Systematic review of psychosocial needs assessment tools for caregivers of paediatric patients with dermatological conditions

Carleen Walsh, Gerard Leavey, Marian McLaughlin

ABSTRACT

Objective To identify validated dermatology-specific and disease-specific psychosocial needs assessment tools for caregivers of paediatric patients with dermatological conditions. A secondary objective was to assess the adequacy of their measurement properties.

Design Systematic review.

Data sources EMBASE, PsycINFO, MEDLINE (in Ovid SP), Cochrane, Cumulative Index to Nursing and Allied Health (EBSCO), U Search and Web of Science were searched (2000–5 October 2021). Grey literature, bibliographies, online databases of QoL tools and several trial registers were searched (2000–5 October 2021).

Eligibility criteria Eligible studies involved adult caregivers caring for a child (no age limit) with any form of skin condition. Predetermined exclusion criteria, as per protocol, were applied to the search results.

Data abstraction and synthesis Title, abstract, full-text screening and data abstraction (standardised forms) were done independently in duplicate. Both’s predefined methodological criteria assessed risk of bias. Narrative synthesis was used to present the findings.

Results 187 full-text articles were examined from a total of 8979 records. Most tools were generic QoL tools, relevant to spouse/partner or based on their child’s perception of the disease or assessed patients’ quality of life. Following quality appraisal, 26 articles were identified, and 11 tools (1 dermatology-specific and 10 disease-specific) were included. Information outcome domains were provided for each tool (study specific, questionnaire specific, adequacy of measurement properties and risk of bias). No literature was found pertaining to the use of these tools within healthcare settings and/or as e-tools.

Discussion With limited evidence supporting the quality of their methodological and measurement properties, this review will inform future dermatological Core Outcome Set development and improve evidence-based clinical decisions. Increasing demand on limited healthcare resources justifies the development of an accessible solution-focused psychosocial needs assessment e-tool to promote caregiver health outcomes.

Strengths and limitations of this study

► The first systematic review to provide a comprehensive overview of psychosocial assessment tools validated for use among dermatological caregivers of paediatric patients.
► This study was conducted with the involvement of a health and life subject-specific librarian and an international multidisciplinary expert group.
► The protocol was registered on the PROSPERO database (CRD42019159956), the COMET database and was conducted according to the recommendations from the Preferred Reporting Items for Systematic Reviews and Meta-Analyses 2020 statement and ENTREQ statement.
► Adequacy of measurement properties was assessed using Both et al’s criteria.
► Included articles were limited to being published in English between 2000 and 2021.

INTRODUCTION

Paediatric dermatology is a unique specialty in that children with lifelong and life-limiting skin disorders are increasingly being cared for by caregivers at home, which requires considerable cognitive, emotional and physical resources. Skin disease is the fourth leading cause of global disease burden with associated prevalence, care requirements and costs comparable with other diseases, such as cardiovascular disease and diabetes. Delayed identification of dermatological caregiver needs and provision of timely supports can seriously compromise the long-term psychosocial well-being of caregivers and particularly undermine the care and treatment of paediatric patients affected by rare or chronic skin disease. Caregivers of skin disease require similar systems of monitoring and integrated biopsychosocial support as other comparable chronic conditions.

The WHO directive, recent international guidelines and reports emphasise the importance of identifying psychosocial needs.
assessment tools for use among long-term caregivers, particularly self-referral models. Timely and appropriate identification of caregivers’ unmet psychosocial needs has the potential to reduce caregiver strain and increase their ability to provide quality care within the home at reduced public health cost. Although a psychosocial needs assessment could be considered preventative in nature, by anticipating caregiver burnout and decreasing the need for emergency interventions, there is a lack of evidence regarding the use of caregiver assessment tools within healthcare settings. To date, no comprehensive review of psychosocial needs assessment tools validated for use among informal dermatological caregivers of paediatric patients has been conducted. With increasing competition for valuable healthcare resources and services, there is an urgent need to reconceptualise global burden within the construct of ‘prevention is better than cure’ by informing evidence-based decisions and promoting caregiver health outcomes within day-to-day clinical practice.

Objectives
This review aimed to improve clinician access to existing dermatology-specific and disease-specific psychosocial needs assessment tools, validated for use among caregivers of paediatric patients with dermatological conditions. Additionally, this review assessed the adequacy of their measurement properties.

METHODS
This review was conducted according to the recommendations from the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) 2020 statement.17 The ENTREQ statement was read and guided in reporting the synthesis of the findings.18

Eligibility criteria
Studies that involved adult caregivers (age 18 years and over) caring for a child (no age limit) with any form of any skin condition were included. Predetermined exclusion criteria were adhered to (see protocol). Included articles were limited to being published in English between 01 January 2000 and 5 October 2021. This ensured that relevant assessment tools developed in the years before publication of the 2017 review were included as that review had limited their search to one database and quality-of-life measures only, which contrasts with the measures recommended by the Cochrane Skin Centre of Evidence Based Dermatology.

Information sources
MEDLINE, PsycINFO and Embase (OVID interface) and Cumulative Index to Nursing and Allied Health Literature (CINAHL) EBSCO were searched (1 January 2000–5 October 2021). Grey literature, bibliographies, online databases of QoL tools and several trial registers were also searched (1 January 2000–5 October 2021). A ‘snowball’ search was carried out to identify additional studies by manually searching the reference lists of all publications eligible for full-text review. The PRISMA flow diagram (figure 1) includes the number of records identified from each source.

Search strategy
One known relevant systematic review19 was used as a starting point to identify records. A draft search strategy was developed by using candidate search terms that were identified in the titles, abstracts and subject indexing of that systematic review. The full search strategy development process is included in online supplemental file 1. This strategy was tailored to the specifications of each of the databases searched and developed in collaboration with a subject-specific librarian (JA) and expert group. Each tailored database search strategy is included in online supplemental file 2. All search terms/categories used to search within the supplementary sources are included in online supplemental file 3.

Selection and data collection process
Title, abstract and full-text screening were conducted manually in duplicate (independently) by two reviewers (CW and GL). Extracted data from full-text articles was processed using three standardised extraction forms: (1) study-specific information included the name of the tool, country of origin, disease of affected patients, sample sizes used in each stage of its development and study setting; (2) questionnaire-specific information included the outcome domains, number of items and subscales, recall period, scoring system, respondent feedback and administration mode and time; (3) adequacy of measurement properties was evaluated using five methodological domains: validity, reliability, structure, interpretability and transferability. At the full-text screening stage, any discrepancies were resolved by discussion and, where necessary, the third author (MM) was consulted.

Risk of bias assessment
Risk of bias in the included studies was assessed independently by two reviewers using Both et al’s20 criteria, made possible by the similarities between the studies. Each methodological domain and item were graded for risk of bias using predefined criteria. Any discrepancies were resolved by consensus discussions (CW and GL) and, where necessary, by deferment to the third author (MM). No overall risk of bias judgement that summarised across domains was given due to the wide variation in assessment across domains within each tool. To improve the robustness of the synthesis and facilitate replicability, an overview of the domain definitions, items, effect measures, grades and criteria used in assessing the risk of bias is provided in online supplemental file 4.

Synthesis methods
In line with synthesis guidelines,22 a narrative approach was used to arrange the results into two categories: dermatology-specific and disease-specific tools psychosocial needs assessment tools. To ease identification of
variability between and within the included tools, results were also tabulated using the subheadings used in each of the three data extraction forms.

**Certainty assessment**
The robust search strategy was validated in MEDLINE when it successfully identified the one known systematic review as part of the search strategy development process (online supplemental file 1). Two authors (CW and GL) independently assessed the certainty of evidence by assessing risk of bias using a predefined checklist of criteria.

**Ethics approval**
Informed consent was obtained from all participating members of the expert group associated with the research project.

**Patient and public involvement (PPI)**
An international multidisciplinary expert group (n=15), including affected adults, clinical psychologists, clinical nurse specialists, consultant dermatologists, health policy advisors and caregivers, was established at the outset of the project (September 2017). Anonymity remains protected due to their ongoing involvement in another follow-on study. The Guidance for Reporting Involvement of Patients and the Public Short Form checklist was used to improve the reporting of PPI in our study. PPI helped identify the research question, guide in terms of review design (search strategy, inclusion and exclusion criteria and data extraction subheadings) and improve the dissemination of findings (invitations to poster and orally
RESULTS
This review identified 8979 records: 8256 records from database searching and 723 records from supplementary sources. After duplicates were removed (n=2577), 6402 records were available. Of the 6402 titles screened, 992 abstracts were screened, and 187 full-text articles were assessed for eligibility. This included 15 records identified from the one known systematic review. Of the 187 full-text articles assessed for eligibility, 161 records were excluded for reasons that met the exclusion criteria (PRISMA flow diagram; figure 1). No full-text records were included after snowballing reference lists (48 screened).

To improve transparency, summaries of the records identified during the initial and updated searches, for both databases and supplementary sources, are included in online supplemental files 3 and 5. PRISMA flow diagrams are included for both the initial search (1 January 2000–1 April 2020) (online supplemental file 5 (figure 1)) and updated search periods (1 April 2020–5 October 2020) (online supplemental file 5 (figure 2)) and provide a breakdown of the number of records identified for each database and supplementary source. The two full-text articles, identified in the updated search, were both excluded when assessed for eligibility. One record contained psychometric data resulting from a biased study design and statistical analysis ('validity was established in a limited range of subjects’, ‘the parents that responded to the survey were all mothers’, ‘single-institution cross-sectional study in Japan targeting parents of first-time patients less than 7 years old’). The other record identified the Family Dermatology Life Quality Index (FDLQ), which was already identified in the initial search.

The majority of existing, validated dermatological assessment tools identified were generic quality of life (QoL) tools and/or assess the patients’ QoL. Of those tools validated for use among caregivers, most were either relevant to spouse/partner or depend on the caregiver to complete but are based on their child’s perception of the disease (figure 1). Very few needs assessment tools were validated for use among caregivers of paediatric patients affected with dermatological disease. In summary, a total of 11 assessment tools were identified from the 26 articles included in this review. Ten disease-specific assessment tools were identified (The Psoriasis Family Index (PFI-15), Family Pso, Quality of life in Primary Caregivers of Children with Atopic Dermatitis (QPCAD), Childhood Atopic Dermatitis Impact Scale (CADIS), Parents’ Index of Quality of Life in Atopic Dermatitis (PIQoL-AD), Dermatitis Family Impact (DFI), Parental Self-Efficacy with Eczema Care Index (PASECI), CareGiver Oncology Quality of Life (CarGOQoL), Epidermolysis Bullosa – Burden of Disease (EB-BoD) and Family Burden of Ichthyosis (FBI) and one dermatology-specific assessment tool was identified (FDLQ). Table 1 provides a summary of study-specific information and includes the name of tool, country of origin, disease of affected patient, sample sizes and study setting. Table 2 summarises questionnaire-specific information under the subheadings outcome domains, subscales, number of items, recall period, scoring system and administration time. Table 3 provides an overview of the adequacy of the measurement properties of the included tools, including transferability, reliability, validity, structural and interpretability. Table 4 provides a graded risk of bias assessment (using the predefined criteria) of each methodological domain and item for each of the 11 tools.

Disease-specific needs assessment tools

The Psoriasis Family Index (PFI-15) is recommended for use alongside a dermatology-specific tool. As it is assessed on current time only, it does not rely on accurate recall. However, due to the small sample size, factor analysis could not be done, and there is a lack of comparison of PFI scores with other generic family QoL scales. In order to achieve its Cronbach’s alpha value (0.86), it was necessary to delete five items. It has a weaker focus on the emotional aspects of living with affected members. Those accompanying patients to the primary care centre and inpatients were not included in the creation of the PFI, which restricts the generalisation of the quantitative findings.

The Family Pso was created from interviews (n=95) with psoriasis patients and their family members. Three experts (no caregiver involvement) decided the generation items for piloting and item reduction. Other limitations include that a small sample was used in its testing and were predominantly female partners of the interviewees. Its advantages include that the wording is more focused on emotional aspects of caregiving as opposed to HR-QoL.

Four tools were found that assess the impact of atopic dermatitis on the family. The QPCAD has a 1-week recall and has been validated for use among primary caregivers of children with AD in the Japanese version only. Convergent validity requires further study, and only caregivers of mild and moderate patients from an urban area were included in the study.

The CADIS is validated for use with both patients and parents of patients younger than 6 years. Rasch analysis reduced the tool to a 45-item version, which is responsive to clinical change in AD.

The Parents’ Index of Quality of Life in Atopic Dermatitis (PIQoL-AD) assesses the impact of AD on caregivers of affected children, aged 8 years or younger. The PIQoL-AD adopts a dichotomous response system, which is less sensitive to subtle changes in HR-QoL and includes only items that consider the negative aspects of psychological well-being.

The DFI tool is the most widely reported in studies, having been used in over 750 clinical trials, although often at longer intervals despite being
validated for use with a 1-week recall period. As most of DFI studies are in secondary care hospitals, there exists the possibility of maximising the chances of the DFI scores showing significant improvements following an intervention. Dodington’s review found that internal consistency and test–retest reliability was adequately demonstrated but highlighted that psychometric measures were less well established due to a lack of validation for use with a 1-week recall period. As most of DFI studies are in secondary care hospitals, there exists the possibility of maximising the chances of the DFI scores showing significant improvements following an intervention. Dodington’s review found that internal consistency and test–retest reliability was adequately demonstrated but highlighted that psychometric measures were less well established due to a lack of validation for use with a 1-week recall period. As most of DFI studies are in secondary care hospitals, there exists the possibility of maximising the chances of the DFI scores showing significant improvements following an intervention. Dodington’s review found that internal consistency and test–retest reliability was adequately demonstrated but highlighted that psychometric measures were less well established due to a lack of validation for use with a 1-week recall period. As most of DFI studies are in secondary care hospitals, there exists the possibility of maximising the chances of the DFI scores showing significant improvements following an intervention. Dodington’s review found that internal consistency and test–retest reliability was adequately demonstrated but highlighted that psychometric measures were less well established due to a lack of validation for use with a 1-week recall period. As most of DFI studies are in secondary care hospitals, there exists the possibility of maximising the chances of the DFI scores showing significant improvements following an intervention. Dodington’s review found that internal consistency and test–retest reliability was adequately demonstrated but highlighted that psychometric measures were less well established due to a lack of validation for use with a 1-week recall period. As most of DFI studies are in secondary care hospitals, there exists the possibility of maximising the chances of the DFI scores showing significant improvements following an intervention. Dodington’s review found that internal consistency and test–retest reliability was adequately demonstrated but highlighted that psychometric measures were less well established due to a lack of validation for use with a 1-week recall period. As most of DFI studies are in secondary care hospitals, there exists the possibility of maximising the chances of the DFI scores showing significant improvements following an intervention. Dodington’s review found that internal consistency and test–retest reliability was adequately demonstrated but highlighted that psychometric measures were less well established due to a lack

Table 1  Study-specific information relevant to included assessment tools

<table>
<thead>
<tr>
<th>References of included publications (first author, year, reference)</th>
<th>Country of origin</th>
<th>Disease of affected patients</th>
<th>Name of measurement instrument</th>
<th>Sample size (n)</th>
<th>Study setting</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mrowietz et al (2017)27 Germany</td>
<td>Psoriasis</td>
<td>Family Pso</td>
<td>Interviews (14) Pilot (96) Validation (96)</td>
<td>Monocentric Outpatient clinic</td>
<td></td>
</tr>
<tr>
<td>Kondo-Endo et al (2009)28 Japan</td>
<td>Atopic dermatitis</td>
<td>QoL in Primary Caregivers of children with Atopic Dermatitis</td>
<td>Interviews (unknown) Pilot (33) Validation (400)</td>
<td>Monocentric Outpatient clinic</td>
<td></td>
</tr>
<tr>
<td>Chamlin et al (2005)29 USA</td>
<td>Atopic dermatitis</td>
<td>Childhood Atopic Dermatitis Impact Scale</td>
<td>Interviews (unknown) Pilot (20) Validation (300)</td>
<td>Two dermatology paediatric practices (San Francisco and Chicago)</td>
<td></td>
</tr>
<tr>
<td>McKenna et al (2005)35 UK, Netherlands, Italy, Spain, USA, Switzerland, Germany, France (simultaneous development)</td>
<td>Atopic dermatitis</td>
<td>Parent’s Index QoL – Atopic Dermatitis</td>
<td>Interviews (140 total) Validation (ranged between countries 45–328)</td>
<td>Monocentric Outpatient clinic</td>
<td></td>
</tr>
<tr>
<td>Lawson et al (1998)31 UK</td>
<td>Dermatitis</td>
<td>Dermatitis Family Impact</td>
<td>Interviews (29) and focus groups (10) Pilot (14) Validation (56)</td>
<td>Monocentric Outpatient clinic</td>
<td></td>
</tr>
<tr>
<td>Minaya et al (2012)33 France</td>
<td>Skin cancer</td>
<td>CareGiver Oncology Quality of Life</td>
<td>Interviews (77) Pilot (837) Validation (unknown)</td>
<td>Monocentric Outpatient clinic</td>
<td></td>
</tr>
<tr>
<td>Dufresne et al (2015)34 France</td>
<td>Epidermolysis bullosa</td>
<td>Epidermolysis Bullosa – Burden of Disease</td>
<td>Complaints (23) informed item generation Pilot (Lionbridge institution) Validation (55)</td>
<td>Monocentric Outpatient clinic</td>
<td></td>
</tr>
<tr>
<td>Dufresne et al (2013)35 France</td>
<td>Ichthyosis</td>
<td>Family Burden Ichthyosis</td>
<td>Interviews (94) Pilot (42) Validation (30)</td>
<td>Monocentric Outpatient clinic</td>
<td></td>
</tr>
<tr>
<td>Basra et al (2008)36 UK</td>
<td>All – general dermatology instrument</td>
<td>Family Dermatology Life Quality Index</td>
<td>Interviews (50) Pilot (20) Validation (14)</td>
<td>Monocentric Outpatient clinic</td>
<td></td>
</tr>
</tbody>
</table>

QoL, quality of life.
<table>
<thead>
<tr>
<th>Name of measurement instrument</th>
<th>Outcome domains measured</th>
<th>Number of items and subscales</th>
<th>Recall period</th>
<th>Scoring system</th>
<th>Respondent feedback</th>
<th>Admin mode (time in minutes)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Psoriasis Family Index (PFI-15)</td>
<td>Social life, leisure activities, sporting activities, people's reactions, worry about future, housework, relationships treatment duration, clothing shopping and sleep</td>
<td>15 items</td>
<td>Now</td>
<td>4-point scale (0-3)</td>
<td>Brief in length, Simple to administer, score and interpret, Weak evidence of alternative forms.</td>
<td>Self-administered (2)</td>
</tr>
<tr>
<td>Family Pso27</td>
<td>Emotional domain – emotional impact. Social domain –impact on daily activities and work/school and treatment. Leisure domain – influence on leisure/personal relationships</td>
<td>15 items</td>
<td>1 month</td>
<td>5-point Likert format (0–4) and ‘Does not apply’</td>
<td>Brief in length, Simple to administer, score and interpret, Weak evidence of alternative forms.</td>
<td>Self-administered (3)</td>
</tr>
<tr>
<td>QoL in Primary Caregivers of children with Atopic Dermatitis28</td>
<td>Achievement (3) Worry (6) Family cooperation (3) Exhaustion (8)</td>
<td>19 items</td>
<td>Past week</td>
<td>5-point scale (none to extremely)</td>
<td>Brief in length, Moderate to administer, score and interpret, Conflicting evidence of alternative forms.</td>
<td>Self-report (unknown)</td>
</tr>
<tr>
<td>Childhood Atopic Dermatitis Impact Scale29</td>
<td>Impact on family (three domains) Sleep and emotions Family and social function</td>
<td>45 items</td>
<td>1 month</td>
<td>5-point scale (never to all the time)</td>
<td>Long in length and problems of acceptability. Moderate to administer, score and interpret, Absent evidence of alternative forms.</td>
<td>Self-administered (6)</td>
</tr>
<tr>
<td>Parent's Index QoL – Atopic Dermatitis30</td>
<td>One domain – needs that can be influenced by a child with a diagnosis of AD.</td>
<td>28 items</td>
<td>Not reported</td>
<td>5-point scale (never to all the time)</td>
<td>Brief in length, Simple to administer, score and interpret, Weak evidence of alternative forms.</td>
<td>Self-administered (3)</td>
</tr>
<tr>
<td>Dermatitis Family Impact31</td>
<td>Personal relationships and helping with treatment, Food and feeding, sleep, housework shopping, financial, leisure tiredness and emotional distress</td>
<td>10 items</td>
<td>1 week</td>
<td>4-point scale (not at all, a little, a lot, very much)</td>
<td>Brief in length, Simple to administer, score and interpret, Weak evidence of alternative forms.</td>
<td>Self-administered (unknown)</td>
</tr>
<tr>
<td>Parental Self-Efficacy with Eczema Care Index32</td>
<td>Managing medications Managing eczema and symptoms Communication with healthcare teams Managing personal challenges</td>
<td>29 items</td>
<td>1 week preintervention and 4 weeks postintervention</td>
<td>11-point Likert Scale</td>
<td>Brief in length, Simple to administer, score and interpret, Weak evidence of alternative forms.</td>
<td>Clinician administered (3)</td>
</tr>
<tr>
<td>CareGiver Oncology Quality of Life questionnaire33</td>
<td>Psychological well-being, burden, relationship with healthcare, administration and finances, coping, physical well-being, self-esteem, leisure time, social support and private life</td>
<td>29 items</td>
<td>1 week</td>
<td>5-point Likert scale (never/not at all, rarely/a little, sometimes/somewhat, often/a lot, always/very much)</td>
<td>Brief in length, Simple to administer, score and interpret, Weak evidence of alternative forms.</td>
<td>Self-administered (3)</td>
</tr>
<tr>
<td>Epidermolysis Bullosa – Burden of Disease34</td>
<td>Economic and social impact (5) Family life (7) Disease and treatment (5) Child's life (3)</td>
<td>20 items</td>
<td>Not stated</td>
<td>7-point scale (always, very often, often, sometimes, rarely, never and not applicable)</td>
<td>Moderate to administer, score and interpret, Absent evidence of alternative forms. Long in length and problems of acceptability.</td>
<td>Self-administered (unknown)</td>
</tr>
<tr>
<td>Family Burden Ichthyosis35</td>
<td>Work and psychological impact, daily life, pain, familial and personal relationships</td>
<td>25 items</td>
<td>Not stated</td>
<td>4-point scale (definitely yes, maybe, definitely not and I don't know)</td>
<td>Long in length and problems of acceptability. Moderate to administer, score and interpret.</td>
<td>Self-administered (3)</td>
</tr>
<tr>
<td>Family Dermatology Life Quality Index36</td>
<td>Housework and expenditure Emotional and physical well-being Impact on study/job, social life burden of care, leisure activities</td>
<td>10 items</td>
<td>1 month</td>
<td>4-point scale (not at all, not applicable, a little, quite a lot and very much)</td>
<td>Brief in length, Simple to administer, score and interpret, Weak evidence of alternative forms.</td>
<td>Self-administered (3)</td>
</tr>
</tbody>
</table>

QoL, quality of life.
Table 3  Adequacy of the measurement properties relevant to included assessment tools with excellent and good methodological quality

<table>
<thead>
<tr>
<th>Name of measurement instrument</th>
<th>Transferability</th>
<th>Reliability</th>
<th>Validity</th>
<th>Structure</th>
<th>Interpretability</th>
</tr>
</thead>
<tbody>
<tr>
<td>Family Psq</td>
<td>Never translated using guidelines. Never analysed in a cultural equivalence study.</td>
<td>Cronbach’s α &gt; 0.70. Retest reliability: k or ICC not reported or correlation coefficient &lt;0.70.</td>
<td>Conceptual – more focused on objective/subjective domains. Construct – no information. Convergent &lt;0.70.</td>
<td>Factor analysis. Weak sensitivity to detect changes. Weak item bias.</td>
<td>Norms: general nor dermatology patients. Categorisation: not reported. MCID: not reported.</td>
</tr>
<tr>
<td>QoL in Primary Caregivers of children with Atopic Dermatitis</td>
<td>Never translated using guidelines. Never analysed in a cultural equivalence study.</td>
<td>Cronbach’s α &gt; 0.70. Retest reliability: k or ICC &gt; 0.70.</td>
<td>Conceptual: more focused on objective/subjective domains. Construct &lt; 75% results in accordance with hypothesis. Convergent &lt;0.70.</td>
<td>Satisfactory response to change in disease severity. Satisfactory test-retest reliability.</td>
<td>Norms: general nor dermatology patients. Categorisation: not reported. MCID: not reported.</td>
</tr>
<tr>
<td>Childhood Atopic Dermatitis Impact Scale</td>
<td>Sometimes translated using guidelines. Never analysed in a cultural equivalence study.</td>
<td>Cronbach’s α &gt; 0.70. Retest reliability: k or ICC &gt; 0.70.</td>
<td>Conceptual: well balanced. Construct &lt; 75% results in accordance with hypothesis. Convergent &lt;0.70.</td>
<td>IRT. Strong sensitivity to detect changes. Weak item bias.</td>
<td>Norms: general nor dermatology patients. Categorisation: not reported. MCID: not reported.</td>
</tr>
<tr>
<td>Parent’s Index QoL – Atopic Dermatitis</td>
<td>Always translated using guidelines. Never analysed in a cultural equivalence study.</td>
<td>Cronbach’s α &gt; 0.70. Retest reliability: k or ICC &gt; 0.70.</td>
<td>Conceptual: more focused on objective/subjective domains. Construct &lt; 75% results in accordance with hypothesis. Convergent &lt;0.70.</td>
<td>IRT. Strong sensitivity to detect changes item bias. Strong item bias.</td>
<td>Norms: general nor dermatology patients. Categorisation: known in heterogeneous sample. MCID: not reported.</td>
</tr>
<tr>
<td>Dermatitis Family Impact</td>
<td>Always translated using guidelines. Sometimes analysed in a cultural equivalence study.</td>
<td>Cronbach’s α &gt; 0.70. Retest reliability: k or ICC &gt; 0.70.</td>
<td>Conceptual: well balanced. Construct &lt; 75% results in accordance with hypothesis. Convergent &gt;0.70.</td>
<td>Factor analysis. Satisfactory response to change in disease severity. Weak item bias.</td>
<td>Norms: General nor dermatology patients Categorisation: not reported. MCID: not reported.</td>
</tr>
<tr>
<td>Parental Self-Efficacy with Eczema Care Index</td>
<td>Always translated using guidelines. Never analysed in a cultural equivalence study.</td>
<td>Cronbach’s α &gt; 0.70. Retest reliability: k or ICC &gt; 0.70.</td>
<td>Conceptual: well balanced. Construct &gt; 75% results in accordance with hypothesis. Convergent &gt;0.70.</td>
<td>Factor analysis. Satisfactory response to change in disease severity. Weak item bias.</td>
<td>Norms: general nor dermatology patients. Categorisation: used distribution-based techniques. MCID: not reported.</td>
</tr>
<tr>
<td>CareGiver Oncology Quality of Life Questionnaire</td>
<td>Sometimes translated using guidelines. Never analysed in a cultural equivalence study.</td>
<td>Cronbach’s α &gt; 0.70. Retest reliability: k or ICC &gt; 0.70.</td>
<td>Conceptual: more focused on objective/subjective domains. Construct &gt; 75% results in accordance with hypothesis. Convergent &gt;0.70.</td>
<td>Factor analysis. Low/moderate sensitivity to changes item bias. Strong item bias.</td>
<td>Norms: general nor dermatology patients. Categorisation: not reported. MCID: not reported.</td>
</tr>
<tr>
<td>Epidermolysis Bullosa – Burden of Disease</td>
<td>Sometimes translated using guidelines. Never analysed in a cultural equivalence study.</td>
<td>Cronbach’s α &gt; 0.70. Retest reliability: k or ICC &gt; 0.70.</td>
<td>Conceptual: well balanced. Construct &lt; 75% results in accordance with hypothesis. Convergent &gt;0.70.</td>
<td>Factor analysis. Weak sensitivity to detect changes. Weak item bias.</td>
<td>Norms: general nor dermatology patients. Categorisation: not reported. MCID: not reported.</td>
</tr>
<tr>
<td>Family Burden Ichthyosis</td>
<td>Sometimes translated using guidelines. Never analysed in a cultural equivalence study.</td>
<td>Cronbach’s α &gt; 0.70. Retest reliability: k or ICC not reported or correlation coefficient &lt;0.70.</td>
<td>Conceptual: well balanced domains Construct &gt;75% results in accordance with hypothesis. Convergent &lt;0.70.</td>
<td>No factor analysis or IRT. Weak sensitivity to detect changes. Weak item bias.</td>
<td>Norms: general nor dermatology patients. Categorisation: not reported. MCID: not reported.</td>
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<tr>
<td>Family Dermatology Life Quality Index</td>
<td>Always translated using guidelines. Never analysed in a cultural equivalence study.</td>
<td>Cronbach’s α &gt; 0.70. Retest reliability: k or ICC &gt; 0.70.</td>
<td>Conceptual: well balanced domains Construct &gt;75% results in accordance with hypothesis. Convergent &gt;0.70.</td>
<td>Factor analysis. Strong sensitivity to detect changes. Weak item bias.</td>
<td>Norms: general nor dermatology patients. Categorisation: not reported. MCID: not reported.</td>
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IRT, item response theory; IC, internal consistency; ICC, intraclass correlation coefficient; MCID, minimal clinically important difference.

of vigour in both the creation and validation processes. No valid score banding descriptors of DFI score meanings are included, and no information to establish the MCID of DFI score is available. No studies demonstrated dimensionality, factor structure or differential item functioning.
Table 4  Evaluation of disease-specific and dermatology-specific tools (risk of bias assessment criteria outlined in online supplemental file 4)

<table>
<thead>
<tr>
<th>Criteria</th>
<th>PFI-15</th>
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<th>CADIS</th>
<th>PiQoL-AD</th>
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<th>PASECI</th>
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Objective and subjective domains are described by Muldoon et al (1998).1

CADIS, Childhood Atopic Dermatitis Impact Scale; CarGOQoL, The CareGiver Oncology Quality of Life; DFI, Dermatitis Family Index; EB-BoD, Epidermolysis Bullosa Burden of Disease; Family Pso, Family Psoriasis; FBI, Family Burden Ichthyosis; FDLQI, Family Dermatology Life Quality Index; MCID, minimal clinically important difference; PASECI, Parental Self-Efficacy with Eczema Care Index; PFI-15, Psoriasis Family Index; PiQoL-AD, Parents’ Index of Quality of Life in Atopic Dermatitis; QoL, quality of life; QPCAD, QoL in Primary Caregivers of Children with Atopic Dermatitis.
The final tool included in this review was the PASEC. It is a generalised self-efficacy scale focusing on the management of four subscales: medication, symptoms, personal challenges and communication with healthcare teams. It has a two-factor structure that considers the performance of routine management tasks and the management of child symptoms and behaviour. There was reliance on self-reported data, potentially affecting the fidelity of the results. More research is needed on banding and categorisation.

Validation of the CarGQoL was carried out using dermatology experts other than caregivers. Several non-optimal indicators of validity are indicated in table 4.

The EB-BoD tool needed to remove non-discriminatory items, such as frustration and guilt, from the original FBI during its creation. It requires further validation in larger EB patient and/or caregiver groups before being revalidated for use in other languages and cultures.

The FBI is the only validated disease-specific questionnaire that measures the concept of burden for ichthyosis caregivers. The monocentric study used parents and their affected children in the creation of verbatim using an unnamed French social assessment, which could not be accessed for this review. Selection bias was a possibility as 40% of participants cared for those affected by severe forms of ichthyosis (severity score 50 or greater). Limitations include that validation of the FBI was carried out using parents of children affected with only the severest forms of ichthyosis (severity score 50 or greater). Although itch is one of the significant challenges named by parents of children affected with ichthyosis (third most significant impact during the validation of the DFI), it does not feature as an item. Similarly, no items relate to pain in the finalised FBI. Verification of its psychometric properties, preferably in a multicentre study, is required. Caregiver feedback included that the finalised generation items were negatively phrased. The original French questionnaire has been linguistically and culturally adopted in Italy.

Dermatology-specific needs assessment tools

The FDLQI is the most used dermatology-specific Health-Related Quality of Life (HR-QoL). The psychosocial impact loaded six items (emotional impact, physical well-being, impact on relationships, leisure, social life and people’s reactions) and the physical impact loaded four items (burden, effect on job/study, household expenditure and housework). Fifty semistructured interviews took place that informed the items generated for testing during piloting. The feedback from these interviews has been termed ‘the greater concept’. Piloting of the 19 items occurred with 20 parents or partners of those originally interviewed, potentially introducing bias. Limitations include that the life course of skin disease is not reflected in the FDLQI and that it depends on recall accuracy. Definitions, such as MID, and the meaning of FDLQI scores are missing and future research is required to show the unidimensionality of the tool. The FDLQI was not tested for responsiveness for clinical change in a hospital or intervention context. Several items cannot discriminate between inflammatory and uninflammatory groups.

One common theme that emerged was the variation in methodological rigour used in measuring informal dermatological caregiver needs. Using the risk of bias assessment, each of the reviewed tools indicated an incomplete psychometric overview meaning that the generalisability and interpretation of results remain limited. Each reviewed tool (11 of 11; 100%) evaluated four or more psychometric properties. They do not comply with the OMERACT filter criteria and consequently are unable to be included in the development of a future Core Outcome Set (COS).

In terms of structure, five tools reported the use of factor analysis. Three tools reported the use of the more recently developed item response theory (IRT) to determine psychometric properties. Other tools neither reported factor analysis or IRT. Apart from two tools reporting strong item bias, the other nine tools reported weak item bias. One tool reported the use of distribution-based categorisation techniques, but the other 10 tools did not report on categorisation. MCID was not reported for any tool other than one.

In terms of reliability, all tools reported a high internal consistency (IC >0.95). Two tools did not report their retest reliability. One reported a weak retest reliability (ICC <0.70), while the other eight tools reported a good retest reliability (ICC >0.70). In terms of conceptual validity, four tools have less well-balanced domains. The other seven tools include well-balanced domains. No information is given regarding the construct validity for one tool. Five tools demonstrate that <75% of results are in accordance with their hypothesis and five tools demonstrate that >75% of results are in accordance with their hypothesis. The majority of tools demonstrate poor convergent validity apart from two (>.70). The PFI-15 provides no information on convergent validity. The other eight tools in this review show a convergent validity value of <0.70.

**DISCUSSION**

This is the first systematic review to address gaps in the existing evidence base around the identification of appropriate psychosocial needs assessment for caregivers of paediatric patients with dermatological conditions. This topic represents an emerging area for which there is a lack of up-to-date good quality synthesised evidence. With increasing numbers of paediatric patients of chronic skin disease being cared for by informal caregivers, often with limited medical training, key international multidisciplinary stakeholders (including clinicians, dermatological caregivers and policymakers) emphasised an urgent need to improve clinician awareness of existing needs assessment tools, to help them make informed evidence-based
decisions relating to assessment. The need to promote caregiver health outcomes within day-to-day clinical practice has become even more significant during COVID-19, a period of enhanced social isolation and increased caregiver hypervigilance and burnout.

This review identified 11 psychosocial needs assessment tools validated for use among caregivers of paediatric patients with dermatological conditions. A narrative approach was used to arrange the reviewed tools into two groups: dermatology-specific and disease-specific tools. To ease identification of risk of bias, study variability and measurement properties between and within the included tools, results were additionally tabulated using the predefined subheadings on the data extraction forms.

Although skin disease may be characterised at times by unpredictable episodes in symptom severity, that requires similar systems of monitoring and integrated biopsychosocial support as other chronic conditions, our review highlights the lack of literature pertaining to the use of these assessment tools in healthcare settings. This review suggests that the mismatch between the recognised impact of caregiving for skin disease and the failure of practitioners to effectively engage with its management may be attributed to the biomedical model of assessment reflected in existing tools.

In contrast to the tools reviewed, which used measures of other constructs as a proxy for caregivers’ need, it appears vital to directly assess informal dermatological caregivers’ needs (at problem area and support level) and plan for how that knowledge will be used to help support these needs.

Similarly, future assessments should use the scope of the International Classification of Functioning, Disability, and Health, to inform their caregiver framework in terms of contextual factors and in terms of functioning and disability. Despite the recognised difficulty of assessing chronic pathologies by clinical or QoL aspects alone, most tools identified in this review were generic QoL tools. The European Academy of Dermatology and Venereology Quality of Life Task Force, Cochrane Skin Centre of Evidence Based Dermatology and the Harmonising Outcome Measures for Eczema initiative reinforce that generic QoL assessments do not encompass the many factors that contribute to the psychosocial burden of skin disease and are not as sensitive, responsive or relevant to individual patients or their caregivers.

We considered appropriate measurement tools to be theoretically driven, rigorously conceptualised with input from caregivers at each stage, consider disease life course, tested for validity and reliability and intended to assess caregiver needs in relevant settings. Conceptual and theoretical work on dermatological caregivers’ needs could have been relatively lacking because of the varying degree by which the tools were informed by caregiver experience, with minimal description of the questionnaire development process, absence of or exclusionary key definitions such as family, caregiver and domain and participants were not asked to clarify their relationship to the patient attending the outpatient clinics. Some of the tools only included items for the negative aspect of psychological well-being.

Healthcare teams require access to validated assessment tools that consider all dimensions along the care continuum and that do not use measures of other constructs as a proxy for caregivers’ needs to provide culturally sensitive care. An international multicentric approach could best address variables including culture, demographics and disease severity. Although none of the reviewed assessment tools allow for the assessment of disease variables, including disease severity, we recommend that future needs assessment tools include disease parameters when designing their assessment framework. Dufresne found that increased disease severity led to increased caregiver burden, suggesting that tools that assess factors relevant to clinical severity of disease could better inform the types of supports needed long term.

Future assessment should be practical and feasible for daily use within busy clinics. A self-reporting psychosocial needs assessment e-tool, developed to identify caregiver needs (at both problem and support level), could best serve to address non-clinical barriers to assessment, including lack of time, support staff and easy tools, to reduce the reported high rates of non-use of validated tools within daily practice.

Research reinforces improved care recipient and caregiver outcomes when caregivers are facilitated to regularly self-report perceived needs enabling clinicians to identify and/or triage unmet psychosocial care needs.

**Strengths and limitations**

Strengths include a published protocol, a multidisciplinary expert group and health science librarian involved in the design of the review, a comprehensive literature search, information provision on study, questionnaire, measurement properties and risk of bias. This review also provides key recommendations for future research. Although time was needed to ensure that members were involved as equal partners in debates and decisions around key issues, benefits of PPI included having experts with lived experience who creatively contributed towards the methodology. Limitations included studies published in the English language between 2000 and 2021.

To enhance the chances of developing a truer set of outcome domains for improved COS uptake, future assessments should adopt a more thorough typology to assess the degree to which deficits in caregivers’ needs are present and to develop transparent conceptual frameworks that include key definitions and that are built on a hybrid model using good quality caregiver frameworks alongside qualitative feedback from large and culturally diverse international cohorts of caregivers. With increased emphasis on e-healthcare, it seems both desirable and practical to conceptualise an accessible and solution-based model of future e-assessment that can address recognised healthcare challenges, including limited clinic time, poor caregiver identification and healthcare communication, allowing for timely identification and/or triage of unmet psychosocial needs by practitioners.
while strengthening a caregiver’s sense of autonomy, coping ability and resilience.59 60 To inform the development of solution-focused assessment e-tools, it is important that research is also conducted into which supports are rated as most important by informal dermatological caregivers.

CONCLUSION

Although no gold standard tool exists for measuring the psychosocial needs of dermatological caregivers, this comprehensive review improves clinician awareness and knowledge of eleven validated psychosocial needs assessment tools for caregivers of paediatric patients with dermatological conditions. It is hoped that this review will inform the development of solution-based models of outcome assessment for improved dermatology care coordination. As dermatological caregiving research moves forward with significant public and private investment, rigorous measurement of caregivers’ needs is essential for the development of social services, public policies and improved COS uptake. These findings have implications for clinical practice, service development and future research and reinforce that attitude towards caregivers is pivotal in developing assessment for the purpose of accessing supports and services.

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