Localized vitiligo occurring in an old biopsy scar: A case report

Kevin Rychik, BS,a Jason Cohen, MD,b and Alan Glass, MDc
New York and Port Chester, New York

Key words: depigmentation; halo nevus; koebner phenomenon; melanocyte; scar; vitiligo.

INTRODUCTION

Vitiligo is a depigmentation disorder of the skin, hair, or mucosa, which reveals white macules or patches caused by destruction of pigment-producing melanocytes.1 A complex interaction between genetic, environmental, and self-destructive factors generates this form of response.2 An initiating environmental event such as severe sunburns is thought to cause substantial stress in an already susceptible melanocyte, which elicits an autoimmune response in genetically predisposed individuals.3 Vitiligo’s increased prevalence in immediate relatives has been established. Vitiligo affects 1% of the general population, whereas the risk of an affected patient’s sibling and monozygotic twin for developing the disease is 6.1% and 23%, respectively. There is an increased risk of developing other autoimmune diseases as well, such as type 1 diabetes, autoimmune thyroiditis, pernicious anemia, and Addison disease. These frequencies suggest a nongenetic component plays an important role in the pathogenesis of the disease, in addition to a genetic predisposition.5

Oxidative stress is an environmental risk factor that could play a role in causing vitiligo. Stress can result from overproduction of pro-oxidant species or creation of antioxidant chemicals via a reduction reaction.6 Melanocytes are constantly being exposed to stressors, including ultraviolet radiation and harmful chemicals and pathologic conditions such as inflammation and cancer, which increase production of reactive oxygen species (ROS). Healthy melanocytes are capable of lessening these stressors, whereas melanocytes from patients with vitiligo are more vulnerable. Decreased catalase, a key enzyme that degrades hydrogen peroxide and catalyzes the creation of other antioxidant agents, glutathione peroxidase and glucose-6-phosphate dehydrogenase, protects melanocytes from self-destructive ROS. Decreased levels of catalase have been found in melanocytes of patients with vitiligo.7 This finding proposes a present oxidation-reduction defect involved in the disease. Studies found patients with vitiligo experience forms of stress, triggering the immune system to attack the body’s own melanocytes.

The pathogenesis of role of the Koebner phenomenon in the development vitiligo is still unknown, but its presence has been established.8 It has been suggested that trauma to the skin may cause hydrogen peroxide to accumulate. This accumulation may cause proteins such as heat shock protein, that are designated to protect the cell from damage, to then become immunogenic, therefore eliciting an autoimmune response.9 Many people report first noticing their vitiligo after traumatic events, periods of stress, or severe sunburns.10 We report the case of localized vitiligo occurring in an old biopsy scar from 25 years prior.

CASE REPORT

A 33-year old man presented with tenderness and rapid depigmentation over several weeks of an old biopsy scar on his back. At age 9, the patient had a mole biopsied in the exact location, which led to scarring. The mole was determined to be benign. The patient denied a change in scar pigmentation until recently when the area turned white. The

From the Sackler School of Medicinea; Dermpath Diagnostics, Port Chesterb; and 57 West Dermatology.c

Funding sources: None.

Conflicts of interest: None disclosed.

Correspondence to: Alan Glass, MD, 57 West Dermatology, 57 West 57th St, Ste 1109, New York, NY, 10019. E-mail: Skindr57@gmail.com.

JAAD Case Reports 2020;6:326-8.
2352-5126
© 2020 by the American Academy of Dermatology, Inc. Published by Elsevier, Inc. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/).
https://doi.org/10.1016/j.jdcr.2020.02.003
change in pigmentation and pain concerned him. He
denied picking or scratching of the area, and denied
pruritus. The patient reported a history of excessive
sun exposure and chronic sunburns in the area but
denied any recent sunburns. The patient denied a
medical history of autoimmune diseases. The patient
also denied a family history of autoimmune disease,
but reported malignant melanoma in an aunt and
basal cell carcinoma in his mother.

Physical examination found a nonscaly depig-
mented patch occurring in an old biopsy scar
involving medial scapular back. No halo nevi or
evidence of vitiligo was noted in the area. A careful
inspection and palpation of the area was performed,
and skin was of normal temperature, turgor, and
texture. There were no suspicious skin lesions,
rashes, or depigmented areas noted on a total skin
examination.

A differential diagnosis of halo nevus was made
initially, and a shave biopsy of the patch was
performed. Bacitracin and a bandage were applied
to the site. The patient was instructed to apply
Bacitracin twice a day for 2 to 3 days and to begin
clobetasol propionate cream twice a day once the
biopsy site healed. The patient was lost to follow-up.

The shave biopsy showed a proliferation of
fibroblasts aligned parallel to the skin surface inter-
posed among linearly arranged thickened collagen
bundles and small blood vessels. Most notably, there
was an absence of melanocytes and melanin
pigment within the epidermis extending over and

slightly beyond a zone of fibrosis. The clinical
diagnosis was most consistent with recent vitiligo
occurring in an old biopsy scar. See Figs 1 through 4.

DISCUSSION

Current evidence suggests vitiligo is an autoim-
mune disease with a genetic predisposition that can
be triggered by nongenetic environmental risk
factors. Increased prevalence among immediate
relatives has been established. In addition, risk
factors including ultraviolet radiation and harmful
chemicals have been found to increase the produc-
tion of harmful ROS, when combined with decreased
catalase in susceptible melanocytes, and have been
found in patients with vitiligo. A study on vitiligo by
Deo et al10 found that many patients first notice
their vitiligo following severe sunburns. This finding
supports the belief that the vitiligo could be caused
by decreased catalase and severe sunburns. A severe

Fig 1. Clinical photo of a nonscaly depigmented patch occurring in an old biopsy scar in the medial scapular back.

Fig 2. Absence of melanin pigment and melanocytes within the epidermis overlying a zone of fibrosis. (Hema-
toxylin-eosin stain; original magnification: ×10.)

Fig 3. The absence of melanin pigment within the epidermis is highlighted with a Fontana-Masson stain for melanin pigment. (Original magnification: ×10.)
sunburn can cause local trauma to the skin, and the development of vitiligo could be explained by the Koebner phenomenon. However, in contrast, an old biopsy from 25 years prior to the incidence of vitiligo has not been reported, to our knowledge, as a possible triggering environmental event. Furthermore, a family history of autoimmune disease was not established in this patient.

Although a genetic predisposition of autoimmune disease or catalase levels could not be established because of a history of excessive sunburns in the area and excessive scar tissue resulting from the biopsy scar, a local autoimmune attack on melanocytes could have occurred from a combination of these factors.

**REFERENCES**

1. Boissy RE, Manga P. On the etiology of contact/occupational vitiligo. *Pigment Cell Res*. 2004;17(3):208-214.
2. Rashighi M, Harris JE. Vitiligo pathogenesis and emerging treatments. *Dermatol Clin*. 2017;35(2):257-265.
3. Shen C, Gao J, Sheng Y, et al. Genetic susceptibility to vitiligo: GWAS approaches for identifying vitiligo susceptibility genes and loci. *Front Genet*. 2016;7:3.
4. Taieb A, Picardo M. Clinical practice. Vitiligo. *N Engl J Med*. 2009;360(2):160-169.
5. Alkhateeb A, Fain PR, Thody A, Bennett DC, Spritz RA. Epidemiology of vitiligo and associated autoimmune diseases in Caucasian probands and their families. *Pigment Cell Res*. 2003;16(3):208-214.
6. Denat L, Kadekaro AL, Marrot L, Leachman SA, Abdel-Malek ZA. Melanocytes as instigators and victims of oxidative stress. *J Invest Dermatol*. 2014;134(6):1512-1518.
7. Schallreuter KU, Wood JM, Berger J. Low catalase levels in the epidermis of patients with vitiligo. *J Invest Dermatol*. 1991;97(6):1081-1085.
8. Chesner J, Levin MK, Marmur ES. Koebnerization phenomenon after broadband light therapy in a patient with cutaneous sarcoidosis. *JAAD Case Rep*. 2017;3(4):306-309.
9. Sanghavi SA, Dongre AM, Khopkar US. Koebnerization and generalized spread of vitiligo following radiotherapy. *Indian Derm Online J*. 2013;4(2):147-148.
10. Deo SS, Bhagat AR, Shah RN. Study of oxidative stress in peripheral blood of Indian vitiligo patients. *Indian Derm Online J*. 2013;4(4):279-282.

**Fig 4.** The absence of melanocytes within the epidermis is highlighted with a Melan-A immunoperoxidase stain. (Original magnification: ×10.)