QUALITY OF LIFE AND MALADJUSTMENT ASSOCIATED WITH HAIR LOSS IN WOMEN WITH ALOPECIA ANDROGENETICA

J. VAN DER DONK, J. A. M. HUNFELD, J. PASSCHIER, K. J. KNEGT-JUNK and C. NIEBOER

Department of Medical Psychology and Psychotherapy, Erasmus University, Rotterdam, Department of Gynaecology, University Hospital, Rotterdam-Dijkzigt, Department of Dermatology and Venereology, University Hospital, Rotterdam-Dijkzigt and Department of Dermatology, Free University, Amsterdam, The Netherlands

Abstract—Quality of life and maladjustment related to hair loss were studied by means of a standardized interview in a group of 58 women with alopecia androgenetica who applied for treatment at the Department of Dermatology. The hair loss was found to have a negative influence on the quality of life on the majority of them. In 88%, hair loss had negative effects on their daily life; in about 75%, the hair problems were manifested in negative self-esteem and about 50% experienced social problems. General psychosocial maladjustment in relation to hair loss was indicated in almost one-third of the women.

Key words—alopecia androgenetica, maladjustment, quality of life

INTRODUCTION

Compared to the gradually increasing number of studies being published on men with alopecia androgenetica [1–3], the lack of attention being paid to women with this affliction is remarkable.

More psychosocial problems can be expected in women than in men because a smaller proportion of women are affected, which emphasizes their deviation from the norm. Venning and Dawber [4] found that 13% out of 564 premenopausal women showed hair loss on the scalp, while in men this percentage was about 45% [5].

A second reason for expecting more psychosocial problems in women with alopecia androgenetica than in men is that our cultural norms emphasize the relevance of physical attractiveness for women in particular.

So far, the psychological and social problems of women with alopecia androgenetica have mainly been described anecdotally [6]. Eckert [7] studied women with diffuse alopecia and found that 20% of them were suffering from severe marital problems or were overtly depressed. van der Donk et al. [8] compared women with alopecia androgenetica to women with a non-visible dermatological disorder and to male subjects with alopecia androgenetica, using standardized psychological instruments. They found elevated scores for self-satisfaction and social inadequacy in comparison with the women with non-visible dermatological complaints. In addition, the women with alopecia androgenetica scored higher on inadequacy, rigidity, injudgedness and general psychological maladjustment and lower on self-evaluation and self-esteem than the men with alopecia androgenetica. Finally, they showed more psychosocial problems, which they attributed to their hair loss, than any of the other groups.

The present study is a continuation of the previous one by van der Donk et al. [8] and explores the character and extent of psychosocial problems related to alopecia androgenetica in the aforementioned 58 women with alopecia androgenetica more deeply, in order to assess the influence of hair loss on their quality of life.

As the sensitivity of standardized generic quality of life instruments, such as the Nottingham Health Profile, is highest mainly in the severe range [9], data were collected via in-depth interviews.

The use of an interview method also gives detailed information on the situations in which the problems are experienced, which can be used for behavioural management.

As this study was concerned solely with problems which can be attributed to alopecia androgenetica, no control group was included.

Our study questions were:

1. What problems exist in various life areas in the women with alopecia androgenetica who applied for treatment?
2. How many women with alopecia androgenetica show general psychosocial maladjustment which is attributable to alopecia androgenetica?

METHOD

Subjects

Subjects who applied for treatment for their hair loss at the Department of Dermatology at the University Hospital Dijkzigt in Rotterdam and the Free University Hospital in Amsterdam were invited
by letter to participate in a double-blind clinical trial. The trial had the purpose of investigating the effect of 2% minoxidil lotion treatment and will be published elsewhere. From the respondents, 70 healthy, dark-haired women, aged between 18–45 years and with a minimum duration of hair loss of 6 months, were selected by dermatologists (K.J.K.J., C.N.), on the basis of the following criteria for female pattern alopecia:

1. diffuse thinning in the fronto-parietal area of the scalp, both with or without temporal recession,
2. grades I and II according the Ludwig scale [10].

Subjects were assigned at random, in a double-blind fashion, to either a 2% minoxidil lotion treatment group or to a placebo treatment group. Of the 70 women who met these criteria,* 58 agreed to participate in our psychological study, resulting in a response rate of 83%. The protocol of the study was reviewed and accepted by the Medical Ethics Committee of the University Hospital Rotterdam-Dijkzigt. The subjects received a small reward for their invested effort (equivalent of U.S. $5).

**Procedure**

The data were collected using an interview with questions related to the significance of hair, the problems resulting from hair loss and psychosocial maladjustment. The interview was held in the 9th month of the medical study and had a retrospective character, i.e. the questions referred to the situation 1 month before the start of treatment. It took about 30 min and was carried out at the Department of Dermatology by a trained female psychologist (J.A.M.H.) in view of the possibility of specific female problems.

**Instruments**

The interview consisted of two parts and was semi-structured. It was based on the experience gained during the explorative interviews with subjects with androgenetica in the pilot phase of the study. The first part consisted of open questions on the main life areas so that the subject could bring forward topics which were related to hair loss. A written report on this part of the interview was made afterwards for later analysis. The second part of the interview consisted of a number of closed questions, in particular about the subject's behaviour and feelings in specific situations. These situations were adopted from the pilot phase of the study and were only presented to the subject if they had not already been discussed in the first part of the interview.

**Data analyses**

To answer the first research question on the effect on quality of life, the answers were analysed descriptively by frequency counts of the psychological problems. For the second research question on the general psychosocial maladjustment of the women, the written records were analysed to review the women's adjustment to their hair loss. Two independent judges (J. vd D., J.P.) classified the adjustment into (1) adaptive and (2) maladaptive adjustment. They used the following operationalizations:

1. **Good adjustment to hair loss** was expressed by the absence of any general effect of the hair loss on the subject's feelings and her participation in daily activities. A number of specific negative aspects related to hair loss could be present. In addition, the adjustment led to positive expectations of the subject about the future.

2. **Maladjustment to hair loss** was expressed by preoccupation with the negative aspects of hair loss, leading to manifold unpleasant and troubled feelings and to the avoidance of normal daily activities. Furthermore, it led to negative expectations about the future.

On the basis of this classification, the subjects were divided into three categories by the judges: (A) predominantly well-adjusted subjects, (B) predominantly maladjusted subjects and (C) subjects who showed both well-adjustment and maladjustment without an overall predominance of either one. The agreement between the judges on the classification of the women according to their adjustment was 86% initially. After sharpening of the criteria, the remaining 14% were discussed again resulting in an agreement of 94%. There was no concordance in 6%.

**RESULTS**

**Subject characteristics**

The characteristics of the subjects are shown in Table 1.

**Quality of life**

Table 2 shows the frequency and percentage of psychosocial problems mentioned during the interview and categorized by life area. In the following
negative feelings during social contact

For 48% of the subjects, the hair loss formed a problem socially. More than a quarter felt worried that others would see the thinning patches: "I always feel nervous in company because I feel like I have a large, round bald spot on my head." One out of every seven women was afraid that others would make remarks about the hair loss: "I keep quiet particularly at work, because the moment they make a remark about my hair, I feel very small." The women felt offended by jokes or remarks about their hair loss and found them very improper, such as: "Hey, old woman" or "I can see your scalp!" For most of the women it was important who made these remarks. A number of them used to crack cynical jokes about themselves, such as: "Any punk would be jealous. I hardly need to do anything to get this hairstyle."
Adjustment

The written reports showed that 36% of the subjects were predominantly well-adjusted regarding coping with their hair loss. This approach was illustrated by one of the women in the following words:

In company I am usually more worried about being in a bad mood than about my hair loss. Of course, suddenly having thoughts about the hair loss can soon change that. As such, the hair loss isn’t that much of a problem to me. It doesn’t automatically mean that I would start thinking about a wig if I was to go bald—I regard my hair as a means of distinguishing myself from others. Neither is it my intention to use some sort of medication for the rest of my life against hair loss. I prefer to spend my money on other things.

The reports also showed that 29% of the women were predominantly maladjusted regarding their hair loss. One of them told us, for example:

In company I always feel that other people are looking at me all the time and I hate that. That is why I prefer to stay at home. I’ve never been what you might call attractive, but now with my hair like this, it’s even worse. A woman without hair isn’t a really a woman anymore. And if I wasn’t already aware of that, the other people around me would soon let me know! So, I am afraid that I will go completely bald. I think that if my hair doesn’t stop falling out, I will go down hill very fast psychologically.

Both well-adjustment and maladjustment were observed in 29% of the women. The general trend in these cases was, for instance:

I always feel worried in company and I am afraid that my wig won’t stay straight. But when I get chatting, the feeling disappears. The looks other people give me make me terribly annoyed and as soon as they start talking about their own hair-styles, I run for cover! I have always felt unattractive and fat even as a child and when I started to lose my hair too, I wouldn’t go out to play anymore. I locked myself in and began to eat. But nowadays I wear a wig and have the courage to go out again. Perhaps I will try hair fusion some time. A new wig really boosts my self-confidence. Luckily, I am a cheerful person.

DISCUSSION

Scalp hair plays a major role in determining physical attractiveness and is an important aspect in interpersonal contact, as it is the first and most directly visible source of non-verbal information [11,12]. Scientific studies have demonstrated that individuals who are rated as being physically attractive are thought to be more successful [13], to enjoy more respect [14], to have more friends, to be more happily married [15] and to be socially better adjusted [16] than less physically attractive individuals. Physical attractiveness also seems to have intrapersonal consequences, as indicated by the positive correlation with self-esteem and the negative correlation with neuroticism [17].

Studies on individuals who think themselves physically unattractive were almost exclusively conducted on clinical populations. van Keep [18] reported that the majority of the subjects in his study with a visible dermatological disorder avoided social contact and lost touch with acquaintances and friends. Baardman [19] found that nearly 20% of the clients at a clinic for neurosis were negatively preoccupied by their appearance, which caused social avoidance behaviour, poor physical hygiene, etc. In brief, feelings of physical unattractiveness seem to be related to many negative psychosocial aspects. These negative aspects were also found in this study on women who applied for treatment for alopecia androgenetica.

A large number of the women in this study experienced a negative effect on their quality of life as a consequence of this affliction. These problems manifested themselves in the effort taken to hide the hair loss, in their taking care where to sit and in avoiding company and activities, in inhibited behaviour, feelings of envy, lost self-confidence and feelings of insecurity in company. Most of them had sought help for their alopecia androgenetica: the majority had consulted their general practitioner or a dermatologist. In general, they considered the care offered to be insufficient. For this reason, many of the women felt powerless and referred to their worries about the cause and possible progressive nature of the hair loss as most unpleasant. A possible case in point was that three women in our study group considered their hair loss to be comparable with a physical impairment, such as the loss of a limb.

The psychosocial problems of the women with alopecia androgenetica in this study appeared to be comparable with the psychological problems of individuals with a (visible) dermatological disorder, such as acne, eczema and psoriasis. The social life of 40% of the individuals with one of these dermatological disorders was affected by the disorder [20], whereas in our group this percentage was 50%.

Adjustment

It was striking that 29% of the women in our study group were unable to cope with their alopecia androgenetica and experienced severe maladjustment. Furthermore, 36% of the women were predominantly well-adjusted and 29% showed both well-adjustment and maladjustment. While general practitioners in The Netherlands reported that 50% of the patients who presented with alopecia androgenetica had psychological problems, they nevertheless used the same policies for patients with and without psychological problems [21]. The authors of this study conclude that many patients with alopecia androgenetica did not receive sufficient psychological support from their general practitioner in a psychological respect. As patients with alopecia androgenetica primarily seek help and comfort in the medical field, psychological support, such as information, emphatic listening and behavioural advice regarding the above problems, ought to be supplied by the general practitioner and the dermatologist.
The question arises as to whether the problems and adjustment of the women with alopecia androgenetica in this study can be extrapolated to the general population of women suffering from this affliction. A further epidemiological study on a non-clinical group of women with alopecia androgenetica might help to find the answer to this question.

Acknowledgements—This study was supported by a grant from The Upjohn Company, Kalamazoo, MI. Mrs Goudsmit provided the administrative support.

REFERENCES