Onychoheterotopia in children

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Abstract

The ectopic nail (EN) is an additional nail located in an abnormal site. It belongs to the onycho-heterotopia, a rare condition whose pathogenesis is indeterminate. This article illustrates the clinical-morphological and dermoscopic points of view, the diagnostic criteria, the possible pathogenesis, and surgical treatment of this pediatric onycho-heterotopia.

Introduction

Ectopic nail (EN) is an additional and independent nail located in an abnormal site. Also known as onycho-heterotopia, it is an extremely rare condition whose pathogenesis remains undetermined.1 EN and most of the nail diseases, which appear in the first decade of life, are congenital; less frequently, EN is described as acquired or post-traumatic. Onychoheterotopia is a single lesion involving the sole or, in most of the reported cases, occurring along with the regular fingernail, on the distal phalanx, and is attributed to digital malformations (e.g., palmar nail syndrome) or bone alterations.2 The appearance of the anomalous nail at diagnosis ranges from 2 to 13 years, and females are more affected than males.

Case Report

An 11-year-old girl, skin phototype 3 of Fitzpatrick, was evaluated for an EN lesion on the right sole. The patient complained of occasional pains while walking and dancing. Family history was not significant for genodermatosis or nail disorders. The lesion appeared at the age of 4 and had slowly increased since then. It was recently diagnosed as a wart and treated with cryotherapy without any benefit.

The clinical presentation showed cylindrical-conical keratotic horns on admission (Figure 1). The surrounding skin was smooth, with no signs of swelling or inflammation. We performed a dermoscopic analysis, which revealed a surface made irregular by transverse grooves at the periangual sulcus base (Figure 2). Treatment of EN consists of its removal. EN was treated under local anesthesia by surgical excision followed by phenol’s application on the nail matrix. No local recurrence was observed in the two-years follow-up.

Discussion

The onycho-heterotopia is a single lesion that usually appears near the fingers and rarely involves the feet.3 The clinical features are dependent on the nail tissue’s anatomical site, and occasional pain may be reported. Histopathological and ultrastructural features are similar to those of a regular nail with a reverse growth pattern.4 Three zones constitute EN: nail matrix, nail bed, nail plate, similar onycholysis, but differently arranged compared to a standard nail.5 The EN has a matrix in the center or back, with a horizontal-vertical growth pattern and a distal or inverted orientation. The neo-nail is generally conical and spatula-shaped at dermoscopic presentation, looking like a rudimentary fingernail with a corrugated surface and transverse furrows.6 The right foot’s little finger is the most commonly involved in EN, followed by the 4th digit and the 2nd digit. Several hypotheses tried to explain the origin of EN: the traumatic inoculation of onycholysis or implantation of nail germ cells and their abnormal growth;6 the variation in the long arm of chromosome 6 with congenital polyonychia;7 the presence of stray germinal cells in ectopic sites;8 the phenomenon of “paleoteı´vnia” (paleo = ancient; teı´v = to verge on) meaning the appearance of a primitive silent character, present in other animal species having a common phylogenetic origin. The pathogenesis remains unclear; it seems most probable that ectopic nails are related to the alterations of particular genes responsible for nail growth and orientation.9 The EN is often confused with acquired digital fibroma, cutaneous horn, clavus, foreign body, split nail deformity, rudimentary polychydactyly associated with callus formation and warts, and treated accordingly. The lesion is often undiagnosed or misdiagnosed, and cryotherapy alone often results in relapses. Surgical treatment should be followed by the phenolization of the nail matrix so that the germinal matrix is completely removed and destroyed.

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Conclusions

This report describes an unusual localization of a pediatric ectopic nail. Although this is a rare condition, its prompt recognition can avoid unnecessary treatments and allows appropriate surgical treatment without relapses.

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