An Overview of Quality of Life in Vitiligo Patients – At Glance

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Abstract
Vitiligo is an acquired depigmenting disorder characterized by white maculae on the skin associated with social stigma. Vitiligo has major impact on the quality of life due to its high cosmetic value, chronic nature of disease and long-term ineffective treatments. Vitiligo patients suffer from low self-esteem and poor body image that affects their emotional and psychological well-being. To investigate quality of life of Vitiligo patients, psychological impact of unpredictable course of disease and long term treatment among South Indian population. A total of 155 Vitiligo patients, undertaking treatment in dermatology clinic at the Central Research Institute of Unani Medicine (CRIUM), who completed the Dermatology Quality of Life Index (DLQI) questionnaire, participated in this study. Patients with Vitiligo averaged at about 27.4% of maximum psychological impairment as measured by the DLQI summary score. Disability in patients was highest in treatment category with 54.6% of impairment followed by symptoms and feelings (41.83%).

Keywords: Vitiligo; Quality of life; DLQI

Introduction
Vitiligo, also known as leukoderma, is an acquired depigmentary disorder characterized by the appearance of white patches resulting from the loss of functional melanocytes and melanin from the skin [1]. Vitiligo is an acquired, idiopathic, and worldwide common depigmentation disorder with an estimated prevalence from 0.1 to 8%. These numbers are based on clinical population studies and field research examining inhabitants of geographically enclosed areas. The often cited prevalence of 8% could not be confirmed after excluding clinical patient populations. Accordingly, the worldwide prevalence of Vitiligo ranges between 0.5 and 2% [2]. Adults and children of both sexes are equally affected; the proportion of patients with a positive family history varies from one part of the world to another, with particularly wide ranges reported in India (25-18%), with reports of up to 40% elsewhere in the world [3]. Some dermatological outpatient records show the incidence of vitiligo to be 3% to 4% in India although an incidence as high as 8.8% has also been reported [4]. The exact aetiology of Vitiligo is unknown, but four main theories exist to explain it: the autoimmune hypothesis, the neural hypothesis, the self-destruct hypothesis, and the growth factor defect hypothesis, but none satisfactory. It is believed that Vitiligo is a polygenic trait and that a convergence theory, combining elements of different theories across a spectrum of expression is the most accurate aetiology [5]. Vitiligo is not a physically damaging disease; other than an increased sensitivity to UV radiation most of the disease effects are social and psychological, especially for dark-skinned races. There are both surgical and non-surgical treatments for Vitiligo but fails to give cosmetically acceptable pigmentation [6].

Vitiligo is an acquired depigmenting disorder characterized by the loss of functional melanocytes from the epidermis [7]. Vitiligo is in most cases diagnosed clinically, and so obvious are the features of a well-developed lesion of this disease. Other investigations are rarely required. However, occasionally Vitiligo especially in its early evolving stage or in some of its localised expressions may pose diagnostic difficulties as the lesions are hypo pigmented rather than depigmented and other differential diagnoses may be considered [8]. Vitiligo is of great cosmetic concern since the change in appearance caused by Vitiligo can affect a person’s emotional and psychological well-being. Since ancient times patients with Vitiligo suffered the same mental abuses as lepers. In India young woman with Vitiligo have little chance of getting married. Appearance of skin can condition an individual self-image, and any pathological alteration can have psychological consequences. Implications of Vitiligo ruin young girls who are almost condemned of spinsterhood and at many occasions these patients are denied of first line jobs [9]. Vitiligo is thus an important skin disease having major impact on the quality of life; therefore we intended to study the psychological impairment in different aspect of life in Vitiligo patients. This analysis describes self-reported dermatological life quality index (DLQI) of patients with segmental and non-segmental Vitiligo.

Methodology
A total of 155 patients undergoing treatment for Vitiligo in Dermatology clinic at the Central Research Institute of Unani Medicine (CRIUM) consented for the study and completed the Dermatology Life Quality Index (DLQI) questionnaire (with permission from Wales College of medicine, Dermatology department). Study subjects were over 16 years of age, of either sex, with Segmental Vitiligo (SV) or Non-Segmental Vitiligo (NSV) types. The study was approved by the CRIUM ethical committee. DLQI questionnaire measures six disease-specific domains, i.e., symptoms and feelings, leisure activities, daily activities, work and school, personal relationships, and treatment. Individual items are summed to generate an overall DLQI score [10]. Scores are expressed as percentages, with higher percentage indicating

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greater impairment in the quality of life. Results were reported for the total group and for subgroups.

**Statistical Analysis**

Data were stored and analysed using SPSS 19 (Statistical Package for Social Science; release 19.0). Chi-square test and Fishers exact test were used and a P-value <0.05 was considered as significant.

**Results**

The mean age of patients is 32.4 ± 13.9 and having a mean chronicity of 3.09 ± 5.01 yrs. The demographic characteristics of the total analysis group and the subgroups are described and DLQI scores are displayed. Patients with Vitiligo averaged at about 27.4% of maximum impairment as measured by the DLQI summary score. Vitiligo patients scored highest in the treatment component of the DLQI, which measures pain, stress and difficulties faced in getting the right kind of treatment, with 56.1% of impairment reported. This was followed by symptoms and feelings, with 41.83% of impairment measuring the embarrassment and suffering associated with the disease. Impairment in work and school, and leisure activities ranged from 9.9% to 17%. Males and females were equally affected (8.25 ± 5.0 vs 8.13 ± 4.69, p=0.8). Patients with non-segmental type consistently scored higher (worse) on all components of the DLQI. Dark skinned individuals with Vitiligo had maximum total impairment of 31.3% when compared to other group. Unmarried Patients showed greater impairment in symptoms and feeling and leisure component of DLQI when compared to married patients. Patients with chronicity greater than 5 yrs averaged 29.6% of maximum disability which is higher (worse) than those below 5 yrs 27.1% and also had the maximum score in treatment component 76.5%. Slow responders (repigmentation less than 40% after one year of treatment) showed greater impairment in treatment section of DLQI 62.3% compared to rapid responders 26.19%.

**Discussion**

Vitiligo can have a significant effect on the psychological well-being of the patient [11]. This is especially true for darker skinned patients as the contrast between the pigmented and depigmented skin can be quite drastic. In some cultures there is a stigma attached to having Vitiligo. Those affected with the condition are sometimes thought to be evil or doomed to certain death. It is often perceived as a source of embarrassment and the frustration of resistant lesions over exposed part of hands and feet can lead to anger and disillusionment.

In the present study scores on the DLQI ranged from 0-22 (mean ± SD,8.21 ± 4.48), indicating poor quality of life in South Indian Vitiligo population when compared to the scores on the DLQI of Persian population that ranged from 0 to 24(mean ± SD,7.05 ± 5.13) [9]. The mean DLQI score (8.21) in this study was higher than that reported in France (7.17), Belgium (6.9) and that of patients with Acne vulgaris (7.45), but lower than Psoriasis (10.53) or Atopic dermatitis (11.20) [14,15]. The mean DLQI score in this study was also higher than that obtained mean 7.3 and Kent and Al-Abadie’s [16] study mean 4.82 but lower than Parsad et al. [17] mean 10.6 this difference is due to the fact that south India has higher literacy rate and different culture as compared to north India). The association between poor QoL and depressive symptoms suggests that healthcare professionals should watch for possible depression in high-risk patients. Frequent complaint of embarrassment, despite low rates of poor self-esteem, suggests that while most patients cope well with their appearance, they continue to encounter unpleasant social interactions. Increasing public awareness about vitiligo to reduce unnecessary attention may benefit more than self-esteem management for vitiligo patients in South India [18].

Patients with non-segmental type of vitiligo reported more impairment than corresponding subgroups because of irregularly spread depigmented patches on the exposed part of the skin. Weiss et al compared the difficulties faced by patients with vitiligo with those with leprosy in India [19]. Salzer and Schallreuter reported that 75% patients found their disfigurement moderately or severely intolerable. These findings also support the relationship between stress and the development of vitiligo. Al-Abadie et al. [20] indicated that psychological stress increases levels of neuroendocrine hormones that affects the immune system and alters the level of neuropeptides, which may be the initial steps in pathogenesis of vitiligo. The chronic nature of disease, long term treatment, lack of uniformity, high cost and unsatisfactory pigmentation of therapy are usually very demoralizing for patients suffering from Vitiligo. Compliance for regular long term visits for PUVA/UVB therapy; side effects of immunosuppressive therapies are other limitation in treatment of Vitiligo patients. Interestingly in this study, an encouragingly huge section (92.46%) of the patients responded to various degrees to the Unani treatment given to them at CRIUM and in most cases, the disease was found to limit itself for about 10 years after onset. Data collected from case records of these patients showed that the response to Unani treatment (with a mean 46% repigmentation) was remarkably better when compared to Ayurveda (12%), Homeopathy (5.6%) and Allopathy (5.8%) treatment.

Our results show that in Vitiligo patient’s quality of life is poor and it is strongly associated with their social well-being. Patients with high DLQI responded less favourably to given treatment indicating that psychological counselling and comfort could obtain a better treatment outcome. Papadopoulos et al. [21] have shown that counselling can help to improve body image, self-esteem and quality of life of patients with Vitiligo, also having positive effect on the course of the disease. A recent study which measured the frequency and number of stressful life events occurring over a specified period among Vitiligo and alopecia areata suggested that such patients endured a significantly higher number of stressful life events than do controls suggesting that psychological distress may have contributed to the onset of Vitiligo and alopecia areata [22]. Hence improving quality of life in this group should be taken as an important task by the dermatologist while treating Vitiligo patients.

**Acknowledgement**

We are grateful to staff and subjects for help in compiling the data and senior dermatologist at CRIUM Hyderabad, Dr. M.A. Waheed for his kind cooperation.
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This article was originally published in a special issue, Vitiligo News handled by Editor(s). Dr. Yan Valle, Vitiligo Research Foundation, USA