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Discovering the hidden benefits of cognitive interviewing in two languages: The first phase of a validation study of the Integrated Palliative care Outcome Scale

Eva K Schildmann¹, E Iris Groeneveld², Johannes Denzel¹, Alice Brown², Florian Bernhardt¹, Katharine Bailey², Ping Guo², Christina Ramsenthaler², Natasha Lovell², Irene J Higginson², Claudia Bausewein¹ and Fliss EM Murtagh²

Abstract

Background: The Integrated Palliative care Outcome Scale is a newly developed advancement of the Palliative care Outcome Scale. It assesses patient-reported symptoms and other concerns. Cognitive interviewing is recommended for questionnaire refinement but not adopted widely in palliative care research.

Aim: To explore German- and English-speaking patients' views on the Integrated Palliative care Outcome Scale with a focus on comprehensibility and acceptability, and subsequently refine the questionnaire.

Methods: Bi-national (United Kingdom/Germany) cognitive interview study using ‘think aloud’ and verbal probing techniques. Interviews were audio-recorded, transcribed verbatim and analysed using thematic analysis and pre-defined categories. Results from both countries were collated and discussed. The Integrated Palliative care Outcome Scale was then refined by consensus.

Setting/participants: Purposely sampled patients from four palliative care teams in palliative care units, general hospital wards and in the community.

Results: A total of 15 German and 10 UK interviews were conducted. Overall, comprehension and acceptability of the Integrated Palliative care Outcome Scale were good. Identified difficulties comprised the following: (1) comprehension problems with specific terms (e.g. ‘mouth problems’) and length of answer options; (2) judgement difficulties, for example, due to the 3-day recall for questions; and (3) layout problems. Combining the results from both countries (e.g. regarding ‘felt good about yourself’) and discussing them from both languages’ perspectives resulted in wider consideration of the items’ meaning, enabling more detailed refinement.

Conclusion: Cognitive interviewing proved valuable to increase face and content validity of the questionnaire. The concurrent approach in two languages – to our knowledge the first such approach in palliative care – benefited the refinement. Psychometric validation of the refined Integrated Palliative care Outcome Scale is now underway.

Keywords

Patient-reported outcome measures, outcome measurement, cognitive interviewing, palliative care

What is already known about the topic?

- Outcome measures need to be specifically developed in palliative care populations to reflect the concerns of advanced illness and ensure minimal respondent burden.
- The Palliative care Outcome Scale (POS) and Palliative care Outcome Scale–symptom module (POS-S) are brief, valid and reliable questionnaires developed for this population.
- However, there is some overlap, and patient and clinician feedback indicates scope for further refinement.
- Cognitive interviewing is recommended for questionnaire development but not adopted widely in palliative care research.

¹Department of Palliative Medicine, Munich University Hospital, Munich, Germany
²Department of Palliative Care, Policy & Rehabilitation, Cicely Saunders Institute, King’s College London, London, UK

Corresponding author:
Eva K Schildmann, Department of Palliative Medicine, Munich University Hospital, Marchioninistr. 15, 81377 Munich, Germany.
Email: Eva.schildmann@med.uni-muenchen.de
What this paper adds:

- In response to patients’ and clinicians’ demands, we present a refined measure (Integrated Palliative care Outcome Scale (IPOS)) that has been shaped by the patients’ perspectives.
- This study demonstrates once more the importance of cognitive interviewing in the refinement of outcome measures and that it can be done effectively in a palliative care population.
- The concurrent use of cognitive interviewing in two languages – for the first time in palliative care – provided additional valuable information, resulted in wider consideration of the meaning and context of each item and led to a better overall refinement of the IPOS questionnaire.

Implications for practice, theory or policy

- IPOS is now available for clinical use; it is one of the few brief measures validated in this population that covers all relevant domains.
- The refined version of the IPOS is currently undergoing phase II of validation in Germany and the United Kingdom to demonstrate the psychometric properties.
- Wider awareness and use of cognitive interviewing in measure refinement in palliative care is crucial, and cognitive interviewing in more than one language can have added value.

Background

Patient-reported outcome measures (PROMs) are questionnaires completed by patients to measure their perceptions of their functional status and well-being. The Integrated Palliative care Outcome Scale (IPOS) is a PROM and a newly developed advancement of the Palliative care Outcome Scale (POS) which assesses symptoms and other palliative care (PC) concerns as perceived by the patient. There are versions for patients and healthcare professionals.

The POS was developed in 1999 with the involvement of patients and has been validated in a variety of settings: hospice in-patient care, day care, home care, hospital in-patient care and community and outpatient services. It has been used to inform clinical practice, to monitor service interventions, to evaluate and improve the quality of care and as a research tool. The POS team has currently endorsed 11 translations, and POS is being used in over 20 countries. To describe the impact of PC-specific symptoms, a symptom module (the POS-S) was developed as an addition to the POS. Requests from clinical colleagues to merge POS and POS-S triggered the development of IPOS (see Figure 1). The IPOS is based on POS and POS-S, with changes to the wording of certain questions and response options. The 10 questions cover patients’ main concerns, common symptoms, patient and family distress, existential well-being, sharing feelings with family, information received and practical concerns.

Adopting a recommended quality criterion for questionnaires, we aimed to involve patients in the questionnaire development through cognitive interviewing to ensure face and content validity and acceptability. Cognitive interviewing is increasingly used as a step in the refinement of survey questions and PROMs. A complex cognitive process takes place when research participants answer questionnaires. This includes comprehension of the question, retrieval of information from memory, judgement processes to estimate an answer and formulation of a response. The European Organisation for Research and Treatment of Cancer (EORTC) guidelines for questionnaire development recommend cross-language pre-testing. Accordingly, we aimed to base the IPOS refinement on views from German- and English-speaking patients and have the refined questionnaires available for concurrent psychometric validation in both countries.
Methods

Design

Cross-sectional, qualitative, bi-national and multicentre study enrolling purposively sampled PC patients in Germany and the United Kingdom, using cognitive interviews.

Setting and participants

In Germany, patients were recruited from a university hospital PC unit and its associated hospital support and community PC teams. UK patients were recruited from a university hospital support team. Inclusion criteria were as follows: age 18 years or above, capacity to give consent and sufficient fluency in written and spoken English or German language to answer the questionnaire. Exclusion criteria were as follows: being too unwell or distressed to participate as judged by the clinical team. Participants were purposively sampled according to the following criteria in order to achieve variation across key characteristics: gender, over and under 65 years, high, medium and low functional status as measured by the Australia-modified Karnofsky Performance Status (AKPS) scale (see Table 1) and malignant/non-malignant main diagnosis. It was not aimed to have equal numbers in all cells, but to include participants with all characteristics. Screening took place at multidisciplinary team meetings or handovers, and a clinical team member asked whether the patients were interested and provided an information leaflet. Interested patients were approached by the researcher – except five who had deteriorated or been discharged. Of the 29 approached patients, 25 gave their informed consent.

Cognitive interviews

Formal translation of the IPOS from English to German had previously been conducted using standard techniques. The interviews were conducted in the participants’ current care setting, that is, on the ward or in the participants’ home. Participants were asked to complete the questionnaire, using a combination of ‘think aloud’ and concurrent verbal probing techniques to elicit problems or comments regarding the completion. ‘Think aloud’ techniques (uttering the thoughts during question answering) were demonstrated by the interviewer at the start of the interview. Probing techniques were both spontaneous, to explore verbal or non-verbal utterances like hesitations or irritation, and standardised according to an interview guide (see Appendix 1). This was based on the four-stage question response model as described by Tourangeau (comprehension, retrieval, judgement and response formulation). Probes were developed for each of the four stages, and additional questions related to layout, missing aspects in the questionnaire and burden associated with its completion. The interviews were audio-recorded and transcribed verbatim.

Analysis of the interviews and refinement of the IPOS

Interviews were analysed by two independent researchers in each country using thematic analysis as described by Willis. In accordance with the interview guide, the transcript was also categorised following Tourangeau’s question response model (comprehension, retrieval, judgement and response formulation). Additional categories included the following: layout, acceptability/questionnaire burden and additional aspects. Results of all participants were compared and aggregated for each question and each category in both countries to identify the reasons for difficulties in answering the questionnaire and other emergent themes.

The German results were translated. Results from both countries were then collated and discussed between the research teams and the IPOS was refined accordingly, based on consensus. The German version was adapted according to the refined English version, taking German-language-specific results into account.

The study was approved by the relevant ethics committees (Ethikkommission of Ludwig-Maximilians-Universität Munich, reference number 169-13; National Research Ethics Service (London–Dulwich Committee) reference number: 13/LO/0573).

Results

Demographics

A total of 15 German and 10 UK interviews (16–54 min long) were completed between July 2013 and July 2014. The median age was 65 years in Germany (range 22–85 years) and 61 years in the United Kingdom (range 43–83 years). In Germany, 10/15 and in the United Kingdom, 7/10 participants were female. The predominant primary diagnosis was cancer. A third of the participants in Germany and two-thirds in the United Kingdom had an AKPS score of 50% or lower, which is defined by requiring ‘considerable assistance and frequent medical care’.

The majority of participants were in-patients. Details are shown in Table 1.

Findings from the interviews and IPOS refinement

The interviews in both countries demonstrated that for the majority of participants, most questions and answer options worked well. The identified difficulties were mainly comprehension problems and some pertained to judgement. No problems were identified with retrieval or response formulation. The interview results and the changes made based on them are shown in Table 2. The revised questionnaire can be found on the POS website.
 Patients were certain about their main problems and concerns (question 1), and many described these as the things that are always on your mind. Positive comments about the symptom list in question 2 included the following:

 Oh, easy to understand. (United Kingdom, female aged 43 years)

and

I think question 2 is good, that it is all dealt with in a very ‘compact’ way. (Germany, female aged 25 years)

Some comprehension problems were identified in both countries. First, difficulties arose with the wording of specific questions. For example, the term ‘mouth problems’ in question 2 was regarded as too vague, and in contrast to the intent of the question, many patients did not think of dry mouth when answering it. Furthermore, comprehension problems concerned answer options which were regarded

Table 1. Demographic and clinical details of participants.

<table>
<thead>
<tr>
<th></th>
<th>Germany</th>
<th>United Kingdom</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total number interviewed</td>
<td>15</td>
<td>10</td>
</tr>
<tr>
<td>Age (years) Median, range</td>
<td>65 (22–85)</td>
<td>61 (43–83)</td>
</tr>
<tr>
<td>≤65</td>
<td>9</td>
<td>7</td>
</tr>
<tr>
<td>&gt;65</td>
<td>6</td>
<td>3</td>
</tr>
<tr>
<td>Sex</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>10</td>
<td>7</td>
</tr>
<tr>
<td>Male</td>
<td>5</td>
<td>3</td>
</tr>
<tr>
<td>Nationality</td>
<td></td>
<td></td>
</tr>
<tr>
<td>German</td>
<td>14</td>
<td></td>
</tr>
<tr>
<td>British</td>
<td></td>
<td>7</td>
</tr>
<tr>
<td>Other</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Not recorded</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>British</td>
<td>7</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Not recorded</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>Ethnicity</td>
<td></td>
<td></td>
</tr>
<tr>
<td>White British</td>
<td>Not recorded</td>
<td>7</td>
</tr>
<tr>
<td>Black African</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Black Caribbean</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Marital status</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Married</td>
<td>7</td>
<td>6</td>
</tr>
<tr>
<td>Partner</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Separated or divorced</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>Widowed</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Single</td>
<td>5</td>
<td>0</td>
</tr>
<tr>
<td>Not specified</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Living situation</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Alone</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td>With others</td>
<td>11</td>
<td>6</td>
</tr>
<tr>
<td>Working situation</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Retired</td>
<td>Not recorded</td>
<td>4</td>
</tr>
<tr>
<td>Working</td>
<td>5</td>
<td></td>
</tr>
<tr>
<td>Unemployed</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Primary diagnosis</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Malignant</td>
<td>13</td>
<td>9</td>
</tr>
<tr>
<td>Non-malignant</td>
<td>2 (cardiomyopathy and motor neuron disease)</td>
<td>1 (chronic obstructive pulmonary disease)</td>
</tr>
<tr>
<td>AKPS score Median, range</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&gt;70%</td>
<td>4</td>
<td>0</td>
</tr>
<tr>
<td>&gt;50%–&lt;70%</td>
<td>6</td>
<td>3</td>
</tr>
<tr>
<td>≤50%</td>
<td>5</td>
<td>7</td>
</tr>
<tr>
<td>Care setting</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Palliative care unit</td>
<td>10</td>
<td></td>
</tr>
<tr>
<td>Other hospital ward with support from hospital palliative care team</td>
<td>2</td>
<td>10</td>
</tr>
</tbody>
</table>

The Australia-modified Karnofsky Performance Status Scale (AKPS) grades functional status, for example, 70% represents ‘cares for self, unable to carry on normal activity or to do active work’ and 50% represents ‘requires considerable assistance and frequent medical care’.12
### Table 2. Issues regarding IPOS completion identified in the cognitive interviews in Germany and the United Kingdom and resultant changes made to the IPOS.

<table>
<thead>
<tr>
<th>Prototype IPOS items</th>
<th>Prototype IPOS answer options</th>
<th>Aspects identified with supporting quotations</th>
<th>Revised IPOS items</th>
<th>Revised IPOS answer options</th>
</tr>
</thead>
<tbody>
<tr>
<td>Q1. What have been your main problems or concerns at the moment?</td>
<td>Open question with three lines for answers</td>
<td>Good comprehension (18/25) of this question. Participants responded with personal problems as well as health-related problems. 2/15 German participants explicitly welcomed that the question asks for main problems AND concerns. Oh yes, that is good. Main problems is of course also associated with concerns. (Germany, male aged 55 years) 3/10 of UK and 4/15 of German patients queried the openness of the question or were at first unsure about the layout with three free lines but consequently answered it.</td>
<td>Q1. What have been your main problems or concerns over the past 3 days?</td>
<td>Open question with three lines for answers</td>
</tr>
<tr>
<td>Q2. Below is a list of symptoms which you may or may not have experienced. For each symptom, please tick one box that best describes how it has affected you over the past 3 days.</td>
<td>Not at all</td>
<td>3 days was not considered long enough by some patients (5/25). Why has it got to be the last three days? You know, like, in the last week, I’ve been really sick for five days, the last two days I’ve been alright. (UK, male aged 53 years) Judging severity of symptom over the time period of 3 days was sometimes difficult when people have had the symptom for one of the days or had a fluctuation of symptom severity (6/25). The subheadings in the answer options were confusing and/or too long (5/25). Poor appetite, well I mean ‘marked effect on activities or concentration’, how does it affect? But activities you just don’t wanna eat that’s all. So I can’t find the right one to tick. (UK, female aged 83 years) In contrast 2/25 of patients found that the explanations helped.</td>
<td>Not at all</td>
<td>Slightly</td>
</tr>
<tr>
<td>Pain</td>
<td>Slightly</td>
<td>Little or no effect on activities or concentration</td>
<td>Difficulty in um not breathing as you normally do. (UK, female aged 47 years)</td>
<td>Pain</td>
</tr>
<tr>
<td></td>
<td>Moderately</td>
<td>Some effect on activities or concentration</td>
<td>Pain</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Severely</td>
<td>Marked effect on activities or concentration</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Overwhelmingly</td>
<td>Unable to think of anything else</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Weakness or lack of energy</td>
<td></td>
<td></td>
<td>Weakness or lack of energy</td>
<td></td>
</tr>
<tr>
<td>Nausea (feeling like you’re going to be sick)</td>
<td></td>
<td></td>
<td>Nausea (feeling like you are going to be sick)</td>
<td></td>
</tr>
<tr>
<td>Vomiting (being sick)</td>
<td></td>
<td></td>
<td>Vomiting (being sick)</td>
<td></td>
</tr>
<tr>
<td>Poor appetite</td>
<td></td>
<td></td>
<td>Poor appetite</td>
<td></td>
</tr>
<tr>
<td>Constipation</td>
<td></td>
<td></td>
<td>Constipation</td>
<td></td>
</tr>
</tbody>
</table>

(Continued)
Prototyp IPOS items | Prototype IPOS answer options | Aspects identified with supporting quotations | Revised IPOS items | Revised IPOS answer options
--- | --- | --- | --- | ---
**Mouth problems** | Too vague and unclear (5/25) | *Mouth problems? It's funny they mention mouth problems, why would they ask about mouth problems? (UK, female aged 83 years)* | *Sore or dry mouth* |  
8/15 of patients in Germany did not necessarily think of dry mouth but of sore mouth/chewing/swallowing problems/tooth ache | *The tongue is sore, there will be thrush or so, and with the teeth they are not all own teeth – and so on there are problems with eating. (Germany, male aged 69 years)*

9/10 of UK patients had good comprehension of the word *drowsiness*. *Um … when just … when your eyes are like going fogy and you just want to sleep … Oh it was easy to understand. (UK, female aged 43 years)* | *Drowsiness* |  
2/15 of German patients were uncertain if it meant tiredness by day or by night | *Problems going to sleep or problems sleeping through the night – or the drowsiness only relates to the daytime. (Germany, female aged 77 years)*

15/15 of German participants had good comprehension | *Poor mobility* |  
2/15 of German participants commented that it was helpful having this question included and 4/10 UK patients wrote other symptoms | *Interpretation of mobility and weakness/fatigue as similar to UK patients (3/10)*

2/15 of German participants felt family/friends was clear; however, 2/10 UK patients were uncertain about who should be included in 'family or friends' | *Please list any other symptoms not mentioned above, and tick one box to show how they have affected you over the past 3 days.*

**Q3. Over the past 3 days, have you been feeling anxious or worried about your illness or treatment?** | No, not at all | Good comprehension by all participants | *Over the past 3 days* |  
Occasionally | Q3. Have you been feeling anxious or worried about your illness or treatment?

Sometimes | Always

Q4. Over the past 3 days, have any of your family or friends been anxious or worried about you? | Yes, always | *Variation in level of insight into family and friends worries and/or anxieties (3/25)* | Q4. Have any of your family or friends been anxious or worried about you?

Q5. Over the past 3 days, have you been feeling depressed? | 8/10 of UK patients had a very good understanding of the word depressed | To be depressed … um … to be in darkness and see no light at all … And your whole being is black. (UK, female aged 70 years) | Q5. Have you been feeling depressed? in German version alternative term for 'niedergeschlagen': 'traurig bedrückt'

15/15 of German participants felt family/friends was clear; however, 2/10 UK patients were uncertain about who should be included in 'family or friends' | *Alternative term for 'niedergeschlagen': 'traurig bedrückt'* |
Prototype IPOS items | Prototype IPOS answer options | Aspects identified with supporting quotations | Revised IPOS items | Revised IPOS answer options
--- | --- | --- | --- | ---
Q6. Over the past 3 days, have you felt good about yourself as a person? | Yes, all the time. | Germany participants interpreted the term used in German mostly as feeling 'physically well' or 'well in surroundings', and if asked did not associate it with a spiritual aspect (9/15). For me feeling good would be standing up, walking … and playing tennis and normal living. … Relatively, I feel good, I do not have pain at the moment, but generally the situation is crap. (Germany, male aged 75 years) | Q6. Have you felt at peace? | Always
Most of the time
Sometimes
Occasionally
No, not at all
3/10 of UK participants found the question strange to be asked and confusing How can you feel good about yourself when you're lying in hospital … everything's crap. (UK, male aged 75 years) | Yes … I've been able to share how I feel … Most of the time. (UK, female aged 65 years) | Q7. Have you been able to share how you are feeling with your family or friends? | Yes, as much as I wanted | Good comprehension by 23/25 participants
Most of the time
Sometimes
Occasionally
Not at all with anyone
1/10 of UK participants found it difficult to answer for both themselves and their family and friends Well I don't know if it was enough for my family I can't answer for them. (UK, male aged 54 years)
Q8. Over the past 3 days, how much information have you and your friends been given? | Full information/as much as I wanted | Questions and answer options considered by 6/25 patients to be too long or complicated Informed 'by whom' and 'about what?' is missing (4/25). Question missed whether information was wanted or not (UK, 1/10) and whether information was understood at the right 'level' (UK, 1/10) 1/10 of UK participants found it difficult to answer for both themselves and their family and friends Well I don't know if it was enough for my family I can't answer for them. (UK, male aged 54 years) | Q8. Have you had as much information as you wanted? | Problems addressed/no problems
Problems mostly addressed
Problems partly addressed
Problems hardly addressed
Problems not addressed
Q9. Over the past 3 days, have any practical matters resulting from your illness, either financial or personal been addressed? | No problems/problems being addressed | Question and answer options confusing/difficult to comprehend (7/25) They're very similar answers aren't they … or is it my brain not working. (UK, female aged 43 years) ‘Practical matters’ not ideal (Germany, 6/15) ‘Problem’ in answer options inappropriate (funeral’ or ‘legacy’ is not a problem) (Germany, 2/15) | Q9. Have any practical problems resulting from your illness been addressed (such as financial or personal)? | Problems addressed/no problems
Problems mostly addressed
Problems partly addressed
Problems hardly addressed
Problems not addressed
Q10. How did you complete this questionnaire? | On my own | IPOS: Integrated Palliative care Outcome Scale. The figures refer to the number of participants who made these comments out of the total number of participants. The remaining participants did not make specific comments and had no difficulties with completion. Changes to the IPOS questions or answer options are typed in **bold**.
With help from a friend or relative
With help from a member of staff
Q 10. How did you complete this questionnaire? | On my own
With help from a friend or relative
With help from a member of staff
as too long or complicated in question 2 (symptom list), question 8 (information received) and question 9 (practical matters). In addition, a third of patients in each country rated the severity of the symptom instead of the degree of being affected by it in question 2 (symptom list).

For some questions, interviewing in the two countries elucidated additional aspects that were important for the refinement of the IPOS: The question ‘Have you felt good about yourself as a person?’ (Q6) had been difficult to translate into German as there is no equivalent phrase that captures the same concept. The translation used for the interviews was understood by most participants as feeling physically well or feeling well in their surroundings. This complemented the comments of UK patients that this question was strange to be asked. A UK patient noted that question 8 (information needs) missed whether the received information was wanted. There were cases when comments made by the German patients diverged with those from the UK patients: Some UK patients interpreted ‘weakness and lack of energy’ and ‘mobility’ as similar, whereas all German patients had a clear understanding of ‘mobility’. Some German patients were unsure about the meaning of ‘drowsiness’, while UK patients had a good understanding of the word. In addition, there were a few translation-related difficulties commented on by the German participants, for example, comprehension problems of the German word used for ‘depressed’.

Judgement. A judgement problem was identified in both countries due to the 3-day time period over which symptoms are sought: some participants said they were unsure how to answer a question when they had a fluctuation of symptom severity; however, all answered the questions. Five participants commented that they would prefer a longer time frame, for example, 7 days. In addition, three participants said that judging the level of insight and anxiety of family and friends was difficult, but all answered the question.

Acceptability/questionnaire burden, missing aspects and layout. Overall, acceptability of the measure was good. Spontaneous comments included:

Well I think it’s a good questionnaire I really do … it lets everybody like staff that they’ve only got to look at it and they know how you feel. (United Kingdom, female aged 74 years)

One patient became distressed when she talked about the worries of her family, but explicitly wanted to continue answering the questions because she thought they were important. One other patient found answering the questionnaire burdensome because she had problems concentrating. All other patients did not find it distressing or burdensome to answer the questions. In the United Kingdom, 8 of the 10 patients stated that they wanted to keep all questions in IPOS, one patient expanded on the importance and variety of all questions:

… when someone’s not well as I say it changes your whole system your whole … routine … and everything needed to be looked into properly. (United Kingdom, female aged 70 years)

One or two patients each suggested the following aspects to be additionally included in the questionnaire: ease of contacting the relevant team, experience of care, general well-being, insomnia and diarrhoea. We did not incorporate these in the refined IPOS for the following reasons: We recognise the importance of measuring patient experience of care, but intended this questionnaire to remain a PROM. The aim was to assess specific symptoms and concerns. Defining insomnia is difficult, often different types of it are described.16 Besides, an open list is provided for patients for not-listed symptoms, for example, insomnia and diarrhoea. The group agreed that this was sufficient, as we wanted to minimise respondent burden.

Finally, layout problems were identified, especially the difficulty of staying in the right line for questions 1, 2, 7 and 8. The layout changed automatically due to the item refinement, especially shortening/aligning of answer options.

Added value of cognitive interviewing in two languages for IPOS refinement. The difficult translation of ‘felt good about yourself as a person’ (question 6) into German and the fact that the chosen translation was mainly understood by patients as physical well-being led to detailed discussions between both research teams about the question’s meaning and intention. As the intended emphasis was on spiritual well-being, the question was replaced by the respective question from the African POS ‘Have you felt at peace?’ This question is a good measure of spiritual well-being in African PC and is highly relevant across diverse cultures and beliefs.17 Discussion in the team also helped to find concise and clear phrasing in both languages for ‘mouth problems’ (question 2) as well as for questions 8 (information received) and 9 (practical matters). The items ‘weakness or lack of energy’, ‘drowsiness’ and ‘poor mobility’ were left unchanged after extensive discussion. For this discussion, the partly conflicting results from the interviews in both languages and considerations for alternatives in both languages – which were deemed inferior to the current items – were paramount.

Discussion

The cognitive interviews conducted simultaneously in Germany and the United Kingdom showed that for the majority of patients, most questions and answer options of the IPOS worked well. The difficulties that were identified were mainly comprehension problems, which were addressed in questionnaire refinement. A judgement problem was reflected in some patients’ replies regarding the 3-day time period over which symptoms were sought. These patients found it hard to judge the score if
symptoms were fluctuating. This corresponds with the findings of Bergh et al.15 in their cognitive interview study of the Edmonton Symptom Assessment System (ESAS), where patients found judging intensity ‘now’ difficult for fluctuating symptoms. The scoring of fluctuating symptoms seems to be a challenge for patients regardless of whether they are asked to score them ‘now’ or over a period of time. In accordance with the wish of some patients for a longer time frame than 3 days, a 7-day IPOS version is now also available. Generally, this problem of scoring fluctuating symptoms on a questionnaire illustrates that any questionnaire can only be one component of a comprehensive clinical assessment.19

Problems with retrieval or response formulation were not identified in this study. The absence of retrieval problems corresponds with the findings of a former cognitive interviewing study in PC and may be due to the short time period for which symptoms were sought.20 However, the absence of retrieval problems is important to note for this population, which was mostly elderly and with low functional status. Another important finding was that except the two patients mentioned earlier, these PC patients with mostly poor functional status did not find it distressing or burdensome to complete the questionnaire, and that they wanted to retain all questions.

Overall, cognitive interviewing showed good comprehension and acceptability of the questionnaire, identified difficulties in its completion and helped to refine it. Enabling patients to shape IPOS greatly benefited the refinement. Cognitive interviewing of patients therefore proved a valuable tool in questionnaire refinement. Problems with retrieval or response formulation were not identified in this study. The absence of retrieval problems corresponds with the findings of a former cognitive interviewing study in PC and may be due to the short time period for which symptoms were sought.20 However, the absence of retrieval problems is important to note for this population, which was mostly elderly and with low functional status. Another important finding was that except the two patients mentioned earlier, these PC patients with mostly poor functional status did not find it distressing or burdensome to complete the questionnaire, and that they wanted to retain all questions.

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PC in-patients. However, these patients have complex PC needs – as this is by definition the reason for a referral to a PC hospital support team or PC unit. Therefore, it seems adequate to base the development of a tool that aims to assess PC concerns mainly on the views of these patients.

Conclusion

Cognitive interviewing of patients for whom the questionnaire is intended proved a valuable tool to increase its face and content validity. The concurrent approach in two languages – to our knowledge the first such approach in PC – was beneficial. The refined version is currently undergoing psychometric validation in Germany and the United Kingdom. Wider awareness and use of cognitive interviewing in measure refinement in PC is crucial.

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References

Appendix I

**Topic guide**

*Integrated Palliative care Outcome Scale (IPOS) pilot survey (Phase I)*

**Objective:**

- To explore the cognitive processes used by respondents when reading, interpreting and responding to items on the IPOS questionnaire.

**Introduction:**

- Study purpose, confidentiality, able to stop at any time, decline questions.
- I’m going to show you a questionnaire and I want you to read and answer the questions one at a time.
- We will stop and talk about each question before moving onto the next.
- Please try to ‘think out loud’ as you read and answer the questions (*DEMONSTRATE: that is, When the question is ‘What did you have for breakfast?’, instead of answering ‘Porridge’ I would say ‘Well I was running late for work this morning so I picked up a croissant and a coffee on the way to work, which I ate at my desk’)*.
- I will also ask you some more specific things about each question.
- Apologies if the questions get repetitive.
- In this study, we are less interested in your answers to the questions, but how you arrive at the answers – what you think the question means, and the things you were thinking about when you chose your answer.
- You can tell me *any* thoughts or views you might have about the questions.

--------------------------------------------------------------------------------------------------------------------------- START RECORDING -----------------------------------------------------------------------------------------------

**General:**

- What were you thinking about when you answered that question?
- I noticed you hesitated before giving your answer – what were you thinking about then?

**Comprehension: what does the respondent believe the question to be asking?**

- What does the question mean to you, in your own words?
- What does the word XXXXXX mean to you? (if certain words are thought to be problematic)
- How easy or difficult was it to understand this question?
- (If problem) How would you change this question?

**Retrieval: could they recall the information required by the question? Was the time frame suitable?**

- How well could you remember your experience when answering this question?
- Was it easy or difficult to think about the past (week) when answering this question?
- Would there be a different time period that would be easier to understand?

**Judgement: is the respondent able to make an evaluation based on the information recalled?**

- What were you thinking about when you answered this question?
- How did you arrive at your answer to that question?
- Was that easy or hard to arrive at your answer? Why do you say that?
- How sure are you of the answer to this question?

**Response: is the respondent able to map their internally generated answer to a response option?**

- How did you choose your answer to this question?
- Was it hard or easy to select an answer from the options given?
- Did all options make sense for this question?

• Other:

  - Is there anything else you would like to say about this question/The questionnaire as a whole?
  - Did you find any of the questions upsetting/embarrassing/inappropriate?
  - Are there any topics/questions that you would leave out of this questionnaire?
  - Are there any topics/questions that you would add to this questionnaire?
  - Do you have any thoughts about the way your answers were captured? (i.e. tablet/paper)

------------------------------------------------------------------- THANKS + STOP RECORDING ---------------------------------------------------

The German topic guide additionally included some questions regarding the translation options of specific words.