Multiple Nevus Achromicus Lesions: A Rare Entity

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ABSTRACT
Nevus achromicus is an uncommon birthmark characterized by a well-defined hypopigmented patch resembling a splash of paint. It is usually noted at birth or in early childhood. They do not progress and are mostly seen on the trunk. We report a 20-year-old young man with multiple nevus achromicus scattered over his trunk and limbs. Nevus achromicus is usually solitary, but sometimes it could be multiple wherein the distribution is segmental. Hence, this is a rare case of generalized multiple nevus achromicus and so worth reporting.

Keywords: Depigmentosus, Generalized multiple nevus achromicus, Hypopigmented macule.

INTRODUCTION
Nevus depigmentosus or achromicus is a congenital pigmentary disorder that can occur in all sexes and races.¹ It presents as a single large irregular area of hypo- or depigmentation in childhood and is static. It is strictly unilateral and usually limited to a single dermatome.¹ Sometimes, the lesions may be small and multiple, but these are still unilateral and segmental.²

Nevus depigmentosus is caused by functionally defective melanocytes and abnormal melanosomes.² Localized solitary lesions need to be differentiated from conditions like nevus anemicus, ash leaf macule, and vitiligo. Generalized lesions should be differentiated from hypomelanosis of Ito.² Treatment is usually not required although cosmetic camouflage, phototherapy, and grafting can be done.¹

CASE DESCRIPTION
We report the case of a 20-year-old young man born to non-consanguinely married couple. He came with multiple white patches on the skin, most of which were present since birth. Some of the patches had developed during his childhood. The patient was asymptomatic and had no history of developmental delay/anomalies. There was no history of similar lesions over the oral or genital mucosa and no history of any other skin lesions,

On examination, multiple well-defined hypopigmented macules and patches distributed over the back, trunk, and upper and lower limbs were seen (Figs 1 to 3).

Based on the clinical findings, a differential diagnosis of ash leaf macules, multiple nevus anemicus, or achronamic was considered.

Diascopy test ruled out nevus anemicus. Absence of angiofibromas, shagreen patch, or history of epilepsy in the patient or family members ruled out tuberous sclerosis.

Hypomelanosis of Ito was ruled out as it is a unilaterally occurring condition.

A Wood’s lamp examination showed no accentuation of the lesions, ruling out vitiligo.

Hansen’s disease/leprosy was ruled out as most of the lesions were asymptomatic (no loss of sensations, sweating, or appendages) and present since childhood.

Dermoscopy showed normal skin markings, presence of well-defined margins, normal looking hair, and yellow and white dots in polarized light. No vessels were seen and loss of pigment network was also observed (Fig. 4).³

Histopathology of the involved skin showed a normal number of melanocytes and was not suggestive of anything specific.

Based on the abovementioned exclusions, the patient was diagnosed with multiple nevus achronamic.

DISCUSSION
Nevus depigmentosus or achronamic or achronic nevus is an uncommon disorder of pigmentation where an infant
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or a child develops a single large irregular area of hypo- or depigmentation. Nevus depigmentosus (ND) has serrated borders and is nonprogressive.

It needs to be differentiated from other similar-looking hypopigmentary skin conditions like vitiligo, nevus anemicus, hypomelanosis of Ito, ash leaf macules of tuberous sclerosis, and indeterminate leprosy. It requires no treatment except when patient asks for it, for a better esthetic appearance.

There are three clinical variants:

- Single well-defined lesion – the most common form
- Segmental (dermatomal) nevus achromicus
- Systematized form with whorls or streaks that are predominantly unilateral resembling hypomelanosis of Ito

Based on the size, the following classification can be made:

- Type I—size less than 10 cm².
- Type II—simplex form—size more than 10 cm².

Our patient did not fit into any of the abovementioned three clinical types. He had multiple lesions that were generalized and asymmetrical. Considering the rarity of this kind of nevus achromicus, it was thought to be worthy of reporting.

REFERENCES