The psychological Burden of Hair Disease

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Abstract

Background: Hair diseases have an important burden on patients’ life, causing significant emotional and psychosocial distress. However, the comparison between the impairment due to different hair conditions, such as alopecia areata (AA) and androgenetic alopecia (AGA), has rarely been done.

Objective: The aim of this study was to assess the psychological burden of sub-group of patients with different hair diseases and to compare them to a healthy population.

Methods: This study is part of a large multicenter study including 3635 adult dermatological out-patients and 1359 controls from 13 European countries.

In the sub-group of patients with hair diseases, we analyzed the sociodemographic characteristics, the stress level, and the impact of hair diseases on quality of life (QoL), anxiety, and depression and we compared them among patients with AA, AGA, and healthy controls.

Results: The study population included 115 patients with hair diseases. Women suffered more from hair diseases than men (77% vs 23%). Patients with hair diseases had a lower education level than healthy controls (medium educational level: 43% vs 28%). Overall, 41% of the patients reported stressful life events during the last 6 months compared with 31% of the controls. Patients with same age, sex, depression level and comorbidities had a worse QoL when suffering from AA than from AGA (DLQI score: 5,8 vs 2,5).

Conclusion: Our results indicate that hair diseases patients are more anxious, depressed and have a worse QoL than controls. According to the results a psychotherapeutic and psychopharmacological approach should be proposed when necessary.

INTRODUCTION:

In humans, hair is predominantly considered as an aesthetic concern, however hair diseases may lead to an important burden on patients’ life, causing significant emotional and psychosocial distress (1). The most common hair diseases are alopecia areata (AA) and alopecia androgenetica (AGA).

It has been observed that AA significantly reduces patients’ quality of life (QoL) (2). In addition, AA patients have a high prevalence of psychiatric comorbities, such as anxiety, depression, social phobia, and personality disorders (3, 4). It is therefore controversial to what extent the impact of this impairment is comparable to other skin diseases.

In an Indian study on a wide range of etiologies including chemotherapy related alopecia, Sellami et al have shown that alopecia patients were more anxious and depressed than controls (5). In 50 AA patients, 62% had symptoms of anxiety and 38% of depression (5). These percentages were significantly higher than in the control group. In patients with AGA a significant impact on QoL was found, particularly when associated with high severity, a longer duration, younger age, having received previous non-medical hair care, and hospital visits for AGA treatment (6).

Compared to other skin conditions, which may be accompanied by itching, pain or other sensations that provide distinctive symptoms, in most hair diseases the patients generally only experience the symptom of “hair loss” and may help the patients to distinguish between different diseases. Considering this mono-symptomatic aspect of hair diseases, it is interesting to investigate other aspects that may differentiate the impact of different hair diseases (AA, AGA, other diseases).

The aim of this study was to assess the psychological burden of patients with hair diseases in a large European sample. For this purpose, we compared different subgroups of patients with hair diseases with a healthy control population, regarding the level of anxiety, depression, stress, and QoL impairment.

MATERIAL AND METHODS:

*Study population*

A multicenter, observational, case-control study was conducted in 13 European countries. Material and methods have been described in detail in a previous paper (7). In brief, in the dermatology clinics of each country, patients with all dermatological diseases were consecutively recruited. Controls were recruited in all centers among the hospital staff who did not currently had or previously have had any dermatological disorder. Both cases and controls had to complete questionnaires on QoL and psychological comorbidities. Data were complete for 3635 patients and 1359 controls. In the present study, we have included patients with hair diseases and healthy controls.

*Questionnaires*

The first part of the questionnaire recorded socio-demographic variables including age, gender, marital status, and self-reported socio-economic status. Ethnicity was self-reported by each participant referring to their own country of birth.

The presence of symptoms of depression and anxiety was assessed with the HADS, a well-validated instrument showing good psychometric properties, aimed to assess symptoms of anxiety disorders and depression in somatic, psychiatric, and primary care patients, as well as in the general population. HADS includes seven items assessing anxiety and seven assessing depression, each with four possible answers. For each dimension of anxiety and depression a score from 0 to 7 is considered normal, from 8 to 10 a borderline case, and from 11 to 21 a case in need of further examination or treatment. This instrument was used in the validated translations relevant to the study countries.

To assess health related quality of life (HRQoL), we used 2 different self-rated questionnaires: the Dermatology Life Quality Index (DLQI) and the EQ-5D-3L. The DLQI is a dermatological specific, ten-item questionnaire used to measure the impact of skin disease on the QoL of an affected person. The ten questions cover the following topics: symptoms, embarrassment, shopping and home care, clothes, social and leisure, sport, work or study, close relationships, sex and treatment. Each question is scored from 0 to 3, giving a possible total score range from 0 (no impact of skin disease on QoL) to 30 (maximum impact on QoL).

The EQ-5D-3L essentially consists of 2 parts: the EQ-5D descriptive system and the EQ visual analogue scale (EQ VAS). The EQ-5D-3L descriptive system comprises the following 5 dimensions: mobility, self-care, usual activities, pain/discomfort and anxiety/depression. Each dimension has 3 levels: no problems, some problems, extreme problems. The respondent is asked to indicate his/her health state by ticking (or placing a cross) in the box against the most appropriate statement in each of the 5 dimensions. The EQ VAS records the respondent’s self-rated health on a vertical, visual analogue 100-degree scale (0= worst imaginable health state”; 100= best imaginable health state”).

This information can be used as a quantitative measure of health outcome as judged by the individual respondents. Stress was assessed with the item ‘’Have you had any stressful life event during the last 6 months?’’ (yes/no).

Cases completed all the questionnaires, while controls did not complete the DLQI.

*Statistical analysis:*

Analyses were performed with R version 3.1.0. Baseline characteristics were compared between groups (controls versus any hair diseases; controls versus AA and AGA; men versus women; different age groups) by Fisher’s exact tests for categorical variables and t-tests or ANOVA for continuous variables (or Wilcoxon tests if needed). The categorical scores were compared by Fisher’s exact tests between groups. The DLQI scores and EQ5D scores, considered on continuous scales, were also compared (separately) by multivariate regressions after a logarithmic transformation for DLQI (to fulfill the normality assumption) with the groups, age, sex, depression and comorbidities as predictors (adjusting variables). The HADS scores, considered on a continuous scale, were compared between groups with t-tests or ANOVA (the normality assumption has been checked with QQ plot). The EQ5D scores were compared by Wilcoxon tests, or Kruskal-Wallis non-parametric ANOVA between gender and categorized age, while their five components were compared by Fisher’s exact tests. The population was adjusted by linear model for age, sex, depression and comorbidities. Some results are presented using box-plots which are a convenient way of graphically depicting groups of numerical data through their [quartiles](https://en.wikipedia.org/wiki/Quartile). Box plots may have lines extending vertically from the boxes (*whiskers*) indicating variability outside the upper and lower quartiles, hence the terms box-and-whisker plot and box-and-whisker diagram.

RESULTS:

Details on participants characteristics have been previously published (7). In this study, we included 115 patients with hair diseases and 1359 controls. In the group of patients with hair diseases, there were 37 patients with AA, 20 patients with AGA, and 58 patients with other hair conditions such as lichen planus of the scalp, telogen effluvium, cicatricial alopecia, frontal fibrosing alopecia and others.

Table 1 shows a comparison between the characteristics of the hair diseases group and the controls. In the hair diseases group, there were more females than in the control group; 46% of the patients were educated to a medium education level (i.e., primary or secondary school) compared with 28% of the controls. Finally, tertiary (university) level education was achieved by a smaller proportion of patients than of the controls (29% vs 43%).

Comparing subgroups of patients with hair diseases, we observed that AA and the subgroup of other hair diseases were more likely to have achieved a medium level education than AGA patients (47% and 49%, respectively, vs. 35%), (*p=0.007*). A higher proportion of AA patients were married (70%) compared to the other hair disease population (49%), (*p=0.021*). Stress was reported by 49% of the AA patients and 35% of the AGA and 31% of controls (*p=0.061*).

HADS mean score in patients (*anxiety 7.9; depression 5.4*) was significantly higher compared to controls (*anxiety 5.6; depression 3.6*). Overall, 41% of the patients reported stressful life events during the last 6 months compared with 31% of the controls.

Patients with same age, sex and comorbidities had a higher DLQI score when suffering from AA than AGA, (Fig. 1). Figure 2 (Fig. 2) shows that patients with AA had a higher HADS score compared to AGA population. An ANOVA Test was performed to compare controls, AA and AGA population and confirmed the results above. *(Anxiety p<0.001; depression p=0.005).*

Females had a poorer quality of life (Fig. 3a), especially in the pain dimension (*p=0.001*) and anxiety and depression dimension (*p=0.012).* Patients aged more that 50 years old had lower EQ-5D scores (*p<0.001*) and EQ VAS scores (Fig. 3b), i.e. a more impaired QoL.

Comparison between controls, AA and AGA patients showed a lower EQ-5D score in both AA and AGA patients when adjusted for age, sex, depression and comorbidities (Fig. 3c). Specifically, EQ-5D activity, pain and anxiety and depression dimensions showed lower scores in all hair patients compared to controls (*p=0.002, p<0.001, p<0.001).* Also, EQ VAS score was lower for all hair patients compared to controls when adjusted for age, sex, depression and comorbidities*.* AA patients reported lower self-evaluated health than AGA patients and controls when using the EQ VAS scale (fig. 3d) and had a lower EQ-5D anxiety and depression score (*p<0.001*). All other comparisons did not show any significant differences.

DISCUSSION:

How do hair diseases affect patients’ lives? In this study, we found that having any hair diseases was associated with more anxiety, a more depressed mood and impaired QoL compared to a control group without skin disease. These differences were most pronounced in AA patients.

The special impact of AA is in line with previous studies. Liu et al. showed that HRQoL experienced by patients with AA is similar to that seen in patients with other chronic skin diseases including atopic dermatitis and psoriasis (8, 9). Rencz et al. have also demonstrated that patients with AA experience significant impairment in HRQoL, especially in the area of mental health (2).

AA patients may suffer more than patients affected by other hair diseases for different reasons. AA is a chronic disease and is well known for having a significant impact on the psychological sphere of patients. Indeed, AA is a burdensome medical problem because concealing the symptoms of that condition is relatively difficult, especially for women. [T. Cartwright](https://onlinelibrary-wiley-com.gate2.inist.fr/action/doSearch?ContribAuthorStored=Cartwright%2C+T) showed that QoL was significantly poorer for women with alopecia, compared with men. There were gender differences in several of the DLQI subscales used by the authors. Women were significantly more likely to feel that alopecia affected their leisure activities and personal relationships, while men scored higher on work interference than women (10)

Also, AA has a major impact on the social life of the patient, leading him/her to feel socially alienated and rejected. Furthermore, AA is associated with mental disorders like depression, anxiety disorders, body dysmorphic disorders, social phobia and suicidal thoughts. These mental disorders could be primary conditions which declare themselves in medical problems associated with the skin or these could be a result of this serious and disfiguring condition (11).

An interesting result was the significant difference at the level of the marital status between the AA patients and other hair diseases patients. In fact, 70% of AA patients were in a relationship compared to 49% of the others. In a previous study, we demonstrated that there are more dysfunctional families in AA patients (12). Those dysfunctions could be fusions and ruptures, generational repetition of behaviors of dependence or vulnerability and unsolved mourning if traumatic events are experienced. We could hypothesize that not resolving these problems could lead to increasing the perceived stress and consequently to more anxiety and depression. Moreover, these patients have a lower education level, which is usually correlated with earlier partnerships (13,14). Further studies should be done to confirm this hypothesis.

About the AGA patients, we also found an impact on HRQoL. Their level of anxiety and depression was also higher than for the controls as well. In previous studies, it has been shown that HRQoL of male and female AGA patients is altered, in particular in younger patients (6,13,15). This should not be underestimated and a psychological support could be helpful for these patients.

The weakness of this study is that several generic and dermatology-specific HRQoL instruments have been used, but no specific instruments for AA were used. The newly developed AA-specific measures seem very promising, however, a more extensive assessment of validity and reliability is needed.

In conclusion, patients with hair diseases experience a considerable impact on their QoL and may benefit from an assessment at psychological level. We recommend the use of anxiety and depression questionnaires like HADS, which is specific and user-friendly. Some caution remains warranted: we must take into account the risk of biases such as underestimation with self-administered questionnaires; moreover, AA patients are often alexithymic (16) and therefore, have problems to articulate their affections. We therefore recommend to let the patient fill the questionnaire on his own and evaluate the answers with the dermatologist. Also the observation of patients’ nonverbal behavior is an invaluable tool to assess the presence of depression. Finally, once an adequate psychological assessment has been performed, psychotherapeutic and psychopharmacological support should be offered to patients whenever such support appears to be conducive to a better outcome.

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Table 1: Comparison between hair diseases groups and controls. Mean values given with standard deviation (SD)

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| VARIABLE | LEVEL | N (%) Hair diseases | N (%) controls | P-value |
| Sex | Male | 27 (23%) | 453 (33%) | P=0,03 |
|  | Female | 88 (77%) | 903 (67%) |  |
|  |  |  |  |  |
| Educational level | Medium | 49 (46%) | 375 ( 28%) | P<0,001 |
|  | High | 26 (25%) | 399 (30%) |  |
|  | University | 31 (29%) | 577 (43%) |  |
| Have you had any  stressful life events during the last 6 months?” (Yes/no) |  | 47 (41%) | 412 (31%) | P=0,02 |
|  |  |  |  |  |
|  |  | Mean values (SD) | Mean values (SD) | P-value |
| Age |  | 41.6 (SD) | 41.1 (SD) | P=0,732 |
| HADS Anxiety  HADS Depression |  | 7.9 (SD)  5.4 (SD) | 5.6 (SD)  3.6 (SD) | P<0,001  P<0,001 |
|  |  |  |  |  |
|  |  | Median | Median |  |
| EQ-5D |  | 0.8 | 1 | P<0,001 |

Figure 1: Median scores of quality of life as assessed with the DLQI in alopecia areata (AA) and androgenetic alopecia (AGA) (*p= 0,022*)



Figure 2: Score of Anxiety and Depression (HADS) in Control - AA–AGA patients (*Anxiety p< 0,001; depression p=0,02*)



# Figures 3a: Median scores for Health Related Quality of Life (HRQoL) as assessed by the (EQ5D) in male versus female patients (*p=0,001)*



Figure 3b: HRQoL as assessed by the EQ5D in different age groups (*p<0.001*)



Figure 3c: HRQoL in general health (EQ5D) in control- AA-AGA patients (*p=0,003*)



Figure 3d: HRQoL with visual analogue score (EQ VAS) in control - AA-AGA patients (*p<0,001*)

