White Fibrous Papulosis of the Neck: A Case Report

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Abstract

White fibrous papulosis of the neck is a rare entity, with a benign course and unknown pathogenesis. It is clinically characterized by the appearance of firm, persistent, usually asymptomatic, non-follicular papules located on the neck. We present the case of a 72-year-old patient who presented pruritic lesions on the neck whose biopsy was compatible with this entity.

Categories: Dermatology
Keywords: wfpn, rare skin disease, progressive disease, elderly onset, white fibrous papulosis of the neck

Introduction

White fibrous papulosis of the neck (WFPN) is a rare entity. The first case reported dates back to 1983 in Japan [1]. Since then, it has been described in other countries such as Italy, Great Britain, Saudi Arabia, France, Spain, Belgium, and China [2-3]. In Latin America, cases have been reported in Argentina, Brazil, and Chile [4-6]. In most cases, the nature of this pathology is asymptomatic, which causes its true incidence to be underestimated. We present a female patient with this rare disease.

Case Presentation

A 72-year-old female patient arrived at the Dermatology outpatient clinic due to pruritic "pimples" on her neck, which progressively increased in number over one year prior to the dermatology appointment. The patient had a medical history of type 2 diabetes mellitus, systemic arterial hypertension, end-stage kidney disease on maintenance hemodialysis, hypothyroidism, and hypersensitivity to trimethoprim-sulfamethoxazole.

Physical examination showed a skin disease localized on the posterior and lateral sides of the neck (Figure 1) consisting of multiple papular lesions measuring 2 mm in diameter, skin-colored, and hard to touch (Figure 2).
FIGURE 1: Papulous eruptions measuring 2-3 mm located on the side of the neck (black arrows)
An incisional biopsy from the posterior region of the neck was taken, which showed a slightly folded epidermis with orthokeratosis, and atrophy of the spiny layer but conserving most of its interpapillary processes; thickening of the collagen fibers was observed in the superficial and medium dermis, with no change in its orientation (Figure 3A). Argentic staining identified a decrease in the elastic fibers and fragmentation (Figure 3B). Thus, the diagnosis of WFPN was made [7]. The patient did not request cosmetic treatment and remains stable at one year of follow-up.
FIGURE 3: A: Histological examination with hematoxylin & eosin staining (10X) revealed a folded epidermis with hyperkeratosis and orthokeratosis with thickening of collagen fibers in the superficial and medium dermis (black arrows). B: Argentic staining for elastic fibers (10X) revealed a decrease in elastic fibers, with fragmentation in some of them (transparent arrow)

Discussion

WFPN was first reported in Japan in 1983 by Shimizu et al. who reviewed 32 patients with asymptomatic papules on the neck. The histological examination of these patients showed fibrosis in the papillary dermis and, in some cases, slight changes in the elastic fibers [1].

In 1992, Riongioletti and Rebora reported two elderly women with similar neck lesions, which they described as “acquired elastolysis of the papillary dermis simulating elastic pseudoxanthoma.” The histological examination revealed papillary elasticity without fibrosis. In 1997, Balus et al. considered there was an overlap between the clinical and pathological characteristics of these entities, and the term "fibroelastolytic papulosis of the neck” was proposed [8].

White fibrous papulosis appears as monomorphic papulose eruptions about 2 to 3 millimeters in diameter, which can be isolated or confluent. These papulose eruptions are firm, non-follicular, and with a smooth surface. The lesions can be round or oval, of normal skin color, ivory white or yellow, and they are asymptomatic and sometimes pruritic [1]. They are mainly located around the neck but can also affect the upper part of the trunk, abdomen, armpits, and limb folds [1,3]. Most of the patients are women over 60 years of age, but there are case reports of younger ages [1,3], the youngest being 26 years old [1]. Lesions appear progressively and do not go away over time.

The pathogenesis of this disease is unknown. Some authors believe that sun exposure is involved, by accelerating the skin’s aging process [1]. Other authors think it is related to chrono-aging, which is why they consider this papulosis a “fibroelastolytic pattern of intrinsic skin aging” [9].

There are no reports of association with drugs, diseases, family history of similar lesions, or racial predilection to date. However, Catacra et al. reported a case of papillary dermal elastolysis type pseudoxanthoma elastic with the involvement of hormonal factors in young women [10].

Histological findings by hematoxylin and eosin stain include superficial dermal fibrosis with little or no change in elastic fibers, as well as a variable thickening of collagen fibers in the superficial dermis. With the Verhoeff-Van Gieson stain, specific for elastic fibers, the content of these fibers is shown to be slightly reduced or absent. In transmission electron microscopy, no significant morphological changes are shown [2-5]. Dermatohistology reveals white, homogeneous, circumscribed areas with thin dotted or short vessels, without follicular involvement [11].

The differential diagnosis must be established with other papular eruptions that appear on the neck. It should first include elastic pseudoxanthoma, which presents ocular and cardiovascular manifestations, with histology showing the absence of fragmentation of elastic fibers and calcium deposits. Acrocerdons, fibrofolliculomas, and trichodiscomas in Birt Hogg-Dube syndrome, Buschke-Ollendorf syndrome, cutis rhomboidalis nuchae, eruptive xanthomas, and perifollicular elastosis should also be considered (Table...
J) [4,12].
| Source: [13-17] |

| TABLE 1: Differential diagnosis of white fibrous papulosis of the neck |
| TGF, transforming growth factor; Er: YAG laser, Erbium-doped Yttrium aluminum garnet laser |

| Definition | Etiology | Physiopathology | Clinical Features | Diagnosis | Treatment | Prognosis |
|------------|----------|-----------------|-------------------|-----------|-----------|-----------|
| Elastic Pseudoxanthoma | A multisystem disorder that primarily affects the skin, eyes, and cardiovascular system, characterized by progressive and degenerative calcification of the elastic fibers. | ABCG2 gene defect on chromosome 16q24.1 with an autosomal recessive inheritance. | Skin signs: yellow calloskin lesions on areas of bending. Ophthalmological manifestations: angiod streaks, paxi of orange, maculopathy. Cardiologically manifestations: intermittent claudication, coronary artery disease, arterial hypertension, angina, myocardial infarction, congestive heart failure, restrictive cardiomyopathy, and valvulopathies. | Histology: calcium deposit with injured, swollen, clustered, and fragmented elastic fibers between the reticular and deep dermis. Electron microscopy: mineralization of the elastic fibers starts at the core, and as calcification progresses, fragmentation of the fibers occurs. | Ophthalmological and cardiothoracic evaluation. Laser photocoagulation to prevent retinal bleeding. Although not curative, adequate cardiovascular risk factor management, moderate dietary calcium intake, and prophylactic anticoagulants. Pathological changes are irreversible. | Usually normal life expectancy. |
| Acrochordons (Skin Tags) | Pedunculated benign cutaneous neoplasms. | Associated with increased risk of diabetes mellitus and hypertension. | Lesions are located in sites of friction, such as the shoulders, neck, and elbows. Hyperpigmentation results from UV exposure. Lesions are symmetrical and indistinguishable from each other on visual examination. | Histology: Lesions consist mainly of fibroblasts, fat, and other types of tissues such as blood, blood vessels, mast cells, Langerhans cells, and dermis. | Aesthetic removal with laser, electrodesiccation, cryosurgery with liquid nitrogen, and dermabrasion. | Lesions rarely reappear after removal. New lesions may develop in areas of previous skin. |
| Fibropodiumomas and Trichodiscomas | Aneurysmal tumors that arise from or around the hair follicles. | Dependent on the underlying pathology. | Peppers of 2-3 mm in the shape of a mushroom. They appear on the face (chin, nose, cheeks, ears, and eyes). Both lesions are asymptomatic and indistinguishable from each other on visual examination. | Histology: Fibropodiumomas: Dilated central follicles, increased papillary dermis, epithelial strands of basaloid cells emanating from the infundibulum of the hair follicle. Histology: Trichodiscomas: Proliferation of connective tissue and fibrous stroma, located near a hair follicle. | Surgical excision and combined CO2/N2: YAG laser. | Depends on the underlying pathology. |
| Buschke-Ollendorff Syndrome | Genodermatosis with connective tissue nevi, osteopoikilosis, and an autosomal dominant inheritance. | Linear cortical hyperostosis is seen on long bones in an X-ray. Connective tissue nevi are seen as papules, plaques, and yellowish or skin-colored nodules on the palms and soles. | Histology of connective tissue nevi: Increased elastic fibers or collagen fibers. | No specific treatment is required. | No specific treatment is required. | Benign and usually asymptomatic (25% of patients experience bone pain and joint edema). |
| Cutis Romboidea Nuchae | Manifestation of prolonged sun exposure and resulting skin damage that occurs at the back of the neck. | Sun-induced skin disorder in which ultraviolet radiation causes the degeneration of elastic and collagen fibers. | Diffuse thickening on the back of the neck with yellowing of the skin, and formation of deep grooves which result in a typical irregular rhomboidal pattern. | Histology: Thickening of the epidermis and atrophy in the composition of the dermis. | Photoprotection, topical retinoids, and fluorouracil. | Association with acral keratosis and basal cell carcinoma. |
There is no established treatment for injury remission. Although the lesions are benign, patients may appreciate these lesions as cosmetically unpleasant, so treatment should focus on flattening the lesions to achieve an aesthetic result acceptable to patients.

The use of 1550-nm fractional non-ablative laser, Erbium-doped Yttrium aluminum garnet laser, and carbon dioxide (CO\textsubscript{2}) laser have been reported. Other treatment options include photoprotection and topical preparations with retinoids [5, 18] or antioxidants that reduce age-induced free radicals [19]. Tacrolimus 0.1% twice daily has demonstrated an improvement in pruritus [20].

Conclusions

WFPN is a rare entity, with a characteristic clinical picture and unknown pathogenesis. There are few cases reported to date, so full documentation and publication are of paramount importance. These will help us know the true prevalence of WFPN, and the factors associated with it, thus resulting in a better characterization of this rare disease. Also, it’s important to highlight that these lesions tend to be pruritic and aesthetically unacceptable, they affect the quality of life of those who suffer from it, so it is necessary to implement new treatment strategies that lead to the well-being of our patients. We hope this case report will call the attention of other health professionals and improve the detection of WFPN.

Additional Information

Disclosures

Human subjects: Consent was obtained or waived by all participants in this study. Conflicts of interest: In compliance with the ICMJE uniform disclosure form, all authors declare the following: Payment/services info: All authors have declared that no financial support was received from any organization for the submitted work. Financial relationships: All authors have declared that they have no financial relationships at present or within the previous three years with any organizations that might have an interest in the submitted work. Other relationships: All authors have declared that there are no other relationships or activities that could appear to have influenced the submitted work.

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