Investigative Dermatology and Venereology Research

A Study of Halo Naevi, Vitiligo and Asymptomatic Autoimmune Thyroiditis

Alin Laurentiu Tatu1*, Victor Gabriel Clatici2, Marius-Anton Ionescu3, Violeta Corina Cristea4

1Department of Dermatology, Faculty of Medicine and Pharmacy, Dunarea de Jos University, Galati
2Department of Dermatology, Elias Emergency Hospital, Bucharest, Romania
3Dermatology Outpatient Clinic, University Hospital “Saint-Louis”, Paris, France
4Carol Davila University and Central Reference Laboratory Synevo, Bucharest, Romania

*Corresponding author: Tatu Alin Laurentiu, Department of Dermatology, Faculty of Medicine and Pharmacy, Dunarea de Jos University, A.I. Cuza street no 39, Galati, Postal code-800101, Romania, Tel: 00-40-728267435; Fax: 00-40-236-415705; E-mail: dralin_tatu@yahoo.com, dr.toni.ionescu@gmail.com


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Background

Halo Naevus (HN) is a dermatologic condition characterized by a typical whitish rim around an existing melanocytic congenital or acquired nevus, it was first described in 1916 by Sutton[1]. Halo naevi are common in children and young adults, with a mean age at onset of 15 years. The incidence in general population is estimated to be approximately 1%[2]. Affected individuals frequently have multiple lesions which are usually localized on the back. HN is an autoimmune response and T lymphocytes are considered to play a key role in the progressive destruction of naevus’ cells. Multiple HN is rather rare and this could suggest a higher risk of vitiligo or of autoimmune disease development comparing to solitary HN lesion risks. The mechanism of halo phenomenon is still non completely defined and is suggested to be induced by a destruction of melanocytes by immune response of cytotoxic T cells[3] or due to IgM autoantibodies induction[4].

Purpose

We sought to assess the association of Halo Naevi (HN) or of HN-vitiligo with thyroiditis by screening markers of thyroiditis in patients presenting Halo Naevi (HN) and HN associated with vitiligo.

Materials and Methods

In this study were included adults patients with halo naevus and / or vitiligo presented in dermatology outpatient clinic between 2010 and 2016. Clinical and dermoscopy examination were performed. The possible association between vitiligo and thyroiditis was assessed by peripheral blood dosage of Anti-Thyroid Peroxidase (ATPO).

Results

Were included 37 patients, 16 female and 21 male. Twenty five patients (67.56%) had one HN, 7 patients(18.91%) had 2 HN and 5(15.31%) of the patients had 3 HN. From 37 patients with halo naevus 10 (27.02%) had associated vitiligo. In all cases vitiligo occurred 3 - 4 years after the onset of HN. In all 10 patients with HN and associated vitiligo, lesions were of non-segmental type: limited, symmetric, small and large lesions of extremities. Among them five had one HN (4 female and 1 male), 2 patients had 2 HN (one male and one female), 3 patients had 3 HN.

In 27 (72.98%) patients with HN ( and no vitiligo): ATPO peripheral blood levels were normal in 26 patients (96.29%), a 13 years old boy had high ATPO (presented in the discussion) (3.71%).

In 10 patients with HN and associated vitiligo: 3 had high ATPO results (33.33% of patients,1 male with 2 HN and 2 female with 3 HN) with a 2/1 female/male ratio. One of female with 3 HN had a history of autoimmune thyroiditis and vitiligo - both occurred after the onset of HN (by order thyroiditis, HN then vitiligo). In 2 others cases with high ATPO clinical appearan-
ance of the conditions was HN, vitiligo and thyroiditis. Vitiligo features were in all cases of type non-segmental with bilateral hypo-chromic, symmetrical small macules (0.3 - 0.8 cm diameter) and Koebner phenomena or large plaques located on extremities (Figure 1).

**Figure 1:** Vitiligo of the hand.

**Discussion**

Thirty eight patients were initially included. One patient was excluded as he presented one lesion that was not a halo nevus but a melanoma with peripheral regression send to surgery and oncology. Vitiligo and halo naevi are both pigmentary disorders of the skin characterized by the acquired loss of functional epidermal melanocytes manifesting as white macules and patches. The cellular mechanisms and biochemical changes that result in the appearance of these two types of achromic lesions and the direct relationship between vitiligo and halo naevi are still subject of discussion. The tendency of different morphologic alterations in mitochondria from perilesional vitiligo skin and from perilesional halo nevi skin reflect heterogeneous backgrounds between the two diseases, revealing that vitiligo and halo naevi may have separate pathogenic mechanisms.

There is strong evidence that immunological mechanisms are at the base of melanocytic destruction: the presence progressive destruction of of an inflammatory lymphocytic infiltrate that leads to progressive destruction of naevi cells; the presence of antibodies against nevi cells[5]. Recent studies highlighted similarities and differences between events that occur in regression of melanoma, vitiligo and in Sutton’s phenomenon[6-8].

The association of vitiligo with autoimmune thyroid diseases and the increased prevalence of autoantibodies including thyroid autoantibodies in vitiligo had been proven by several recent studies[9-11]. In Sutton’s nevi was usually described an inflammatory cellular infiltrate surrounding symmetrically the naevus, consisting mainly of T-lymphocytes. B-lymphocytes, Langerhans cells and macrophages are absent or rarely observed[8]. Thyroiditis is defined histologically by the presence of a diffuse lymphocytic infiltrate in the thyroid gland, with the formation of lymphoid follicles. Cell mediated and antibody mediated immune responses lead to the destruction of thyroid cells[10]. In patients with multiple HN, the risk of vitiligo and other autoimmune diseases seems to be higher than in pediatric patients with a single HN. The female/male ratio is 5 - 10/1 in autoimmune thyroiditis and the onset of disorder usually takes place between the third and sixth decade of life. Due the fact that halo naevus’ onset is usually earlier than vitiligo, clinicians should pay particular attention even to patients presenting children/boys with one or several HN. The follow-up of HN is clinical and by dermoscopy, very high photo-protection is necessary (SPF 50+) for HN of photo exposed areas. Non-segmental, stable, limited vitiligo management is most frequently made by potent topical steroid therapy, topical calcineurin inhibitors, or narrow-band UVB 311 nm or associations phototherapy by UVA and psoralen (PUVA) (phototherapy is contraindicated in areas where vitiligo is associated with HN), excimer laser 308 nm.

A 13 years old boy with congenital melanocytic naevi that was referred to dermatology presented a recent white patch around one naevus. Clinical evaluation revealed a whitish rim around an congenital melanocytic nevus from the lumbar area, without cutaneous induration, resembling a halo nevus (Figure 2).

**Figure 2:** Halo nevus. Clinical image.

Dermoscopy revealed a mixed pattern composed of a central network, structureless light brown-gray area and some small brown globules in periphery (Figure 3)[12]. There was no family or personal history of vitiligo or autoimmune diseases. Anti-Thyroid Peroxidase (ATPO) levels were 107.6UI/mL (normal values 0 - 34UI/mL), Thyroid Stimulating Hormone (TSH) was 3,2 uUI/mL (normal values 0,4 - 4 uUI/mL), free thyroxine (fT4) was 1,14 ng/dL (normal values 0,98 - 1,63 ng/dL). Thyroid ultrasound showed heterogeneous thyroid echo texture. Anti-thyroglobulin and fT4 were not performed but the patient was referred to the endocrinologist for specialty investigations and monitoring.

**Figure 3:** Halo nevus. Dermoscopy 10 x magnification.
Conclusions

In a series of 37 halo naevi, ten cases (27, 2%) had an associated vitiligo, among them 3 (33.33%) had peripheral blood levels of Anti-Thyroid Peroxydase (ATPO). One case from 27 patients with HN without vitiligo (3.70%) had high levels of ATPO and ultrasound sign of thyroiditis. In patients with one or several HN, a systematic dermatoscopy and endocrine assessments can reveal associated autoimmune diseases as vitiligo and thyroiditis but halo naevi and vitiligo patients have more frequent high levels of ATPO (33.33%) than those with just halo naevi without vitiligo (3.70%).

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References