Sebaceous Hyperplasia en plaque: A Rare Variant

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Abstract

Observation: We present a rare variant of sebaceous hyperplasia in the postauricular area presenting with yellowish papules coalescing to form a plaque. We would like to present this unique case because of its plaque-like formation of the sebaceous papules as well as its rare localization in the retroauricular region.

Introduction

Sebaceous hyperplasia (SH) occurs frequently as solitary or multiple yellowish papules on the face in adults of middle age and older [1]. We present here a rare variant of sebaceous hyperplasia in the postauricular area presenting with yellowish papules coalescing to form a plaque.

Case Report

A 20- year- old female presented with asymptomatic skin lesions in the postauricular area for 2 years. She was otherwise healthy. Dermatological examination revealed multiple yellowish papules coalescing to form plaque on the left postauricular region (Figure 1). A 4-mm- punch biopsy was made from one of the yellowish papules which revealed hyperplastic sebaceous glands (Figure 2). The diagnosis was sebaceous hyperplasia en plaque.

Discussion

Sebaceous hyperplasia occurs frequently as solitary or multiple yellowish papules on the face in adults of middle age and older [1]. Recently, several cases in adolescence and
young adult patients have been described and identified as premature sebaceous gland hyperplasia [2]. The pathogenesis of this early clinical variant seems to be different from the senile type and it is poorly understood. Some reports have documented a family history [3]. In addition to this early clinical variant of sebaceous hyperplasia, several more clinical variants have been reported. There are patients with a giant form [4] on the cheek and also with linear arrangement [5]. Similar to our case, patients who had grouped hyperplastic sebaceous glands in a plaque pattern were rarely described [6].

The most frequent localizations of the senile and premature type of SH are the face, neck and upper thorax [2]. However, reports of SH on atypical localizations such as areola [7], vulva [8] and penis [9] have been described. To the best of our knowledge, the only report with SH forming a plaque around auricular region apart from our case is the one described by Kim et al [6].

However, SH is thought to be associated with decreased androgen levels in aging individuals and the pathogenesis of these early and unusual variants remain unknown. In conclusion, we would like to present this unique case because of its plaque like formation of the sebaceous papules as well as its rare localization in the retroauricular region.

References