Letter

Rapid Melanoma Death of an Adult Male with Congenital Bathing Trunk Nevus despite Initiation of Combination Immunotherapy

Thilo Gambichler 1,2,* , Kathrin Noldes 1 , Yousef Arafat 1 , Matthias Neid 3 , Arno Rütten 4 and Stefanie Boms 1

1 Department of Dermatology, Christian Hospital Unna, 59423 Unna, Germany
2 Skin Cancer Center, Department of Dermatology, Ruhr-University Bochum, 44791 Bochum, Germany
3 Institute of Pathology, Ruhr-University Bochum, 44789 Bochum, Germany
4 MVZ Dermatopathology, 88048 Friedrichshafen, Germany
* Correspondence: t.gambichler@klinikum-bochum.de

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Dear Editors: Giant congenital melanocytic naevus (GCMN)-associated melanoma in adults is very rare [1]. We here report an adult male with fatal GCMN-associated melanoma despite the initiation of combined immunotherapy (Figure 1).

Figure 1. An adult male with congenital giant melanocytic nevus presenting as huge brownish-black partly hairy bathing trunk nevus including several tumorous nodules (arrow head, (a,b)). Moreover, there were countless satellite nevi on the entire body.

A 28-year-old male refugee from Syria presented with increasing abdominal pain over a three-week period. The patient as well as his family had no history of cancers. Dermato-
logical examination revealed a huge CGMN presenting as huge brownish black, partly hairy bathing trunk nevus including several tumorous nodules. There were countless satellite nevi on the entire body. Moreover, there were huge skin folds with neurofibromatosis-like appearance (Figure 2) [1].

Abdominal and thoracic computed tomography revealed a partly necrotic abdominal/retroperitoneal tumor conglomerate with displacement of the urinary bladder. Evidence for metastatic disease was also detected intrapulmonal, mediastinal bihilar, hepatic, and osseal. Cranial magnetic resonance imaging, including axial pre- and post-contrast T1-weighted images, did not reveal signs for neurocutaneous melanosis or metastatic disease. However, serum lactate dehydrogenase and S100B were significantly elevated with 486 U/L (135–225) and 2.66 µg/L (<0.15).

Histopathology of two nodules excised from the lower back and right buttock revealed similar findings, including dermal monomorphic small epithelioid and spindle cell shaped melanocytes and melanophages, reaching down to the subcutaneous tissue. Underneath there was a partly necrotic nodule including large melanocytes with atypia, prominent nucleoli, and increased and atypical mitoses (Figure 3a–c).

The entire lesion was strongly positive for S100B and Melan-A, whereas the deep atypical part also showed in Ki-67 positive melanocytes loss of 5-hydroxymethylcytosine and H3K27me3 expression (Figure 4).
Figure 3. Histopathology of two nodules excised from the lower back and right buttock revealed similar findings, including dermal monomorphic small epithelioid and spindle cell shaped melanocytes and melanophages, reaching down to the subcutaneous tissue (a). Underneath there was a partly necrotic nodule including large melanocytes with atypia, prominent nucleoli, and increased and atypical mitoses (b,c).

PD-L1 expression was <1%. Mutation analysis did reveal wildtypes for BRAF, NRAS, and KIT genes. Immunotherapy with ipilimumab (3 mg/kg body weight) and nivolumab (1 mg/kg body weight) every three weeks was recommended. However, 14 days following the first immunotherapy cycle the patient deceased from rapid tumor progression and septic complications.
A nevus with a projected adult size greater than 40 cm is classified as GCMN, as in our case. The clinical risk factors associated with the development of melanoma in patients with GCMN include childhood, truncal involvement, numerous satellite nevi, and nevi located over the posterior axis [2]. Scard et al. [3] recently showed in a systematic review that most melanomas in patients with congenital melanocytic nevi occur early in life. In fact, a higher risk of melanoma onset was observed in the first year of life, in the groups defined by small or medium as well as large or GCMN. For the large and giant categories, a higher risk before 3 years of age was already known [2]. Hence, melanoma development in adult patients with GCMN is considered a rarity [2,3]. Moreover, melanomas arising in congenital melanocytic nevi appear as a dermal/deep-seated melanoma with histopathologic features difficult to distinguish from cutaneous melanoma metastasis or proliferative nodules which often represent a diagnostic challenge given a close resemblance to malignant melanoma [3,4]. In the present case, however, it was not difficult to diagnose melanoma in the deep part of the GCMN using routine histopathology. Moreover, the proliferating melanocytes in the deep nevus part showed loss of 5-hydroxymethylcytosine and H3K27me3 expression which is a hallmark for melanoma [5–7]. According to recent research, NRAS gene mutation is the main driving factor in GCMN [8]. Due to the absence of druggable mutations and tumoral PD-L1 expression, we initiated combined immunotherapy which could unfortunately not prevent the early fatal outcome in the present case.

Conclusively, we reported this tragic case because of the rarity of adult-onset melanoma in GCMN and to highlight the need for melanoma assessments and rigorous follow-up in adult patients with GCMN as well.

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References

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