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Cultural adaptation of the German Integrated Palliative care Outcome Scale for neurological symptoms

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Keywords:	Patient-related measure, neurological patients needing palliative care, cultural adaptation, cognitive interviewing, German
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	(n=11) confirmed that the IPOS Neuro-S8 is an intelligible tool that is well accepted (phase 5), some linguistic and cultural differences were found between the original English and German versions. These mainly concerned the items mouth problems and spasms. Significance of Results: The German version of the IPOS Neuro-S8 has demonstrated face and content validity and captures relevant symptoms of neurological patients needing palliative care. Its psychometric properties, including construct and criterion validity, will be investigated next.

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Cultural adaptation of the IPOS Neuro-S8

Original Article

Cultural adaptation of the Integrated Palliative care Outcome Scale for neurological symptoms

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Cultural adaptation of the IPOS Neuro-S8

Abstract

Objectives: Standardised measures for assessing neurological patients needing palliative care remain scarce. The Integrated Palliative care Outcome Scale for neurological patients in its short form (IPOS Neuro-S8) helps assess and identify patient's symptom burden and needs early but has not yet been validated in German. The aim was to culturally adapt and translate the IPOS Neuro-S8 to the German healthcare context and evaluate its face and content validity.

Methods: Cultural adaptation study following the first six out of eight phases of the Palliative care Outcome Scale measures manual: 1) conceptual definition, 2) forward translation to German, 3) backward translation to English, 4) expert review, 5) cognitive debriefing, and 6) proof reading. Complex neurological patients in need of palliative care of the Department of Palliative Medicine or Department of Neurology of the University Hospital of Cologne, clinical staff with extensive experience in either palliative care or neurology, actively working in either department were included. Data were analysed using thematic content analysis and descriptive statistics.

Results: A total of 13 patients and 16 clinical staff members participated in this six-phase study. The expert review panel (phase 4) consisted of eleven additional members. While patients (n=9) and clinical staff (n=11) confirmed that the IPOS Neuro-S8 is an intelligible tool that is well accepted (phase 5), some linguistic and cultural differences were found between the original English and German versions. These mainly concerned the items mouth problems and spasms.

Significance of Results: The German version of the IPOS Neuro-S8 has demonstrated face and content validity and captures relevant symptoms of neurological patients needing palliative care. Its psychometric properties, including construct and criterion validity, will be investigated next.

Keywords: Patient-related measure, neurological patients needing palliative care, cultural adaptation, cognitive interviewing, German

Cultural adaptation of the IPOS Neuro-S8

Introduction

Patients with severe neurological diseases typically face problems with mobility, neuropsychological and cognitive disability, communication problems and / or increased care needs (Allen et al., 2020). These neurological conditions are largely incurable, reduce life expectancy and may thus require palliative care (Boersma et al., 2014). End of life for these patients is usually approaching with the onset of swallowing problems, frequent infections like aspiration pneumonia, significant functional and cognitive decline reflected in high caregiver burden and weight loss suggesting that a palliative care approach should be initiated (Ebke et al., 2018; Oliver et al., 2016). Palliative care aims to improve the quality of life of patients and their family members providing a holistic approach. Physical symptoms as well as psychological, social and spiritual concerns are addressed with the help of an interdisciplinary and multiprofessional team specifically trained in palliative and hospice care (World Health Organization, 2020). Integration of palliative care for patients with severe neurological diseases improves prognosis estimation, symptom management, patients' quality of life and family satisfaction (Oliver et al., 2016). Although the number of patients with severe neurological diseases cared for in German palliative and hospice care structures has increased from 0.8% in 2005 to 4.8% in 2017, these patients are still underrepresented in palliative and hospice care structures considering their prevalence, morbidity and mortality (Dillen et al., 2019). Unsurprisingly, the most common place of death of neurological patients is the hospital and not an in- or outpatient palliative or hospice care setting (Boersma et al., 2014; Dasch & Lenz, 2021). However, little is known on how to integrate palliative care for neurological patients best as they present unique challenges that a palliative care approach developed primarily for oncological patients cannot sufficiently meet (Boersma et al., 2014; Turner-Stokes et al., 2007). Yet, such information on a combined approach is crucial to improve care (Saleem et al., 2007) and can be ensured by a proper and valid outcome tool.

Cultural adaptation of the IPOS Neuro-S8

Patient-reported outcome tools (PROMs) are used to capture patients' perception of their health and psychological, social and spiritual concerns by means of standardized, validated questionnaires. This can help clinical staff to focus on the patients' main concern (Bausewein et al., 2016; Dawson et al., 2010). The Palliative care Outcome Scale (POS) is one of the few palliative outcome measures that captures the main concerns of patients with an incurable life-limiting disease (Hearn & Higginson, 1999). Although the POS is widely used in research studies and in the clinical routine, a more refined version was needed that would incorporate more symptoms and refine spiritual or existential issues (Higginson et al., 2012; Murtagh et al., 2019), the Integrated Palliative care Outcome Scale (IPOS). Four versions were created, i.e., patient self-report and staff proxy report both with a timeframe of three and seven days, respectively. These have been cognitively tested and validated in various languages, including English and German (Murtagh et al., 2019; Schildmann et al., 2016). The IPOS has proven very valuable in research studies to assess and measure patients' symptoms and palliative care needs (Evans et al., 2021; Golla et al., 2020; Golla et al., 2022; Schloesser et al., 2022; Schunk et al., 2020). In the clinical care of palliative care patients, outcome measures are becoming increasingly important as well (Bausewein et al., 2016; Bausewein et al., 2018). For example, the German Comprehensive Cancer Centers (CCCs) are currently planning to implement the IPOS as palliative outcome measure which is in agreement with the IPOS being one of the few recommended palliative outcome measures in the German guideline for palliative care to screen for palliative care needs (Leitlinienprogramm Onkologie, 2020). The IPOS is mainly used as self-report by the patient itself but proxy reports filled out by clinical staff also exist and gain in importance once patients cannot sufficiently assess their symptoms themselves anymore due to the progression of their disease. It is important for palliative outcome measures to capture the full range of concerns of patients with progressive incurable diseases. When applied to specific diseases such as neurological conditions they might not be sensitive enough to detect key symptoms that require palliative care in that specific patient population. For this reason the

Cultural adaptation of the IPOS Neuro-S8

POS-MS (Sleeman & Higginson, 2013) and the IPOS Neuro (Gao et al., 2016; Wilson et al., 2019) have been developed. The latter is a reliable and valid psychometric instrument, thus far only available as a self-report version, and allows for identification of problems at an early stage and, if indicated, consultation of palliative care structures. Its full version comprises 45 items covering symptom experience, information needs, practical concerns, anxiety, and feeling at peace, which are scored on a 5-point Likert scale. There are two shorter adaptations containing eight and 24 carefully selected symptom specific items, which have both demonstrated satisfactory to good psychometric properties (Gao et al., 2016; Wilson et al., 2019). As a rather short tool, the IPOS Neuro-S8 (see supplementary file 1) seems to be suitable in the clinical routine irrespective of being applied in an outpatient, inpatient, or semi-inpatient setting such as a private practice, hospitals, nursing homes, hospices, rehabilitation or day clinics, in general or specialized palliative care settings as well as research studies. It is easy and quick to administer which is a prerequisite for patients with a severe or terminal disease as it will minimize their time constraints. It is also a validated measure for use in English-speaking populations but translation and validation in non-English speaking populations has not yet been done. Prior to its use in a specific country, measures must be translated, culturally adapted, and validated to create a reliable and relevant measure reflecting care concepts that are applicable in the target culture with its particular population (Bausewein et al., 2016).

This study aimed to translate and cross-culturally adapt the IPOS Neuro-S8 into the German healthcare context and evaluate its face and content validity. Both are comparable forms of validity assessing whether a test covers all relevant parts of the construct with face validity being more subjective. This was done as groundwork before assessing its validity and reliability so that it can be used in Germany as a brief and feasible instrument in the clinical routine in various outpatient, inpatient, and semi-inpatient settings and research studies. In the

Cultural adaptation of the IPOS Neuro-S8

current study, we used cognitive interviewing as the method of choice to ensure that the instructions and items were accurately expressed and to indicate face and content validity.

Methods

Design

Cultural adaptation study following the first six phases of “The Palliative care Outcome Scale (POS) Family of Measures Manual for Translation, Cross-Cultural Adaptation and Psychometric Testing” (Antunes et al., 2012).

Setting and Participants

The study was conducted at the Departments of Palliative Medicine and Neurology of the University Hospital Cologne. The Ethics Commission of Cologne University’s Faculty of Medicine (#20-1086, 28-May-2020) approved the study, which was registered at the German Clinical Trials Register (#DRKS00021783, 30-June-2020). The study followed the Declaration of Helsinki (World Medical Association, 2014). All study participants provided written informed consent.

Complex neurological patients suffering from a largely incurable disease that reduces their life expectancy and are thus in need of palliative care of at least 18 years of age were recruited from the Departments of Palliative Medicine and Neurology. If a patient was unable to give written informed consent due to physical disabilities, a legal representative who had full command of the German language and could give written informed consent was allowed to act on behalf of the patient. Both had to be native German speakers, patients additionally needed to have basic knowledge of the English language (understanding). Clinical staff with extensive

Cultural adaptation of the IPOS Neuro-S8

experience in either palliative care or neurology were recruited from both departments. All clinical staff needed to be 18 years or older, native German speakers with basic knowledge of the English language (understanding), and able to give informed consent. Participants were screened by a clinical team member (YG, HG, CW) and, if interested, approached by a researcher (KD).

Measure

The IPOS Neuro was developed for people with progressive, long-term neurological conditions and has been adapted into two shortened forms, i.e., IPOS Neuro-S8 (Gao et al., 2016) and IPOS Neuro-S24 (Wilson et al., 2019). Both of these shortened adaptations contain three key questions with instructions of which the second question lists a selection of core symptoms from the full 45-item version. The IPOS Neuro-S8 covers eight physical symptoms, i.e., pain, nausea, vomiting, mouth problems, sleeping difficulties, breathlessness, spasms, and constipation over the past three days. The response categories range from 0 (not at all) to 4 (overwhelmingly). The total score is obtained by summing the item scores, that is, 0-32. The English version of both the IPOS Neuro-S8 and IPOS Neuro-S24 have been validated and evaluated using data from patients severely affected by multiple sclerosis, idiopathic Parkinson's disease, multiple system atrophy, and progressive supranuclear palsy (Gao et al., 2016; Wilson et al., 2019). For the validation of the IPOS Neuro-S24 patients with motor neurone disease were also included (Wilson et al., 2019).

Phase 1. Conceptual Definition

The conceptual definition clarifies the concepts underlying each item which is crucial to ensure that the new measure reflects the care concepts of the target culture (Antunes et al., 2012). This

Cultural adaptation of the IPOS Neuro-S8

was done by following a three-step process: i) literature search on health-related quality of life issues relevant for palliative care and neurological patients, ii) identification, analysis, and definition of critical concepts underscoring each item using semi-structured interviews with clinical staff possessing knowledge in palliative care or neurology, iii) discussion of the concepts defined in step two with palliative care and neurological patients. Both interview guides were developed by the first author (KD) following the POS measures manual (see supplementary files 2, 3) (Antunes et al., 2012).

Phase 2 and 3. Forward and Backward Translation

First, the original IPOS Neuro-S8 was translated into German (forward translation). This forward translation involved two independent translators with complementary backgrounds. One had clinical knowledge and was familiar with palliative care and neurology terminology; the other had no clinical or medical background and therefore used a language spoken by the general population. Both were native German speakers and proficient in English (i.e., they were very skilled in reading, writing, speaking, and listening). Discrepancies were identified by a third person who served as a mediator to reach a consensus. The mediator was knowledgeable with palliative care concepts but had no medical background and was not involved in the preceding forward translation. Next, this version was translated back to English (backward translation), ensuring the German version reflected the item content of the original English version. This check was done by a native English speaker who had no clinical or medical background, no conceptual knowledge of the IPOS Neuro-S8 and was blind to the original English version.

Cultural adaptation of the IPOS Neuro-S8

Phase 4. Expert Review

The translated intermediary versions were then evaluated, revised, and consolidated by a multi-disciplinary panel of experts, including members from the research team and clinical staff with palliative care or neurological background through a one-time online video conference coordinated and chaired by the first author (KD) who was taking minutes. These were then discussed with the principal investigator of this study (HG) before the pre-final version of the measure was created by KD. Using the conceptual elements from the first phase, this was done to achieve conceptual, semantic, experiential, and content equivalence.

Phase 5. Cognitive Debriefing

By conducting cognitive interviews verbal information related to the responses are additionally collected while administering a questionnaire (Beatty & Willis, 2007). This can be ensured by the think-aloud technique during which interviewees are asked to verbalize their thoughts while answering and by using specific probes (Beatty & Willis, 2007; Schildmann et al., 2016). The think-aloud technique in particular can be useful in designing a questionnaire (Beatty & Willis, 2007), we thus opted for cognitive interviewing or debriefing as it relates to the qualitative pre-testing phase of a measure in the target language. Clinical staff and patients were interviewed separately using two different semi-structured interview guides to evaluate the measures' comprehension, acceptability, clarity, relevance, and length (see supplementary files 4, 5). This procedure allowed assessing content and face validity. All interviews started with a question about the interviewees' overall impression and relevance followed by specific questions related to the test instructions and each item, each with probing questions to generate verbal information and record cognitive processes (think-aloud technique) and ended with an open-ended question for additional remarks. Each interview was conducted via online conferencing,

Cultural adaptation of the IPOS Neuro-S8

recorded digitally, timed and performed by the first author (KD), who is experienced in conducting qualitative interviews and did the verbatim transcription.

Phase 6. Proof Reading

Upon completing all five phases, all required documents (i.e., forward translation, backward translation, records, final report, pre-final measure) and completed templates as requested by the POS Development Team were sent as aggregate data to the POS Development Team (MH) for final proof reading.

Data Analysis

Data collected during phase 1 and 5 were analysed by the first author (KD) following the cultural adaptation phases (Antunes et al., 2012). Thematic content analysis was used to categorise and identify central themes following cognitive interviewing (Anderson, 2007). Each interview was read thoroughly. Responses for each element and item were listed on a coding sheet. These were aggregated and compared for each item and finally compared between patients and clinical staff. Statements to the last question about additional remarks concerned similar issues so they were clustered together. Ambiguous text segments were discussed with the last author (HG) until a consensus was found. Sample characteristics were described by medians and ranges or absolute frequency. Time to completion was calculated descriptively by medians and ranges.

Results

Demographical Data

Cultural adaptation of the IPOS Neuro-S8

Five clinical staff members and four patients participated in an interview for phase 1. These were conducted between May and August 2020. Two more patients were approached but declined participation. The expert review panel (phase 4) consisted of four physicians, three nurses, one researcher, and all three translators from phases 2 and 3. Out of 12 patients who agreed to participate in a cognitive interview for phase 5, three had to be excluded due to physical deterioration at the time of the interview. Eleven eligible clinical staff members were identified, all of whom agreed to be interviewed. These interviews (phase 5) were completed between October and November 2020. Demographical characteristics for interviewees of phases 1 and 5 can be found in table 1. A summary of all participants is shown in figure 1.

<< insert table 1 and figure 1 about here >>

Phase 1. Conceptual Definition

This phase aimed to define and discuss key concepts that underscore each item. A summary is shown in table 2. In general, there was a high consensus between the concepts as defined by clinical staff (n=5) and patients (n=4), although clinical staff expressed clear, objective concepts while patients spoke from their own subjective experience. A few minor challenges were highlighted in defining the concepts for all but one item. When asked about the underlying concept of *pain*, both clinical staff and patients differentiated between physical and psychological pain.

On the one hand, I understand by *pain* physical pain that can be expressed neuropathically as a burning, stabbing, agonizing, sudden, spasmodic, or permanent sensation, but I also understand by *pain* a psychological pain component, i.e., it cannot be assigned to anything at all, well, I'm aware of the term "total pain", so simply the feeling of a comprehensive psychological pain that cannot be treated with pain medication alone. (K05)

Cultural adaptation of the IPOS Neuro-S8

They expected the psychological component of *pain* and other items, including *shortness of breath* and *nausea*, in other countries and cultures to be neglected in favour of the physical component. Regarding the items *nausea* and *vomiting*, clinical staff explained that the English parenthesis *to be sick* is not associated with either of the symptoms mentioned earlier in the German language.

... sometimes this is related to language elaboration, for example *vomiting (being sick)*, you wouldn't say "I feel quite sick", but "I feel nauseous", "I have to throw up soon". (K02)

All interviewees had difficulties defining *mouth problems* and listed more examples than a solid definition. The item *spasms* was difficult to define, especially the distinction of cramps vs spasticity. *Difficulty in sleeping* was also a controversial item pending between difficulties, problems and disturbances. Potential differences regarding the importance of sleep in different cultures and countries were mentioned.

<< insert table 2 about here >>

Phase 2. Forward Translation

There were minor linguistic and content differences between both forward translations for the items *vomiting (being sick)*, *mouth problems*, *spasms*, *difficulty in sleeping*. As one translator was a clinician who knew which terms are easiest understood by patients, her suggested translations were used for the two items *vomiting* [Erbrechen] and *difficulty in sleeping* [Schlafstörungen]. These were adopted into the final version of the measure. The parenthesis *being sick* after *vomiting* was deleted as it was found to be idiomatic to the English language. The other two items (*mouth problems*, *spasms*) were discussed with the mediator until a consensus was found. For *mouth problems*, the literal translation [Mundprobleme] was chosen, not leaving room for interpretation. For *spasms*, five German words were initially considered

Cultural adaptation of the IPOS Neuro-S8

by both translators [Spastiken, Spastik, Krämpfe, Spasmen, Verkrampfungen], with one being mutual and generally understandable [Krämpfe]. To differentiate it from another disease-specific symptom, it was further narrowed down [Muskelkrämpfe] in mutual agreement with the mediator and both translators. However, these two items were further discussed during the subsequent expert review (phase 4) and cognitive interviews (phase 5) and eventually rewritten in the final version (see below for details).

Phase 3. Backward Translation

This intermediary version of the preceding forward translation was then used for the backward translation. However, there were a few discrepancies in the backward translation compared to the original measure. The instructions and Likert response options of question 2 were particularly different, as were the following four items: *nausea (feeling like you are going to be sick)*, *constipation*, *spasms*, *difficulty in sleeping*. Two of these items were already considered problematic during the forward translation (spasms, difficulty in sleeping). All inconsistencies were discussed with the mediator, and a protocol was kept for further debate within the expert review.

Phase 4. Expert Review

Both, the intermediary forward and backward translations were then discussed within the online expert review (n=11). The instructions and Likert response options of question 2 and two items already considered challenging during the translation phases were particularly discussed. The distinction between symptom severity and impact as expressed in question 2 was clarified. During the backward translation, this phrase was translated as *how severe the symptoms were*, which did not match the original sub-sentence. Consequently, both the forward and backward

Cultural adaptation of the IPOS Neuro-S8

translations were revised accordingly [wie sehr Sie sich dadurch ... beeinträchtigt gefühlt haben – that best describes how the symptoms have affected you...]. Similarly, the backward translation for the Likert response option *slightly* was translated differently [leicht - light], i.e., it did not describe how a symptom can affect a person, so both the forward and backward translations were updated [ein wenig – a little]. The translations were inconclusive for two items already discussed during the translation phases. First, the item *constipation* was translated as *congestion* [Verstopfung – congestion]. However, expert review members associated this with the sinuses rather than the gastrointestinal tract. Therefore, the forward and backward translation were modified for clarification [Verstopfung (Darmträgheit) – blockage]. The other item *spasms* was discussed at length, specifically the difference between spasticity and muscle cramps. Eventually, both intermediary translations of the preceding phases 3 and 4 were rewritten to reflect the concept of spasticity [Spastik] and not muscle cramps [Muskelkrämpfe] as agreed upon during the forward translation. The remaining two items that differed from the original measure *difficulty in sleeping* and the parenthesis *feeling like you are going to be sick* were not altered as experts felt they were not to be misunderstood or misinterpreted by patients in the German healthcare context.

Phase 5. Cognitive Debriefing

The pre-final version created after the expert review by the first author (KD) was then used in the qualitative pre-testing phase using cognitive debriefing or interviewing. Both patients (n=9) and clinical staff (n=11) found the measure comprehensible and well-structured, especially the Likert response options were commended. Herein, only one response option was rewritten as suggested by two clinical staff members. The phrasing of the first two questions itself were criticized and modified accordingly in the final version.

Cultural adaptation of the IPOS Neuro-S8

Q1: I would replace the word problems with (medical) complaints [Beschwerden], problems are rather broad by definition, as in “my account was frozen”, these are problems but a (medical) complaint I would associate with the body and health. (K15)

Q2: This "Or not at all", well I believe you could delete this, because either you have experienced it then you tick it off or you just tick "not at all", then it is "or not at all"... that's just the same thing twice, but I think that is a result of the English translation, because it is a standard phrase in the English language. (K3)

The length and time to completion was deemed acceptable (median time patients: 3:11 min, range 02:09 - 09:51, median time clinical staff: 1:22 min, range 1:02 – 2:05), only one patient felt that the time for completion was too long. The length of the measure was also commended for its brevity.

Well, I think the physical symptoms are well covered... I would not make it longer, in no way longer, it should remain concise and clear. (K13)

Three patients considered the symptoms irrelevant for themselves. The items *mouth problems* and *spasms* posed some minor problems for both patients and clinical staff. Both groups felt that *mouth problems* were too general and nonspecific and proposed to add some specific examples, e.g., dry mouth, which was considered highly relevant for the involved patient population. As a result, two of the suggested examples “dry mouth and sores” were incorporated in the final version. Comprehension difficulties were identified for *spasms* and solved by adding *muscle cramps* in parenthesis.

I stumbled across two things that might not be clear to some patients, *mouth problems* I wasn't quite sure what was meant by that, I can imagine as a [occupation] what is meant by it, but I don't know if a patient would think of it right away, I could imagine that *dry mouth* and things like that are meant by that or maybe *swallowing problems*.... and with *spasms* I'm also not sure

Cultural adaptation of the IPOS Neuro-S8

if every patient knows what is meant by it, if s/he never had it... if you could actually just add muscle cramps, then it is not isolated to *spasms*, but refers to a broader context (K08)

All other items and elements of the measure were left unchanged after extensive discussion. Conceptual elements underlying each item were adjusted accordingly. For a detailed overview, see table 3.

<< insert table 3 about here >>

Phase 6. Proof Reading

The final version (see supplementary file 2) and all aggregated documentation were reviewed and approved by the POS Development Team, facilitated by MH. The external reviewer of the POS Development Team suggested minor editorial changes to the instructions of the first two questions and the instructive prompt, which we have incorporated into the final version.

Discussion

In this study, we culturally adapted and translated the IPOS Neuro-S8 for the first time and demonstrated acceptability and content validity through cognitive interviews with patients and clinical staff within the Department of Palliative Medicine and the Department of Neurology of the University Hospital Cologne. However, the translation for some items was too vague and needed cultural adaptation, emphasising the importance of cognitive debriefing. This finding is in good agreement with other translations and cultural adaptations of the parent measure (Antunes & Ferreira, 2020; Sterie & Bernard, 2019; Veronese et al., 2019).

We followed the POS measures manual (Antunes et al., 2012) to create a version of the IPOS Neuro-S8 conceptually equivalent to the original measure. Although there was a

Cultural adaptation of the IPOS Neuro-S8

consensus for all eight items, the concepts of four items were slightly adjusted after cognitive interviewing, which attests to the significance of cognitive interviews as part of a cultural adaptation study of an outcome measure. Interestingly, for the three items *pain*, *shortness of breath* and *nausea*, both patients and clinical staff mentioned psychological aspects during the conceptual definition. This might be considered controversial in other cultures, especially when it might be unusual to acknowledge psychological influences on physical sensations. In Germany, however, patients appreciated the psycho-somatic aspects of illnesses and complaints and expected clinical staff and researchers to think within this dimension, too.

Overall, our results confirm the measure's acceptability and comprehension, despite some minor problems with comprehension of two items questioning its conceptual equivalence, i.e., whether the concepts of given items in both cultures actually exist and are equal (Antunes et al., 2012). Herein, agreeing on an appropriate term for *spasms* was a significant hurdle already during the translation process. Different wordings and parentheses were discussed while reviewing its concept. Ultimately, conceptual equivalence was ensured by adding *muscle cramps*. The second questionable item was *mouth problems*. It was considered too vague, and although some patients reported thinking of *dry mouth*, which is the intention of the item (Veronese et al., 2019), not every patient did. This misperception has already been described in the translation and cultural adaptation study of the parent IPOS (Schildmann et al., 2016). However, as the original item *sore or dry mouth* was intentionally changed to *mouth problems* in the neurological version (Veronese et al., 2019), we only added *dry mouth* as an explanatory addition for clarification as proposed by many interviewees. While two clinical staff members additionally proposed a change of wording, we considered the addition of two specific examples sufficient as explanation and to confirm conceptual equivalence.

Content and face validity were also revealed through cognitive debriefing. Overall, interviewees found the measure valuable and intelligible, confirming its clinical applicability.

Cultural adaptation of the IPOS Neuro-S8

Nevertheless, two patients and one clinical staff member doubted the relevance and therewith content equivalence of three items (*shortness of breath, nausea, vomiting*) whilst all other symptoms were considered essential. Also, nine clinical staff members and one patient suggested the addition of the following symptoms: psychological symptoms such as depression or anxiety, weakness, loss or change of appetite, hallucinations, difficulty speaking, tingling, restlessness, and tiredness. However, although we recognise the relevance of these symptoms, we did not incorporate them in the refined IPOS Neuro-S8 as the included eight items are considered core items of the IPOS Neuro (Gao et al., 2016) and the intent of the current study was not to add or remove given items but to remain as close as possible to the original measure.

One clinical staff member criticised the period of three days (asked in questions 1 and 2) as neurological patients are often diagnosed with a progressive, long-term disease, and instead suggested asking for *changes*. Similarly, the period of the past three days was considered too short by one patient who would have preferred a more prolonged time reference. This is in line with the findings of another translation and cultural adaptation study (Schildmann et al., 2016). While the IPOS Neuro was developed as self-report specifically for people with long-term neurological conditions (Gao et al., 2016; Wilson et al., 2019), the three-day recall version does not incorporate the fluctuating symptoms this patient population might be affected by. As there are versions of the IPOS with a seven-day recall period, our results suggest developing a seven-day IPOS Neuro version, which is more reasonable for neurological patients. Shall such a version with an extended recall period be developed, it seems appropriate to use both versions in various clinical settings providing palliative care including hospitals, private practices, rehabilitation or day clinics, hospices, nursing homes and at home by specialized or general palliative home care **depending on the patients' condition, i.e., for neurological patients with a progressive, long-term disease**. It may also be used in non-palliative care settings, such as neurological units, to trigger the referral process (Gao et al., 2016). It

Cultural adaptation of the IPOS Neuro-S8

might aid in the early initiation of palliative care for patients with severe neurological diseases approaching end of life.

Additionally, there were some discrepancies with the phrasing of question 2. Herein, the severity of a symptom was rated instead of the degree of being affected by it (Schildmann et al., 2016). This is one of the major differences between the HOPE+, already available in the German language, and the IPOS Neuro-S8, with the former evaluating the incidence and intensity of symptom burden while the IPOS Neuro-S8 assesses explicitly the *impact* of symptoms on a patients' everyday life within the past three days (Dillen et al., 2019). Our own experience suggests that patients tend to rate the presence or incidence and intensity of a symptom rather than the impact it has on them. It might be easier for patients to indicate whether they have a specific symptom than how much they are affected by it as this requires a higher-level cognitive function as well as the capacity of emotional reflection skills. It is, however, clinically relevant to differentiate between the impact of a symptom and the incidence and intensity as treatment plans might be different; however, both views are important for a comprehensive understanding.

In general, both patients and clinical staff found the wording clear and understandable and felt that the Likert response options, length, and time to completion were appropriate. Clinical staff also commended the measure's conciseness. This is crucial for a measure developed specifically for terminally ill patients. They even seemed to appreciate the opportunity to talk about certain aspects related to the symptoms (Beck et al., 2017). Therefore, our results suggest that the instructive prompt at the end of the measure is of essential importance. Instead of being left alone after filling out an outcome measure which might have triggered something in a patient, this prompt offers a follow-up consultation. Thus, while a measure must be easy and quick to administer, their wish to talk about it should also be considered.

Cultural adaptation of the IPOS Neuro-S8

Strengths and Limitations

In our study, we used both the “think aloud” and probing technique to optimise unintended and specific, detailed information flow (Willis, 2005). To enhance credibility, the interview guide was meticulously discussed with the last author (HG). We also ensured that the first author (KD) had the required knowledge and training to perform the study. A major strength of our study relates to the heterogeneity of our sample. We were able to cognitively interview a broad range of persons with various progressive neurological conditions at different stages of their disease, so we even included severely affected patients.

There are also some limitations that need to be discussed. The first caveat relates to the recruitment setting. All except one patient were recruited from the Department of Neurology. However, we carefully selected severely affected patients who were considered palliative care patients. Another limitation is the small sample size, which limits the generalizability. However, small sample sizes of 5-15 interviewees have been recommended for cognitive interviews (Beatty & Willis, 2007), also by the POS team itself (Antunes et al., 2012), and our sample size is comparable to other translation and cultural adaptation studies (Beck et al., 2017; Gerlach et al., 2020; Schildmann et al., 2016; Sterie & Bernard, 2019).

Conclusion

The German IPOS Neuro-S8, a patient-reported measure used to assess and treat patient-related problems in clinical practice, is well accepted by severely affected neurological patients and clinical staff and demonstrated face and content validity. The cross-cultural adaptation and translation process resulted in changes for the items *vomiting*, *constipation*, *spasms* and *mouth problems*. As a translated measure must stay as close as possible to the original measure, other items remained unchanged, although there were some inconsistencies. This is the first measure

Cultural adaptation of the IPOS Neuro-S8

for neurological patients in need of palliative care that assesses the *impact* of symptoms on patients' everyday life and can be used longitudinally to treat problems and direct conversations in routine clinical practice, which is essential for patients with severe neurological disease with fluctuating symptoms. It is also appropriate for an international audience, so our results suggest cultural adaptations to other non-English speaking populations and might already have raised awareness for the importance of such a tool. The tool is now available for download in German on the POS website (<https://pos-pal.org/>) for routine clinical assessments, clinical trials, and education to capture patient-centred needs of neurological patients. Next, we will investigate its psychometric properties, including construct and criterion validity.

Conflicts of Interest: None

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Cultural adaptation of the IPOS Neuro-S8

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For Peer Review

Cultural adaptation of the IPOS Neuro-S8

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Cultural adaptation of the IPOS Neuro-S8

Figures

Figure 1. Summary of study participants

Tables

Table 1. Characteristics of both patients and clinical staff who participated in phases 1 and 5

Table 2. Summary of conceptual definition for each item

Table 3. Issues identified within cognitive debriefing by all interviewees (n=20)

Supplementary Files

Supplementary file S1. IPOS Neuro-S8 (original English version)

Supplementary file S2. Interview guide for clinical staff - phase 1

Supplementary file S3. Interview guide for patients – phase 1

Supplementary file S4. Interview guide for clinical staff – phase 5

Supplementary file S5. Interview guide for patients – phase 5

Supplementary file S6. IPOS Neuro-S8 (translated German version)

Cultural adaptation of the IPOS Neuro-S8

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Cultural adaptation of the IPOS Neuro-S8

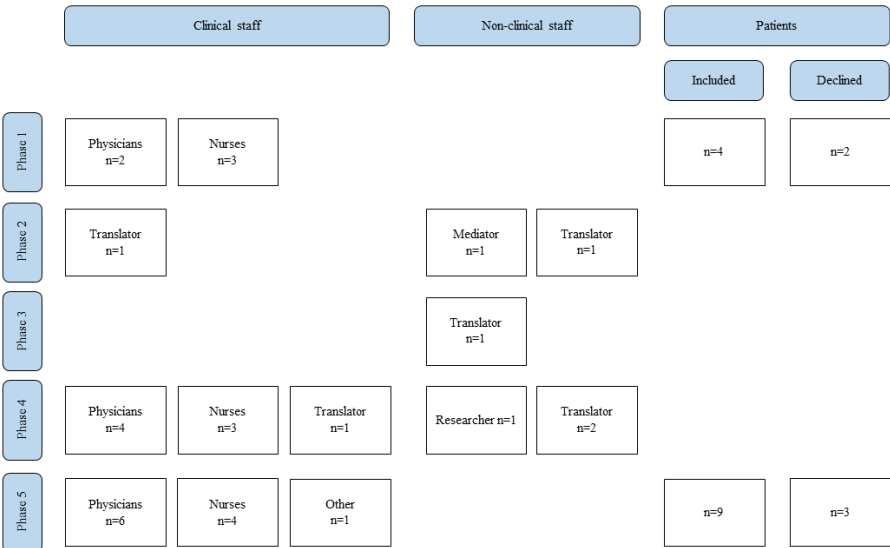
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Summary of study participants

97x54mm (300 x 300 DPI)

Table 1. Characteristics of both patients and clinical staff who participated in phases 1 and 5

	Phase 1		Phase 5	
	Patients	Clinical staff	Patients	Clinical staff
	(n=4)	(n=5)	(n=9)	(n=11)
Age (years)				
Median	60.5	36	58	36
Range	31-74	32-41	31-84	27-64
Gender (n)				
Female	2	5	6	7
Male	2	0	3	4
Patients' primary diagnosis (n)*				
Amyotrophic lateral sclerosis	1		1	
Spinal muscle atrophy	1		1	
Glioblastoma	1		1	
Parkinson's disease	1			
Guillain-Barre syndrome with rapidly progressive paraparesis			1	

Thoracal myelitis		1	
Cervical dystonia with deep brain stimulation		1	
Stroke		1	
Multiple sclerosis		2	
Patients' care setting (n)			
Department of Neurology	3	8	
Department of Palliative Medicine	1	1	
English proficiency of patients (n)			
Sufficient	2		
Good	2		
Very good	0		
Business fluent	0		
Mother tongue	0		
Occupation of staff (n)			
Physician		2	6

Nurse	3	4
Other	0	1
Workplace of staff (n)		
Department of Neurology	3	6
Department of Palliative Medicine	2	5
English proficiency of staff (n)		
Sufficient	0	
Good	2	
Very good	3	
Business fluent	0	
Mother tongue	0	

* Patients were carefully screened by a clinical team member (YG, HG, CW). Included patients either had problems with mobility, communication problems, increased care needs, or significant functional decline.

Table 2. Summary of conceptual definition for each item

Pain	• Strong unpleasant feeling or perception that impedes the quality of life, restricts thinking and acting, and can be physical or psychological • Physical pain can be burning, agonizing, stinging, sudden, spasmodic, permanent, pressing, pulsating, and due to an illness • Psychological pain cannot be assigned to a particular body part, so pain medication cannot provide relief, this kind of pain arises from the soul instead, e.g., grief, shortness of breath
Shortness of breath	• Air or respiratory distress, poorer breathing of any form, does not allow deep ventilation of the lungs • Gasping for breath • Objective: oxygen content in blood decreases, heart beats faster, accompanied by physical symptoms, one has to take breaks while talking or walking • Subjective: pressure on the chest, difficulty to breathe due to weakness of the muscular system, feeling strongly overburdened because something is taking one's breath away, although it cannot be objectified, goes beyond objectifiable measurements, e.g., oxygen saturation • Is often accompanied by panic, fear of suffocation, hyperventilating because of the feeling of not getting enough air

Nausea (feeling like you are going to be sick)	• Feeling of needing to vomit, one may start gagging or have the feeling that stomach content is coming back up even though it is not • More than just physical nausea, also feeling sick, although you cannot name it (psychological aspect), not knowing what your body needs • Associated with reluctance to eat and drink • Affects the whole body, often accompanied by a pale face, belching, swallowing, slower movement, increased salivation, sweating, trembling
Vomiting (being sick)	• Ejection of stomach content through the oesophagus via the mouth (or nose) or, if the stomach is empty, of bile • Great effort for the body: cold sweats, shaky, afterwards possible pain, burning sensation, tiring but also relieving • Often previous nausea, accompanied by retching, which cannot be stopped but can also come out of nowhere, without previous nausea
Constipation	• Intestinal content cannot be excreted rectally • Medical: lack of bowel movement for more than three days, but very individual • Subjective: abdomen full and bloated, may cause cramps and pain • Cause: insufficient fluid intake or side effect of medication

Mouth problems	• Problems with outer and inner mouth area, oral mucosa, oral health, oral cavity, lip, dental and pharyngeal health, oral hygiene • E.g., dry mouth, inflammation, salivation, impaired taste, aphthae, tartar inflammation, lesions, fungi, caries, periodontitis, difficulty swallowing, speech disorder, herpes, swollen tongue, irritated gums
Spasms	• Severe, persistent muscle stiffness, cramping, or twitching, muscular system hardens • Increased muscle tension • Shortening of the muscles or tendons • Stiffness of the extremities, cramped extremities • Uncomfortable, exhausting, painful, restricted mobility
Difficulty sleeping	in • Problems sleeping: e.g., difficulty falling asleep or sleeping through the night, little restful sleep, generally sleeping too little, early awakening, superficial sleep, no deep sleep • Consequence: fatigue, concentration problems, dizziness, circulatory problems • Cause: ruminating thoughts, psychological, anxiety/restlessness, external circumstances such as noise, body- or disease-related (e.g., sleep apnea, pain)

Table 3. Issues identified within cognitive debriefing by all interviewees (n=20)

Question / item (in the original English version)	Interviewees' comprehension and acceptability	Question / item revised
<i>Heading</i>		
IPOS Neuro-S8 Patient Version	No suggested changes.	no
<i>Questions</i>		
1. What have been your main problems or concerns over the past 3 days?	Ten patients and clinical staff found the first question too general and wished for a change of wording that indicated the relation to the disease. One clinical staff commended the phrasing of the first question. The time frame of three days was perceived as too short by one patient.	yes no
2. 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.	Good comprehension but sub-sentence was deemed redundant by three clinical staff members.	yes

3. How did you complete this questionnaire? No suggested changes. One clinical staff member commended the inclusion of this question.

Symptoms in question 2

Pain No suggested changes. no

Shortness of breath Two interviewees found the item irrelevant. no
Five clinical staff members suggested changing the German translation to a more subjective and acute word while another five clinical staff members and eight patients wished to leave the translation as is.

Nausea (feeling like you are going to be sick) One patient found the item irrelevant, three clinical staff members suggested combining nausea and vomiting, and four interviewees felt that the parenthesis could be removed while five clinical staff members and seven patients wished to leave the translation as is. no

Vomiting (being sick) One patient found the item irrelevant, no suggested changes. no

Constipation Six interviewees felt that the parenthesis could be removed while five clinical staff no

members and four patients wished to leave the translation as is. One clinical staff member commended the addition of the parenthesis.

Mouth problems Thirteen interviewees found this item too general and recommended the addition of specific examples in brackets, e.g., dry mouth and sores.

Two clinical staff proposed a change of wording.

Spasms This item proved to be the most challenging. Ten interviewees found this item too specific and difficult to understand and advised for a change of wording or at least the addition of an explanatory definition or examples in brackets.

Difficulty in sleeping Two interviewees recommended the use of a parenthesis while five clinical staff and six patients wished to leave the translation as is.

Response options

Not at all No suggested changes. no

Slightly	No suggested changes.	no
Moderately	No suggested changes.	no
Severely	One clinical staff suggested a change of wording.	no
Overwhelmingly	Two clinical staff suggested a change of wording.	yes
On my own	No suggested changes.	no
With help from a friend or relative	No suggested changes.	no
With help from a member or staff	No suggested changes.	no

Instructive prompt

If you are worried about any of the issues raised on this questionnaire please speak to your doctor or nurse	One clinical staff suggested a change of wording. One clinical staff member commended the inclusion of this prompt.	no
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IPD 2017 2018 2019 2020 2021 2022 2023 2024 2025 2026 2027 2028 2029 2030

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Q1. What have been your main problems or concerns over the past 3 days?

- 1.....
- 2.....
3.

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.

	Not at all	Slightly	Moderately	Severely	Over- whelmingly
Pain	0 ☐	1 ☐	2 ☐	3 ☐	4 ☐
Shortness of breath	0 ☐	1 ☐	2 ☐	3 ☐	4 ☐
Nausea (feeling like you are going to be sick)	0 ☐	1 ☐	2 ☐	3 ☐	4 ☐
Vomiting (being sick)	0 ☐	1 ☐	2 ☐	3 ☐	4 ☐
Constipation	0 ☐	1 ☐	2 ☐	3 ☐	4 ☐
Mouth problems	0 ☐	1 ☐	2 ☐	3 ☐	4 ☐
Spasms	0 ☐	1 ☐	2 ☐	3 ☐	4 ☐
Difficulty in sleeping	0 ☐	1 ☐	2 ☐	3 ☐	4 ☐

	On my own	With help from a friend or relative	With help from a member of staff
Q3. How did you complete this questionnaire?	1 ☐	2 ☐	3 ☐

If you are worried about any of the issues raised on this questionnaire please speak to your doctor or nurse



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Interview guide for the project „cross-cultural adaptation and translation of the IPOS Neuro-S8“ - phase 1

Introductory question

You have been invited to this personal interview because of your outstanding expertise in hospice and palliative care and/or neurology. Today I would like to go through symptom specific items of a palliative and neurological outcome measure and discuss the underlying definitions with you. Since this questionnaire has not yet been used in Germany, we need your help to find out whether the definitions of the individual symptoms in Germany are similar to those used in the English-speaking community or whether they should possibly be adapted culturally.

Transition question

Subjects are given a copy of the original version of the IPOS Neuro-S8

The IPOS Neuro-S8 is already in use in English-speaking countries. As indicated by its name, it comprises eight symptoms, which I will now name of after the other in the original language, thus in English. You also have a copy of the original version in front of you. I kindly ask you to define those symptoms in your own words.

Key questions (5 minutes per symptom, resulting in a total of 40 minutes) The narrative prompt is used separately for each of the eight symptoms		
Guiding question (narrative prompt)	Memo for possible follow-up questions – only to be asked if not addressed by itself	Specific questions – to be asked verbatim
What do you understand by <xy>?	• Identification • Analysis • Definition	• What does the symptom <xy> mean to you and how would you describe or define it in your own words? • What were you thinking about when describing the symptom <xy>?
Final question (5 minutes)		
Is there anything else you would like to add regarding the cultural adaptation of this questionnaire in Germany that we have not yet addressed?		



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Interview guide for the project „cross-cultural adaptation and translation of the IPOS Neuro-S8“ - phase 1

Introductory question

You have been invited to this personal interview because you are currently receiving or have received medical care in the past. Today I would like to go through symptom specific items of a questionnaire with you and discuss the underlying ideas that various healthcare professionals had about them before. We need your help to find out whether you think the healthcare professionals' ideas make sense or whether you think differently about them.

Transition question

Patients are given a copy of the original version of the IPOS Neuro-S8 along with a summary of the items as defined by interviewed healthcare professionals

You now have eight items including a definition in front of you. We will now go through these items and you tell me whether you find the definition for each item convincing and correct. Otherwise, I kindly ask you to describe in your own words what you understand by the respective item.

Guiding question (narrative prompt)	Memo for possible follow-up questions – only to be asked if not addressed by itself	Specific questions – to be asked verbatim
What do you understand by the given definition for item <xy>?	Is the basic concept for item <xy> fully captured?	- How would you rephrase the item <xy>? - What does the item <xy> mean to you? - What were you thinking about when hearing the symptom <xy>?

Final question (5 minutes)

Is there anything else you would like to add regarding the definitions of the items of the questionnaire that we have not yet addressed?



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Interview guide for the project „cross-cultural adaptation and translation of the IPOS Neuro-S8“ - phase 5

Introductory question

You have been invited to this personal interview because of your outstanding expertise in hospice and palliative care and/or neurology. Today I would like to go through and discuss symptom specific items of a palliative and neurological outcome measure that has not yet been used in Germany and was therefore translated by us. We now need your help to find out if this translated questionnaire is well understood.

Transition question (5-10 minutes)

Subjects are given a copy of the translated version of the IPOS Neuro-S8 along with a pen. Time to completion is written down.

You now have the translated version of the questionnaire in front of you. Please read it carefully and fill it out at your convenience. When you are done, we will discuss any queries you may have.

Key questions (15-20 minutes each)

Guiding question (narrative prompt)	Memo for possible follow-up questions – only to be asked if not addressed by itself	Specific questions – to be asked verbatim
1. How did you find the questionnaire in general, in terms of language and content?	• Test instructions and questions understandable? • Difficulty understanding and answering the questions? • Length? • Overall relevance for assessing health problems in the neurological, palliative care setting?	• What was your linguistic understanding of the test instructions, questions and the individual symptoms? • How did you feel about the time needed to complete the questionnaire? • What can you say about the significance of the listed symptoms in relation to the expected health problems of your neurological palliative care patients?
2. Let us now go through the individual symptoms together. What do you understand by the symptom <xy> (<i>the symptoms are all named one after the other</i>) and how do you justify your answer?	• Reasons why individual symptoms were difficult to understand or answer • Suggestions on how to rewrite unclear or inappropriate symptoms	• Can you elaborate a bit, why did you find the symptom <xy> difficult to understand or have difficulty responding to? • How could we rewrite the symptom <xy>? • How did you come with this?
Final question (5 minutes)		
Is there anything else you would like to add that we have not yet addressed?		



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Interview guide for the project „cross-cultural adaptation and translation of the IPOS Neuro-S8“ - phase 5

Introductory question

You have been invited to this personal interview because you are currently receiving or have received medical care in the past. Today I would like to go through and discuss symptom specific items of a questionnaire with you that has not yet been used in Germany. We now need your help to find out if this translated questionnaire is well understood.

Transition question (5-10 minutes)

Subjects are given a copy of the translated version of the IPOS Neuro-S8 along with a pen. If the patient is physically not able to fill in the questionnaire, the interviewer will fill in the questionnaire on behalf of the patient. Time to completion is written down.

You now have the translated version of the questionnaire in front of you. Please read it carefully and fill it out at your convenience or tell me your answers and I will gladly write down your answers for you. When you are done, we will discuss any queries you may have.

Key questions (15-20 minutes each)		
Guiding question (narrative prompt)	Memo for possible follow-up questions – only to be asked if not addressed by itself	Specific questions – to be asked verbatim
1. How did you find the questionnaire in general, in terms of language and content?	• Test instructions and questions understandable? • Difficulty understanding and answering the questions? • Length? • Overall relevance for assessing health problems?	• What was your linguistic understanding of the test instructions, questions and the individual symptoms? • How did you feel about the time needed to complete the questionnaire? • What can you say about the significance of the listed symptoms in relation to your health problems?
2. Let us now go through the individual items together. What do you understand by the item <xy> (the symptoms are all named one after the other) and how do you justify your answer?	• Reasons why individual symptoms were difficult to understand or answer • Suggestions on how to rewrite unclear or inappropriate symptoms	• Can you elaborate a bit, why did you find the symptom <xy> difficult to understand or have difficulty responding to? • How could we rewrite the symptom <xy>? • How did you come with this?
Final question (5 minutes)		
Is there anything else you would like to add that we have not yet addressed?		


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IPOS NEURO S8 - Palliative & Supportive Care - IPOSNEUROS8-V1-DE 10/02/2021

F1. Welche Beschwerden oder Anliegen haben/hatten Sie in den letzten 3 Tagen?

1.
2.
3.

F2. Im weiteren Verlauf befindet Sie eine Liste an Symptomen, die Sie ggf. haben. Bitte kreuzen Sie für jedes Symptom jeweils ein Kästchen an und beurteilen Sie, wie stark Sie sich durch das Symptom in den letzten 3 Tagen beeinträchtigt gefühlt haben.

	Gar nicht	Ein wenig	Mäßig	Schwer	Sehr schwer
Schmerzen	0 ☐	1 ☐	2 ☐	3 ☐	4 ☐
Kurzatmigkeit	0 ☐	1 ☐	2 ☐	3 ☐	4 ☐
Übelkeit (das Gefühl, erbrechen zu müssen)	0 ☐	1 ☐	2 ☐	3 ☐	4 ☐
Erbrechen	0 ☐	1 ☐	2 ☐	3 ☐	4 ☐
Verstopfung (Darmträgheit)	0 ☐	1 ☐	2 ☐	3 ☐	4 ☐
Symptome im Mund (z.B. Mundtrockenheit, Apthen)	0 ☐	1 ☐	2 ☐	3 ☐	4 ☐
Spastik (Muskelkrämpfe)	0 ☐	1 ☐	2 ☐	3 ☐	4 ☐
Schlafstörungen	0 ☐	1 ☐	2 ☐	3 ☐	4 ☐

	Alleine	Mit Hilfe eines Freundes/einer Freundin oder eines Angehörigen/einer Angehörigen	Mit Hilfe eines Mitarbeiters/einer Mitarbeiterin
F3. Wie haben Sie den Fragebogen ausgefüllt?	1 ☐	2 ☐	3 ☐

Wenn Sie sich über einen der in diesem Fragebogen angesprochenen Punkte Sorgen machen, sprechen Sie bitte mit Ihrem Arzt/Ihrer Ärztin oder Ihrem Pflegepersonal.