Symmetrical cutaneous bilateral appendage – a case study

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Keywords
skin diseases; skin neoplasms

Question 1
What is the differential diagnosis?

Question 2
What is the most likely diagnosis?

Case study
A 13-year-old male was referred to our dermatology department for the presence of congenital asymptomatic verruciform lesions on the hands. Past medical history was unremarkable. No previous treatments were proposed.

On examination, two normocoloured papules were observed on the ulnar surface of the proximal phalanx of the fifth finger of both hands (Figure 1). The patient did not present any foot malformations, psychomotor delay or organ abnormalities. There were no skeletal components on radiographic evaluation.

Surgical removal of both lesions by excision and high ligation of the accompanying digital accessory nerve was performed. Hematoxylin-eosin staining showed the presence of a well circumscribed dermal nodular tumor comprised marked neural proliferation and large numbers of Wagner-Meissner bodies in the papillae (Figure 2). After two years of follow-up, recurrence or postsurgical neuroma has not been observed.

Question 3
How many types of this condition are we likely to find in our daily practice?

Question 4
Is it necessary to rule out any extra-cutaneous disease?

Question 5
What are the treatment options for this disease?

Answer 1
The main differential diagnosis is common warts, dermatofibroma, neurofibroma and dermal naevus. Common warts are clinically hyperkeratotic and dome-shaped papules with punctate black dots that represent thrombosed capillaries. Dermatofibromas are hyperpigmented papules, sometimes normocoloured, minimally elevated and they may seem attached to subcutaneous tissue. Neurofibromas are skin-colored papules, usually solitary and soft. Dermal naevus feels soft to the touch. They could be hypopigmented or normocoloured but rarely are located on fingers. None of these entities are congenital.
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Answer 2
Ulnar type B polydactyly.

Answer 3
All classifications have been performed with academic purposes. Swanson1 and Buck-Gramcko2 have classified polydactyly as radial/tibial affecting the thumb or great toe. When the affected digit is the little finger, the polydactyly is referred to as ulnar/fibular and cases in which the three central digits are affected are referred to as central polydactyly, the least common of the three types. This classification has replaced previous classifications into preaxial and postaxial and polydactylies. This has the advantage of avoiding confusion between the central morphological axis in the human hand, running along the third digit, and the more posterior metapterygial axis.3 Temtamy and McKusick4 prefers to divide polydactyly on type-A, when it is fully developed and type-B when it is not, including bilateral cases as the present case. A radiological study may help discern whether there is an associated skeletal component.

Answer 4
This is the most frequent congenital hand anomaly in people of African descent and the second most common, after syndactyly, in those of Caucasian descent. Isolated polydactyly is frequently inherited as an autosomal dominant trait of variable expression and its prevalence is approximately 1 in 531 live births. Although it often appears as a bilateral isolated feature usually underdiagnosed and misdiagnosed, an association with other birth defects such as mental retardation, underdevelopment, craniofacial dysmorphism, type 1 neurofibromatosis, hypospadias or hydrourerotonephrosis is present in 6.6% of cases.5

Answer 5
Ulnar type B polydactyly is often successfully treated using suture ligation in the delivery room with absorbable sutures6 or surgery. We consider excision with high ligation of the accompanying digital accessory nerve (digital nerve in the supernumerary digit) to be safer and more effective. It avoids the risk associated with anesthesia for the newborn and the possible occurrence of post-surgical neuromas.7 Other ablative techniques such as shaving and cautery, cryotherapy or CO2 lasers have also been used.

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