

A 23-year-old male patient was admitted to the Clinical Center Kragujevac in Serbia for surgical treatment of multiple skin tumours that were localized between the tragus and the corner of the mouth (Figure 1). The largest tumour had a fistulous channel and secreted cerumen. There were no...
openings of the channels into the mouth. Ophthalmologists and otorhinolaryngologists were consulted, and the patient had no other anomalies. All of the tumours were completely removed together with the fistulous channel to the level of the facial muscles. Histopathology confirmed that this was an inherited anomaly (Figure 2). He was discharged from the hospital on the 3rd postoperative day with no complications, and at the last follow-up examination three months after surgery, there were no signs of recurrence.

CASE REPORT 2

A one-year-old male patient underwent surgery at the Clinical Center Kragujevac in Serbia because of a growth on the left side of the neck that was medial to the anterior edge of the sternocleidomastoid muscle (Figure 3). Ophthalmologists and otorhinolaryngologists were consulted, and the patient had no other anomalies. During surgery, the fistulous channel, which extended to the common carotid artery bifurcation, was completely removed. Histopathological analysis confirmed that the growth was an accessory auricle. The postoperative period was uncomplicated, and the patient was discharged from the clinic on the 3rd postoperative day. There was no recurrence during the three-month observation period.

DISCUSSION

Congenital anomalies of the ear are numerous but relatively rare. Their aetiology is not known. Data in the literature indicate that the incidence of these malformations is 1:6000 and is most prevalent among the Japanese population and the Navajo Indians (4). It is thought that ear malformations occur due to improper fusion of the six auricular hillocks during auricle development. One of the most common congenital anomalies of the ear is the accessory auricle. In most cases, it is a mild malformation that manifests as a supplementary tragus in front of the normal tragus or at the ascending crus of the antihelix. The more severe anomalies are less common, and those with a morphologically formed auricle are referred to as polyotia (5-9). This is a very rare form of malformation and is defined as an accessory auricle that, by its size and morphology, resembles a normal auricle. Accessory auricles are more common in male patients and are unilateral in 90% of cases (10). This anomaly most frequently occurs on the face, but it can be found on other locations such as the lateral side of the neck (11, 12), suprasternal region (13), middle ear (14), Eustachian tube (15), nasopharynx (16, 17, 18), oropharynx (19) and nasal vestibule (20).

Many genetic studies of accessory auricle have indicated that it is an autosomal, dominantly inherited disorder, although there have also been reports of it being acquired from X-linked recessive and autosomal recessive inheritance. One study analysed gene maps of 11 families for the autosomal dominant accessory auricular anomaly (ADAA) and showed that it was an autosomal dominant abnormality with complete penetration, and the isolated gene locus was 14q11.2-q12 (21). According to numerous studies, accessory auricle may be associated with other congenital anomalies. Goldenhar’s syndrome is the most common form and is also known as oculo-auriculo-vertebral syndrome (OAV) (22-24). This is a rare syndrome with incomplete ear, nose, soft palate, lip, and lower jaw development, and often occurs with spinal scoliosis and lordosis. Sometimes, there are anomalies of internal organs such as the heart, lungs and kidneys. The incidence of this syndrome in the UK is 1/3,500 to 1/26,000. These anomalies are unilateral in 90% of cases. They are frequently accompanied by limbal dermoid of the eye, accessory tragus, and strabismus. The following syndromes are much rarer: Wolf-Hirschhorn syndrome (WHS) (25, 26), also known as chromosome 4p deletion syndrome; and Pitt-Rogers-Danks syndrome (PRDS) or Pitt’s syndrome. The main features are microcephaly, micrognathia, hypertelorism, accessory auricle, short philtrum, prominent glabella, psychophysical retardation, muscular hypotonia, and abnormalities of the heart. Less common features are hypospadias, iris
coloboma, renal abnormalities, and IgA deficiency. Multiple accessory tragus and aplasia cutis congenita are part of the rare Preston Delleman’s syndrome (oculocerebrocutaneous syndrome) (27). Townes-Brocks syndrome (TBS) is extremely rare and occurs in only 200 people worldwide. It is the result of mutations in the SALL1 gene and is autosomal dominantly inherited. It is characterized by accessory tragus, inner ear anomalies, and malformations of the anorectal region, heart, kidneys, hands and feet.

Accessory auricle is diagnosed by clinical examination and histopathologic analysis. It is usually treated surgically and usually involves a radical excision into the periphery and below the growth (2, 5). The operative duration is dependent on the severity of the anomaly and the presence of associated anomalies. Occasionally, preoperative fistulography is required. In addition to traditional excision, other surgical methods have been described such as the application of special clips (28), which may be useful for milder forms of the anomaly and for those without a fistula.

The diagnosis of accessory auricle is usually based on clinical examination and sometimes requires fistulography; a consultation to rule out the possibility of a skin tumour is also warranted. Therefore, definitive diagnoses are possible only after histopathological analysis of the completely excised skin lesion. It is important to exclude the associated anomalies, especially in the middle and inner ear as well as the eye. Accessory tragus is a feature of Goldenhar’s syndrome, and therefore, otorhinolaryngological and ophthalmological examination is recommended. Occasionally, there is a need for paediatric or genetic consultation. Treatment of this anomaly is surgery (30). Since the rudimentary canal may spread to the deeper facial and neck neurovascular structures, a plastic surgeon and/or otorhinolaryngologist should be consulted.

DECLARATION OF CONFLICTS OF INTEREST

The authors declare that they have no conflicts of interest.

REFERENCES