Clinical, Dermoscopic and Microscopic Features of a “Collision Tumour” Ultimately Confirmed as a Regressing Melanoma – Lessons Learnt from a Chance Diagnosis

Lawrence Chukwudi Nwabudike1, Ana Maria Oproiu2,3, Ioana Mihaela Dogaru2,3,*, Mariana Costache3,4,5, Alin Laurentiu Tatu5–7

1Outpatient Dermatology, N. Paulescu National Institute of Bucharest, Bucharest, Romania; 2Department of Plastic Surgery, Emergency University Hospital, Bucharest, Romania; 3Carol Davila University of Medicine and Pharmacy, Bucharest, Romania; 4Department of Pathology, Emergency University Hospital, Bucharest, Romania; 5Clinical Medicine Department, Faculty of Medicine and Pharmacy, Dunarea de Jos University of Galati, Galati, Romania; 6Dermatology Department, St Parascheva Clinical Hospital of Infectious Diseases, Galati, Romania; 7Multidisciplinary Integrated Center for Dermatologic Interface Research (MIC-DIR), Galati, Romania

*These authors contributed equally to this work

Correspondence: Lawrence Chukwudi Nwabudike, N. Paulescu Institute of Bucharest, 5-7 Ion Movila Street, Sector 2, Bucharest, Romania, Email Chukwudi.nwabudike@live.com; Ana Maria Oproiu, Carol Davila University of Medicine and Pharmacy, Bucharest, Romania, Email anamariaoproiu@gmail.com

Abstract: Spontaneous regression of malignant melanoma was first reported over a century ago. Clinically, areas of blue or grey colouration in lesions may be indicative of regression. Dermoscopy is a very useful tool for diagnosing regression. An important criterion is the blue-white scar. About 10–35% of excised melanomas show features of regression histopathologically. We present a case of regressing melanoma, with clinical and dermoscopic features suggesting a collision tumour, diagnosed histopathologically. This case might improve our knowledge of the potential clinical manifestations, and the biology, of regressing melanoma.

Keywords: regressing melanoma, collision lesion, blue-white scar, pepper-like granules, milia-like cysts, pseudofollicular openings

Introduction

Spontaneous regression in malignant melanoma was first reported in 1953 by Sumner in a remarkable case.1 Although the patient had several histopathologically proven tumour foci, the presumed primary was not biopsied and could only be conjectured based on history and cutaneous depigmentation.

Areas of depigmentation, blue or grey colouration may suggest regression clinically.2 The blue-white scar-like structure (BWS) is considered an important dermoscopic criterion for regression in pigmented lesions.3 Regression occurs in 10–35% of melanomas excised and examined histologically.1 Often regressing melanomas are diagnosed histologically, ie post factum, when the process has been completed.

We present a case of regressing melanoma, showing clinical and dermoscopic features that suggest a collision tumour. Clinical and dermoscopic manifestations suggestive of collision are not normally recognised criteria for regression in melanomas. This case has the potential to offer us a window also into other characteristics that could suggest a melanoma in the process of regression to clinicians, as well as into the biology of this phenomenon.

Case Report

A 75-year-old male patient presented for a varicose ulcer on his left leg. He received a dressing for the ulcer and was instructed on how to care for it. During the instruction, the attending physician noticed a black spot on the left side of his
neck. Closer examination revealed a black, approximately 1 cm × 1cm, triangular-shaped plaque, which was macular and hypopigmented on one half and nodular as well as pigmented on the other (Figure 1).

Overall, the lesion showed a multicomponent pattern. Dermoscopy showed an abrupt cut-off between two areas of differing pigment intensity. Atypical grey network irregular thick lines with incomplete rhomboids on the left side (oval shape). Irregular black blotches. Irregular streaks at 12 and 6 O’clock. White scar-like depigmented areas (hexagon), blue–gray pepper-like granules and blue-white veil. On the right side, blue-white veil, milia like cysts (circle), comedo-like openings as in seborrheic keratosis (rectangle), at 6 O’ clock thick pigmented lines around appendageal openings (arrows), which are sometimes called rhomboidal structures similar to those of lentigo maligna (Figure 2).

Histopathology (HE, 10x, Figure 3) shows an area of regression in the left part of the image (inflammatory infiltrate, which is predominantly lymphocytic and perivascular, encompassing also melanophages; some capillaries are newly formed; no “junctional activity” tumour cells can be identified, ie no residual tumor epidermal component is present). On the right, there are elongated cells with large hyperchromatic and atypical nuclei that are arranged in nests of varying dimension at the dermoeidermal junction, surrounded by a rich perivascular, predominantly lymphocytic infiltrate that also contains melanophages and extends along the whole of the base of the lesion. The melanoma cells involve all layers of the epidermis. The Clark level was IV and Breslow thickness 0.85 mm. No ulceration, neoplastic emboli or perineural infiltration was observed. Regression level was less than 75%.

Surgical excision was carried out with appropriate margins, the patient recovered without complications and the excised tissue was sent for histopathology, with the results elaborated above.

The patient was subsequently referred to the oncologists.

This case report has been presented according to the CARE guidelines for case reporting. A summary of the timeline of this case is presented in Table 1.

Written informed consent for the publication of the patient’s clinical details, including photographs and laboratory results was obtained.

![Figure 1](https://doi.org/10.2147/CCID.S361793)

**Figure 1** Polymorphous plaque, with flatter depigmented area and slightly raised pigmented area on the left side of the neck, suggesting melanoma or collision lesion.
Figure 2 (A) Dermoscopy showing abrupt cut-off between areas of stronger and milder pigmentation. Blue-whitish veil, pepperling, multicomponent pattern, suggesting melanoma, with milia-like cysts and pseudofollicular openings suggesting seborrheic keratosis. (B) containing marked up photo.

Figure 3 Histopathology, (HE x 40) with features of regression and absence of junctional activity (left side of image) and nests of elongated tumour cells at the dermoepidermal junction (right side of image).
Discussion
Regression is not an infrequent feature of melanocytic lesions, including melanoma. Histopathologically, it is usually reported after the fact. Early regression is characterised by patchy disappearance of melanoma cells and their replacement by T-lymphocytes, and by fibrosis in the late phase. CD4+ cells predominate in regressing melanoma, as well as a milieu rich in fibrogenetic cytokines, such as IL-6, platelet-derived growth factor and transforming growth factor-β. This may explain some of the features seen in regressing primary melanomas. Certain features have been found to correlate with regression in melanoma such as depigmentation, blue or grey colours clinically as well as BWS, multicomponent pattern and peppering dermoscopically, and these were seen in our case. Structures typical of seborrheic keratosis (milia-like cysts, comedo-like openings) are rarely found in melanoma. Confocal microscopy promises to offer a non-invasive method of differentiating between melanomas and other skin cancers. It may also help differentiate areas of regression from active melanoma. Round or dendritic pagetoid cells located at the dermoepidermal junction can be seen using confocal microscopy to examine melanoma tissue. Other features that may be indicative of malignancy include epidermal disarray, pagetoid infiltration, deranged dermoepidermal junction architecture and atypical nests. There are limited data on fibrosis studied using confocal microscopy. In classic frontal fibrosing alopecia and lichen planopilaris, confocal microscopy showed perifollicular lichenoid inflammatory infiltrate in 25% of cases, extensive perifollicular fibrosis in 75% of cases, as well as an increased number of white, ill-defined, coarse collagen fibres at the superficial dermis in 58.3% of cases and dilated blood vessels in 58.3% of cases. The overall features, suggestive of collision clinically and dermoscopically, seen in our case, together with the histopathology suggestive of an earlier phase of regression of malignant melanoma, may offer us a window into the evolution of regressing melanomas.

The original case by Sumner is reminiscent of the phenomenon of field cancerisation, which may be triggered in cases of immunosuppression. Our case is a patient with diabetes, which may predispose to cancer. Further, diabetics are at high risk for cardiovascular disease. Drugs such as hydrochlorothiazides, beta-blockers and statins have been associated with increased risk of cutaneous carcinogenesis. This may indeed accentuate the field cancerisation. While field cancerisation is a predisposition to developing cancer, regression in melanoma is not. It may signal an immune response to the tumour, but it may also signal a poorer prognosis, including the existence of distant metastases, as it is often seen in advanced melanomas.

From a clinical perspective, when facing such patients, the first difficult task may be to accurately observe and diagnose the lesions. Indeed, one of the reasons that regressing melanoma is traditionally associated with a poor prognosis is the delayed discovery and inaccurate initial evaluation of the tumour thickness. Such lesions evolve in two stages, first the melanocytic, usually pigmented proliferation appears; it can grow in size, darken in colour or bleed, and, left untreated, develops secondary regressive changes such as depigmentation and atrophy — flattening. These changes occur gradually, so lesions like the one presented in this case can be observed. Frequently, the patient seeks care
for other symptoms such as enlarged lymph nodes, and a diagnosis of metastatic disease is established in the absence of a visible primary tumour. Sometimes a lesion resembling an atrophic scar can be observed in the case of a completely regressed melanoma, thus differential diagnosis is all the more difficult for such patients.

The immunologic mechanism of regression is not yet fully understood. Some correlation between regressing melanoma and vitiligo-like depigmentation has been described in the literature, and it is believed to be caused by an immune response to similar antigens shared by normal melanocytes and melanoma. Similar discoloration has been observed in patients treated for melanoma with immunomodulators. This evidence may suggest that regression is based on the host immunologic response to the tumour and has led some specialists to believe that it could indicate a favourable prognosis. However, that view is not shared by all authors and the data are inconclusive. Other studies have linked regression with a poor prognosis in thin lesions, but unchanged prognosis in thicker tumours.

The patient’s associated type II diabetes is a risk factor for cancers. Diabetes has been shown to increase the risk of developing several types of cancer, particularly liver and pancreatic cancers. Several possible links between diabetes and carcinogenesis have been described, most important of which are hyperinsulinemia, hyperglycemia and insulin resistance, however the mechanism of carcinogenesis and its correlation with elevated glucose levels is still unclear. The use of antidiabetic medication is also believed to influence carcinogenic potential by inducing different insulin concentrations, thus metformin and thiazolidinediones are potentially safer than sulfonylureas and exogenous insulin, though results have varied.

Surgical treatment in these cases can be an issue if the primary tumour has regressed completely. Our patient presented with a partially regressed melanoma and thus the pigmented part of the lesion was easier for the examining physician to notice. When the lesion is completely regressed, it is often the regional metastases that are biopsied first. Generally, in the presence of nodal dissemination, complete lymph node dissection is recommended for all patients, because it reduces local recurrence rates, it provides valuable information on prognosis and it also aids in the selection of an adjuvant treatment. Managing metastatic melanoma almost always involves some kind of systemic therapy. Newer agents that modulate the immune response such as Ipilimumab or BRAF enzyme inhibitors such as Vemurafenib significantly prolong survival in the presence of distant metastases. Surgical excision of the metastases can prolong survival in some cases, but characteristics such as their number, site, resectability and also the biological aggressiveness of the disease should be carefully considered. The 5-year survival rate for locally advanced melanoma is still only 63%, and drops to 18% in the presence of metastatic disease. It is therefore extremely important to emphasise the role of periodic examination and self-examination, especially knowing that in the presence of regression, diagnosis can be challenging even for the experienced specialist. Fortunately, this patient did not suffer metastases.

**Conclusion**

Some features of melanomas in the process of regression have been reported, though not for the same lesion simultaneously. Our case, with features of a collision tumor, offers a rare view into potential clinical, dermoscopic and histopathologic features of a melanoma in the earlier phases of regression, thus contributing to the understanding of the biology of the phenomenon of melanoma regression.

**Ethics and Consent Statements**

Institutional approval was not required as this is a case report and the patient had already given written informed consent for the publication of photographs and all the test results. Written informed consent for the publication of this case was provided by the patient for the publication of photographs and all the test results.

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