Management of Stahl’s ear: a case report
Manejo da deformidade auricular de Stahl: um relato de caso

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ABSTRACT
Stahl’s ear deformity was first described in the 19th century and it consists of a rare auricular deformity characterized by hypoplasia of the antihelix crus with enlargement of its base and a third crus of the antihelix connected to posterior portion of helix crus, which deforms the posterolateral wall of the external auditory canal. Reconstructive surgery is the definitive treatment, however, because of the diversity of clinical presentations, no standard technique exist for all cases. The method described in this report is another treatment option and entails the resection of the third crus and reconstruction of superior crus of the antihelix.

Keywords: Ear deformities acquired; Ear; Reconstructive surgical procedures.

RESUMO
Descrita no século XIX, a deformidade auricular de Stahl consiste em uma má formação auricular rara, caracterizada por hipoplasia da raiz da anti-hélice, com o alargamento de sua base e uma terceira raiz da anti-hélice conectando-a à parte posterior da hélice, deformando a porção posterossuperior do pavilhão auditivo. A correção cirúrgica é o tratamento definitivo, porém, pela diversidade de apresentações clínicas, não há uma técnica padrão para todos os casos. O método descrito neste relato é mais uma opção de tratamento e consiste na ressecção da terceira crus e confecção da raiz superior da anti-hélice.

Descritores: Deformidades adquiridas da orelha; Orelha; Procedimentos cirúrgicos reconstrutivos.

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INTRODUCTION

In the 19th century, Stahl described and classified different auricular malformations into three types: “helix transversus spleniformis”, “crus antihellicis trifurcata” and “crus superium turgidum”. Currently, the term Stahl’s ear refers to the second type, a hypoplasia of the antihelix crus, with enlarged base and a third crus of the antihelix extending from the antihelix to the helix crus, deforming the posterolateral wall of the external auditory canal.

This anomaly has aesthetic and social consequences, but it does not affect hearing. It is more common among eastern people and rarely seen in white people, however, its real incidence has not been established so far. The deformity is most commonly unilateral, and it can be bilateral in 20% of cases. In general, other anomalies are also present, such as narrowing of the helix, hypoplasia or absence of the superior crus of antihelix and enlargement of the triangular fossa.

Stahl’s ear would be the consequence of dysgenesis of the intrinsic atrial muscle during the third month of embryogenesis. Embryonic muscular dysgenesis leads to hypoplasia of the superior crus of antihelix, and development of a third crus of the antihelix. The third crus that connects the antihelix to the posterolateral wall of the helix is responsible for causing the defect on the curvature of the auditory canal. The solution for the deformity is reconstructive surgery, but there is no consensus regarding the technique to be used.

CASE REPORT

We report a case of an 18-year-old white man with Stahl’s ears deformity (Figure 1) who underwent reconstructive surgery under general anesthesia. An infiltration of the incisions was performed using 0.75% ropivacaine with epinephrine in a 1: 100,000 proportion. A retroauricular access was performed in the concha-Antihelix transition with detachment to anterior portion, enabling degloving of the ear and exposure of the third crus (Figure 2).

A resection of the third crus was performed with primary approximation of the defect. To reconstruct the superior crus of the antihelix, prior cartilage weakening was done with metal scraping and Mustarde 4-0 polyglactin suture. To reconstruct a uniform curvature of the helix, a subsequent cartilage weakening was carried out with a scalpel blade no.15 and x-suture for eversion with 4-0 polyglactin.

Closing of the posterior incision was performed using a Greek bar suture with 4-0 polyglactin. Ear dressings were made with sterile bandages and maintained for 48 hours. The same procedure was performed in both ears. The result after the surgery is shown in the late postoperative period (Figure 3) in which little changes in size of the ear is observed associated with absence of third crus, and presence of helix and superior crus.

DISCUSSION

Congenital auricular deformities are defined as malformations (microtia, cryptotia). Deformities are characterized as a normal chondrocutaneous component, but with an abnormal architecture categorized as constricted, prominent, and Stahl deformity, as described in this study.

A number of techniques for surgical reconstruction of Stahl’s ear have been described. These techniques range from simple procedures as zetaplasty, which is a realignment of the third crus, the wedge resection of cartilage, the local grafting after its reversion to more complex procedures such as temporal periosteal flap for auricular support described by Nakayama et al.
REFERENCES


CONCLUSION

Stahl’s ear is a rare auricular malformation that can bring social stigma to the patient. The method described constitutes another alternative for treating this disease, especially in cases where the deformity is more significant, and may produce satisfactory aesthetic results.

COLLABORATIONS

**JM** Statistical analysis: final approval of the manuscript to be published, conception and design of the study, operations and/or experiments, drafting the manuscript or critical review of the content.

**ACPO** Final approval of the manuscript to be published, conception and design of the study, operations and experiments.

**LMP** Analysis and/or interpretation of data, conception and design of the study, drafting the manuscript or critical review of the content.

**JB** Analysis and/or interpretation of data, drafting the manuscript or critical review of the content.

**MVMC** Final approval of the manuscript to be published, operations and/or experiments.

Figure 3. Late postoperative period. A: Right ear; B: Left ear.

There are reports of non-operative treatment using splints, which are used to shape the auricle, however more satisfactory results are seen if this treatment is performed in the neonatal period. However, there are different degrees of clinical presentation of the syndrome, therefore it is difficult to achieve consistent results using a standard surgical technique.

Currently, the modified Chongchet technique are the most commonly used for the treatment of this anomaly. There is a trend to use Mustarde sutures in mild deformities associated with cartilage weakening maneuvers. Severe cases require the use of previous technique associated with excision of cartilage and skin.

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