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The Role of Therapy in Impairing Quality of Life in Dermatological Patients: A Multinational Study

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Skin disease and its therapy affect health-related quality of life (HRQoL). The aim of this study was to measure the burden caused by dermatological therapy in 3,846 patients from 13 European countries. Adult outpatients completed questionnaires, including the Dermatology Life Quality Index (DLQI), which has a therapy impact question. Therapy issues were reported by a majority of patients with atopic dermatitis (63.4%), psoriasis (60.7%), prurigo (54.4%), hidradenitis suppurativa (54.3%) and blistering conditions (53%). The largest reduction in HRQoL attributable to therapy, as a percentage of total DLQI, adjusted for confounders, was seen in blistering conditions (10.7%), allergic/drug reactions (10.2%), psoriasis (9.9%), vasculitis/immunological ulcers (8.8%), atopic dermatitis (8.7%), and venous leg ulcers (8.5%). In skin cancer, although it had less impact on HRQoL, the reduction due to therapy was 6.8%. Treatment for skin disease contributes considerably to reducing HRQoL: the burden of dermatological treatment should be considered when planning therapy and designing new dermatological therapies.

Key words: quality of life; HRQoL; DLQI; dermatological therapy; burden of skin disease; therapy burden.

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Topical and other dermatological therapies can add to the burden of skin disease, as they may be time-consuming, messy, intervene with clothing choice, and impact on health-related quality of life (HRQoL) in ways that are unique to the skin (1, 2). This contrasts with the relatively low burden of oral therapy in other diseases (3) where, for most, oral medication becomes routine.

However, even systemic dermatological medications, such as cytotoxic drugs, corticosteroids, retinoids, intravenous or injected biologics, may have an associated burden. Topical and injection routes of drug administration have the lowest levels of convenience and global satisfaction (4).

Impairment of HRQoL due to dermatological therapy is little explored, even though the burden caused by skin disease treatment is very important, both to patients and because it contributes to poor adherence (5).

Most generic measures of HRQoL were developed without including skin diseases. It is therefore unsurprising that they miss the burden experienced by dermatological patients. In measures designed for use across skin diseases, only the Dermatology Life Quality Index (DLQI) includes a question concerning the impact of treatment on everyday life (6).

The aim of this study was to measure how therapy for skin disease contributes to reducing HRQoL in outpatients across Europe.

SIGNIFICANCE

Treatments for skin diseases differ from those used for other diseases. They may be messy, time-consuming, affect clothing or be painful. Some diseases are burden-some (psoriasis, eczemas, itching) and their therapy causes extra impairment, which should be appreciated. Others showed little impact from therapy, although the diseases themselves were serious (hidradenitis suppurativa, psychodermatological conditions, acne). Adequate therapy should be sought to alleviate symptoms without adding further impairment. Lastly, some skin diseases stood out as more burdened by therapy than by the disease itself (cancer, allergies, scars). For these patients, choice of therapy is most important for providing optimal help.
METHODS

Data were obtained from a cross-sectional multicentre study on patients recruited from 15 dermatological outpatient clinics in 13 European countries: details have been previously reported (7). The study was approved by the Regional Committee for Medical Research Ethics in Norway. Separate ethical approvals were obtained where necessary. The study was conducted in accordance with the Declaration of Helsinki.

Consecutive patients, age over 18 years, understanding the local language and not having severe mental disease were invited to participate on random days, giving written consent. Participants completed questionnaires on sociodemographics (sex, age, ethnicity, education, marital and socioeconomic status), the DLQI and other questionnaires (7–11).

Patients were examined by the dermatologist, who recorded comorbidities: diabetes mellitus, cardiovascular, chronic respiratory, rheumatological or other disease. Workers from each hospital’s service division were invited to participate as controls.

The DLQI, a 10-item questionnaire, was used to assess impairment in HRQoL. Question 10, which concerns the impact of therapy, was used to assess how treatment impaired HRQoL: “How much of a problem has the treatment for your skin been, for example by making your home messy, or by taking up time?” with possible answers “very much” (scored 3), “a lot” (2), “a little” (1) or “not at all/not relevant” (0).

The DLQI was not designed for use by healthy individuals. Patients with naevi (n=192) served as “healthy” controls, since there were no significant differences between the patients with naevi and healthy controls (7, 8).

Statistical analysis

Data from all centres were merged. Diagnoses were organized in 35 disease groups (8, 12).

SPSS 24 software was used for statistical analysis. Frequencies and means for patient and control characteristics were calculated.

The answers to DLQI question 10 were dichotomized into “no impairment” (0) or “impaired” (1, 2 or 3) when calculating frequencies of positive answers.

For each diagnosis the mean scores for question 10 and total DLQI were calculated. Their relationship was calculated as the mean total DLQI score × 100, denoted as Q10%.

Comparisons between patients with naevi and healthy controls were performed with the t-test for continuous variables (age) and the χ² test for categorical variables (sex, marital status, socioeconomic status, comorbidities, economic difficulties, stress, depression and anxiety (7)) and linear (EQ-VAS) and logistic regressions (EQ5D) for comparing HRQoL outcomes (8).

Linear regression was performed to analyse Q10% for each diagnosis, adjusting for age, sex, socioeconomic status and comorbidity with “naevi” as controls.

A search for publications on therapy issues in dermatology using DLQI or other instruments was performed using MEDLINE, EBASE and Cochrane Library following standard search strategies. Search terms and medical descriptors (MeSH) included skin disease, dermatosis, dermatoses, quality of life, DLQI, skin therapy, topical therapy, photodynamic therapy, cryotherapy, cryosurgery, cryoablation, laser, phototherapy, photochemotherapy, ultraviolet B (UVB), UVA, UV A1, psoralen plus UVA (PUVA), retinoid plus PUVA (RePUVA), topical drug administration, parenteral administration, biological therapy, tumour necrosis factor (TNF)-α inhibitors, infusion therapy, skin cancer therapy, and surgical dermatological therapy.

RESULTS

Participants

There were 4,010 participants and 1,359 healthy controls. Comparative details have been published previously (7–11) and are given briefly in Table S1.

Dermatology Life Quality Index data

There were 3,846 (96%) valid answers to DLQI. 5.2% of which had a DLQI >20 (extremely large effect on HRQoL). One-fifth (20.3%) experienced at least a very large effect (DLQI >11) and 44.9% had a DLQI >6, mea-

Table I. Frequencies of Dermatology Life Quality Index (DLQI) scores (n=3,846)

| DLQI score band descriptors (ref. 6, 13) | Valid % | Number | Cumulative % |
|-----------------------------------------|---------|--------|--------------|
| Extremely large (21–30)                 | 5.2     | 200    | 5.2% >20     |
| Very large (11–20)                      | 20.3    | 782    | 25.5% >11    |
| Moderate (6–10)                         | 19.4    | 745    | 44.9% >6     |
| Small (2–5)                             | 26.6    | 1,023  | 71.5% >2     |
| No (0–1)                                | 28.5    | 1,096  |              |
| Total                                   | 3,846   |        |              |

Mean DLQI (SD). n=Valid number of patients

| Sex          | Male: 6.4 (6.7) n=1,686 | Female: 7 (6.8) n=2,168 |
|--------------|--------------------------|--------------------------|
| Age groups   | 18–35 years: 7.34 (6.8) n=1,247 | 36–65 years: 6.94 (7) n=1,880 |
|              | >66: 5.06 (5.9) n=652       |                           |
| Socioeconomic status | Low: 8.22 (7.2) n=720 | Middle: 6.44 (6.6) n=2,844 |
|              | High: 5.88 (5.9) n=327       |                           |
| Comorbidity  | None: 6.74 (6.7) N=2,573 | Any: 6.89 (6.9) N=1,033   |

| Country | BE | DK | FR | GER | HU | IT* | NL | NO* | PL | RUS | ES | TR | UK |
|---------|----|----|----|-----|----|-----|----|-----|----|-----|----|----|----|
| n       | 338 (3.9) | 579 (6.7) | 458 (5.3) | 714 (7.4) | 726 (7.3) | 8.01 (6.7) | 5.09 (5.7) | 6.99 (6.7) | 10.89 (7.8) | 10.38 (7.0) | 2.26 (3.6) | 7.35 (5.6) | 5.23 (6.8) |

*Padua and Rome. †Oslo and Stavanger.

BE: Belgium; DK: Denmark; FR: France; GER: Germany; HU: Hungary; IT: Italy; NL: The Netherlands; NO: Norway; PL: Poland; RUS: Russia; ES: Spain; TR: Turkey; UK: United Kingdom.
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Fig. 1. The percentage of positive answers to having therapy issues (Question 10 of the DLQI) for each diagnosis. Diagnoses represented by fewer than 20 valid answers (hyperhidrosis (12), nail diseases (17) and granuloma annulare (13) excluded). NMSC: non-melanoma skin cancer.

The total patient population (n = 3,846) had a mean ± standard deviation (SD) DLQI score of 6.7 ± 6.8, meaning moderately impaired HRQoL. Except for naevi, no skin disease had a mean score < 2, so all had at least a small effect on patients’ HRQoL. Twenty-seven of the 35 (77%) skin conditions had mean DLQI scores > 5, indicating at least a moderate effect on a patient’s life (Table SII1).

Higher DLQI values, indicating higher impairment, were seen in females, younger age groups, patients with comorbidities or those of low socioeconomic status.

Therapy impact data (DLQI question 10)

Question 10 in the DLQI addresses therapy-related issues. The numbers of patients answering with “a little”, “a lot” or “very much”, i.e. other than “no impact/not relevant”, are given in Fig. 1. More than half of the patients with atopic dermatitis (AD) (63.4%), prurigo (60.7%), psoriasis (54.4%), hidradenitis suppurativa (HS) (54.3%) or blistering disorders (53%) answered positively. Fifteen of 32 skin conditions had >33.3% patients scoring positively.

The mean scores with SD for question 10 and Q10% for each diagnosis are presented in Table II. There are no existing cut-off values for interpreting results from single questions of the DLQI, and isolated values may not give a clear perspective as to how large the impact is. Q10% is not a standardized method for interpreting DLQI data, but does provide perspective on how therapy issues relate to the total HRQoL impairment. Table II lists the diseases in descending values according to Q10%, adjusted for age, sex, socioeconomic status and comorbidity. The positive standardized β coefficients for all diseases denote influence of therapy on HRQoL even when adjusted. For many diseases the β coefficient was relatively high, indicating robustness of the presented results.

When assessing Q10%, males and older patients showed more impairment, the reverse of what was seen for total mean DLQI. The impairment was highest in patients with comorbidities or those of low socioeconomic status.

When considering the impact of therapy on HRQoL, highest mean scores and most positive answers to question 10 were seen in diseases that commonly affect large areas of the skin (e.g. AD, psoriasis, allergic/drug/phototoxic conditions, prurigo, papulosquamous diseases, eczemas, connective tissue disease and vitiligo), as well as diseases accompanied by blisters/erosions, ulceration or crusting (blistering diseases, venous leg ulcer, vasculitis, immunological ulcers and oral diseases) and pruritic dermatoses (prurigo, urticaria and pruritus) (Table II, Fig. 1).

Q10% reveals which diagnostic groups are most affected by therapy relative to their total HRQoL impairment. Blistering conditions showed the highest value (10.7), followed by allergic, drug, phototoxic/-allergic reactions (10.2) and psoriasis (9.9), a ranking that differs from total mean DLQI values (Table SII1). This gives insight into the true extra burden of therapy for different diseases.

HS, prurigo, pruritus and urticaria show the highest impairment when mean DLQI scores are evaluated, but drop in ranking when therapy is assessed. Likewise, acne, rosacea and psychodermatological conditions, scoring among the average impaired as measured by mean DLQI scores, were some of the least affected by therapy. Conversely, blistering conditions, non-melanoma skin cancer (NMSC), actinic keratoses (AK), allergic/drug reactions, vasculitis and venous leg ulcers rank higher when evaluated according to therapy-related impairment.
Table II. Effect of treatment on Dermatology Life Quality Index (DLQI). Ranking according to the percentage of Question 10 of the DLQI (therapy issues) to the mean total DLQI (Q10%) for diagnoses with at least 20 valid answers (hyperhidrosis (12), nail diseases (17) and granuloma annulare (13) excluded). Linear regression (standardized β) with “naevi” as a “healthy” control group, adjusting for age, sex, socioeconomic status and comorbidity (diabetes mellitus, cardiological, respiratory, rheumatological or other disease).

| Diagnosis                                                                 | Q10%a | Standardized β | Question 10 Mean ± SD | DLQI Mean ± SD | Valid n/Valid b |
|---------------------------------------------------------------------------|-------|---------------|------------------------|----------------|----------------|
| All patientsc                                                             | 7.73  | 0.06          | 0.52 ± 0.8             | 6.73 ± 0.8     | 3,846/3,553    |
| 1 Blistering conditions                                                   | 10.71 | 0.47          | 0.92 ± 1.0             | 8.59 ± 7.4     | 66/60          |
| 2 Allergic, drug, phototoxic/-allergic reactions                         | 10.21 | 0.39          | 0.54 ± 0.8             | 5.29 ± 4.3     | 24/21          |
| 3 Psoriasis                                                               | 9.85  | 0.19          | 0.90 ± 1.0             | 9.14 ± 7.6     | 660/615        |
| 4 Vasculitis and immunological ulcersd                                   | 8.78  | 0.28          | 0.62 ± 0.9             | 7.06 ± 6.1     | 67/60          |
| 5 Atopic dermatitis                                                       | 8.67  | 0.33          | 1.0 ± 0.9              | 11.53 ± 7.2    | 172/150        |
| 6 Vitiligo                                                                | 8.62  | 0.36          | 0.33 ± 0.6             | 3.83 ± 3.7     | 24             |
| 7 Venous leg ulcers                                                       | 8.47  | 0.27          | 0.80 ± 1.0             | 9.45 ± 7.3     | 113/87         |
| 8 Other hair disorders                                                    | 8.38  | 0.27          | 0.42 ± 0.8             | 5.01 ± 5.4     | 82/76          |
| 9 Prurigo                                                                 | 8.13  | 0.33          | 0.93 ± 0.9             | 11.44 ± 8.2    | 27/24          |
| 10 Scars, fibrosis of the skin, morphea                                   | 8.11  | 0.24          | 0.43 ± 0.9             | 5.3 ± 4.8      | 27             |
| 11 Papulosquamous skin diseasesd                                          | 7.69  | 0.21          | 0.49 ± 0.8             | 6.37 ± 6.4     | 113/103        |
| 12 Connective tissue disease                                              | 7.43  | 0.20          | 0.58 ± 0.9             | 7.81 ± 7.0     | 91/74          |
| 13 Oral conditionsg                                                       | 7.39  | 0.24          | 0.50 ± 0.8             | 6.77 ± 6.6     | 26             |
| 14 Eczema                                                                 | 7.36  | 0.21          | 0.62 ± 0.9             | 8.42 ± 7.2     | 234            |
| 15 Urticaria                                                              | 7.09  | 0.35          | 0.68 ± 0.9             | 9.59 ± 6.7     | 69/60          |
| 16 Hand eczema                                                           | 7.05  | 0.18          | 0.60 ± 0.9             | 8.51 ± 7.2     | 156/146        |
| 17 Alopecia areata                                                        | 6.99  | 0.18          | 0.39 ± 0.8             | 5.58 ± 6.8     | 31/30          |
| 18 Pruritus                                                               | 6.84  | 0.24          | 0.75 ± 1.0             | 10.97 ± 7.1    | 60/58          |
| 19 Non-melanoma skin cancer and actinic keratosis                         | 6.75  | 0.11          | 0.16 ± 0.5             | 2.37 ± 5.0     | 401/372        |
| 20 Genital (non-venerereal)b                                              | 6.36  | 0.22          | 0.56 ± 0.8             | 8.81 ± 6.4     | 32/30          |
| 21 Otherf                                                                | 6.15  | 0.20          | 0.39 ± 0.7             | 6.34 ± 6.6     | 96/67          |
| 22 Hidradenitis suppurativa                                               | 6.14  | 0.24          | 0.78 ± 0.8             | 12.7 ± 7.6     | 46/44          |
| 23 Infections of the skin                                                | 6.09  | 0.21          | 0.38 ± 0.8             | 6.24 ± 5.8     | 253/244        |
| 24 Benign skin tumours                                                   | 5.51  | 0.09          | 0.15 ± 0.5             | 2.72 ± 3.7     | 159/154        |
| 25 Lichen planus                                                         | 5.42  | 0.07          | 0.33 ± 0.7             | 6.09 ± 5.4     | 46/41          |
| 26 Seborrhoeic dermatitis                                                | 5.41  | 0.23          | 0.34 ± 0.6             | 6.28 ± 4.4     | 75/74          |
| 27 Psychodermatological conditions                                       | 5.41  | 0.14          | 0.46 ± 0.8             | 8.5 ± 7.1      | 34             |
| 28 Acne                                                                  | 4.99  | 0.16          | 0.31 ± 0.6             | 6.21 ± 5.2     | 234/228        |
| 29 Rosacea                                                               | 4.66  | 0.09          | 0.25 ± 0.6             | 5.37 ± 5.3     | 75/68          |
| 30 Naevi                                                                 | 4.61  | –             | 0.07 ± 0.3             | 1.52 ± 2.9     | 186            |
| 31 Malignant melanoma                                                    | 4.41  | 0.03          | 0.12 ± 0.4             | 2.72 ± 4.4     | 86/75          |
| 32 Melasma, pigment disorders                                            | 2.01  | 0.16          | 0.10 ± 0.3             | 4.97 ± 4.7     | 32/30          |

DISCUSSION

Using a dermatology-specific measure this study identified the extent of the reduced HRQoL associated with therapy. For several diseases, patients experience a high burden associated with therapy (blistering conditions, allergic/drug reactions, psoriasis, vasculitis, vitiligo and venous leg ulcers). Ranking the diseases according to what percentage of the burden is caused by therapy gives new insight into this specific impairment for the separate diagnoses.

Most skin diseases are treated with topical therapy. However, dermatological treatments include oral therapy, phototherapy, photodynamic therapy, lasers, cryotherapy, intraleisional and surgical procedures and parenteral administrations, which may be painful, time-consuming or cause infusion reactions. The use of these specific dermatological medications and therapeutic approaches presents issues and challenges unique to skin disease.

Generic HRQoL measures have been developed without specific reference to the impact of therapy for skin disease (Table III). Assessment may therefore be inaccurate if this burden experienced by dermatological patients is missed. There are no questions related to the impact of therapy in the most commonly used generic measures. However, the generic measures Treatment Satisfaction with Medicines Questionnaire (SATMED-Q) (3) and Treatment Satisfaction Questionnaire for Medication (TSQM) (4) are designed to address issues with medication, but are little used in dermatology. The DLQI is the only non-disease-specific dermatological measure
Role of therapy in impairing QoL in dermatological patients

that addresses therapy burden (Table III), although the DLQI is the most widely used measure in dermatology (14) the issue of therapy is little explored.

There are very few studies evaluating the contribution of therapy to impairment of HRQoL. In 3 studies (15–17) the generic instrument Short Form Health Survey (SF-36) was used in random samples of the population. A large proportion of patients reported dermatological problems of therapy to impairment of HRQoL. In 3 studies (15–17) the issue of therapy is little explored.

Table III. Overview of dermatology-specific, disease-specific and generic instruments assessing quality of life with comments on whether the impact of therapy is addressed in the questionnaire.

| Type and name of instrument | Therapy impact | Authors, year |
|-----------------------------|----------------|---------------|
| Dermatology-specific instruments | | |
| DLQI | Yes | Finlay & Khan 1994 (6) |
| Skindex, Skindex-29, Skindex-16, Skindex-17 | No | Chren et al. 1996 (28), Nijsten et al. 2006 (29) |
| DSQOL | No | Anderson & Rajagopal 1997 (30) |
| DQOLS | No | Morgan et al. 1997 (31) |
| Disease-specific instruments | | |
| PDI | Yes | Finlay & Kelly 1987 (32) |
| PSORIQoL | Yes | McKenna et al. 2003 (33) |
| RosaQoL | Yes | Nicholson et al. 2007 (34) |
| QOLHEQ | Yes | Ofenloch et al. 2014 (35) |
| VEINES-QOL | Yes | Bland et al. 2015 (36) |
| CADI | No | Motley & Finlay 1992 (37) |
| DSQOL – Contact dermatitis | No | Anderson & Rajagopal 1997 (30) |
| DSQOL – Acne version | No | Anderson & Rajagopal 1997 (30) |
| WAA Questionnaire | No | Dole et al. 2000 (38) |
| Acne-QoL Questionnaire | No | Martin et al. 2001 (39) |
| PSS-AD in adults | No | Ando et al. 2006 (40) |
| SCI | No | Rhee et al. 2006 (41) |
| AAQ | No | Endo et al. 2012 (42) |
| MELASQoL Scale | No | Lieu & Pandaya 2012 (43) |
| AKQoL questionnaire | No | Esman et al. 2013 (45) |
| QA-QLI | No | Fabbrocini et al. 2013 (44) |
| FQL index | No | Heisterberg et al. 2013 (45) |
| VitQoL | No | Lilly et al. 2013 (46) |
| ABQoL | No | Sebaratnam et al. 2013 (47) |
| Generic instruments* | | |
| TSQM | Yes | Atkinson et al. 2004 (4) |
| SATMED-Q | Yes | Ruiz et al. 2008 (3) |
| EuroQol (EQ5D) | Yes | EuroQolGroup 1990 (48) |
| Medical Outcome Study (MOS) | No | Ware & Sherbourne 1992 (49) |
| MOS Short-Form 36 (SF-36) | No | | |
| WHOQoL-100 | No | WHO 1996 (50) |
| WHOQoL-BREF | No | WHO 1998 (51) |

*Only the most commonly used generic instruments that do not address therapeutic issues are shown here.

Studies of the same data-set rank HS patients with some of the lowest HRQoL (8), highest risk for psychiatric comorbidity (7, 20) and impairment in sexual life (9). Despite very high impairment of HRQoL, therapy contributes little to this burden.

AD and psoriasis rank highly when mean DLQI, positive answers to therapy issues or Q10% are evaluated, suggesting that these patients are equally adversely affected by all aspects of HRQoL, including therapy.

Diseases affecting small areas of the body, such as facial dermatoses (seborrhoeic dermatitis, rosacea and acne), as well as psychodermatological conditions rank lower on therapy relative to the total DLQI than might be expected, demonstrating that it is the disease itself and not the therapy that is the driving cause of HRQoL impairment. Treating these conditions adequately should alleviate the patient’s experienced burden without additional improvement.

In contrast, patients with AK, NMSC, allergic/drug reactions, scars/fibrosis and morphea, who do not report severe impairment of HRQoL as measured by the mean DLQI, rank highly in impairment when assessing therapy as a percentage of this total score. AK and NMSC do not apparently have a high impact on HRQoL, nor psychiatric comorbidity (7, 8, 20), but score relatively worse when therapy is assessed, ranking them higher than HS and several other diseases.

Studies evaluating the burden caused by AK and/or NMSC have shown low impact on HRQoL of these diseases (21–24), raising the possibility that currently available measures may be missing therapy issues and that there may be a need for a skin-cancer-specific HRQoL measure. Existing disease-specific instruments do not include therapy questions (22, 25) (Table III).

Burdensome treatments have a negative effect on adherence to therapy (5) and can be the reason for undertreatment and relapse of disease. Measuring HRQoL without taking into account therapy issues may not represent the true extent of suffering that dermatological patients experience. On the other hand, knowing which diseases have the highest potential to cause therapy issues can alert clinicians to which patients need a different approach, by giving them better information, providing a variety of options, offering training in therapy application, or at least acknowledging the issue.

When developing clinical guidelines in dermatology, optimization of therapy and minimizing the burden of treatment should be considered. Developers of HRQoL instruments should pay attention to therapy issues when measuring HRQoL in some specific diagnoses, such as...
skin cancer, as this burden may go undetected using currently available measures (7, 8, 20–23).

**Strengths and limitations**

The high number of patients in this study, the unbiased selection of participants and adjusting for confounding factors resulted in robust data on therapy as a factor contributing to impairment in HRQoL. Similar studies on therapeutic issues are lacking and studies using DLQI typically have no healthy control group.

One potential limitation is in the detail of the wording of DLQI question 10: “(...by making your home messy, or by taking up time)”, which may bias the respondents into only considering topical therapy. However, the main question itself is neutral on this point “…how much of a problem has the treatment for your skin been…”.

Detailed information on all treatments used by our patients was not obtained systematically. The presented data evaluate therapy issues on a general basis. Further studies evaluating specific dermatological treatments are warranted.

Although we refer to data from each country, the data was based on 1 centre from each country (apart from Italy and Norway). The recruitment centres may not have been representative of clinical practice across each country. There were large differences between countries in scores assessing impairment, which cannot be readily explained. The cross-cultural issue is one that is of relevance to all HRQoL measures (26). The same limitation may apply when comparing diseases (27). The cultural and language factors leading to these differences are not fully understood, though they should be taken into account when making any cross-cultural comparisons and when using HRQoL data as a guide to optimal health policies and creating optimal treatment guidelines. Analysis of the source for country differences may be able to serve as a guide to optimal health policies and creating optimal treatment guidelines.

**Conclusion**

Treatments for skin diseases contribute to the burden on HRQoL. For some diagnoses, therapy may have a larger impact than was previously known, but we also identify diseases that are affected by therapy to a lesser degree. Older, male patients with lower socioeconomic status and comorbidities experience more adverse issues with therapy. This study highlights new aspects to HRQoL. For some diagnoses, therapy may have a larger impact than was previously known, but we also identify diseases that are affected by therapy to a lesser degree.

The high number of patients in this study, the unbiased selection of participants and adjusting for confounding factors resulted in robust data on therapy as a factor contributing to impairment in HRQoL. Similar studies on therapeutic issues are lacking and studies using DLQI typically have no healthy control group.

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Although we refer to data from each country, the data was based on 1 centre from each country (apart from Italy and Norway). The recruitment centres may not have been representative of clinical practice across each country. There were large differences between countries in scores assessing impairment, which cannot be readily explained. The cross-cultural issue is one that is of relevance to all HRQoL measures (26). The same limitation may apply when comparing diseases (27). The cultural and language factors leading to these differences are not fully understood, though they should be taken into account when making any cross-cultural comparisons and when using HRQoL data as a guide to optimal health policies and creating optimal treatment guidelines. Analysis of the source for country differences may be able to serve as a guide to optimal health policies and creating optimal treatment guidelines.

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**References**

1. Ruiz MA, Heras F, Alomar A, Conde-Salazar L, de la Cuadra J, Serra E, et al. Development and validation of a questionnaire on ‘Satisfaction with dermatological treatment of hand eczema’ (DermASat). Health Qual Life Outcomes 2010; 8: 127.
2. Jubert-Estève E, Del Pozo-Hernando LJ, Izquierdo-Herce N, Bauza-Alonso A, Martín-Santiago A, Jones-Caballero M. Quality of life and side effects in patients with actinic keratosis treated with ingenol mebutate: a pilot study. Actas Dermosifiliogr 2015; 106: 644–650.
3. Ruiz MA, Pardo A, Rejas J, Soto J, Villasante F, Aranguren JL. Development and validation of the “Treatment Satisfaction with Medicines Questionnaire” (SATMED-Q). Value Health 2008; 11: 913–926.
4. Atkinson MJ, Sinha A, Hass SL, Colman SS, Kumar RN, Brod M, et al. Validation of a general measure of treatment satisfaction, the Treatment Satisfaction Questionnaire for Medication (TSQM), using a national panel study of chronic disease. Health Qual Life Outcomes 2004; 2: 12.
5. Feldman SR, Vrijens B, Gieler U, Piaserico S, Puig L, van de Kerkhof P. Treatment adherence intervention studies in dermatology and guidance on how to support adherence. Am J Clin Dermatol 2017; 18: 253–271.
6. Finlay AY, Khan GK. Dermatology Life Quality Index (DLQI) – a simple practical measure for routine clinical use. Clin Exp Dermatol 1994; 19: 210–216.
7. Dalgard FJ, Gieler U, Tomas-Aragones L, Lien L, Poot F, Jemec GBE, et al. The psychological burden of skin diseases: a cross-sectional multicenter study among dermatological out-patients in 14 European countries. J Invest Dermatol 2015; 135: 984–991.
8. Balieva F, Kupfer J, Lien L, Gieler U, Finlay AY, Tomas-Aragones L, et al. The burden of common skin diseases assessed with the EQSD: a European multicentre study in 13 countries. Br J Dermatol 2017; 176: 1170–1178.
9. Sampogna F, Abeni D, Gieler U, Tomas-Aragones L, Lien L, Titeca G, et al. Impairment of sexual life in 3,485 dermatological outpatients from a multicentre study in 13 European countries. Acta Derm Venereol 2017; 97: 478–482.
10. Szabo C, Allmayer A, Lien L, Poot F, Gieler U, Tomas-Aragones L, et al. Attachment styles of dermatological patients in Europe: a multi-centre study in 13 countries. Acta Derm Venereol 2017; 97: 813–818.
11. Lesner K, Reich A, Szepietowski JC, Dalgard FJ, Gieler U, Tomas-Aragones L, et al. Determinants of psychosocial health in psoriatic patients: a multi-national study. Acta Derm Venereol 2017; 97: 1182–1188.
12. Rea JN, Newhouse ML, Halli T. Skin disease in Lambeth. A community study of prevalence and use of medical care. Br J Prev Soc Med 1976; 30: 107–114.
13. Hongbo Y, Thomas C, Harrison M, Salek MS, Finlay AY. Translating the science of quality of life into practice: what do dermatology life quality index scores mean? J Invest Dermatol 2005; 125: 659–664.
14. Ali FM, Cueva AC, Vyasa J, Atwan AA, Salek MS, Finlay AY, et al. A systematic review of the use of quality-of-life instruments in randomized controlled trials for psoriasis. Br J Dermatol 2017; 176: 577–593.
15. Bingeefors K, Lindberg M, Isacson D. Self-reported dermatological problems and use of prescribed topical drugs correlate with decreased quality of life: an epidemiological survey. Br J Dermatol 2002; 147: 285–290.

www.medicaljournals.se/acta
16. Lindberg M, Isacson D, Bingefors K. Self-reported skin diseases, quality of life and medication use: a nationwide pharmaco-epidemiological survey in Sweden. Acta Derm Venereol 2014; 94: 188–191.

17. Bingefors K, Lindberg M, Isacson D. Quality of life, use of topical medications and socio-economic data in hand eczema: a Swedish nationwide survey. Acta Derm Venereol 2011; 91: 452–458.

18. Riis PT, Vinding GR, Ring HC, Jemec GB. Disutility in patients with hidradenitis suppurativa: a cross-sectional study using EuroQol-5D. Acta Derm Venereol 2016; 96: 222–226.

19. Matusiak L, Bieniek A, Szepietowski JC. Psychophysical aspects of hidradenitis suppurativa. Acta Derm Venereol 2010; 90: 264–268.

20. Balieva F, Lien L, Kupfer J, Halvorsen JA, Dalgard F. Are common skin diseases among Norwegian dermatological outpatients associated with psychological problems compared with controls? An observational study. Acta Derm Venereol 2016; 96: 227–231.

21. Rhee JS, Matthews BA, Smith TL, Burzynski M, Nattinger AB. Skin cancer and quality of life: assessment with the Dermatology Life Quality Index. Dermatol Surg 2004; 30: 525–529.

22. Tennvall GR, Norlin JM, Malmberg I, Erlendsson AM, Hadersdal M. Health related quality of life in patients with actinic keratosis—an observational study of patients treated in dermatology specialist care in Denmark. Health Qual Life Outcomes 2015; 13: 111.

23. Blackford S, Roberts D, Salek MS, Finlay AY. Basal cell carcinomas cause little handicap. Qual Life Res 1996; 5: 191–194.

24. Lee EH, Klassen AF, Nehal KS, Cano SJ, Waters J, Pusic AL. A specific measure of quality of life designed for use in clinical practice and trials. Br J Dermatol 2003; 149: 323–331.

25.花生 mono, 黄梅 G, 手术 GB, 评估影响的皮肤癌对患者‘‘品质生活’’的品质：皮肤的 AKQOL 问题。Br J Dermatol 2013; 168: 277–283.

26. Nijsten T, Meads DM, de Korte J, Sampogna F, Gelfand JM, Ongenae K, et al. Cross-cultural inequivalence of dermatology-specific health-related quality of life instruments in psoriasis patients. J Invest Dermatol 2007; 127: 2315–2322.

27. Twiss J, Meads DM, Preston EP, Crawford SR, McKenna SP. Can we rely on the Dermatology Life Quality Index as a measure of the impact of psoriasis or atopic dermatitis? J Invest Dermatol 2012; 132: 76–84.

28. Chren MM, Lasek RJ, Quinn LM, Mostow EN, Zyzanski SJ. SkinQ16, a quality-of-life measure for patients with skin disease: reliability, validity, and responsiveness. J Invest Dermatol 1996; 107: 707–713.

29. Nijsten TE, Sampogna F, Chren MM, Abeni DD. Testing and reducing Skindex-29 using Rasch analysis: Skindex-17. J Invest Dermatol 2006; 126: 1244–1250.

30. Anderson RT, Rajagopalan R. Development and validation of a quality of life instrument for cutaneous diseases. J Am Acad Dermatol 1997; 37: 41–50.

31. Morgan M, McCreedy R, Simpson J, Hay RJ. Dermatology quality of life scale – a measure of the impact of skin diseases. Br J Dermatol 1997; 136: 202–206.

32. Finlay AY, Kelly SE. Psoriasis – an index of disability. Clin Exp Dermatol 1987; 12: 8–11.

33. McKenna SP, Cook SA, Whalley D, Doward LC, Richards HL, Griffiths CE, et al. Development of the PSORIqol, a psoriasis-specific measure of quality of life designed for use in clinical practice and trials. Br J Dermatol 2003; 149: 323–331.

34. Nicholson K, Abramova L, Chren MM, Yeung J, Chon SY, Chen SC. A pilot quality-of-life instrument for acne rosacea. J Am Acad Dermatol 2007; 57: 213–221.

35. Ofenloch RF, Weisshaar E, Dumke AK, Molin S, Diepgen TL, Apfelbacher C. The Quality of Life in Hand Eczema Questionnaire (QOLHEQ): validation of the German version of a new disease-specific measure of quality of life for patients with hand eczema. Br J Dermatol 2014; 171: 304–312.

36. Bland JM, Dumville JC, Ashby RL, Gabe R, Stubbs N, Ad- derley U, et al. Validation of the VEINES-QOL quality of life instrument in venous leg ulcers: repeatability and validity study embedded in a randomised clinical trial. BMC Cardiovasc Disord 2015; 15: 85.

37. Motley RJ, Finlay AY. Practical use of a disability index in the routine management of acne. Clin Exp Dermatol 1992; 17: 1–3.

38. Dolte KS, Girman CJ, Hartmaier S, Roberts J, Bergfeld W, Waldstreicher J. Development of a health-related quality of life questionnaire for women with androgenetic alopecia. Clin Exp Dermatol 2000; 25: 637–642.

39. Martin AR, Lookingbill DP, Botek A, Light J, Thiboutot D, Girman CJ. Health-related quality of life among patients with facial acne – assessment of a new acne-specific questionnaire. Clin Exp Dermatol 2001; 26: 380–385.

40. Ando T, Hashiro M, Noda K, Adachi J, Hosoya R, Kamide R, et al. Development and validation of the psychosomatic scale for atopic dermatitis in adults. J Dermatol 2006; 33: 439–450.

41. Rhee JS, Matthews BA, Neuburg M, Logan BR, Burzynski M, Nattinger AB. Validation of a quality-of-life instrument for patients with nonmelanoma skin cancer. Arch Facial Plast Surg 2006; 8: 314–318.

42. Endo Y, Miyachi Y, Arakawa A. Development of a disease-specific instrument to measure quality of life in patients with alopecia areata. Eur J Dermatol 2012; 22: 531–536.

43. Liu SJ, Pandy A. Melasma quality of life measures. Dermatol Clin 2012; 30: 269–280.

44. Fabbrocini G, Panariello L, De Vita V, Vincenzi C, Lauro C, Nappo D, et al. Quality of life in alopecia areata: a disease-specific questionnaire. J Eur Acad Dermatol Venereol 2013; 27: e276–281.

45. Heisterberg MV, Menné T, Johansen JD. Fragrance allergy and quality of life – development and validation of a disease-specific quality of life instrument. Contact Dermatitis 2014; 70: 69–80.

46. Lilly E, Lu PD, Borovicka JH, Victorderon M, Kwasny MJ, West GP, et al. Development and validation of a vitiligo-specific quality-of-life instrument (VitiQoL). J Am Acad Dermatol 2013; 69: e11–e18.

47. Sebaratnam DF, Hanna AM, Chee SN, Frew JW, Venugopal SS, Daniel BS, et al. Development of a quality-of-life instrument (VitiQoL). J Am Acad Dermatol 2013; 69: e11–e18.

48.EuroQolGroup, Euroqol – a new facility for the measurement of health-related quality of life. Health Policy 1990; 16: 199–208.

49. Ware JE, Jr., Sherbourne CD. The MOS 36-item short-form health survey (SF-36). I. Conceptual framework and item selection. Med Care 1992; 30: 473–483.

50. Nordmeyer Group. What quality of life? World Health Organization Quality of Life Assessment. World Health Forum 1996; 17: 354–356.

51. The-WHOQOL-Group. Development of the World Health Organization WHOQOL-BREF quality of life assessment. Psychol Med 1998; 28: 551–558.