Acromicric dysplasia and hearing loss: 
A case report

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ABSTRACT
Introduction: Acromicric dysplasia is an extremely rare autosomal dominant bone dysplasia characterized by progressive growth retardation, short hands and feet, mild facial dysmorphism and generalized joint limitation. Association with ear, nose and pharyngolaryngeal alterations has been previously reported. However, little is known about the otolaryngologic aspects that may accompany this disease. We pretend to raise awareness to the otolaryngologic aspects that may accompany acromicric dysplasia and to the possible interventions that improve these patients quality of life.

Clinical Case: We present the case of a ten-year-old girl referred to the otorhinolaryngology consultation with complaints of progressive hearing loss. Hearing evaluation confirmed bilateral severe mixed hearing loss and the patient was successfully rehabilitated with bone-anchored hearing aids.

Discussion/Conclusions: Evaluation by an otolaryngologist should be part of the management of children with bone dysplasias, especially if they present with associated symptoms, to allow an early diagnosis and proper intervention.

Keywords: Bone dysplasias; hearing loss, hearing impairment rehabilitation

DISPLASIA ACROMÍCRICA E ALTERAÇÕES AUDITIVAS:
A PROPÓSITO DE UM CASO CLÍNICO

RESUMO
Introdução: A displasia acromicrárica é uma displasia óssea extremamente rara, de transmissão autossómica dominante, caracterizada por atraso progressivo do crescimento, mãos e pés curtos, dismorfismo facial ligeiro e limitação generalizada do movimento das articulações. A associação com alterações nos ouvidos, nariz, faringe e laringe foi descrita em estudos prévios. No entanto, pouco se sabe sobre os aspectos otorrinolaringológicos que podem acompanhar esta doença. Pretende-se sensibilizar para os aspectos otorrinolaringológicos que podem acompanhar a displasia acromicrárica e para as possíveis intervenções que melhoram a qualidade de vida destes doentes.

Caso Clínico: Apresentamos o caso de uma criança do sexo feminino, de dez anos de idade, referenciada à consulta de otorrinolaringologia por noção de diminuição progressiva da audição. A avaliação audiológica confirmou a existência de uma hipoacusia de mista severa bilateral e a doente foi reabilitada com sucesso com um implante osteointegrado.

Discussão/Conclusões: A observação por um otorrinolaringologista deve fazer parte da avaliação de crianças com displasias ósseas, especialmente se apresentam sintomas associados, para permitir um diagnóstico precoce e uma intervenção adequada.

Palavras-Chave: Doenças do Desenvolvimento Ósseo; hipoacusia; reabilitação auditiva
INTRODUCTION

The evaluation of children with bone dysplasias is often directed to the associated morphological and radiological alterations. However, many of these patients present with otolaryngological complaints that should be taken into account. Acromicric dysplasia (AD) is an extremely rare autosomal dominant bone dysplasia (prevalence <1/1 000 000) first described by Maroteaux et al. in 1986. The genetic cause of AD was shown to be heterozygous mutations in the transforming growth factor (TGF) β-binding protein-like domain 5 of the fibrillin 1 gene (FBN1).

AD is characterized by a postnatal and progressive growth retardation resulting in short stature with a pseudomuscular build, short hands and feet, mild facial dysmorphism, generalized joint limitation and skin thickening. It is associated with some distinctive skeletal and radiologic features, such as delayed bone age, short metacarpals and phalanges, cone-shaped epiphyses, ovoid vertebral bodies and internal notch of femoral head. Less than sixty cases have been reported in the literature to date. In these cases, association with ear, nose and pharyngeal-laryngeal alterations has been reported. However, little is known about the otolaryngologic aspects that may accompany this disease.

CASE DESCRIPTION

A ten-year-old girl, diagnosed with acromicric dysplasia, was referred to the otolaryngology consultation with complaints of progressive hearing loss. The diagnosis of AD was clinically suspected by the distinctive clinical and radiological features that the patient presented (growth retardation with short stature and short limbs, brachydactyly with broad thumbs, partial syndactyly of 3rd and 4th fingers of both feet and short metacarpals and phalanges in x-ray) and was genetically confirmed by molecular analysis of the FBN1 gene. There was no familiar history.

Physical examination showed a mild facial dysmophia (figure 1) and low implantation of both ears with bilateral moderate stenosis of the external auditory canals. The audiogram revealed a bilateral severe mixed hearing loss (figure 2). She had a history of adenoidectomy and myringotomy with placement of transtympanic ventilation tubes at five years of age, with no hearing improvement. A CT scan was performed and revealed a class two middle ear malformation (Cremers classification).

She was successfully rehabilitated with a bone-anchored hearing aid (BAHA®) in the right ear at age 11. A few years later the parents and the patient decided to place a bone-anchored hearing aid (BAHA®) in the left ear, which resulted in additional gain (figure 3).

Figure 1 – Mild facial dysmophia

Figure 2 – Audiogram shows a bilateral severe mixed hearing loss
DISCUSSION

Acromicric dysplasia is characterized by a short stature and distinctive skeletal abnormalities with normal intellectual development. Otolaryngological alterations, such as a hoarse voice and recurrent ear, tracheal and respiratory infections (such as otitis and other upper respiratory infections) have been associated with this disorder.

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The pathogenesis of this disease is related with heterozygous FBN1 mutations that are all located in exons 41 and 42 and encode TGFβ-binding-protein-like domain 5 of FBN1 gene. Microfibrillar network disorganization and enhanced TGFβ signaling were consistent features in AD fibroblasts. This mutation is also associated with other disorders that present with short stature phenotypes (like geleophysic dysplasia) and with Marfan syndrome.2,5

Diagnosis is based on the typical skeletal and radiological alterations and is confirmed by molecular study of the FBN1 gene.2-5

Life expectancy of these patients is normal and no major complications, such as cardiac disease or major orthopedic problems, were described in the course of the disease.4 Nevertheless long-term follow-up of AD requires an orthopedic surveillance and annual clinical and radiological assessment is recommended.3

In terms of hearing loss rehabilitation, the bone anchored hearing aids and the conventional hearing aids are good options. In this case, with the bilateral external canal stenosis, the bone anchored hearing aid was decided and had excellent audiological results. This intervention significantly improved this patient quality of life, allowing her to continue her potential for a normal cognitive development.

In this case the authors expect to raise awareness to the otolaryngologic aspects that may accompany this rare disease and to the possible interventions that undoubtedly improve these patients quality of life.

REFERENCES


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