Pigmented Bowen’s disease:  
a case report of an unusual variant

Doença de Bowen pigmentada: relato de caso de uma variante rara

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ABSTRACT

Bowen’s disease is a squamous cell carcinoma in situ with the potential to turn into invasive carcinoma. Pigmented Bowen’s disease is an unusual variant, with few reported in the literature, and becomes a problem in the differential diagnosis with other pigmented tumors. We reported the case of a white female patient, 73-year-old, with a brownish asymmetric plaque, with more than one color, in the right upper limb, with 2-years of follow up.

Key words: squamous cell carcinoma; Bowen’s disease; oncology.

INTRODUCTION

Bowen’s disease (BD) is a squamous cell carcinoma in situ that is typically presented as asymptomatic erythematous plaque, scaly, well-defined borders, and slow growth1,2; it affects any area of the body, most frequently in sun-exposed areas3,4. It is more prevalent in the elderly, affecting both sexes3,5. It has a rare pigmented variant, with few reports in the literature and difficult diagnosis1-3. We report a case of pigmented BD in a 73-year-old female patient.

CASE REPORT

A 73-year-old white female patient, presenting brownish hyperpigmented plaque, asymmetrical, irregular borders, with more than one color, approximately 2.5 cm in diameter in the right upper limb, two years ago (Figure 1). There was growth and darkening of lesion in recent months. Personal history of squamous cell carcinoma (SCC) in anterior chest region excised two years ago. The lesion biopsy was performed, and histopathology revealed remarkable keratinocytes atypia with loss of cell polarity and presence of some dyskeratotic cells, as well as irregular and pronounced increase in melanin pigmentation in keratinocytes, both in basal and in scaly layer, characterizing pigmented BD (Figures 2 and 3). Dermoscopy was not performed, because before medical follow up the patient died due to acute myocardial infarction.

FIGURE 1 - Hyperpigmented brownish plaque, asymmetrical, irregular border, more than one color, approximately 2.5 cm in diameter in the right upper limb
Discussion

The pigmented BD is a rare variant that accounts for less than 2% of BD cases, and is considered difficult to diagnose\(^2\). It is characterized by the typical alterations of BD and by melanin increase in the epidermis and dermis\(^2\); clinically it appears as hyperpigmented, well defined, with a velvety surface, flat or verrucous plaque\(^1\). Desquamation and ulceration may be present, as well as itching or burning, although most lesions are asymptomatic\(^2\). Risk factors are the same as BD's: exposure to sunlight, radiotherapy, arsenic, trauma, and infection with human papillomavirus (HPV)\(^4\). Our patient had typical clinical lesion of slow growth and had a history of exposure to solar radiation, with SCC history.

Pigmented BD should be considered in the differential diagnosis of melanocytic lesions, pigmented basal cell carcinoma, pigmented actinic keratosis, seborrheic keratosis, and solar lentigo – often being associated with the last two\(^7\).

Dermatoscopy assists diagnosis, demonstrating the absence of criteria for melanocytic lesion, brownish amorphous pigment (if any epidermal melanin) or greyish (if any dermal melanin), while brownish spots in a regular arrangement on the periphery of lesion and dotted or glomerular vessels in linear arrangement are more specific\(^7\), but these features are not present in all cases; histopathology is the gold standard for diagnosis\(^7\).

By histopathology we found acanthotic epidermis with keratinocytes, showing atypia and loss of usual pattern of maturation\(^9\), abundant presence of melanin pigments in the cytoplasm of atypical keratinocytes and a slight increase in the number of melanocytes, which are typical\(^2\). There is diffuse hyperpigmentation of epidermis, best observed in Fontana-Masson staining, melanophages in papillary dermis, and transepidermal elimination of melanin, which focally accumulates in the horny layer\(^9\).

Little is known about the mechanism of pigmentation of BD, and several theories attempt to explain its origin\(^8\). The theory of “colonization” asserts that the pigment occurs due to the presence of an increased number of melanocytic hyperplasia with of hypertrophic dendritic processes dispersed through the tumor\(^8\). Another theory suggests that, in pigmented BD case without scaling and keratosis signals, atypical keratinocytes are still in a well-differentiated stage, having, therefore, melanin\(^8\). In later stages, atypical keratinocytes lose this ability, resulting in loss of pigmentation, keratosis, and scaling. A third hypothesis suggests that specific growth factors or cytokines produced by cancer cells may stimulate the proliferation of melanocytes and melanin production\(^8\). Also, the association and overlapping of pigmented BD with seborrheic keratosis and solar lentigo, some authors argue that in such cases, the preexisting hyperpigmentation spreads with the expansion of neoplastic...
lesion and, as generally seborrheic keratoses and solar lentigo are not excised and subjected to histopathological routine examination, it is believed that the incidence of pigmented BD may be underestimated.8,9

The pigmented BD presents clinical similarity with other melanocytic lesions and should always be considered in the differential diagnosis of pigmented lesions. Although under-diagnosed, it may not be so rare, and its diagnosis is still a challenge.

RESUMO

A doença de Bowen é um carcinoma espinocelular in situ com potencial para transformar-se em invasivo. Ela é pigmentada, uma variante rara, possui poucos relatos na literatura e torna-se um problema no diagnóstico diferencial com outros tumores pigmentados. Relatamos o caso de uma paciente de 73 anos, branca, com uma placa acastanhada, assimétrica, com mais de uma cor, em membro superior direito, com evolução de dois anos.

Unitermos: carcinoma de células escamosas; doença de Bowen; oncologia.

REFERENCES


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