Implementing patient-reported outcome measures in palliative care clinical practice: A systematic review of facilitators and barriers

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Abstract

Background: Many patient-reported outcome measures have been developed in the past two decades, playing an increasingly important role in palliative care. However, their routine use in practice has been slow and difficult to implement.

Aim: To systematically identify facilitators and barriers to the implementation of patient-reported outcome measures in different palliative care settings for routine practice, and to generate evidence-based recommendations, to inform the implementation process in clinical practice.

Design: Systematic literature review and narrative synthesis.

Data Sources: Medline, PsycInfo, Cumulative Index to Nursing and Allied Health Literature, Embase and British Nursing Index were systematically searched from 1985. Hand searching of reference lists for all included articles and relevant review articles was performed.

Results: A total of 3863 articles were screened. Of these, 31 articles met the inclusion criteria. First, data were integrated in the main themes: facilitators, barriers and lessons learned. Second, each main theme was grouped into either five or six categories. Finally, recommendations for implementation on outcome measures at management, health-care professional and patient levels were generated for three different points in time: preparation, implementation and assessment/improvement.

Conclusions: Successful implementation of patient-reported outcome measures should be tailored by identifying and addressing potential barriers according to setting. Having a coordinator throughout the implementation process seems to be key. Ongoing cognitive and emotional processes of each individual should be taken into consideration during changes. The educational component prior to the implementation is crucial. This could promote ownership and correct use of the measure by clinicians, potentially improving practice and the quality of care provided through patient-reported outcome measure data use in clinical decision-making.

Keywords

Palliative care, hospice, patient-reported outcome measures, narrative synthesis, clinical practice, end-of-life care

Introduction

Measurement is a fundamental component of evidence-based medicine and provides the necessary information clinicians require to make decisions in patient management. However, patients are increasingly active in their own treatment; therefore, collecting biological, laboratory information and mortality rates only is no longer adequate. Other outcomes such as health status, level of disability and quality of life are now common in the literature. It is commonly asserted that the patient’s perspective should inform in clinical decision-making and direct collection of outcome data from the patient should be incorporated in clinical practice. Patient-reported outcome measures (PROMs) are standardised, validated questionnaires that are completed by patients to measure their perceptions of their own functional status and wellbeing. In the past two decades, many PROMs have been developed, and increasingly, their role in clinical practice has been emphasised. PROMs are considered to be the gold standard for outcome measurement of subjective experiences, because the information

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captured comes directly from the patient and well-validated tools reflect the concerns and main problems of the clinical population.\textsuperscript{1,9} In palliative care settings, systematic collection of patient-reported outcomes using validated questionnaires may benefit clinical practice both at the population and individual patient level. This may be achieved by the following mechanisms: (a) facilitating identification and screening of physical, psychological, spiritual and social unmet needs that might otherwise be overlooked; (b) providing information on disease progression and impact of treatment prescribed; (c) facilitating patient/family/carer–clinician communication, promoting the model of patient-centred care by shared decision-making and advanced care planning, establishing common priorities and expectations regarding outcomes of treatment and disease progression, which does not mean using a PROM to act as a substitute for the therapeutic relationship, but rather to complement it; and (d) monitoring outcomes by performing audits as a strategy for improvement of the quality of care provided and its costs.\textsuperscript{10,11} Finally, if funding health-care institutions were to be allocated depending on performance and outcomes of the services, having those data available would be critical.

Models have been developed to fit health-related quality of life (HRQoL) and other PROMs data collection into clinical practice.\textsuperscript{12} Building on Wilson and Cleary’s\textsuperscript{12} conceptual model of patient outcomes, Osoba developed a model for HRQoL assessment, describing how to incorporate HRQoL measures into clinical practice, throughout the disease trajectory. Outcome measures are used like a conventional laboratory test, that is, to inform the differential diagnosis and monitor a patient’s progress. These types of data can be helpful in several phases of a condition or disease management: during the initial history and physical examination, laboratory and imaging procedures of investigation leading to a diagnosis, all phases of treatment and follow-up.\textsuperscript{13} However, despite recommendations, the routine use of PROMs in palliative care clinical practice has been slow to implement, so their optimal role in assisting decision-making and improving quality of care is yet to be achieved.\textsuperscript{2,14}

In other areas, for example, psychiatry, work has been developed on implementing PROMs in clinical practice where it has been suggested that to implement outcome measurement in a clinical unit and have clinicians conduct it as part of their clinical routine, takes at least 1 year.\textsuperscript{15} Despite the evidence indicating that routine outcome assessment has benefits for mental health patients, there are a number of reasons as to why it has not been adopted in most services.\textsuperscript{16} These include lack of appropriate instruments, time and incentives (financial and professional); lack of consensus regarding what outcome domains to include and what assessment measures to use; and clinicians remaining unconvinced of the benefits of routinely monitoring outcomes. Furthermore, barriers to successful screening for depression in cancer settings involve patient and clinician factors: acceptability of screening is critical to implementation, clinician confidence/skills influence screening success and training may improve confidence but effects upon long-term outcomes are modest.\textsuperscript{17} The latter seems to be a consequence of not providing clinicians with interpretation of scores, no mandating follow-up, not linking screening with training or other types of clinician support, and screening being generally ineffective without aftercare.\textsuperscript{17}

Two frameworks that have been developed are relevant for this review. One pertains to the implementation of clinical practice guidelines that describes five steps: assessment of clinician’s stage of readiness to change, assessment of specific barriers to guideline use, determination of appropriate level of intervention, design of dissemination and implementation strategies and evaluation of the strategies implemented.\textsuperscript{18} The other, ‘Promoting Action on Research Implementation in Health Services’ (PARIHS) framework was developed to incorporate research into practice.\textsuperscript{19–21} It suggests that successful implementation is a function of the relation between the nature of the evidence, the context in which the proposed change is to be implemented and the mechanisms by which the change is facilitated. It is advocated that (a) implementing research into practice is an organisational issue rather than an individual one; (2) there is a need to carefully plan implementation strategies such as interventions that address the need for education, audit and the management of change and (c) criteria for evaluating the impact of the intervention must be identified and agreed upon before implementing any change. The role of the facilitator seems to be paramount as the facilitation is considered to be an intervention for getting research into practice. Finally, the facilitation process is also emphasised in a study, which aimed at implementing two end-of-life care tools in care homes.\textsuperscript{22} A model of high facilitation was used. This included an experienced palliative care nurse facilitating the project and visiting the sites multiple times and key champions attending a 4-day facilitative course and cascading this training down to their own staff.

Systematic reviews on the use of outcome measurement in clinical practice have focused on the impact and effectiveness of PROMs in clinical practice but none of them covered the process of implementing the measures.\textsuperscript{23–25} There is a brief review exploring assessment of only quality of life in one particular setting, that is, oncology practice.\textsuperscript{26} Understanding the facilitators and barriers to the implementation of PROMs in palliative care settings could potentially inform the process of their implementation. Therefore, this review aimed to systematically identify facilitators and barriers to the implementation process of routine use of PROMs in different palliative care settings.
and generate evidence-based recommendations, to inform the implementation process in clinical practice. The objectives of this review were to (a) identify barriers and facilitators to the systematic implementation of PROMs in palliative care clinical practice, (b) identify needs and other comments of clinical teams regarding the routine use of PROMs and (c) identify lessons learned on the process of implementation of PROMs in clinical practice.

**Methods**

**Design**

This systematic review follows the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) recommendations.27

**Study identification**

Studies were identified using a systematic search of electronic databases with additional hand-searching of reference lists of included articles.27

**Search strategy**

Five databases (Medline, PsycInfo, Cumulative Index to Nursing and Allied Health Literature (CINAHL), Embase and British Nursing Index (BNI)) were searched and included the literature published between 1985 and August 2011. It was decided to exclude all the literature before 1985 because PROMs only start to emerge in the literature in the late 1980s. The searches were conducted between 29 August 2011 and 2 September 2011. The search was updated on 19 March 2012. Using Medical Subject Heading (MeSH) terms and free text, keyword groups, including ‘palliative care’, ‘PROMs’, ‘clinical practice’ and ‘facilitators and barriers’, were combined in several ways. Slight changes were made according to each database. Appendix 1 has the full search strategy for BNI. Data search was supplemented by hand search of reviews and relevant articles.27

**Study selection**

Inclusion criteria are as follows: (a) primary studies reported in English, Portuguese, Spanish, Italian, German and French because these languages were spoken in the wider study team; (b) studies using a PROM alongside the clinical care of adult patients (18 years old or older) with advanced disease in palliative care settings; and (c) studies reporting barriers and/or facilitators of the implementation of the PROM. Articles had to meet all the inclusion criteria, and no exclusion criteria to be included. All study designs were included.

Exclusion criteria are as follows: (a) published literature other than primary studies (i.e. review articles, books and conference articles); (b) studies reporting exclusively on development and feasibility phases of new measures, which did not include comments of users; and (c) studies of PROMs completed by proxies.

**Selection procedure**

All titles and abstracts were reviewed by one reviewer (B.A.) to assess eligibility for inclusion in the review. Full texts were retrieved if the inclusion criteria were met or the abstract contained insufficient information. Any uncertainty about eligibility after assessing full text was assessed by a second reviewer (I.J.H.).

**Data extraction**

Data were extracted by one reviewer (B.A.) and assessed by a second reviewer (N.C.). Tabulation was used as a first step. The author, country, aim, population, study design, measures used, settings, facilitators and barriers were extracted. Clauses were taken and kept as reported.

**Analysis**

Incorporating qualitative data into systematic reviews has been recognised as an important contribution for providing and analysing non-numerical research data by answering questions that are not easily addressed exclusively by experimental methods.28,29 Hence, a narrative synthesis was conducted following a framework of four elements to ensure robustness of the findings as described in the Guidance on the Conduct of Narrative Synthesis in Systematic reviews.30 Element 1 concerns the role of theory in evidence synthesis, Element 2 refers to the development of a preliminary synthesis, Element 3 pertains to exploring relationships within and between studies and in Element 4, the robustness of the synthesis is assessed. This method has proved to be useful in synthesising different types of data from different study designs.31 There is more than one technique to choose from in order to conduct Elements 2–4, depending on the data extracted, and the following were the chosen techniques for each element in this review:

**Element 1: the role of theory in evidence synthesis.** We present a theory as suggested in the guidance, which will contribute to the interpretation of the findings and the assessment of how widely applicable the findings might be.

**Element 2: developing a preliminary synthesis.** Element 2 was achieved by (a) grouping the data by tabulation and (b) performing thematic analysis. This allowed for the systematic identification of the main themes.

**Element 3: exploring relationships within and between studies.** Element 3 was achieved by (a) exploring the
influence of heterogeneity by focusing attention on the characteristics of the different studies and their potential relationships to the findings, (b) reciprocal translation by identifying common concepts across studies and (c) conceptual mapping was performed linking multiple pieces of evidence extracted from the different studies representing the relationships between them. Categories were first developed for each main theme. Data were then organised in a timeline, by implementation steps and sequenced actions. Any discrepancies were resolved by discussion (B.A. and N.C.).

**Element 4: assessing robustness of the synthesis.** There are two steps to carry out described in the guidance, as follows: (a) assessing the methodological quality of included studies and (b) being rigorous in the methodology used to synthesise findings and being critical regarding limitations of methods used. The latter will be included in the ‘Strengths and limitations’ section given under the ‘Discussion’ section. There are no universally accepted guidelines for either assessing the quality of specific qualitative methods nor an accepted set of criteria to assess the quality of qualitative and quantitative studies in order to incorporate their findings in a systematic review.\(^{28–30,32–36}\) We used the modified Harden criteria to assess the methodological quality of the included studies (Table 1).\(^{37,38}\) Two reviewers (B.A. and V.S.) assessed the quality of the studies independently. Discrepancies were resolved by discussion. Studies with a qualitative design were assessed according to 16 criteria subdivided into two dimensions: clarity of reporting and robustness of the study methods.

| Clarity of reporting                                                                 | Qualitative | Quantitative |
|-----------------------------------------------------------------------------------|-------------|--------------|
| 1. Was the context of the study clearly described?\(^{a}\)                         | X           | X            |
| 2. Were the goals of the study clearly described?                                  | X           | X            |
| 3. Was the research question clearly defined?                                      | X           | X            |
| 4. Was the design adequate for the study goal/question?                           | X           | X            |
| 5a. Was the identification and the recruitment of the sample clearly described and justified? | X           |              |
| 6. Were the data collection methods clearly described?                             | X           | X            |
| 7. Were the data analysis methods clearly described?                               | X           | X            |
| 8. Were the findings clearly described?                                            | X           | X            |

**Robustness of the study methods**

| Robustness of the study methods                                                                 | Qualitative | Quantitative |
|-----------------------------------------------------------------------------------------------|-------------|--------------|
| 9a. Was the sampling strategy comprehensive to ensure the generalisability of the results?  | X           |              |
| 9b. Was the size of the study population sufficient to ensure the generalisability of the results? | X           |              |
| 9c. Was the response rate sufficient to ensure the generalisability of the results?           | X           |              |
| 10a. Were methods used to establish the reliability of the data collection methods?          | X           |              |
| 10b. Were reliable measurement instruments used?                                             | X           |              |
| 11a. Were methods used to establish the validity of data collection?                         | X           |              |
| 11b. Were valid measurement instruments used?                                                 | X           |              |
| 12a. Were methods used to establish the reliability of the data analysis?                   | X           |              |
| 12b. Were adequate analysis techniques used?                                                  | X           |              |
| 13a. Were methods used to establish the validity of the data analysis?                       | X           |              |
| 13b. Were methods used to establish the validity of the data analysis?                       | X           |              |
| 14. Did the research move logically from a description of the data, through quotations or examples, to an analysis and interpretation of the meanings and their significance? | X           |              |
| 15. Was evidence of reflexivity in the process reported (interim data analyses guides further data collection and analyses)? | X           |              |
| 16. Were the findings really rooted in the perspectives of the population studied?          | X           |              |

Number of positive criteria

| Qualitative | Quantitative |
|-------------|--------------|
| 16          | 13           |

The original 10 criteria from Harden are printed in italic format.

\(^{a}\)Modified Harden criteria in Slort et al.\(^{37,38}\)
of the respondents, logically proceeding from data to interpretation and reflexivity, which has a maximum score of 8. Regarding the studies with a quantitative design, 13 criteria were used, also subdivided in those two dimensions. The maximum possible score for robustness of study methods was 5. If the study used a mixed-methods approach, then the methodology that provided the data of interest for this review, was rated. Each criterion was rated as ‘yes’ or ‘no’, and if there was insufficient information, the score was ‘no’. Studies were rated by both reviewers as ‘high’, ‘medium’ or ‘poor’ quality. However, no studies were excluded based on the quality score because there is no consensus on the minimum quality assessment score required for inclusion in a review.29

Results

Identification and inclusion of studies

The search strategy yielded a total of 3863 hits. Following screening of titles and abstracts (Figure 1), 421 articles were reviewed in full detail and 393 were excluded for not having comments on PROMs use, implementation of the measure(s), nor use in clinical practice. Twenty-six studies were included within 31 articles.

Eleven studies were conducted in the United Kingdom, seven in the United States, two in the Netherlands and one in each of the following countries: Australia, Canada, Israel, Italy, Malaysia and Vietnam (Table 2). There were no intervention studies testing the introduction of facilitators to overcome barriers when implementing PROMs in clinical practice. Eight studies used qualitative methods and provided structured, in-depth information about the facilitators, barriers and lessons learned when implementing PROMs in palliative care clinical practice. These studies used a range of designs (n = 4) including web-based online surveys, semi-structured interviews with patients and clinicians either face to face or by phone and meetings with clinical teams and focus groups. Seventeen studies used quantitative methods (n = 7) and none looked at facilitators or barriers to the implementation of PROMs as a primary outcome. One randomised controlled trial evaluated the efficacy of standardised HRQoL assessments used in facilitating patient–physician communication. The PROMs used in the studies include seven measures on pain, seven on symptoms, eight on quality of life and six measuring other concepts (Table 2).

Narrative synthesis framework

Element 1: the role of theory in evidence synthesis. We draw on work conducted in other fields, including an adaptation of the Slade70 model, and on the early social and behavioural psychology theories, which are the base for organisational theories of change.15,18 The model (Figure 2) takes into consideration that implementation is an ongoing dynamic process, and that staff from any setting with particular characteristics and rules are individuals with ongoing cognitive and emotional processes. These influence how they relate and work with colleagues, patients and families and how they react to change.

Element 2: developing a preliminary synthesis. Tabulation allowed us to find, at an early stage of the analysis, that there were no intervention studies of implementing facilitators to overcome barriers when using a PROM in palliative care clinical practice. This was an important finding, which allowed for the integration and analysis of the data in a way that was most informative but still answered our question. Then, thematic analysis was performed. We first applied our a priori three main themes: facilitators, barriers and lessons learned (see Appendix 2 for all data extracted). Most data extracted from the included studies were in the form of narrative observations located in the ‘Discussion’ and ‘Conclusion’ sections. In-depth data explicitly reported in the ‘Results’ section were also extracted, mainly from the focus groups studies and the web-based online survey study.

Element 3: exploring relationships within and between studies. After integrating the data in the main themes, in-depth inductive analysis led to the creation of five different categories for facilitators and lessons learned and six categories for barriers, namely, management, education, tool specific, clinical utility, financing and attitudes. All categories are presented in Table 3 accompanied by an illustrative quote per category.

In addition, as analysis continued, a different typology emerged. This led us to reorganise the data in a timeline. We suggest three sequential steps at different levels (see Appendix 3 for detailed data): (a) preparation, with different actions put in place simultaneously at management level, health-care professional level and patient level; (b) implementation, in which the measure is taken into clinical practice, and different tasks are fulfilled at different levels and (c) assessment and improvement, which begins with the implementation step and involves discussion and potential changes of actions previously implemented. Hence, data were organised by those three sequential steps including barriers, facilitators and lessons learned for each. Similarities between studies emerged as some facilitators, barriers and lessons learned appear to be common to all sequential steps. Furthermore, by categorising the data as described (Figure 3), it seemed that some barriers discussed in some studies were being answered/resolved by facilitators and lessons learned mentioned in other studies. By exploring these relationships, we were able to link different concepts and synthesise them in the form of recommendations for implementing PROMs in palliative care clinical practice (Table 4).
Element 4: assessing the robustness of the synthesis. We applied our quality criteria and categorised six qualitative studies as high quality because these scored 13 or more, two as medium quality having scored between 10 and 12 and one as poor quality since it scored below 10. Regarding the quantitative studies, 10 were rated high quality for scoring 11 or higher, and a score of 10 was rated as medium quality; however, there were no studies with this score, and
| Reference, country | Aim | Study design, population and setting | Type of data reported | Measures | Clarity of reporting | Robustness of the study methods | Quality assessment sum score |
|-------------------|-----|--------------------------------------|-----------------------|----------|---------------------|-----------------------------|-----------------------------|
| Bouvette et al.,39 Canada | To determine the feasibility of implementing the PSAR in a variety of settings | Qualitative exploratory; focus groups with nurses; chart audits Palliative care institutions in Ottawa | In-depth data explicitly reported in results Facilitators and barriers extracted | PSAR | 8/8 | 7/8 | 15/16 |
| Bourbonnais et al.,40 Canada | To focus on the importance of the use of tools for pain and symptom management, issues around implementing them and sustaining their use in the clinical setting | Article reporting lessons learned from conducting the earlier study Narrative opinions related to the earlier study Facilitators, barriers and lessons learned extracted | – | – | – | – | |
| Dunckley et al.,41 UK | To identify facilitators and barriers to implementing outcome measures and to identify and facilitate methods of overcoming those barriers | Qualitative: action-research approach. Semi-structured interviews with clinicians; diary completion; monthly meetings Hospice | In-depth data explicitly reported in results Mainly facilitators extracted | POS | 8/8 | 7/8 | 15/16 |
| Bausewein et al.,42 UK | To describe the use and experiences with PROMs of professionals working in palliative care in Europe and Africa | Qualitative: web-based online survey to clinicians Multiple settings, both in Europe and in Africa | In-depth data explicitly reported in results Mainly barriers extracted | N/A | 8/8 | 7/8 | 14/16 |
| Davson et al.,43 UK | To examine and compare doctors’ and nurses’ views and experiences regarding outcome measurement in palliative care, including PROMs | Qualitative: web-based online survey to clinicians Multiple settings, both in Europe and in Africa | In-depth data explicitly reported in results Mainly barriers extracted | N/A | – | – | – |
| Harding et al.,44 UK | To identify the outcome tools currently in use in end-of-life care (both clinically and for research) across Europe and investigate the preferred features of outcomes tools from the perspective of those who select and apply them | Qualitative: web-based online survey to clinicians Multiple settings, both in Europe and in Africa | In-depth data explicitly reported in results Mainly barriers extracted | N/A | – | – | – |
| Schulman-Green et al.,45 USA | To develop and pilot test performance measures and a data collection system that hospices could use in partial fulfilment of CMS requirements | Qualitative: semi-structured telephone interviews Hospice staff | In-depth data explicitly reported in results Mainly lessons learned extracted | ESAS | 7/8 | 7/8 | 14/16 |
| Schwartz et al.,46 USA | To evaluate the feasibility and utility of the MVQOLI-R during over 6 weeks of use; explore the utility of the MVQOLI-R as a research and/or clinical tool (i.e. a psychometric versus clinimetric instrument) | Qualitative: semi-structured interviews to clinicians Hospice, home and Pall Care settings | In-depth data explicitly reported in results Mainly lessons learned extracted | MVQOLI-R, MSAS | 8/8 | 6/8 | 14/16 |
| Hughes et al.,47 UK | To describe the implementation of a palliative care outcome measure in non-specialist palliative care settings and to understand the implementation of the measure | Qualitative: semi-structured interviews to staff and patients 15 non-specialist palliative care settings | In-depth data explicitly reported in results Mainly barriers and lessons learned extracted | POS | 7/8 | 6/8 | 13/16 |
Table 2. (Continued)

| Reference, country | Aim | Study design, population and setting | Type of data reported | Measures | Clarity of reporting | Robustness of the study methods | Quality assessment sum score |
|--------------------|-----|------------------------------------|-----------------------|----------|---------------------|-------------------------------|-----------------------------|
| Cox et al.,48 UK   | To focus on the clinical acceptability of a computerised assessment tool and present the difficulties in evaluating the support the tool provided to patients | Qualitative: semi-structured interviews by telephone or in person to clinicians and patients Hospice | In-depth data explicitly reported in results | Mainly barriers extracted | ESAS, EQ-5D | 7/8 | 4/8 | 11/16 |
| Hughes et al.,49 UK| This study investigated professionals’ views of using outcome measures with special reference to one, the POS | Qualitative: semi-structured telephone interviews to health-care professionals | In-depth data explicitly reported in results | Mainly barriers and lessons learned extracted | POS | 6/8 | 4/8 | 10/16 |
| Hughes et al.,50 UK| Assessing palliative care outcomes for people with MND living at home | Qualitative: semi-structured interviews to MND patients | In-depth data explicitly reported in results | Mainly facilitators and barriers extracted | – | – | – | – |
| Barret,51 UK       | To develop an assessment tool in wound care that can be used by the practitioner Establishing a national system (the PCOC) to measure outcomes and quality of specialist palliative care services and to benchmark services across the country | Qualitative: meeting with district nursing team | Narrative opinions | Mainly lessons learned extracted | Pain monitoring Aid | 4/8 | 4/8 | 8/16 |
| Eagar et al.,52 Australia | Establishing a national system (the PCOC) to measure outcomes and quality of specialist palliative care services and to benchmark services across the country | National system to measure outcomes in palliative care services | Narrative opinions | Mainly lessons learned extracted | PCOC tool kit (Palliative Care Phase, SAS, RUG-ADL, Australian-modified KPS, PCPSS and some items of the POS) | 8/8 | 5/5 | 13/13 |
| Rawlings et al.,53 Australia | Using PACA tools to influence and enhance clinical practice | National system to measure outcomes in Palliative care services | Narrative opinions related to the earlier study | Mainly lessons learned extracted | – | – | – | – |
| Detmar et al.,54 the Netherlands | To evaluate the efficacy of standardised HRQoL assessments in facilitating patient–physician communication and increasing physicians’ awareness of their patients’ HRQoL-related problems | Quantitative: prospective, randomised crossover trial | Narrative opinions | Mainly lessons learned extracted | EORTC QLQ-C30, COOP, WONCA, SF-36, ECOGS, PSQ C | 8/8 | 5/5 | 13/13 |
| Caraceni et al.,55 Italy | To assess the compliance of hospitalised patients with chronic cancer pain, referred to an inpatient palliative care consultation service, with self assessment of pain intensity by means of a daily pain form | Quantitative: prospective longitudinal Hospitalised patients Tertiary oncological referral centre | Narrative opinions | Facilitators, barriers and lessons learned extracted | 0-10 NRS | 7/8 | 5/5 | 12/13 |
| Ellershaw et al.,56 UK | To evaluate the effectiveness of a hospital palliative care team in the provision of symptom control, patients’ and relatives’ awareness of the diagnosis and outcome regarding the patients’ placements | Quantitative: prospective longitudinal study Cancer patients referred to a hospital-based palliative care team | Narrative opinions | Mainly lessons learned extracted | PACA tool | 8/8 | 4/5 | 12/13 |
| Reference, country | Aim                                                                 | Study design, population and setting                                                                 | Type of data reported                                                                 | Measures                                      | Clarity of reporting                  | Robustness of the study methods | Quality assessment sum score |
|-------------------|----------------------------------------------------------------------|-----------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------|-----------------------------------------------|---------------------------------------|---------------------------------|-------------------------------|
| Escalante et al.,57 USA | To separately profile fatigue in cancer patients with solid tumours or haematological malignancies who were being served in the EC | Quantitative: retrospective chart review Emergency centre                                             | Narrative opinions Mainly lessons learned extracted                                       | NRS for fatigue and pain                      | 7/8                                   | 5/5                             | 12/13                         |
| Ewing et al.,58 UK | To investigate the agreement on symptom assessments between patients at home and GPs and district nurses | Quantitative: cross-sectional prospective study Adult palliative care patients estimated to be in their last year of life | Narrative opinions Mainly lessons learned extracted                                       | CAMPAS-R (VAS), CAMPAS-P (VAS)               | 8/8                                   | 4/5                             | 12/13                         |
| Hoekstra et al.,59 Netherlands | To achieve symptom relief through systematic and regular symptom reporting by patients themselves | Quantitative: RCT Hospital and GP practice                                                           | Narrative opinions Mainly barriers and lessons learned extracted                         | Symptom monitor                               | 8/8                                   | 4/5                             | 12/13                         |
| Kamel et al.,60 USA | To investigate the effect of utilising a combination of three easily administered pain assessment instruments on the frequency of diagnosing pain among elderly nursing home residents | Quantitative: cross-sectional pain assessment + chart review Nursing homes                           | Narrative opinions Mainly lessons learned extracted                                       | VAS for pain, pain descriptive scale and behaviour scale | 7/8                                   | 5/5                             | 12/13                         |
| De Rond et al.,61 the Netherlands | To assess the feasibility of daily pain assessment from nurses’ and patients’ perspective in multiple settings | Quantitative: feasibility study 2 general hospitals and 1 university hospital. In each, 2 surgical and 1 medical wards | In-depth data explicitly reported in results Mainly lessons learned extracted             | NRS                                           | 7/8                                   | 4/5                             | 11/13                         |
| Jette et al.,62 USA | To determine (a) the extent of the use of standardised outcome measures and (b) perceptions regarding their benefits and barriers to their use | Quantitative: observational. Paper survey Physical therapists                                        | In-depth data explicitly reported in results Mainly barriers and lessons learned extracted | N/A                                           | 8/8                                   | 3/5                             | 11/13                         |
| Defilippi and Cameron,63 UK | To assess the impact of a model of introducing a palliative care component and professional supervision of community caregivers on the quality of care given to people living with HIV/AIDS and their families | Quantitative: prospective longitudinal pilot study Hospice based but home care program             | Narrative opinions Mainly barriers and lessons learned extracted                         | APCA POS                                      | 5/8                                   | 4/5                             | 9/13                          |
| Hardy et al.,64 UK | To determine whether a QoL instrument could be used as a primary end point or outcome measure in palliative care | Quantitative: prospective longitudinal study Palliative care unit                                    | Narrative opinions Mainly barriers and lessons learned extracted                         | RSCL                                          | 5/8                                   | 4/5                             | 9/13                          |
| Bercovitch et al.,65 Israel | To present preliminary experience with the MCPAC | Quantitative: prospective longitudinal Hospice terminal cancer patients                              | Narrative opinions Mainly barriers and lessons learned extracted                         | MCPAC                                         | 5/8                                   | 4/5                             | 9/13                          |
| Chang et al.,66 USA | To evaluate the acceptability of the computer-administered program to patients and the most useful and appropriate presentation of patients' QoL information to treating oncologists | Quantitative: feasibility study, prospective longitudinal Oncology clinic                           | Narrative opinions Mainly barriers and lessons learned extracted                         | Electronic FACT-L                             | 6/8                                   | 2/5                             | 8/13                          |
Clark et al., 67  
USA  
To describe the implementation of the touch screen technology as an effective psychosocial screening tool with immediate clinical utility  
Quantitative: prospective longitudinal study of Cancer centre  
In-depth data explicitly reported in results  
Mainly facilitators extracted  
‘How can we help you and your family’ screening instrument provided useful data as secondary outcomes or process data  
6/8  
2/5  
8/13

Devi and Tang, 68  
Malaysia  
To study the feasibility of modifying the use of a pain assessment tool  
Quantitative: feasibility study of Hospital, oncology ward  
Narrative opinions of Mainly facilitators extracted  
Short form of the Brief Pain Inventory, Wong-Baker FACES Scale  
5/8  
35/2  
8/13

Green et al., 69  
Vietnam  
To evaluate the impact of integrating palliative care services within a district HIV outpatient ART clinic setting by assessing changes in pain, other symptoms, depression, anxiety and perceived social support, and exploring intervention accessibility, acceptability and feasibility  
Quantitative: non-randomised controlled trial of HIV outpatient clinics  
Narrative opinions of Mainly lessons learned extracted  
10 point symptom severity scales  
5/8  
2/5  
7/13

PROM: patient-reported outcome measure; CMS: Centers for Medicare and Medicaid Services; PCOC: Palliative Care Outcomes Collaboration; MND: motor neurone disease; HRQoL: health-related quality of life; GF: general practitioner; RCT: randomised controlled trial; QoL: quality of life; ART: antiretroviral treatment; PSAR: The Ottawa pain and symptom assessment record; POS: Palliative care Outcome Scale; N/A: not applicable; ESAS: Edmonton Symptom Assessment Scale; MVQOLI-R; Missoula-VITAS Quality of Life Index- Revised; MSAS: Memorial Symptom Assessment Scale; EQ-5D: European Quality of Life-5 Dimensions; SAS: symptom assessment scale; RUG-ADL: Resource Utilization Group/Activities of Daily Living; KPS: Karnofsky Performance Score; PCPS: Palliative Care Problem Severity Score; EORTC QLQ-C30: European Organization for Research and Treatment of Cancer, Quality of Life Questionnaire Core 30; COOP: Dartmouth Primary Care Cooperative Information Functional Health Assessment; WOONCA: World Organization Project of National Colleagues and Academics; SF-36: Medical Outcomes Study 36-Item Short-Form Health Survey; ECOGS: Eastern Cooperative Oncology Group scale; PSQ C: Patient Satisfaction Questionnaire C; NRS: numeric rating scale; PACA: Palliative Care Assessment; CAMPAS-R: comprehensive measure for palliative care for patients; VAS: visual analogue scale; CAMPAS-P: comprehensive measure for palliative care for professionals; APCA POS: African Palliative Care Association African Palliative care Outcome Scale; RSC: Rotterdam symptom checklist; MCPAC: multidimensional continuous pain assessment chart; FACT-L: Functional Assessment of Cancer Therapy-Lung; EC: Emergency Centre.

Articles ordered by quality assessment sum score. Articles, which do not present scores, describe the same study as the article immediately above.

seven studies scored 9 or below and so were rated as poor quality (Table 2).

Discussion

To our knowledge, this is the first systematic review that specifically identifies and appraises facilitators and barriers and draws recommendations of implementation of PROMs in clinical practice in palliative care. Although there were no intervention studies identified in this review, the qualitative data provided structured, in-depth information about the facilitators, barriers, needs and lessons learned when implementing PROMs in palliative care clinical practice. The quantitative studies identified did not meet our inclusion criteria as primary aims but provided useful data as secondary outcomes or process data from the original studies.

Our main findings are in line with our initial theory based on work conducted in other fields. First, during the implementation process, there is the need to acknowledge interpersonal relationships between the clinical team members and the ongoing emotional and cognitive processes that occur in each individual. Fear of change, feeling that one is being assessed and that one's work is open to criticism due to the results that PROMs might show, and fear of added work are a few of the concerns that may lead to behaviour, which opposes change. Therefore, a firm but sensitive leadership seems to be needed to motivate individuals and reassure them that the use of PROMs is beneficial and ultimately aims to improve the quality of care provided to patients (Appendix 2, management category). Allowing individuals to take ownership of the measures rather than mandating their use may be important to avoid conflict. Providing feedback to clinicians appears to be a powerful tool to influence beliefs and attitudes towards the use of PROMs in clinical practice (Appendix 2, clinical utility). Some authors have used organisational theories as a backdrop to inform and design indicators to measure the quality of delivery of health care in general practice. 71
A clinical setting is a dynamic system with its own rules, ranks, values and beliefs. It is formed by individuals that may or may not share those values and beliefs and hence may be a driving force or an opposite force to change. Each individual has ongoing cognitive and emotional processes. The transtheoretical model of change, considers behaviour to be a continuous process made up of five stages: pre-contemplation, contemplation, preparation, action and maintenance. Moving from the 1st to the 2nd implies changing knowledge and attitudes, going from the 2nd to the 3rd and 4th stages involves changes in emotional processes such as positive beliefs about self-efficacy and having or developing the necessary skills and finally, evolving to the final stage implies restructuring the environment or system in which the behaviour occurs and providing support.

Decisions to be made prior to implementation of a PROM:
- Selection of outcomes of interest
- Selection of outcome measure(s)
- Educational component about measure and how to use results
- Selection of one coordinator/facilitator
- Who applies the measure and its periodicity

Second, implementing a PROM in palliative care clinical practice appears to benefit from careful planning of three distinct phases or steps (Appendix 3). The preparation phase involves (a) ensuring all health-care professionals are comfortable with changes planned and having a coordinator who will be in charge, (b) an education component and (c) selection of the measure(s) to be used. The latter is becoming more complex due to the high number of existing measures, many used only in research and/or just used in one study. This may not provide enough information about the measure in terms of using it in practice. Hence, the educational component seems to be key because if met, it will allow for clinicians to (a) understand why a measure is needed and how it could potentially benefit their practice; (b) learn about the measure(s), which will be implemented, namely, measurement properties, design and scoring system(s); (c) provide role play on how to explain PROMs usage to patients; (d) explore interpretation of results at
different levels to benefit not only the individual patient but also the population, which the setting serves; (e) discuss the best options for storing and managing the collected data; and (f) understand the evaluation, which will follow. This seems to be the most time-consuming step. During the actual implementation, the results of each assessment point (previously decided, for example, daily, weekly, at each appointment) provide feedback on monitoring the patient’s progress and disease management. Having timely feedback on patient outcomes aids clinical decision-making and may enhance communication not only between patient and clinician but also within the multidisciplinary team. Finally, acknowledging what could be changed throughout the process is important in order to make the necessary enhancements with the overall aim of improving practice and the quality of care.

**Implications for research**

There are potential implications for future research that can be drawn from the findings, namely, whether the introduction of the measure(s) is an intervention in itself, regardless of the primary outcome of the study. Perhaps, there is an adjustment phase that needs to be taken into account before the actual main data collection takes place. It might be important to consider whether study protocols should include this period. There may be an impact on timing, that is, will the baseline be measured immediately after implementing the use of the PROM(s) or will there be a period of adjustment for the users to become more familiar with the tool(s). The same could be said about analysis: thought must be put into whether or not to include the data collected in that period of time in the analysis, considering that

| Table 3. Categories of extracted facilitators, barriers and lessons learned. |
|---|
| **Facilitators – 5 categories** |
| (a) Management/organisational/setting specific |
| (b) Education (of all actors involved/training) |
| (c) Tool specific |
| (d) Clinical utility and relevance of a PROM |
| (e) Psychosocial theories/psychology of work |
| **Quotes** |
| ‘Prior meeting to explore feasibility of implementation of the measure: to implement measure, planning and evaluating are essential; coordinator is identified to undertake overall responsibility for implementation’41 |
| ‘Educational program prior to implementation of the measure: the importance of training in all aspects of use to help staff to become familiar and more comfortable with administering the tool’51 |
| ‘Burden of measure completion on patient/residents is considered’51 |
| ‘Measure asks about issues that are relevant to clinical care’51 |
| ‘Persistence and encouragement by both the research assistant and the unit head nurse on the wards were necessary to ensure the implementation was successful’41 |
| **Barriers – 6 categories** |
| (a) Management/time |
| (b) Education |
| (c) Tool specific |
| (d) Financing |
| (e) Illness specific |
| (f) Motivation/personality/attitudes/beliefs |
| **Quotes** |
| ‘Not enough staff: time constraints and fear of added work’42 |
| ‘Lack of training and guidance about how to use tools’42 |
| ‘Tools are too burdensome for patients & families and staff’42 |
| ‘Cost constraints (e.g. fees for tools)’42 |
| ‘Overall severity of the disease condition’55 |
| ‘Fear of change’57 |
| **Lessons learned – 5 categories** |
| (a) Management/setting level |
| (b) Education specific |
| (c) Tool specific |
| (d) Individual patient level |
| (e) Carer level |
| **Quotes** |
| ‘Integrating a new routine into daily clinical practice takes time and effort’61 |
| ‘Education regarding the tool will be tailored to suit the individual locality needs. This is to allow the staff to take ownership and understand the benefits to its use’51 |
| ‘Whatever scale is chosen, administration must follow specific guidelines, and administration modalities must be appropriate to the clinical or research needs and practical enough to obtain adequate patient compliance’55 |
| ‘Reporting symptoms on a regular basis by patients was valuable because of the patients’ apparent awareness of their own symptoms’59 |
| ‘Caregivers reported increased confidence in caring for people with advanced disease’63 |

PROM: patient-reported outcome measure.

One quote is provided as an example for each category.
misuse of the measure(s) may have an impact on the strength and validity of results.

**Strengths and limitations**

One strength of this review is the heterogeneity of the included studies. These come from a large number of different countries as well as from different settings, which suggests that issues related with implementing PROMs are common and universal to clinical practice in palliative care. Furthermore, the fact that different measures were used in the included studies helps to clarify that challenges in implementing PROMs are not exclusive to the characteristics of the chosen measure, but actually depend on an
Table 4. Recommendations for implementation of PROMs in clinical practice per phases and at different levels.

1. Preparation step

**Management level (13)**
- Initial meeting to explore feasibility of implementation of the measure: planning and evaluating are essential; a coordinator is identified to undertake overall responsibility
- Establish clear boundaries at the outset to avoid unrealistic expectations from all actors involved
- Appraise characteristics and resources of the setting and the requirements of the proposed innovation: that is, it is necessary to attune the implementation protocol to the needs of the specific setting and think about good documentation: will allow staff to take ownership and understand the benefits of using the chosen measure, which needs to be adaptable to local circumstances
- Discussions in this step should include assessment of current data collected, how data are collected, what new data items are to be extracted, how data can be extracted and provided to who will analyse it, and, which measure(s) to use, especially due to the high number of existing measures and the fact that many are used in research or were used in one study only. A measure broadly used will allow for comparisons. The aim should be to embed the collection of standardised clinical assessment and other clinically relevant data into daily clinical practice with a view to improving clinical care
- Visualisation of results should allow easy and quick interpretation by health-care providers, patients and their families
- Consider investing in training of clinical key staff: improves data quality and demonstrates that data can be used (a) on a daily basis as part of clinical practice and (b) to manage and improve services, that is, data can be used in case conferencing and interdisciplinary team meetings and at staff handovers for patient care, discharge planning and discharge and transfer of patients as well as audit for quality assurance
- Education and training sessions: consider your setting and organisational needs, when planning, that is, timing of sessions
- Consider implementation strategies: consistency is paramount. Frequency of use of the measure; burden of completion on patient; measures that can be analysed using existing resources; a measure that is easy and quick to use; establish link with a research/audit group
- Consider establishing a program evaluation system for the first month of each phase as well as every 6 months during the first year of implementation. Should include feedback from patients, staff members, physicians, nurses (and ITS team if applicable) as well as cost savings to the setting
- Organisational support is needed to maximise the tool’s impact: mechanisms for sharing the information with caregivers need to be developed
- Investment in computerised systems for quick data entry and analysis should be considered. Ways of visualising, storing, retrieving and backing up the data should be discussed
- Initiate implementation in stages to improve acceptability
- Coordinator maintains good relationships with all involved by having both an awareness of the extra time and effort needed to implement new outcome measures and providing the appropriate resources and practical support to use the measures and carry out data analysis. Cascade management style is adopted

**Health-care professional level (3)**
- Education and training sessions: understanding outcome measurement and why it is important. Administration of chosen measure must follow specific guidelines so it is used in a valid way: staff will feel comfortable using it and should obtain adequate patient compliance. Standardisation of implementation procedures is critical to the effective use of the measure for quality monitoring but it must be balanced with the priority of individualised care
- Organisational support is needed to maximise the tool’s impact: mechanisms for sharing the information with caregivers need to be developed
- Choosing a measure: efforts should be made to ensure readability and interpretability by patients: reading level, font size and general appearance. Should reflect patient situation, quick and easy to use and interpret, validated for population and setting

2. Implementation step

**Management level (7)**
- Maintain strategies of reminders to incorporate the use of the measures in clinical practice, for example, daily assessment easily fits with daily routine and does not take additional time, that is, if it is incorporated with moment of control of vital signs: better insight into patients’ situation
- Education and motivation of patients could improve compliance
- Timely feedback of results is always done
- Data that are more time consuming to collect but important for quality reporting are collected periodically, rather than, for example, daily
- Persistence and encouragement by the coordinator are necessary to ensure the implementation is successful and that communication among all actors involved is clear
- Space and time are created to discuss how implementation is going: problems and benefits of using outcome measures are easily and rapidly noticeable (what is working and what is not)
- Coordinator investigates all complaints/issues and addresses them immediately, at the time of the complaint/issue (negative views from both patients and professionals tend to be at the outset of implementation when they have not familiarised with the measure)

**Health-care professional level (4)**
- Every member of the team who should be using the outcome measures does so

(Continued)
additional number of factors. The same can be said with regard to the different populations in which the included studies were conducted. This suggests that the findings from this review are applicable to a variety of settings. In addition, we originally aimed at identifying facilitators and barriers to the implementation of PROMs in clinical practice in palliative care only. Using the narrative synthesis approach allowed us to go further and explore the relationships between the different studies. This led to reorganise and synthesise the data in the form of recommendations.

Nevertheless, there are some limitations, starting with the quality assessment, which was not an exclusion criterion, and therefore, articles were not excluded even if rated as poor quality. Second, there was some discussion between reviewers on how to categorise and present these data, mainly because some clauses fit more than one theme, and repeating those clauses would be more beneficial and accurate. However, the size of the appendices was already substantial, and there was no information lost by not using repetition. Third, the wording in the literature is somewhat confusing: the ‘benefits/advantages’ of using PROMs is not the same as facilitators of implementation and ‘disadvantages’ of usage of PROMs is not the same as barriers to implementation. This is subjective and open to personal interpretation. Another potential bias in this review is that there was only one reviewer including/excluding articles, although a second reviewer would be consulted if there were doubts after assessing the full text. Furthermore, only one reviewer developed the categories to synthesise the data, although a second reviewer appraised those and commented on them. In addition, the grey literature was not searched.

**Conclusion**

Implementing PROMs in palliative care clinical practice is an ongoing interactive and continuous process. There is a need to identify and address potential barriers to a successful implementation of PROMs in clinical practice, using appropriate facilitators, tailored to the characteristics of each setting. A key facilitator has to do with the role of a coordinator/facilitator throughout the implementation process. It is important to recognise the ongoing cognitive and emotional processes of individuals when change is being planned and implemented. The educational component for health-care professionals prior to the implementation is also paramount. This could promote ownership and correct use of the measure selected. Ideally, this measure should be short and screen for different palliative needs. Online resources and training courses on outcome measurement and PROMs for clinicians and researchers are starting to emerge. Collaborations such as the Outcome Assessment and Complexity Collaborative (OACC) project are also a step forward in implementing PROMs is clinical practice in palliative care. In conclusion, there is a need for guidance on implementing

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**Table 4. (Continued)**

- Maintain strategies of reminders to incorporate the use of the measures in clinical practice. Daily assessment easily fits with daily routine and does not take additional time
- Interpretation of results is used in practice
- Every member of the team contributes to discussing how implementation is going

**Patient level (1)**

- Education and motivation of patients could improve compliance

**3. Assessment and improvement (this step will start within the implementation step)**

**Management level (7)**

- Collected items are reviewed with the aim of being clinically relevant and not burdensome to collect
- Assess if the measure generated valuable information, without an increase in paperwork, potentially freeing up some time to deliver more patient-centred care
- Assess if practice improved initially as a result of just ‘planting the seed’ of the patient’s needs management. Continue to encourage its use, which will improve confidence with the tool
- Assess if there is a benefit to both patient and practitioner in achieving better outcomes, improved concordance and potentially reducing the cost and effort of that management
- Assess if collation of data generated by the use of the measure allows continuous, accurate collection of information, which should reflect the activity of the palliative care team. It should also identify areas of potential future development
- Continue to refine the process to make it more understandable and acceptable to patients and caregivers
- Assess if changes will be made in practice based on the results of implementing and using the measure

**Health-care professional level (2)**

- Collected items are reviewed with the aim of being clinically relevant and not burdensome to collect
- Assessing and improving documentation will potentially improve practice and quality of care by highlighting needs

**Patient level (1)**

- Assess if there is a benefit to patient in achieving better outcomes

PROM: patient-reported outcome measure.
PROMs in palliative care clinical practice, which could potentially improve practice and the quality of care provided by assisting in clinical decision-making.

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