A systematic review



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outcome measures in palliative care:

Psychometric properties of carer-reported

Abstract

Background: Informal carers face many challenges in caring for patients with palliative care needs. Selecting suitable valid and reliable outcome measures to determine the impact of caring and carers' outcomes is a common problem.

Aim: To identify outcome measures used for informal carers looking after patients with palliative care needs, and to evaluate the measures' psychometric properties.

Design: A systematic review was conducted. The studies identified were evaluated by independent reviewers (C.T.J.M., M.B., M.P.). Data regarding study characteristics and psychometric properties of the measures were extracted and evaluated. Good psychometric properties indicate a high-quality measure.

Data sources: The search was conducted, unrestricted to publication year, in the following electronic databases: Applied Social Sciences Index and Abstracts, Cumulative Index to Nursing and Allied Health Literature, The Cochrane Library, EMBASE, PubMed, PsycINFO, Social Sciences Citation Index and Sociological Abstracts.

Results: Our systematic search revealed 4505 potential relevant studies, of which 112 studies met the inclusion criteria using 38 carer measures for informal carers of patients with palliative care needs. Psychometric properties were reported in only 46% (n=52) of the studies, in relation to 24 measures. Where psychometric data were reported, the focus was mainly on internal consistency (n = 45, 87%), construct validity (n = 27, 52%) and/or reliability (n = 14, 27%). Of these, 24 measures, only four (17%) had been formally validated in informal carers in palliative care.

Conclusion: A broad range of outcome measures have been used for informal carers of patients with palliative care needs. Little formal psychometric testing has been undertaken. Furthermore, development and refinement of measures in this field is required.

Keywords

Palliative care, carers, outcome measures, psychometrics, systematic review

What is already known about the topic?

- The involvement of informal carers is essential for the provision of palliative care, but informal caregiving can have a major ٠ impact on carers' outcomes.
- Studies of informal carer outcomes use a wide range of endpoints.
- Selecting suitable and appropriate carer outcome measures seems problematic.

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What this paper adds?

- An increasing number of studies are conducted in informal carers looking after patients with palliative care needs.
- Only four outcome measures have been formally developed and validated within this population, and limited psychometric information is available on most measures.
- While there has been an increasing trend since 2008 in the use of outcome measures for informal carers in palliative care
 research, most measures used in these studies were developed more than 20 years earlier and may not adhere to current
 standards for measure development.

Implications for practice, theory or policy

- Existing carer outcome measures need to be validated for the palliative care setting and new measures need to be developed in accordance with current guidelines in order to meet the requirements of the growing number of studies, including intervention studies, of informal carers looking after patients with palliative care needs.
- When using an existing outcome measure, the authors should report their rationale for selecting it and should refer to the publications that report the original development of the measure.
- Interventions for supporting informal carers should be evaluated using outcome measures for which appropriate psychometric properties have been reported before they are implemented as policy.

Introduction

The World Health Organization (WHO)¹ defines palliative care as an approach that focuses on the quality of life of patients and their relatives facing problems associated with life-threatening illness, through prevention and relief of suffering. Annually, around 20 million people worldwide need palliative care,² and an ageing population and increases in long-term conditions mean that need is likely to continue to rise.^{3,4}

Informal carers make an important contribution in the provision of palliative care and are regarded as integral to its delivery.^{5,6} Informal carers are defined as carers who are not financially compensated for their services typically spouses, children, siblings or friends.⁷ In 2011, the contribution of approximately six million informal carers in the United Kingdom was estimated at the equivalent of £119 billion a year.⁸ About half a million people are caring for patients during the end-of-life phase and this number is expected to increase to 3.4 million in the coming 30 years.⁹

Palliative care has become an important component of health care, and policy makers are putting more emphasis on informal carers.¹⁰ Informal caregiving may provide emotional benefits and togetherness for carers,^{11,12} but it also involves considerable challenges including adverse psychological, physical, social and financial consequences.^{13,14} Studies indicate that informal caregiving affects carers' wellbeing and their own health resulting in isolation, fatigue, sleeping problems, exhaustion, weight loss, depression and anxiety.^{15–19} It is therefore important that carer outcomes are assessed in order to be able to provide effective support and to reduce negative consequences of caregiving. Carer outcomes refer to a range of concepts including quality of life, burden and strain. While these terms are not well defined and frequently get

used interchangeably, it is generally accepted that they comprise multiple dimensions such as physical impact, mental strain and social functioning. It has been argued that quality of life is a broader concept as it assesses a wider spectrum of wellbeing, whereas burden and strain suggest a more direct measure of duty of care.²⁰

Evidence on effective strategies to reduce the burden of caring and improve their quality of life of informal carers is limited.^{21,22} Although interventions have been developed that aim to improve outcomes for informal carers, their results are difficult to compare as studies focus on a wide range of endpoints.²³ One systematic review identified 62 questionnaires used among informal family carers in various palliative care settings.²⁴ These questionnaires included instruments on carer satisfaction, experience (of health services and support), needs bereavement and outcomes. Previous reviews on interventions for informal carers concluded that it was unclear what kind of support was beneficial, partly due to the lack of appropriate outcome measures.^{21,25}

In order to assess the impact of the caring role on carers, an appropriate choice of outcome measures is required; however, selecting suitable and appropriate measures seems a common problem.^{25–27} This requires reliable and valid measures with robust psychometric properties, which are appropriate for a palliative care context, as this forms the foundation for evaluating caregiver interventions.

This systematic review aimed to identify and evaluate outcome measures that have been used for informal carers in palliative care studies. The measures used in palliative care are described and their psychometric properties (e.g. reliability, validity, feasibility and precision), when available, are evaluated.

Main search terms	Search terms (PubMed database)
Palliative care	palliative care[Mesh Terms] OR hospice care[Mesh Terms] OR hospices[Mesh Terms] OR palliative*[title/abstract] OR terminal care[title/abstract] OR terminal ill[title/abstract] OR hospice*[title/ abstract] OR end-of-life care[title/abstract] OR end-of-life care[title/abstract] OR end-stage[title/abstract] AND
Caregivers	caregivers[Mesh Terms] OR family[Mesh Terms] OR spouses[Mesh Terms] OR volunteers[Mesh Terms] OR (family[title/abstract] AND (caregiver*[title/abstract] OR care giver*[title/abstract] OR caregiving[title/abstract] OR care giving[title/abstract] OR carer*[title/abstract])) OR (informal[title/ abstract] AND (caregiver*[title/abstract] OR care giver*[title/abstract] OR caregiving[title/abstract] OR care giving[title/abstract] OR carer*[title/abstract])) OR (volunteer*[title/abstract] AND (caregiver*[title/ abstract] OR care giver*[title/abstract] OR caregiving[title/abstract] OR caregiver*[title/ abstract] OR care giver*[title/abstract] OR caregiving[title/abstract] OR care giving[title/abstract] OR carer*[title/abstract])) OR (unpaid[title/abstract] AND (caregiver*[title/abstract] OR carer*[title/abstract])) OR (unpaid[title/abstract] AND (caregiver*[title/abstract] OR care giver*[title/ abstract] OR caregiving[title/abstract] OR care giving[title/abstract] OR carer*[title/abstract])) OR spouse*[title/abstract] OR husband*[title/abstract] OR wife*[title/abstract] OR family[title/abstract] OR volunteer*[title/abstract] OR unpaid[title/abstract] OR informal[title/abstract] AND
Outcomes	quality of life[Mesh Terms] OR quality of life[title/abstract] OR QOL[title/abstract] OR anxiety[title/ abstract] OR benefit*[title/abstract] OR burden[title/abstract] OR competence*[title/abstract] OR coping[title/abstract] OR confidence[title/abstract] OR impact[title/abstract] OR need*[title/abstract] OR preparedness[title/abstract] OR satisfaction[title/abstract] OR self-assurance[title/abstract] OR strain*[title/abstract] OR stress[title/abstract] OR support[title/abstract] OR wellbeing[title/abstract] AND
Questionnaires	questionnaires[Mesh Terms] OR self-report[Mesh Terms] OR outcome assessment (health care) [Mesh Terms] OR psychometrics[Mesh Terms] OR assessment*[title/abstract] OR instrument*[title/ abstract] OR measure*[title/abstract] OR outcome*[title/abstract] OR psychometric*[title/abstract] OR psychometry[title/abstract] OR tool*[title/abstract] OR questionnaire*[title/abstract] OR reliability[title/ abstract] OR reliable [title/abstract] OR reproducibility[title/abstract] OR scale*[title/abstract] OR self- report[title/abstract] OR survey [title/abstract] OR validated[title/abstract] OR validation[title/abstract] OR validity[title/abstract]

 Table I. Search strategy employed in systematic review of studies on psychometric properties of carer-reported outcome measures in palliative care.

Methods

Search strategy

We conducted a systematic review of carer outcome measures used in palliative care, according to Cochrane guidelines.²⁸ The databases, Applied Social Sciences Index and Abstracts (ASSIA), the Cochrane Library, Cumulative Index to Nursing and Allied Health Literature (CINAHL), EMBASE, PubMed, PsycINFO, Social Sciences Citation Index and Sociological Abstracts, were searched using four main terms: palliative care, informal carers, outcomes and measures. The search strategy is presented in Table 1 and further detailed search histories are available from the corresponding author on request. All identified citations were imported into the bibliographic database of EndNote, version X5 (Thomas Reuters, New York, NY). Reference lists of the retrieved articles were screened for additional studies.

Study selection

All types of multidimensional measures (generic, carerspecific for any condition and carer-specific for patients with a specific condition) were eligible for inclusion. The study focused on multidimensional measures as we were interested in measures that assess the overall impact of caring in palliative care rather than measures that assess one specific dimension of outcome or impact. A study was included if all of the following were fulfilled: (1) the study used a self-reported multidimensional measure that assessed caregiver outcomes (i.e. burden, strain or quality of life), (2) measures were directed at unpaid informal carers (e.g. spouse, relatives, siblings, friends or neighbours), (3) the patients they supported were diagnosed with an advanced progressive illness or were receiving palliative care (end-of-life care, terminal care or hospice care), (4) both carers and patients were ≥ 18 years old and (5) the study was reported in English.

A study was excluded if any of the following were fulfilled: (1) only unidimensional measures were used; (2) only subscales or individual items and not the full measure were included; (3) only clinician-assessed measures or patient-reported measures were used; (4) all measures completed by carers were on behalf of the patient or (5) it was a qualitative study, comment, editorial, protocol, conference article or grey literature. There were no restrictions regarding publication date and research methods.



Figure I. PRISMA flow diagram of study selection.

Data extraction and analysis

After retrieving all records, the duplicates were removed. All studies were initially screened on the basis of title and abstract, and then on the basis of full-text. Three authors (C.T.J.M., M.B. and M.P.) independently assessed the eligibility of studies: C.T.J.M. assessed all articles, M.B. and M.P. each assessed half of the articles. Any uncertainties were discussed with the other two authors (A.A. and B.W.) and resolved by consensus. C.T.J.M. extracted the data on study characteristics (publication year, country, sample size, research setting, type of disease, intended outcome measure and information on measure) and psychometric characteristics. The following information on psychometrics was collected: content validity, internal consistency, construct validity, reproducibility (agreement and reliability), responsiveness, floor or ceiling effects, acceptability and feasibility. As guidance, we used the definitions given by Terwee et al.²⁹ and Fitzpatrick et al.³⁰ Additionally, when an included study did not report any psychometric information but referred to other articles regarding a measure or its

psychometric values, we assessed these additional articles in order to evaluate the evidence they provided.

Results

Our electronic search, performed on 4 September 2014, identified 8569 studies. Figure 1 provides an overview of the number of studies identified at each stage of the search. After duplicates were removed, 4505 studies were screened on the basis of titles and abstracts, and 231 studies were screened on the basis of full text. This identified 112 studies using 38 different measures for informal carers in palliative care.

Study and measure characteristics

A total of 112 studies (18 randomized controlled trials (RCTs), 78 observational studies and 16 methodological studies) were included. The methodological studies included translation, development and validation studies about an outcome measure for informal carers in palliative

care. The patient population mainly consisted of cancer patients (n=67, 60%) or a mixture of conditions (n=29, 26%). Of the studies, 37% were conducted in the United States. Most studies included a mix of spouses, children, parents or friends (n=99, 88%) and a small number of studies included only spouse carers (n=4, 3%).

Most studies used only one outcome measure that fit our selection criteria (n=91, 81%) and 19% of the studies administered two outcome measures to carers. Studies mainly used carer-specific measures only (n=69, 62%), a quarter used a generic measure (n=29, 26%), and 14 studies used both types (i.e. generic and carer-specific). In total, 38 measures were identified, including 25 carer-specific measures and 13 generic measures. The main study characteristics are presented in Table 2 and in detail in Supplement 1.

The most frequently used generic measure was the SF-36 (n=16, 14%). The most frequently used carer-specific measures were the Caregiver Reaction Assessment (n=21, 19%), Caregiver Quality of Life Index–Cancer (n=14, 13%) and the Zarit Burden Inventory (n=10, 9%). The primary focus of studies using a carer-specific measure was burden (n=14, 13%), followed by quality of life (n=8, 7%) and strain (n=3, 2.6%). An overview of the identified measures and their frequency of use are presented in Table 3.

Psychometrics of measures

More than half of the 112 (n=60, 54%) studies reported no information on psychometric properties. The 52 (46%) studies that did included 33 observational studies, 15 methodological studies and 4 RCTs. Psychometric data were available for only 23 of the 38 measures including 7 generic measures (i.e. McGill Quality of Life Questionnaire,142 World Health Organization Quality of Life,143 Quality of Life Scale,144 Quality of Life Index,94 SF-36,145 SF-12145 and Swedish Health-Related Quality of life¹⁴⁶) and 17 carer-specific measures. These measures consisted of 4-64 items, with a median of 16 items. Table 4 presents an overview of the 24 measures with the available psychometric information. This consisted mainly of information on the Cronbach's alpha (n=45, 40%), construct validity (n=27, 24%), reliability (n=14, 12%), content validity (n=8, 7%), responsiveness (n=8, 7%) and acceptability and feasibility (n=8, 7%).

Of the 24 measures, four were originally developed in a palliative care context, that is, the Quality of Life in Life-Threatening Illness–Family Carer Version (QOLLTI-F),³⁴ the Family Appraisal of Caregiving Questionnaire for Palliative Care (FACQ-PC),³⁵ the Caregiver Burden Scale in end-of-life-care (CBS-EOLC)³⁶ and the Caregiver Quality of Life Index (CQOLI).⁹⁴ The content validity (which examines the extent to which the concepts of interest are represented by the items²⁰¹), internal consistency

(which measures the extent to which items in a scale are inter correlated²⁹) and construct validity (the extent to which scores relate to other similar measured concepts²⁹) were adequate in all four measures. Interpretability (the degree to which one can assign qualitative meaning to quantitative scores) was not reported in all four studies. The reliability (which concerns the degree to which repeated measurements in stable persons provide similar answers²⁹) was positive in two measures^{34,94} and negative for FACQ-PC.³⁵ Floor and ceiling effects (considered to be present if more than 15% of respondents achieved the lowest or highest possible score, indicating that it is likely that extreme items are missing in the lower or upper ends of the scale²⁰²) was negative for QOLLTI-F³⁴ and not reported for the other three measures.^{35,36,94}

For studies (n=60) that did not report psychometric properties but referred to previous publications about the measure, C.T.J.M. additionally extracted psychometric information from the referenced articles (see Supplement 2). An additional 139 references were assessed for study type, study population and psychometric properties. Although this provided information on how the measures were originally developed, it did not result in additional psychometric information for the measures in the context of carers in a palliative care setting.

Discussion

The aim of this systematic review was to identify and evaluate the psychometric properties of self-reported measures used in informal carers in palliative care studies. A total of 112 studies were found, which used 38 different outcome measures for informal carers. The most commonly used generic measure was the SF-36 (n=27) and the most commonly used carer-specific measure was the Caregiver Reaction Assessment (n=21). Psychometric information was available for only 24 carer outcome measures (52 studies). We identified only four measures that were formally tested in a palliative care context.

Measures were mainly used in descriptive studies (n=78) and the overall study sample sizes tended to be quite small. This could be due to methodological and structural challenges in palliative care research.²⁰³ For example, uncertainties in patients' prognosis, heterogeneity of the palliative care population, relatively small palliative care centres, ethical concerns or attrition of patients during the study could inhibit research in palliative care.

We noted an increasing trend in the use of measures in informal carers in palliative care. The majority of the included studies were published relatively recently, with more than 70% published since 2008. However, the majority of measures were developed much longer ago, including the most frequently used such as the Caregiver Reaction Assessment¹⁶⁵ or the Zarit Burden Interview.¹⁵⁵ It is therefore unclear whether measures adhere to the current

I anie z. suug characteri	series of the inicidated studies $(n - 1)^{\tau}$	c).	
Study characteristic		Number of studies (%)	References
Type of study	Methodological Observational	16 (15%) 78 (70%)	31–46 17,47–123
Country	KC I United States Australia Canada Norway	18 (16%) 37 (33%) 12 (11%) 11 (10%) 8 (7%)	124–141 46,47,51,59–61,65,66,68,70,84,86–88,90,94–96,98,102,105–108,117,120,121,124,125,128,129,131,132,134–136,141 17,35,45,50,56,57,82,83,89,123,133,138 34,36,39,52,53,55,62,63,77,78,139 72–76,103,118,119
Study population	United Kingdom Other (e.g. Brazil, China, Germany, Spain, The Netherlands, Japan, Korea, Sweden and Taiwan) Mixture of informal carers	7 (6%) 37 (33%) 99 (88%)	37,64.80,100,126,137,140 31—33,38,40—44,48,49,54,58,67,69,71,79,81,85,91—93,97,99,101,104,109—116,122,127,130 17,31—37,39—47,49,51—63,65—74,76—81,83—92,94,96—120,122—125,127,129,130,132,134—141
-	(e.g. spouse, child and parent) Not reported Spouse	9 (8%) 4 (3%)	38,64,75,82,95,121,126,128,133 48,50,93,131
Patient population (disease)	Cancer Mixture of various diseases Other (e.g. ALS, ESRD, dementia, heart failure, MND and MS) Not reported	67 (60%) 29 (26%) 10 (9%) 6 (5%)	7,3 ,34,35,38,4 -48,5 ,54,55,59,62,63,68–70,72,73,75–78,80,82,84,86–88,91,92,94,97–104,109 –116,118,119,122,124,129,131–135,138–140 32,37,39,40,52,53,56–58,66,67,71,74,81,83,85,89,90,95,96,105,106,108,120,123,127,128,130,136 33,49,50,64,79,93,107,121,126,137 36,60,61,117,125,141
Sample size study population	N < 50 N = 50 N=101-200 N > 200 Not reported	22 (20%) 30 (27%) 31 (28%) 29 (25%) 1 (1%)	4748,50,51,55–57,59,60,63,65,79,90,95,96,100,107–109,121,125,137 17,31,38,41,45,52–54,62,71–73,75,77,80,84,86–88,92–94,97,106,117,120,122,123,129,138 333,55,39,42,46,58,61,64,66,69,70,76,78,82,85,89,98,99,101,104,110,111,115,118,119,124,127,130,133,135,139 32,34,36,37,40,43,44,49,67,68,74,81,83,91,102,103,105,112–114,116,126,128,131,132,134,136,140,141 126
Type of measure	Generic measure only Carer-specific measure only Both generic and carer-specific measure	29 (26%) 69 (62%) 14 (12%)	37,40,42,55,63,64,67–69,72,73,76,89,90,93,97,101,103,107,109,110,119–121,127,130,132,137,141 17,31,32,34–36,38,39,41,43–48,50–54,56–62,66,70,74,75,78–88,91,92,94–96,98,100,104–106,111–117,122, 124–126,129,131,133,134,138,140 33,49,65,71,77,99,102,108,118,123,128,135,136,139
Number of measures asked in study	One outcome measure Two outcomes measures	91 (81%) 21 (19%)	7,3 ,34 40,42,43,45 48,50–64,66–69,72–74,76,78–95,97,98,100,101,103–107,109–113,115–117,119–122,124– 27,129–134,137,138,141 32,33,41,44,49,65,70,71,75,77,96,99,102,108,114,118,123,128,135,136,139,140
RCT: randomized controlled t	rial; ALS: amyotrophic lateral sclerosis;	ESRD: end-stage	renal disease; MND: motor neurone disease; MS: multiple sclerosis.

Table 2. Study characteristics of the included studies (n = 112).

Measures			Number of studies	References
Generic measures	SF-36		16	33,49,64,68,73,76,77,89,90,99,102,103,119,123,128,139
	SF-12		3	63,65,136
	SF-8*		3	69,97,121
	EORTC QLQ-0	230*	3	37,72,118
	EQ-5D*		3	40,110,137
	QOLS		3	67,72,108
	Other (i.e. MS* WHOQOL and	, MQOL, SWED-QOL, OQOLI*, QOLI, I WHOQOL-BREF*)	9	42,93,101,107–109,127,128,130
Carer-specific measures	Burden	CRA	21	41,47,62,74,75,82,83,99,106,111–116,118,122–124,133,138
		ZBI (including 4 item, 6 item, 8 item, 12 item, 22 item 29 item version)	10	50,53,58,66,77,80,81,91,102,126
		CBS	4	33,49,70,135
		Other (i.e. BASC*, BCOS, BSFC, CBS-EOLC, CBI, CIS, FACS*, HP*, MBCBS, RCAS* and BIC)	15	17,31,32,36,38,41,52,54,55,65,71,79,95,105,135
	Quality of life	CQOLI–Cancer	14	44,46,56,57,59,78,92,96,114,117,125,131,139,140
		CQOLI–Revised	5	60,61,120,136,141
		QOLLTI-F	4	34,39,43,71
		Other (i.e. AQOL-EOL*, CH-QOL-F*, FACT, HQOLI* and QOL–Family*)	8	48,51,70,86,94,98,132,134
	Strain	CSI	7	32,84,87,96,100,129,140
		FACQ-PC	4	35,45,85,88
		FSQ*	I	104

Table 3. Identified outcome measures and frequency of use in the included studies.

SF: short form; EORTC QLQ-C30: European Organization for Research and Treatment of Cancer quality-of-life–30-item questionnaire; EQ-5D: EuroQol–5 dimensions; QOLS: Quality of Life Scale; MS: Montgomery Scale; MQOL: McGill Quality of Life Questionnaire; SWED-QOL: Swedish Health-Related QOL Survey; OQOLI: Overall Quality of Life Index; QOLI: Quality of Life Index; WHOQOL: World Health Organization Quality of Life; WHOQOL-BREF: World Health Organization Quality of Lifebrief form; CRA: Caregiver Reaction Assessment; ZBI: Zarit Burden Inventory; CBS: Caregiver Burden Scale; BASC: brief assessment scale for caregivers; BCOS: Bakas Caregiving Outcomes Scale; BSFC: Burden Scale for Family Caregivers; CBS-EOLC: Caregiver's Burden Scale in end-of-life care; CBI: Caregiver Burden Inventory; CIS: Caregiver Impact Scale; FACS: Feelings about Caregiving Scale; HP: Hausliche Pflegeskala; MBCBS: Montgomery Borgatta Caregiver Burden Scale; RCAS: Revised Caregiving Appraisal Scale; BIC: burden index of caregivers; CQOLI: Caregiver Quality of Life Index; QOLLTI-F: Quality of Life in Life-Threatening Illness–Family Carer Version; AQOL-EOL: Assessment Quality of Iife–Spouses; CH-QOL-F: City of Hope–QOL Scale–Family Version; FACT: Functional Assessment of Cancer Therapy; HQOLI: Hospice Quality of Life Index; QOL: quality of Iife; CSI: Caregiver Strain index; FACQ-PC: Family Appraisal of Caregiving Questionnaire for Palliative Care; FSQ: Family Strain Questionnaire.

*No information reported and available on psychometric properties.

development guidelines, such as those set by the Food and Drug Administration for patient-reported outcome measures.²⁰⁴ Evaluating publications on the development of these outcome measures was beyond the scope of our review, and the information would have been of limited value as the measures were mainly developed in other carer populations.

Due to the wide range of identified carer outcome measures and the variety of versions of the measures (e.g. Zarit Burden Interview; Table 3), it is difficult to draw overall conclusions about psychometric properties. The most commonly reported psychometric information was Cronbach's alpha (n=45, 40%), which is a psychometric property that is commonly used, relatively easy to calculate and easy to interpret. In all, 60 did not report any psychometric information. It was not expected that all studies would contain psychometric information, as the lack of psychometrics was not an exclusion criterion. For studies that did not report psychometric properties but referred to previous publications about the measure, we screened an additional 139 references for information on

psychometrics. However, these resulted in limited extra psychometric data, and none of the studies met the inclusion criteria of this systematic review.

Although psychometric information was generally limited, it was even more limited in relation to some psychometric properties such as responsiveness. Responsiveness (or sensitivity to change) is particularly important to highlight as carer-reported outcome measures may be used to assess the effectiveness of interventions. Interventions to support carers in palliative care settings are likely to be complex and require measures that are able to detect change following the intervention.

We identified only four carer-specific measures that were formally developed and tested in this population: QOLLTI-F,³⁴ FACQ-PC,³⁵ CBS-EOLC³⁶ and CQLI.⁹⁴ These four measures were used less frequently than either the Caregiver Reaction Assessment or the Zarit Burden Interview that have not been validated in this population.

Regarding the generic measures, none have been formally validated in this carer population but we found psychometric information on seven^{94,142–146} measures. As

Table 4	Identified psy	chometric	inform	ation in studie	es identified	from the search (n=52).					
Measure	Study	Type of	No. of	Psychometric	information					Measure	Original validation
		study	Items	Content Validity	Internal consistency	Construct validity	Reliability	Responsiveness	Acceptability and feasibility	reterences cited by study	studies of measure (study population)
BCOS	Buscemi et al ⁵⁴	OBS	15 items	NR	α=0.75	NR	NR	NR	NR	Bakas 1999 ¹⁴⁷ , +Bakas 2000 ¹⁴⁸ , +Bakas 2005 ¹⁴⁹	Bakas 2006 ¹⁵⁰ (15 items, caregivers of stroke survivors)
	Govina et al [:] (Greek version)	³⁸ MES	15 items	R	α =0.83	BCOS-LASA (r=0.7), BCOS-G-HADS (r=-0.52), Criterion validity r=0.57	ICC=0.985 ITC=0.47-0.76	91% sensitivity 86% specificity	10–15 minutes	NR	
BSFC	Brogaard et al ³¹ (Danish version)	MES	28 items	X	α=0.91	Social isolation (ρ =0.33, p=0.01), Dyspnoea (ρ =0.32, p=0.01), Self-reported health (ρ =0.03, p=0.80)	ITC=0.02-0.72	Tendency towards floor effect.	X	Graesel 1995 ¹⁵¹ , *Graesel 1998 ¹⁵² , *\$Graesel 2001 ¹³³ , Hecht 2003 ⁷⁹ , \$Holz 1999 ¹⁵⁴ , #Zarit 1980 ¹⁵⁵	Graesel 1995 ¹⁵¹ (Caregivers of patients with various illnesses: no end-of- life diseases)
BIC	Misawa et al	⁴¹ MES	 items	R	AR	"CRA subscales strongly correlated with supposed subscales of BIC"	х	R	Х	Miyashita 2006 ¹⁵⁶	Miyashita 2006 ¹⁵⁶ (Informal caregivers of patients with neurological rroke)
CBS	Akinci and Pinar ³³ (Turkish version)	MES	22 items	R	α= 0.91	CFA=0.43-0.81 (All CBS factors correlated with each other in positive direction. All sub dimensions scores were negatively correlated with SF-36 (-0.58).	ICC=0.985 ITC=0.37-0.70	R	30 minutes	Elmstahl 1996 ¹⁵⁷	Elementary of a construction of the second s
CBI	Merluzzi et al ⁹⁵	OBS	24 items	R	α= 0.88	Correlations CBI-CGI factors 2, 3 and 4 (p<0.05). Strongest correlations for both PSS-CBI were with factors 3+4 of the CGI.	NR	R	10–15 minutes	Novak 1989 ¹⁵⁸	Novak 1989 ¹⁵⁸ (Caregivers of confused/disoriented elderly)
EOLC EOLC	Dumont et al ³⁶	MES	I 6 items	Focus group + qualitative interviews	α=0.95	Construct validity= Most inter-item associations were consistent with conceptual framework qualitative study. Divergent validity=Interscale- correlations ZBI=0.72 (p<0.01), POMS (fatigue)=0.69 (p<0.01), POMS (vigour)=0 0.27(p<0.05). Explaining overall variance = 6.4.8%	X	Sensitivity showed consistent associations with EGOG and unmet needs.	X	X	Dumont 2008 ³⁶ (Family caregivers of terminal cancer patients)
CIS	Cameron et al ⁵⁵	OBS	I4 items	R	α= 0.87	NR	ĸ	R	X	R	Cameron 2002 ⁵⁵ (Caregivers of advanced cancer patients), Devins 1983 ¹⁵⁹ (ESRD patients)

Table 4.	(Continued)										
Measure	Study	Type of	No. of	Psychometric	information					Measure	Original validation
		study	Items	Content Validity	Internal consistency	Construct validity	Reliability	Responsiveness	Acceptability and feasibility	by study	studies of measure (study population)
cQoLI	McMillan and Mahon ⁹⁴	MES	4 items	Expert panel	α =0.76–0.88	Comparison to control group. Correlation between 4 items (r=0.43-0.64)	No test-retest differences	No significant differences found	Might be too short for reliability	R	Weitzner 1999 ¹⁶⁰ (Caregivers of cancer)
CQOLI-C	Connell et al ⁵⁷	OBS	35 items	NR	α=0.90	ZR	Test-retest =0.95	NR	NR	Weitzner 1999 ¹⁶⁰	Weitzner 1999 ¹⁶⁰ (Caregivers of cancer)
	Delgado- Guay et al ⁵⁹	OBS	35 items	NR	α= 0.9 Ι	NR	Test-retest =0.95	NR	NR	Weitzner 1999 ¹⁶⁰	
	Leow et al ⁹²	OBS	35 items	Expert panel (89%)	α =0.87–0.90	NR	NR	NR	NR	#Axelsson 1998 ⁴⁸ , Weitzner 1999 ¹⁶⁰	
	Meyers and Gray [%]	OBS	35 items	NR	α=0.9I	NR	Test-retest =0.95	NR	NR	Weitzner 1999 ¹⁶⁰	
	Tang et al ¹¹⁴	OBS	35 items	NR	α= 0.9 Ι	Х, К	R	NR	NR	^Edwards 2002 ¹⁶¹ (Refers to study Weitzner 1999 ¹⁶⁰), Weitzner 1999 ⁴⁶	
	Tang et al ⁴⁴ (Mandarin version)	μes	35 items	Translated	α=0. 87	EFA showed 7 underlying factors explaining 48.15% of the variance. Caregivers' QOL was inversely related to both patients' (F=0.90, p=0.008) and caregivers' pain (t=-4.22, p<0.001). Correlation CQOLC-M- MOS-SS scores (r=0.26, $P<(0.01)$. CQOLC-M- SVBS scores (r=0.30, $P<(0.01)$.	щ	Ч	٣	Ч	
	Tang 2009 ¹¹⁷	OBS	35 items	NR	α =0.90	NR	NR	NR	NR	Rhee 2005 ¹⁶² [Korean version], Weitzner 1997 ¹⁶³	
	Weitzner I 999 ⁴⁶	MES	35 items	Expert panel	α= 0.9	Correlations with mental health (r=0.64), emotional distress (r=-0.52), burden (r=-0.65) and patient's performance (r=-0.47), physical health (r=0.13), social support (r=0.03). SF-36-CQOLC correlations social support and social desirability were low range (range=0.08-0.20) Correlated with SF-36, BDI, ECOG, STAI, CBS, MSPSS, and MCSDS.	Test-retest =0.95	Correlation CQOLC- Performance status r=−0.46, p<0.0001	10 minutes	R	
											(Continued)

Table 4. (Continued)										
Measure	Study	Type of	No. of	Psychometri	c information					Measure	Original validation
		fenas	Items	Content Validity	Internal consistency	Construct validity	Reliability	Responsiveness	Acceptability and feasibility	by study	studies of measure (study population)
CQOLI-R	Wittenberg- Lyles et al ¹⁴¹	RCT	4 items	R	α=0.77	NR	Reliability=0.94	NR	R	Courtney 2005 ¹⁶⁴	McMillan 1994 ⁹⁴ (Caregivers of hospice cancer patients)
CRA	Andrews ⁴⁷	OBS	24 items	ĸ	ĸ	Correlated with CES-D scale.	R	NR	Х	Given 1992 ¹⁶⁵ , Given 1993 ¹⁶⁶ , #Radloff 1997 ¹⁶⁷	current products) (Caregivers of persons with physical impairments or dementia)
	Grov et al ⁷²	OBS	24 items	NR	α =0.57–0.85	NR	NR	NR	NR N	Given 1992 ¹⁶⁵ , Grov 2006 ⁷²	
	Hudson and Hayman- White ⁸²	OBS	24 items	х	α=0.76-0.83	The initial rotation indicated that a number of items (3, 5, 7, 10, 16, 18) had high loadings. Items with high loading on more than one component were removed. A third and final Principal Component Analysis (PCA) was performed on the remaining items indicating extraction of 5 components, which accounted for 23.9%, 16.7%, 10.1%, 7.9%, and 7.2% of the total variance.	ĸ	ж	ĸ	Given 1992 ¹⁶⁵ , ^Kinsella 1998 ¹⁶⁸ , Nijboer 1999 ¹⁶⁹	
	Misawa et al ⁴¹ (Japanese version)	MES	24 items	"Face and content validity checked."	α=0.73-0.89	EFA= 5 factors were extracted, and the cumulative proportion of variance explained was 76.6%. Correlations Impact on schedule-time dependent burden (r=0.75), Caregiver's self-esteem-emotional burden (r=0.66), Caregiver's self- esteem-Existential burden (r=0.54), Impact on health- provisial hurden (r=0.75)	щ	No ceiling effect or floor effect.	٣	Ř	
	Morishita and Kamibeppu ⁹⁹	OBS	24 items	NR	α=0.83–0.91	NR	NR	NR	NR	Given 1992 ¹⁶⁵ , Misawa 2009 ⁴¹	
	Stein et al ¹³⁸	RCT	NR	R	α=0.82	NR	NR	NR	NR	Given 1992 ¹⁶⁵	
	Tang ^{III}	OBS	NR	NR	α =0.90	NR	NR	NR	NR	Given 1992 ¹⁶⁵ ,	
	Tang et al ¹¹⁵	OBS	NR	NR	α =0.88	NR	NR	NR	NR	Given 1992 ¹⁶⁵	

Table 4.	(Continued)										
Measure	Study	Type of	No. of	Psychometri	c information					Measure	Original validation
		study	Items	Content Validity	Internal consistency	Construct validity	Reliability	Responsiveness	Acceptability and feasibility	relerences cited by study	studies of measure (study population)
	Tang and Li ¹¹³ (Chinese version)	OBS	24 items	R	α =0.88	NR	ĸ	R	R	Given 1992 ¹⁶⁵ , Tang 2007 ¹¹⁵	
	Tang et al ¹¹⁴	OBS	NR	NR	α =0.68–0.85	NR	NR	NR	NR	Given 1992 ¹⁶⁵	
	Tang et al ¹¹⁶	OBS	24 items	R	α =0.88	NR	NR	NR	NR	Given 1992 ¹⁶⁵	
	Utne et al ^{l I8}	OBS	24 items	R	R	Variance in each domain of CRA was explained by different factors, with total explained variance=[55 % (lack of family support)-31.8% (impact on daily schedule)]	NR	ĸ	ĸ	Fletcher 2009 ¹⁷⁰ , Mazanec 2011 ¹⁷¹ , Nijboer 1999 ¹⁶⁹ , Park 2012 ¹⁷²	
	Yoon et al ¹²² (Korean version)	OBS	24 items	R	α =0.63-0.85	NR	R	NR	R	*Rhee 2009 ¹⁷³ , Grov 2006 ¹⁷⁴	
CSI	Hwang et al ⁸⁴	OBS	N. N.	Ř	α=0. 84	Rotated factor patterns showed that items from PPUN, FIN unmet need subscale, CSI, FAMCARE clearly loaded on different factors and suggested that these scales were measuring separate conceptual constructs. CSI correlated with PPUN (r=0.61, p<0.01), FIN (r=0.39,p=0.01)	Х	Ř	ж	Robinson, 1983 ¹⁷⁵	Robinson, 1983 ¹⁷⁵ (Caregivers for people who have arteriosclerotic heart disease or hip operation/ replacement)
	Meyers and Gray ⁹⁶	OBS	I 3 items	N R	α =0.86	NR	NR	NR	NR	Robinson, 1983 ¹⁷⁵	
	Chan and Suen ³² (Chinese version)	E	I 3 items	Ř	α=0.9I	Factor analysis yielded a single factor as the original M-CSI, which explained 49% of variance. Factor of scale was constructed from item loading of at least 0.59 and was not subjected to any rotation. Higher scores in total score of C-M-CSI were substantially associated with high scores of C-CBI and its subscales.	۳	ž	NR Good range of responses to items (Mean= 0.73–1.26; SD=0.72–0.80)	Robinson, 1983 ¹⁷⁵ , Thornton & Travis 2003 ¹⁷⁶ (Modified CSI)	
FACQ-PC	Cooper et al ³⁵	MES	26 items	Experts	α=0.73-0.86	FA=Presence of 6 initial factors with eigenvalues exceeding 1, explaining 25, 14, 8, 7, 5, and 4% of the variance, respectively. Caregiver strain subscale the strongest correlation with subjective burden.	I TC= Strain [0.41-0.69], Distress [0.28-0.65], Family wellbein, [0.37-0.78]	× Z ₩	X	X	Cooper 2006 ³⁵ (Caregivers of palliative care patients)
											(Continued)

Table 4. (Continued)										
Measure	Study	Type of	No. of	Psychometric	information					Measure	Original validation
		donas	Items	Content Validity	Internal consistency	Construct validity	Reliability	Responsiveness	Acceptability and feasibility	by study	studies of measure (study population)
	Keir et al ⁸⁸	OBS	25 items	NR	R	Correlation with distress r=-0.245, p=0.04	NR	NR	<15 minutes	Cooper 2006 ³⁵	
FACT	Northouse et al ¹³⁴	RCT	R	R	α =0.81 –0.84	NR	NR	NR	NR	Northouse 2002 ¹⁷⁷	Cella 1993 ¹⁷⁸ (cancer patients)
мдог	Sherman et al ¹⁰⁸	OBS	17 items	R	α =0.76-0.73	Correlations between subscales MQOL $r < 0.39$, though psychological symptoms and support are significantly correlated to existential wellbeing ($p < 0.0015$, $p < 0.005$).	R	N	R	Cohen Hassan 1996 ¹⁷⁹ , Cohen Mount 1996 ¹⁴²	Cohen Mount 1966 ¹⁴² (cancer patients)
MBCBS	O'Hara et al ¹³⁵	RCT	14 items	R	α =0.76–0.90	N N N N N N N N N N N N N N N N N N N	NR	NR NR	R	\$Montgomery 2000 ¹⁸⁰	Montgomery 1989 ¹⁸¹ (Caregivers of elderly persons)
QOL	Paiva et al ⁴²	MES	26 items	ž	α=0.80	Correlation HCQ-c -WHOQOL-Bref = Overall quality of life (r=0.688, p <0.01), Physical domain (r=0.415, p<0.01), psychological domain (r=0.570, p <0.01), social domain (r=0.561, p <0.01), environmental domain (r=0.619, p <0.01), and global spirituality (r=0.639, p <0.01).	ž	X	ž	Berlim 2005 ¹⁸² (Brazilian version), Chachamovich 2007 ¹¹⁸³ (WHOQOL- BREF)	WHOOOL Group 1998 ¹⁸⁴ (general population)
QOLLTI-F	Cohen et al ³⁴	MES	items	Expert panel	α= 0.86	Factor loadings, only one regarding amount of control the carer has over his/her life remained problematic. 7 domain scores were created with items that loaded most heavily on each factor. Correlations 7 components r≤0.36. The 16 items predicted 55% of variance in global QOL and 53% in 7 domain scores. QOLLTI-F Total score predicted 43%.	г=0.77–0.8 г=0.77–0.8	All scores significantly different between days, with exception of <i>financial.</i> 2 sub measures limited by ceiling effect.	ž	Ϋ́	Cohen 2006 ³⁴ (Caregivers of palliative cancer patients)
	Schur et al ⁴³ (German version)	MES	16 items	Group of various stakeholders (relatives, clinicians)	α= 0.85	Correlations: QOLLTI-F-HIS (integrative hope scale) r=0.40 (p=0.000), explained variance 16.2%	Test-retest r=0.92	R	All items showed a rate of missing responses below 5%	R	

Table 4.	(Continued)										
Measure	Study	Type of	No. of	Psychometri	ic information					Measure	Original validation
		study	Irems	Content Validity	Internal consistency	Construct validity	Reliability	Responsiveness	Acceptability and feasibility	reierences cited by study	studies of measure (study population)
						QOLT1-F –HADS Stress r=-0.41 (p=0.000), explained variance 16.26%. QOLLT1-F –HADS Anxiety r=-0.51 (p=0.000), explained variances 26%. QOLLT1-F –HADS Depression r=-0.52 (p=0.000), explained variances 26%. QOLLT1-F – Subjective burden r=-0.55 (p=0.000), explained variances 29.1%. FA= problematic cross-loading above 0.32 for items 1, 2, 3, 8, 11, 14 and 16. Scale showed a stable 4-factor structure, instead of initial 7.					
QOLS	Grov et al ⁷²	OBS	l 6 items	NR	α =0.88	NR	NR	R	NR	^Burckhardt 2003 ¹⁸⁵ , Wahl 1998 ¹⁸⁶	Flanagan 1978 ¹⁴⁴ (General population)
GOLI	Scott ¹⁰⁷	OBS	36 items	R	α=0.70	N.R.	Total reliability r=0.92–0.96	NR	NR	Ferrans 1992 ¹⁸⁷	McMillan 1994% (Caregivers of hospice cancer patients)
SF-36	Ringdal et al ¹⁰³	OBS	36 items	ĸ	α=0.70-0.94	Correlations from subscales ranges r=0.50–0.89	ĸ	XX	ĸ	Ware 1992 ¹⁴⁵ , Loge 1998 ¹⁸⁸ , Jenkinson 1997 ¹⁸⁹ , McHorney 1993 ¹⁹⁰ , Jenkinson 1993 ¹⁹¹	Ware 1992 ¹⁴⁵ (General population)
	Weitzner et al ⁴⁶	MES	35 items	Х Х	NR	r= 0.52 to 0.78 for subscales with other QoL measures.	Test-Retest r=0.68–0.93	NR	NR	Ware 1992 ¹⁴⁵ , McHorney 1993 ¹⁹⁰ , McHorney 1994 ¹⁹²	
SF-12	Duggleby et al ⁶³ (SFI2v2)	OBS	12 items	NR	NR	√ SF-36 r=0.95 and r=0.96	NR	R	R	*Ware 1998 ¹⁹³	
SWED- QOL	Persson et al ^{ioi}	OBS	64 items	NR	α =0.68-0.93	R	R	NR	R	#Brown, 2005 ¹⁹⁴ , Eriksson 2005 ¹⁹⁵ , Lindqvist 2000 ⁹³ , *jönsson 1999 ¹⁹⁶	Brorsson 1993 ¹⁴⁶ (General population)
ZBI	Bentley et al ^{s,}	° OBS	NR	NR	α=0.87	ЛЯ	NR	R	NR	Bedard 2001 ¹⁹⁷	Zarit 1980 ¹⁵⁵ (Caregivers of dementia patients)
											(Continued)

Table 4.	(Continued)										
Measure	Study	Type of	No. of	Psychometri	ic information					Measure	Original validation
		Annis		Content Validity	Internal consistency	Construct validity	Reliability	Responsiveness	Acceptability and feasibility	reterences cited by study	(study population)
	Brink et al ⁵³	OBS	12 items	ĸ	α=0.78-0.90	CFA examined factor structure of 12 abridged items. Correlation 12 items-22 items r=0.92-0.97. Mean scores were high 19.89, compared to Bedard 2001 ¹⁹⁷ = 11.20, O'Rourke 2003 ¹⁹⁸ =8.29, Higginson 2010 ⁸¹ =12.0	R	NR	X	X	
	Higginson et al ⁸¹	OBS	22 items	NR	α =0.69–0.93	ZBI- I 2 ρ=0.95-0.97, ZBI-8 ρ=0.86-0.93, ZBI-7 ρ=0.90- 0.95, ZBI-6 ρ=0.89-0.95, ZBI-4 ρ=0.88-0.92, ZBI-1 ρ=0.63- 0.78	AR A	Я	X	Bedard 2001 ¹⁹⁷ (ZBI-4), *Gort 2003 ¹⁹⁹ (ZBI-7), *Arai 2003 ²⁰⁰ (ZBI-8), Bedard 2001 ¹⁹⁷ (ZBI-12)	
	Prigerson et al ¹⁰²	OBS	9 items	NR	NR	Mean caregiver burden score was 8.17 for with MDD versus 5.71 for without MDD (p<0.002).	NR	NR	NR	Zarit 1980 ¹⁵⁵	
#Measure nc BCOS=Baka EOLC=Care Quality of Li ogy Group, I Hospital Am Assessment, of Life Queat Threatening Life, ZBI=ZaI	tu used in study, ⁺ (s. Caregiving Outco: giver's Burden Sca fe Index, CQOLI-(FA=Exploratory f ciety and Depressi diety and Depressi ciety and Depressi filmess-Family Care fillness-Family Care it Burden Invento	Conference a omes Scale, B le in End-or-1 Le in End-or-1 C=Caregiver actor analysis actor analysis on Scale, IHS on Scale, IHS nery Borgatta Multidimensic rr Version, Q	Tricle, ^{\$} Gi 3D1=Beck Life Care, Quality o s, ESRD=f =Integrat 1 Caregive 20LS=Qu	rey literature, ⁴ Depression In CES-D=Cente ff Life Index-Ca End stage renal Erve Hope Scale rive Hope Scale of Perceived S ality of Life Sca	'Non English, ^Revi ventory, BIC=The I r for Epidemiologic ncer, CQOLI-R=C, disease, FA=Factoi e, MCGD-C= Holisti, e, MCSDS=Marlow, le, SF=Short, Form, le, SF=Short Form,	ew, NR=Not Reported, MES=Methoc Burden Index of Caregivers, BSFC=Bu Studies Depression Scale, CFA=Con aregiver Quality of Life Index-Revised Analysis, FACQPC=Family Appraisal - Comfort Questionnaire – Caregiver e-Crowne Social Desirability Scale, MI Performance Status Rating, PSS=Perr STAI= State-Trait Anxiety Inventory,	dological study. O urden Scale for Fa di firmatory factor a d. CRA=Caregiveir d. CC angring Qi i of Caregiving Qi i CC antra-class; i CC antra-class; d. CC antra-cla	BS=Observational study mily Caregivers, CBI= C inalysis, CGI= Clinical G - Reaction Assessment, - sestionnaire for Palliativ - Correlation, ITC=Con sisve disorder, MOS-SS- ss, OQL=Quality of life, vedish Health-Related G	 RCT=Randomize Jaregiver Burden In Jabal Impression, C CSI=Caregiver Stra e Care, FACT=Fun rected Item-Total O medical Outcome @OUI=Quality of I 20L Survey, WHO 	d Controlled Trial. ventory, CBS=Caregiv, CS= Caregiver Impact. In Index, ECOG=Easte ctional Assessment of correlation, LASA= Lin correlation, LASA= Lin s Study- Social Suppor frei Index, QOLLTT-For -QOL=World Health (rr Burden Scale, CBS- scale, CQOLI=Caregiver rn Cooperative Oncol- Cancer Therapy, HADS= ear Analogue Scale er Analogue Scale er MOOL=McGill Quality Duality of Life in Life Drganization Quality of

these have been widely validated in a large number of different populations, it seems reasonable to assume that they are applicable for carers in a palliative context as well.

It is interesting that limited psychometric information was reported for the most widely used carer-specific measure, the Caregiver Reaction Assessment.¹⁶⁵ This suggests that psychometric properties of the measures may not be the key factor in researchers' choice of outcome measures. It would be worthwhile exploring in further studies what considerations researchers take into account when selecting their measures and why some carer-specific measures are used more frequently than others, particularly those developed specifically for carers in a palliative context.

Choosing the right measure for a particular study can be challenging because there may be a number of relevant measures from which to choose.²⁰⁵ A systematic review would be appropriate valuable method to identify the most suitable measure, but it may not always be feasible to conduct a systematic review. Alternatively, as our systematic review highlights, no measure may seem entirely appropriate due to a lack of psychometric information. Additionally, measures may include items irrelevant to the study population, but developing new measures is costly and time consuming. Measure listings such as the Mapi research trust²⁰⁶ and published systematic reviews can assist in selecting an appropriate measure.²⁰⁵ Studies in this review did not always reference the measures used or when a reference was provided, it was frequently not the reference of the development of the measure. We encourage authors to reference the original development paper(s) of the measure(s) used and to justify their choice of instrument.

The findings of this systematic review are in line with previously published reviews. Hudson et al.²⁴ identified 62 tools covering a range of topics including satisfaction, experience, bereavement, needs, preparedness, family functioning and outcomes. Hudson et al.24 identified a larger number of tools than we did as they included instruments, which we specifically excluded. The review concluded that appropriate tools were lacking but the authors only gave a broad critical appraisal across substantially different types of instruments. In 2009, Whalen and Buchholz²⁰⁷ identified 74 caregiver burden screening tools for children or adults providing informal care, not specific to a palliative care context. Whalen and Buchholz²⁰⁷ reported that burden measures might seem appropriate for informal carers but many are lacking psychometric information. Deeken et al.²⁰⁸ searched MEDLINE and PubMed from 1966 to 2002 and identified 28 tools on burden (n=17), needs (n=8) and quality of life (n=3). Neither Whalen and Buchholz nor the Deeken et al. reviews focused on palliative care. In contrast, our systematic review was conducted in a broader range of databases, specifically focused on self-reported multidimensional carer outcome measures in a palliative care context.

A strength of this systematic review is the comprehensive search of eight databases using four main search terms and no date restrictions, which meant we could collate and examine the variety of outcome measures that have been used with informal carers in a palliative care context. This review shows that although there is an increasing number of studies of informal carers in palliative care, most of the outcome measures used have not been formally validated within this carer population.

Another strength of the review is the care that was taken with regard to the inclusion criterion of palliative care. Palliative care is a complex process and involves a broad spectrum of health care services and treatments. Not all palliative care studies are labelled as such but refer to 'hospice care' or 'end-of-life care'. These search terms were included but provided some challenges. For example, endstage renal failure is for some patients a chronic disease but when dialysis or treatment is no longer effective, patients need a palliative approach. Two palliative care experts (A.A. and B.W.) independently assessed each study where there was uncertainty to determine whether or not it was in a palliative care population.

A limitation of the review is the exclusion of the grey literature and literature in languages other than English. It is likely that this meant we missed measures published outside the standard academic field or validation studies of translated measures, which might have provided further psychometric information.

A second limitation is rooted in the limitations of literature itself. Limited psychometric information was available, as more than half of the studies (n=60) did not report any psychometric data. We included all studies that used multidimensional outcome measures in informal carers in palliative care, rather than only development or validation studies, as this corresponded to our study aims. We did not intend to include only development or validation studies, but this may be more appropriate for assessing psychometrics. However, if our inclusion criteria had been limited to development or validation studies alone, only four studies^{34–36,94} would have been identified. Trends regarding the increasing number of publications on carer outcomes in palliative care would have been missed. As most of the studies did not include psychometric information, we could not critically assess the quality of most of the measures.

Conclusion

Support for patients receiving care is likely to continue to be devolved to informal carers. The WHO has called for health care provision to be extended to families, ensuring their needs, coping and outcomes are addressed alongside those of patients receiving health care services at the end of life.²⁰⁹ As more interventions are developed to support carers, carers' outcomes will increasingly be assessed in palliative care context. Although a wide range of measures have already been used in this context, very limited formal psychometric testing has been undertaken. The frequently used measures contain limited psychometric information, while the outcome measures developed or validated in this context are not frequently used in research. Hence, further development and refinement of measures for informal carers in palliative care is required in order to be able to sufficiently support informal carers.

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References

- World Health Organization (WHO). Definition palliative care, http://www.who.int/cancer/palliative/definition/en/ (accessed 21 August 2014).
- World Health Organization (WHO) and World Palliative Care Alliance. *Global atlas of palliative care at the end of life*, http://www.who.int/nmh/Global_Atlas_of_Palliative_ Care.pdf (2014, accessed August 2014).
- 3. Ferlay J, Soerjomataram I, Ervik M, et al. *GLOBOCAN* 2012 v1.0: Cancer incidence and mortality worldwide. IARC CancerBase no. 11. Lyon: International Agency for Research on Cancer, http://globocan.iarc.fr (2013, accessed August 2014).
- 4. Gomes B and Higginson IJ. Where people die (1974–2030): past trends, future projections and implications for care. *Palliative Med* 2008; 22: 33–41.
- 5. Claxton-Oldfield S, MacDonald J and Claxton-Oldfield J. What palliative care volunteers would like to know about the patients they are being asked to support. *Am J Hosp Palliat Care* 2006; 23: 192–196.
- 6. Department of Health. End of life care: promoting high quality care for all adults at the end of life. Report, Department of Health, London, July 2008.
- Vitaliano PP, Zhang J and Scanlan JM. Is caregiving hazardous to one's physical health? A meta-analysis. *Psychol Bull* 2003; 129: 946–972.
- Buckler L and Yeandle S. Valuing Carers 2011: calculating the value of carers' support. Carers UK, London, May 2011,

http://circle.leeds.ac.uk/files/2012/08/110512-circle-carers-uk-valuing-carers.pdf

- National Council for Palliative Care (NCPC). Who cares? Support for carers of people approaching the end of life. London: NCPC, 2012. http://www.ncpc.org.uk/sites/ default/files/Who_Cares_Conference_Report.pdf
- Hudson P and Payne S. Family caregivers and palliative care: current status and agenda for the future. *J Palliat Med* 2011; 14: 864–869.
- Hudson PL, Aranda S and Hayman-White K. A psychoeducational intervention for family caregivers of patients receiving palliative care: a randomized controlled trial. J Pain Symptom Manage 2005; 30: 329–341.
- Stoltz P, Willman A and Uden G. The meaning of support as narrated by family carers who care for a senior relative at home. *Qual Health Res* 2006; 16: 594–610.
- Dillehay RC and Sandys MR. Caregivers for Alzheimer's patients: what we are learning from research. *Int J Aging Hum Dev* 1990; 30: 263–285.
- Payne S. European Association for Palliative Care (EAPC) Task Force on Family Carers White Paper on improving support for family carers in palliative care: part 1. *Eur J Palliat Care* 2010; 17(5): 238–245.
- Angelo J and Egan R. Family caregivers voice their needs: a photovoice study. *Palliat Support Care* 2015; 13: 701–712.
- Barg FK, Pasacreta JV, Nuamah IF, et al. A description of a psychoeducational intervention for family caregivers of cancer patients. *J Fam Nurs* 1998; 4: 394–413.
- Hudson P, Quinn K, Kristjanson L, et al. Evaluation of a psycho-educational group programme for family caregivers in home-based palliative care. *Palliat Med* 2008; 22: 270– 280.
- Pinquart M and Sorensen S. Correlates of physical health of informal caregivers: a meta-analysis. J Gerontol B Psychol Sci Soc Sci 2007; 62: P126–P137.
- Ramirez A, Addington-Hall J and Richards M. ABC of palliative care. The carers. *BMJ* 1998; 316: 208–211.
- 20. Morley D, Dummett S, Peters M, et al. Factors influencing quality of life in caregivers of people with Parkinson's disease and implications for clinical guidelines. *Parkinson's Disease* 2012; 2012: 190901.
- Harding R and Higginson IJ. What is the best way to help caregivers in cancer and palliative care? A systematic literature review of interventions and their effectiveness. *Palliat Med* 2003; 17: 63–74.
- McMillan SC. Interventions to facilitate family caregiving at the end of life. *J Palliat Med* 2005; 8(Suppl. 1): S132– S139.
- 23. Harding R, List S, Epiphaniou E, et al. How can informal caregivers in cancer and palliative care be supported? An updated systematic literature review of interventions and their effectiveness. *Palliat Med* 2012; 26: 7–22.
- Hudson PL, Trauer T, Graham S, et al. A systematic review of instruments related to family caregivers of palliative care patients. *Palliat Med* 2010; 24: 656–668.
- 25. Candy B, Jones L, Drake R, et al. Interventions for supporting informal caregivers of patients in the terminal phase of a disease. *Cochrane Database Syst Rev* 2011; 6: CD007617.
- 26. Field D, Clark D, Corner J, et al. *Researching palliative care*. Buckingham: Open University Press, 2001.

- McGuire DB, Grant M and Park J. Palliative care and end of life: the caregiver. *Nurs Outlook* 2012; 60: 351–356.e20.
- Higgins J and Green S. Cochrane Handbook for Systematic Reviews of Interventions (version 5.1.0). The Cochrane Collaboration, www.cochrane-handbook.org (Updated March 2011) (2011, accessed in August 2014).
- 29. Terwee CB, Bot SD, de Boer MR, et al. Quality criteria were proposed for measurement properties of health status questionnaires. *J Clin Epidemiol* 2007; 60: 34–42.
- Fitzpatrick R, Davey C, Buxton M, et al. Evaluating patientbased outcome measures for use in clinical trials. *Health Technol Assess* 1998; 2(14).
- Brogaard T, Neergaard MA, Guldin MB, et al. Translation, adaptation and data quality of a Danish version of the Burden Scale for Family Caregivers. *Scand J Caring Sci* 2013; 27: 1018–1026.
- 32. Chan CL and Suen M. Validation of the Chinese version of the Modified Caregivers Strain Index among Hong Kong caregivers: an initiative of medical social workers. *Health Soc Work* 2013; 38: 214–221.
- Cil Akinci A and Pinar R. Validity and reliability of Turkish Caregiver Burden Scale among family caregivers of haemodialysis patients. *J Clin Nurs* 2014; 23: 352–360.
- Cohen R, Leis AM, Kuhl D, et al. QOLLTI-F: measuring family carer quality of life. *Palliat Med* 2006; 20: 755–767.
- 35. Cooper B, Kinsella GJ and Picton C. Development and initial validation of a family appraisal of caregiving questionnaire for palliative care. *Psychooncology* 2006; 15: 613–622.
- Dumont S, Fillion L, Gagnon P, et al. A new tool to assess family caregivers' burden during end-of-life care. *J Palliat Care* 2008; 24: 151–161.
- Ewing G, Brundle C, Payne S, et al. The Carer Support Needs Assessment Tool (CSNAT) for use in palliative and end-of-life care at home: a validation study. *J Pain Symptom Manage* 2013; 46: 395–405.
- Govina O, Kotronoulas G, Mystakidou K, et al. Validation of the revised Bakas Caregiving Outcomes Scale in Greek caregivers of patients with advanced cancer receiving palliative radiotherapy. *Support Care Cancer* 2013; 21: 1395– 1404.
- 39. Heyland DK, Jiang X, Day AG, et al. The development and validation of a shorter version of the Canadian Health Care Evaluation Project Questionnaire (CANHELP Lite): a novel tool to measure patient and family satisfaction with end-oflife care. J Pain Symptom Manage 2013; 46: 289–297.
- Krevers B and Milberg A. The sense of security in carerelatives' evaluation instrument: its development and presentation. *J Pain Symptom Manage* 2015; 49: 586–594.
- 41. Misawa T, Miyashita M, Kawa M, et al. Validity and reliability of the Japanese version of the Caregiver Reaction Assessment Scale (CRA-J) for community-dwelling cancer patients. *Am J Hosp Palliat Care* 2009; 26: 334–340.
- 42. Paiva BS, de Carvalho AL, Kolcaba K, et al. Validation of the Holistic Comfort Questionnaire-caregiver in Portuguese-Brazil in a cohort of informal caregivers of palliative care cancer patients. *Support Care Cancer* 2015; 23: 343–351.
- 43. Schur S, Ebert-Vogel A, Amering M, et al. Validation of the 'Quality of Life in Life-Threatening Illness–Family Carer Version' (QOLLTI-F) in German-speaking carers of

advanced cancer patients. *Support Care Cancer* 2014; 22: 2783–2791.

- Tang WR, Tang ST and Kao CY. Psychometric testing of the Caregiver Quality of Life Index-Cancer on a Taiwanese family caregiver sample. *Cancer Nurs* 2009; 32: 220–229.
- 45. Ugalde A, Krishnasamy M and Schofield P. Development of an instrument to measure self-efficacy in caregivers of people with advanced cancer. *Psychooncology* 2013; 22: 1428–1434.
- Weitzner MA, McMillan SC and Jacobsen PB. Family caregiver quality of life: differences between curative and palliative cancer treatment settings. *J Pain Symptom Manage* 1999; 17: 418–428.
- Andrews SC. Caregiver burden and symptom distress in people with cancer receiving hospice care. Oncol Nurs Forum 2001; 28: 1469–1474.
- Axelsson B and Sjoden PO. Quality of life of cancer patients and their spouses in palliative home care. *Palliat Med* 1998; 12: 29–39.
- Belasco A, Barbosa D, Bettencourt AR, et al. Quality of life of family caregivers of elderly patients on hemodialysis and peritoneal dialysis. *Am J Kidney Dis* 2006; 48: 955–963.
- 50. Bentley B, O'Connor M, Breen LJ, et al. Feasibility, acceptability and potential effectiveness of dignity therapy for family carers of people with motor neurone disease. *BMC Palliat Care* 2014; 13: 12.
- Borneman T, Chu DZ, Wagman L, et al. Concerns of family caregivers of patients with cancer facing palliative surgery for advanced malignancies. *Oncol Nurs Forum* 2003; 30: 997–1005.
- 52. Brazil K, Kaasalainen S, Williams A, et al. Comparing the experiences of rural and urban family caregivers of the terminally ill. *Rural Remote Health* 2013; 13: 2250.
- 53. Brink P, Stones M and Smith TF. Confirmatory factor analysis of the burden interview of the caregivers of terminally ill home care clients. *J Palliat Med* 2012; 15: 967–970.
- Buscemi V, Antoni F and Viladrich C. Focus on relationship between the caregivers unmet needs and other caregiving outcomes in cancer palliative care. *Psicooncologia* 2010; 7(1): 109–125.
- 55. Cameron JI, Franche RL, Cheung AM, et al. Lifestyle interference and emotional distress in family caregivers of advanced cancer patients. *Cancer* 2002; 94: 521–527.
- Connell T, Fernandez RS, Tran D, et al. Quality of life of community-based palliative care clients and their caregivers. *Palliat Support Care* 2013; 11: 323–330.
- 57. Connell T, Griffiths R, Fernandez RS, et al. Quality-of-life trajectory of clients and carers referred to a community palliative care service. *Int J Palliat Nurs* 2011; 17: 80–85.
- Costa-Requena G, Cristofol R and Canete J. Caregivers' morbidity in palliative care unit: predicting by gender, age, burden and self-esteem. *Support Care Cancer* 2012; 20: 1465–1470.
- Delgado-Guay MO, Parsons HA, Hui D, et al. Spirituality, religiosity, and spiritual pain among caregivers of patients with advanced cancer. *Am J Hosp Palliat Care* 2013; 30: 455–461.
- Demiris G, Oliver DP, Wittenberg-Lyles E, et al. Use of videophones to deliver a cognitive-behavioural therapy to hospice caregivers. *J Telemed Telecare* 2011; 17: 142–145.

- Demiris G, Parker Oliver D, Wittenberg-Lyles E, et al. A noninferiority trial of a problem-solving intervention for hospice caregivers: in person versus videophone. *J Palliat Med* 2012; 15: 653–660.
- Dudgeon DJ, Knott C, Eichholz M, et al. Palliative Care Integration Project (PCIP) quality improvement strategy evaluation. *J Pain Symptom Manage* 2008; 35: 573–582.
- 63. Duggleby W, Williams A, Holstlander L, et al. Evaluation of the living with hope program for rural women caregivers of persons with advanced cancer. *BMC Palliat Care* 2013; 12: 36.
- 64. Fan SL, Sathick I, McKitty K, et al. Quality of life of caregivers and patients on peritoneal dialysis. *Nephrol Dial Transplant* 2008; 23: 1713–1719.
- 65. Fleming DA, Sheppard VB, Mangan PA, et al. Caregiving at the end of life: perceptions of health care quality and quality of life among patients and caregivers. *J Pain Symptom Manage* 2006; 31: 407–420.
- 66. Fried TR, Bradley EH, O'Leary JR, et al. Unmet desire for caregiver-patient communication and increased caregiver burden. J Am Geriatr Soc 2005; 53: 59–65.
- Friethriksdottir N, Saevarsdottir T, Halfdanardottir SI, et al. Family members of cancer patients: needs, quality of life and symptoms of anxiety and depression. *Acta oncol* 2011; 50: 252–258.
- Garrido MM and Prigerson HG. The end-of-life experience: modifiable predictors of caregivers' bereavement adjustment. *Cancer* 2014; 120: 918–925.
- 69. Gotze H, Brahler E, Gansera L, et al. Psychological distress and quality of life of palliative cancer patients and their caring relatives during home care. *Support Care Cancer* 2014; 22: 2775–2782.
- Grant M, Sun V, Fujinami R, et al. Family caregiver burden, skills preparedness, and quality of life in non-small cell lung cancer. *Oncol Nurs Forum* 2013; 40: 337–346.
- 71. Groh G, Vyhnalek B, Feddersen B, et al. Effectiveness of a specialized outpatient palliative care service as experienced by patients and caregivers. *J Palliat Med* 2013; 16: 848–856.
- 72. Grov EK, Dahl AA, Fossa SD, et al. Global quality of life in primary caregivers of patients with cancer in palliative phase staying at home. *Support Care Cancer* 2006; 14: 943– 951.
- 73. Grov EK, Dahl AA, Moum T, et al. Anxiety, depression, and quality of life in caregivers of patients with cancer in late palliative phase. *Ann Oncol* 2005; 16: 1185–1191.
- 74. Grov EK and Eklund ML. Reactions of primary caregivers of frail older people and people with cancer in the palliative phase living at home. *J Adv Nurs* 2008; 63: 576–585.
- 75. Grov EK, Fossa SD, Sorebo O, et al. Primary caregivers of cancer patients in the palliative phase: a path analysis of variables influencing their burden. *Soc Sci Med* 2006; 63: 2429–2439.
- 76. Grov EK and Valeberg BT. Does the cancer patient's disease stage matter? A comparative study of caregivers' mental health and health related quality of life. *Palliat Support Care* 2012; 10: 189–196.
- 77. Grunfeld E, Coyle D, Whelan T, et al. Family caregiver burden: results of a longitudinal study of breast cancer patients and their principal caregivers. CMAJ 2004; 170: 1795–1801.

- Hannon B, Swami N, Krzyzanowska MK, et al. Satisfaction with oncology care among patients with advanced cancer and their caregivers. *Qual Life Res* 2013; 22: 2341–2349.
- Hecht MJ, Grasel E, Tigges S, et al. Burden of care in amyotrophic lateral sclerosis. *Palliat Med* 2003; 17: 327–333.
- Higginson IJ and Gao W. Caregiver assessment of patients with advanced cancer: concordance with patients, effect of burden and positivity. *Health Qual Life Outcomes* 2008; 6: 42.
- Higginson IJ, Gao W, Jackson D, et al. Short-form Zarit Caregiver Burden Interviews were valid in advanced conditions. *J Clin Epidemiol* 2010; 63: 535–542.
- Hudson PL and Hayman-White K. Measuring the psychosocial characteristics of family caregivers of palliative care patients: psychometric properties of nine self-report instruments. J Pain Symptom Manage 2006; 31: 215–228.
- Hudson PL, Thomas K, Trauer T, et al. Psychological and social profile of family caregivers on commencement of palliative care. *J Pain Symptom Manage* 2011; 41: 522–534.
- 84. Hwang SS, Chang VT, Alejandro Y, et al. Caregiver unmet needs, burden, and satisfaction in symptomatic advanced cancer patients at a Veterans Affairs (VA) medical center. *Palliat Support Care* 2003; 1: 319–329.
- Janssen DJ, Spruit MA, Wouters EF, et al. Family caregiving in advanced chronic organ failure. *J Am Med Dir Assoc* 2012; 13: 394–399.
- Juarez G, Ferrell B, Uman G, et al. Distress and quality of life concerns of family caregivers of patients undergoing palliative surgery. *Cancer Nurs* 2008; 31: 2–10.
- Keefe FJ, Ahles TA, Porter LS, et al. The self-efficacy of family caregivers for helping cancer patients manage pain at end-of-life. *Pain* 2003; 103: 157–162.
- Keir ST, Margaret F, Lipp ES, et al. Family appraisal of caregiving in a brain cancer model. *J Hosp Palliat Nurs* 2009; 11(1): 60–66.
- Kenny PM, Hall JP, Zapart S, et al. Informal care and homebased palliative care: the health-related quality of life of carers. *J Pain Symptom Manage* 2010; 40: 35–48.
- Kilbourn KM, Costenaro A, Madore S, et al. Feasibility of a telephone-based counseling program for informal caregivers of hospice patients. *J Palliat Med* 2011; 14: 1200–1205.
- 91. Kim SY, Kim JM, Kim SW, et al. Determinants of a hopeful attitude among family caregivers in a palliative care setting. *Gen Hosp Psychiatry* 2014; 36: 165–171.
- 92. Leow MQ, Chan MF and Chan SW. Predictors of change in quality of life of family caregivers of patients near the end of life with advanced cancer. *Cancer Nurs* 2014; 37: 391–400.
- 93. Lindqvist R, Carlsson M and Sjoden PO. Coping strategies and health-related quality of life among spouses of continuous ambulatory peritoneal dialysis, haemodialysis, and transplant patients. *J Adv Nurs* 2000; 31: 1398–1408.
- McMillan SC and Mahon M. The impact of hospice services on the quality of life of primary caregivers. *Oncol Nurs Forum* 1994; 21: 1189–1195.
- 95. Merluzzi TV, Philip EJ, Vachon DO, et al. Assessment of self-efficacy for caregiving: the critical role of self-care in caregiver stress and burden. *Palliat Support Care* 2011; 9: 15–24.
- 96. Meyers JL and Gray LN. The relationships between family primary caregiver characteristics and satisfaction with

hospice care, quality of life, and burden. *Oncol Nurs Forum* 2001; 28: 73–82.

- 97. Miyashita M, Misawa T, Abe M, et al. Quality of life, day hospice needs, and satisfaction of community-dwelling patients with advanced cancer and their caregivers in Japan. *J Palliat Med* 2008; 11: 1203–1207.
- Moody LE and McMillan S. Dyspnea and quality of life indicators in hospice patients and their caregivers. *Health Qual Life Outcomes* 2003; 1: 9.
- 99. Morishita M and Kamibeppu K. Quality of life and satisfaction with care among family caregivers of patients with recurrent or metastasized digestive cancer requiring palliative care. *Support Care Cancer* 2014; 22: 2687–2696.
- 100. Payne S, Smith P and Dean S. Identifying the concerns of informal carers in palliative care. *Palliat Med* 1999; 13: 37–44.
- 101. Persson C, Ostlund U, Wennman-Larsen A, et al. Healthrelated quality of life in significant others of patients dying from lung cancer. *Palliat Med* 2008; 22: 239–247.
- 102. Prigerson HG, Cherlin E, Chen JH, et al. The Stressful Caregiving Adult Reactions to Experiences of Dying (SCARED) Scale: a measure for assessing caregiver exposure to distress in terminal care. *Am J Geriatr Psychiatry* 2003; 11: 309–319.
- 103. Ringdal GI, Ringdal K, Jordhoy MS, et al. Health-related quality of life (HRQOL) in family members of cancer victims: results from a longitudinal intervention study in Norway and Sweden. *Palliat Med* 2004; 18: 108–120.
- 104. Rossi Ferrario S, Cardillo V, Vicario F, et al. Advanced cancer at home: caregiving and bereavement. *Palliat Med* 2004; 18: 129–136.
- 105. Salmon JR, Kwak J, Acquaviva KD, et al. Transformative aspects of caregiving at life's end. *J Pain Symptom Manage* 2005; 29: 121–129.
- 106. Sautter JM, Tulsky JA, Johnson KS, et al. Caregiver experience during advanced chronic illness and last year of life. J Am Geriatr Soc 2014; 62: 1082–1090.
- 107. Scott LD. Caregiving and care receiving among a technologically dependent heart failure population. ANS Adv Nurs Sci 2000; 23: 82–97.
- 108. Sherman DW, Ye XY, McSherry C, et al. Quality of life of patients with advanced cancer and acquired immune deficiency syndrome and their family caregivers. *J Palliat Med* 2006; 9: 948–963.
- 109. Smeenk FW, de Witte LP, van Haastregt JC, et al. Transmural care of terminal cancer patients: effects on the quality of life of direct caregivers. *Nurs Res* 1998; 47: 129– 136.
- 110. Song JI, Shin DW, Choi JY, et al. Quality of life and mental health in family caregivers of patients with terminal cancer. *Support Care Cancer* 2011; 19: 1519–1526.
- 111. Tang ST. Predictors of the extent of agreement for quality of life assessments between terminally ill cancer patients and their primary family caregivers in Taiwan. *Qual Life Res* 2006; 15: 391–404; discussion 405–409.
- 112. Tang ST, Cheng CC, Lee KC, et al. Mediating effects of sense of coherence on family caregivers' depressive distress while caring for terminally ill cancer patients. *Cancer Nurs* 2013; 36: E25–E33.
- 113. Tang ST and Li CY. The important role of sense of coherence in relation to depressive symptoms for Taiwanese

family caregivers of cancer patients at the end of life. J Psychosom Res 2008; 64: 195–203.

- 114. Tang ST, Li CY and Chen CC. Trajectory and determinants of the quality of life of family caregivers of terminally ill cancer patients in Taiwan. *Qual Life Res* 2008; 17: 387–395.
- 115. Tang ST, Li CY and Liao YC. Factors associated with depressive distress among Taiwanese family caregivers of cancer patients at the end of life. *Palliat Med* 2007; 21: 249–257.
- 116. Tang ST, Liu TW, Tsai CM, et al. Patient awareness of prognosis, patient-family caregiver congruence on the preferred place of death, and caregiving burden of families contribute to the quality of life for terminally ill cancer patients in Taiwan. *Psychooncology* 2008; 17: 1202–1209.
- 117. Tang WR. Hospice family caregivers' quality of life. J Clin Nurs 2009; 18: 2563–2572.
- 118. Utne I, Miaskowski C, Paul SM, et al. Association between hope and burden reported by family caregivers of patients with advanced cancer. *Support Care Cancer* 2013; 21: 2527–2535.
- 119. Valeberg BT and Grov EK. Symptoms in the cancer patient: of importance for their caregivers' quality of life and mental health? *Eur J Oncol Nurs* 2013; 17: 46–51.
- 120. Wilder H, Parker Oliver D, Demiris G, et al. Informal hospice caregiving: the toll on quality of life. J Soc Work End Life Palliat Care 2008; 4: 312–332.
- Williams MT, Donnelly JP, Holmlund T, et al. Family caregiver needs and quality of life. *Amyotroph Lateral Scler* 2008; 9: 279–286.
- 122. Yoon SJ, Kim JS, Jung JG, et al. Modifiable factors associated with caregiver burden among family caregivers of terminally ill Korean cancer patients. *Support Care Cancer* 2014; 22: 1243–1250.
- 123. Zapart S, Kenny P, Hall J, et al. Home-based palliative care in Sydney, Australia: the carer's perspective on the provision of informal care. *Health Soc Care Community* 2007; 15: 97–107.
- 124. Bowman KF, Rose JH, Radziewicz RM, et al. Family caregiver engagement in a coping and communication support intervention tailored to advanced cancer patients and families. *Cancer Nurs* 2009; 32: 73–81.
- 125. Choi YK. The effect of music and progressive muscle relaxation on anxiety, fatigue, and quality of life in family caregivers of hospice patients. *J Music Ther* 2010; 47: 53–69.
- 126. Edmonds P, Hart S, Wei G, et al. Palliative care for people severely affected by multiple sclerosis: evaluation of a novel palliative care service. *Mult Scler* 2010; 16: 627–636.
- 127. Fegg MJ, Brandstatter M, Kogler M, et al. Existential behavioural therapy for informal caregivers of palliative patients: a randomised controlled trial. *Psychooncology* 2013; 22: 2079–2086.
- 128. Hughes SL, Weaver FM, Giobbie-Hurder A, et al. Effectiveness of team-managed home-based primary care: a randomized multicenter trial. *JAMA* 2000; 284: 2877–2885.
- 129. Keefe FJ, Ahles TA, Sutton L, et al. Partner-guided cancer pain management at the end of life: a preliminary study. J Pain Symptom Manage 2005; 29: 263–272.
- 130. Kogler M, Brandstatter M, Borasio GD, et al. Mindfulness in informal caregivers of palliative patients. *Palliat Support Care* 2013; 1–8.

- McMillan SC, Small BJ, Weitzner M, et al. Impact of coping skills intervention with family caregivers of hospice patients with cancer: a randomized clinical trial. *Cancer* 2006; 106: 214–222.
- 132. Meyers FJ, Carducci M, Loscalzo MJ, et al. Effects of a problem-solving intervention (COPE) on quality of life for patients with advanced cancer on clinical trials and their caregivers: simultaneous care educational intervention (SCEI): linking palliation and clinical trials. *J Palliat Med* 2011; 14: 465–473.
- 133. Mitchell GK, Del Mar CB, O'Rourke PK, et al. Do case conferences between general practitioners and specialist palliative care services improve quality of life? A randomised controlled trial (ISRCTN 52269003). *Palliat Med* 2008; 22: 904–912.
- 134. Northouse LL, Mood DW, Schafenacker A, et al. Randomized clinical trial of a brief and extensive dyadic intervention for advanced cancer patients and their family caregivers. *Psychooncology* 2013; 22: 555–563.
- 135. O'Hara RE, Hull JG, Lyons KD, et al. Impact on caregiver burden of a patient-focused palliative care intervention for patients with advanced cancer. *Palliat Support Care* 2010; 8: 395–404.
- 136. Oliver DP, Washington K, Kruse RL, et al. Hospice family members' perceptions of and experiences with end-of-life care in the nursing home. *J Am Med Dir Assoc* 2014; 15: 744–750.
- 137. Sampson EL, Jones L, Thune-Boyle IC, et al. Palliative assessment and advance care planning in severe dementia: an exploratory randomized controlled trial of a complex intervention. *Palliat Med* 2011; 25: 197–209.
- 138. Stein RA, Sharpe L, Bell ML, et al. Randomized controlled trial of a structured intervention to facilitate end-of-life decision making in patients with advanced cancer. *J Clin Oncol* 2013; 31: 3403–3410.
- 139. Wadhwa D, Burman D, Swami N, et al. Quality of life and mental health in caregivers of outpatients with advanced cancer. *Psychooncology* 2013; 22: 403–410.
- 140. Walsh K, Jones L, Tookman A, et al. Reducing emotional distress in people caring for patients receiving specialist palliative care. Randomised trial. *Br J Psychiatry* 2007; 190: 142–147.
- 141. Wittenberg-Lyles E, Kruse RL, Oliver DP, et al. Exploring the collective hospice caregiving experience. *J Palliat Med* 2014; 17: 50–55.
- 142. Cohen SR, Mount BM, Tomas JJ, et al. Existential wellbeing is an important determinant of quality of life. Evidence from the McGill Quality of Life Questionnaire. *Cancer* 1996; 77: 576–586.
- 143. World Health Organization (WHO) Group. The World Health Organization Quality of Life Assessment (WHOQOL): development and general psychometric properties. *Soc Sci Med* 1998; 46: 1569–1585.
- 144. Flanagan JC. A research approach to improving our quality of life. *Am Psychol* 1978; 33: 138–147.
- 145. Ware JE, Jr and Sherbourne CD. The MOS 36-item shortform health survey (SF-36). I. Conceptual framework and item selection. *Med care* 1992; 30: 473–483.
- 146. Brorsson B, Ifver J and Hays RD. The Swedish Health-Related Quality of Life Survey (SWED-QUAL). *Qual Life Res* 1993; 2: 33–45.

- 147. Bakas T and Champion V. Development and psychometric testing of the Bakas Caregiving Outcomes Scale. *Nurs Res* 1999; 48: 250–259.
- 148. Bakas T, Barr J, Croner D, et al. Psychometric analysis of the Bakas Caregiving Outcomes Scale in the lung cancer population. *Paper presentation Midwest Nursing Research Society 24th annual research conference*, Dearborn, MI, 31 March–3 April 2000.
- 149. Bakas T, FC and Williams LS. Psychometric testing of the revised 15-item Bakas Caregiving Outcomes Scale. Poster presentation American Stroke Association thirtieth international stroke conference, New Orleans, LA, 31 January–3 February 2012.
- 150. Bakas T, Champion V, Perkins SM, et al. Psychometric testing of the revised 15-item Bakas Caregiving Outcomes Scale. *Nurs Res* 2006; 55: 346–355.
- 151. Grasel E. Somatic symptoms and caregiving strain among family caregivers of older patients with progressive nursing needs. *Arch Gerontol Geriatr* 1995; 21: 253–266.
- 152. Grasel E. Home care of demented and non-demented patients. II: health and burden of caregivers. *Z Gerontol Geriatr* 1998; 31: 57–62.
- 153. Grasel E. Hausliche-Pflege-Skala zur Erfassung der Belastung bei betreuenden oder pflegenden Personen. 2nd ed. Ebersberg: Vless Verlag, 2001.
- 154. Holz P. Kuren für pflegende Angehörige. Evaluation einer Modellmaßnahme des Projekts "Hilfen für Pflegende", Teil III der Langfassung des Abschlussberichtes [Rehabilitation for family caregivers. Evaluation of a model intervention within the framework of the project "Help for Caregivers", Part III of the unabridged version of the final report]. Baden-Württemberg, Stuttgart: AOK, 1999.
- 155. Zarit SH, Reever KE and Bach-Peterson J. Relatives of the impaired elderly: correlates of feelings of burden. *Gerontologist* 1980; 20: 649–655.
- 156. Miyashita M, Yamaguchi A, Kayama M, et al. Validation of the Burden Index of Caregivers (BIC), a multidimensional short care burden scale from Japan. *Health Qual Life Outcomes* 2006; 4: 52.
- 157. Elmstahl S, Malmberg B and Annerstedt L. Caregiver's burden of patients 3 years after stroke assessed by a novel caregiver burden scale. *Arch Phys Med Rehabil* 1996; 77: 177–182.
- Novak M and Guest C. Application of a multidimensional caregiver burden inventory. *Gerontologist* 1989; 29: 798–803.
- 159. Devins GM, Binik YM, Hutchinson TA, et al. The emotional impact of end-stage renal disease: importance of patients' perception of intrusiveness and control. *Int J Psychiatry Med* 1983; 13: 327–343.
- 160. Weitzner MA, Jacobsen PB, Wagner H, Jr, et al. The Caregiver Quality of Life Index-Cancer (CQOLC) scale: development and validation of an instrument to measure quality of life of the family caregiver of patients with cancer. *Qual Life Res* 1999; 8: 55–63.
- 161. Edwards B and Ung L. Quality of life instruments for caregivers of patients with cancer: a review of their psychometric properties. *Cancer Nurs* 2002; 25: 342–349.
- 162. Rhee YS, Shin DO, Lee KM, et al. Korean version of the caregiver quality of life index-cancer (CQOLC-K). *Qual Life Res* 2005; 14: 899–904.

- 163. Weitzner MA, Meyers CA, Steinbruecker S, et al. Developing a care giver quality-of-life instrument. Preliminary steps. *Cancer Pract* 1997; 5: 25–31.
- 164. Courtney K, Demiris G, Oliver DP, et al. Conversion of the Caregiver Quality of Life Index to an interview instrument. *Eur J Cancer Care* 2005; 14: 463–464.
- 165. Given CW, Given B, Stommel M, et al. The caregiver reaction assessment (CRA) for caregivers to persons with chronic physical and mental impairments. *Res Nurs Health* 1992; 15: 271–283.
- 166. Given CW, Stommel M, Given B, et al. The influence of cancer patients' symptoms and functional states on patients' depression and family caregivers' reaction and depression. *Health Psychol* 1993; 12: 277–285.
- 167. Radloff LS. The CES-D Scale: a self-report depression scale for research in the general population. *Appl Psychol Meas* 1977; 1: 385–401.
- 168. Kinsella G, Cooper B, Picton C, et al. A review of the measurement of caregiver and family burden in palliative care. J Palliat Care 1998; 14: 37–45.
- 169. Nijboer C, Triemstra M, Tempelaar R, et al. Measuring both negative and positive reactions to giving care to cancer patients: psychometric qualities of the Caregiver Reaction Assessment (CRA). Soc Sci Med 1999; 48: 1259–1269.
- 170. Fletcher BA, Schumacher KL, Dodd M, et al. Trajectories of fatigue in family caregivers of patients undergoing radiation therapy for prostate cancer. *Res Nurs Health* 2009; 32: 125–139.
- 171. Mazanec SR, Daly BJ, Douglas SL, et al. Work productivity and health of informal caregivers of persons with advanced cancer. *Res Nurs Health* 2011; 34: 483–495.
- 172. Park CH, Shin DW, Choi JY, et al. Determinants of the burden and positivity of family caregivers of terminally ill cancer patients in Korea. *Psychooncology* 2012; 21: 282–290.
- 173. Rhee YS, Han IY, Lee IJ, et al. Cross cultural validation of the caregiver reaction assessment (CRA) for cancer patient's caregiver. *Korean J Health Promot Dis Prev* 2009; 9: 189–198.
- 174. Grov EK, Fossa SD, Tonnessen A, et al. The caregiver reaction assessment: psychometrics, and temporal stability in primary caregivers of Norwegian cancer patients in late palliative phase. *Psychooncology* 2006; 15: 517–527.
- 175. Robinson BC. Validation of a Caregiver Strain Index. J Gerontol 1983; 38: 344–348.
- 176. Thornton M and Travis SS. Analysis of the reliability of the modified caregiver strain index. J Gerontol B Psychol Sci Soc Sci 2003; 58: S127–S132.
- 177. Northouse LL, Mood D, Kershaw T, et al. Quality of life of women with recurrent breast cancer and their family members. *J Clin Oncol* 2002; 20: 4050–4064.
- 178. Cella DF, Tulsky DS, Gray G, et al. The Functional Assessment of Cancer Therapy scale: development and validation of the general measure. *J Clin Oncol* 1993; 11: 570–579.
- 179. Cohen SR, Hassan SA, Lapointe BJ, et al. Quality of life in HIV disease as measured by the McGill quality of life questionnaire. *AIDS* 1996; 10: 1421–1427.
- 180. Montgomery RJV, Borgatta EF and Borgatta ML. Societal and family change in the burden of care. In: Liu WT and

Kendig H (eds) *Who should care for the elderly*? Singapore: Singapore University Press, 2000, pp. 27–54.

- 181. Montgomery RJ and Borgatta EF. The effects of alternative support strategies on family caregiving. *Gerontologist* 1989; 29: 457–464.
- 182. Berlim MT, Pavanello DP, Caldieraro MA, et al. Reliability and validity of the WHOQOL BREF in a sample of Brazilian outpatients with major depression. *Qual Life Res* 2005; 14: 561–564.
- 183. Chachamovich E, Trentini C and Fleck MP. Assessment of the psychometric performance of the WHOQOL-BREF instrument in a sample of Brazilian older adults. *Int Psychogeriatr* 2007; 19: 635–646.
- 184. WHOQOL Group. The World Health Organization Quality of Life Assessment (WHOQOL): development and general psychometric properties. *Soc Sci Med* 1998; 46: 1569–1585.
- 185. Burckhardt CS and Anderson KL. The Quality of Life Scale (QOLS): reliability, validity, and utilization. *Health Qual Life Outcomes* 2003; 1: 60.
- 186. Wahl A, Burckhardt C, Wiklund I, et al. The Norwegian version of the Quality of Life Scale (QOLS-N). A validation and reliability study in patients suffering from psoriasis. *Scand J Caring Sci* 1998; 12: 215–222.
- 187. Ferrans CE and Powers MJ. Psychometric assessment of the Quality of Life Index. *Res Nurs Health* 1992; 15: 29–38.
- 188. Loge JH and Kaasa S. Short form 36 (SF-36) health survey: normative data from the general Norwegian population. *Scand J Soc Med* 1998; 26: 250–258.
- 189. Jenkinson C, Layte R and Lawrence K. Development and testing of the Medical Outcomes Study 36-Item Short Form Health Survey summary scale scores in the United Kingdom. Results from a large-scale survey and a clinical trial. *Med Care* 1997; 35: 410–416.
- 190. McHorney CA, Ware JE, Jr and Raczek AE. The MOS 36-Item Short-Form Health Survey (SF-36): II. Psychometric and clinical tests of validity in measuring physical and mental health constructs. *Med Care* 1993; 31: 247–263.
- Jenkinson C, Coulter A and Wright L. Short form 36 (SF36) health survey questionnaire: normative data for adults of working age. *BMJ* 1993; 306: 1437–1440.
- 192. McHorney CA, Ware JE, Jr, Lu JF, et al. The MOS 36-item Short-Form Health Survey (SF-36): III. Tests of data quality, scaling assumptions, and reliability across diverse patient groups. *Med Care* 1994; 32: 40–66.
- 193. Ware JE, Jr and Gandek B. Overview of the SF-36 Health Survey and the International Quality of Life Assessment (IQOLA) Project. J Clin Epidemiol 1998; 51: 903–912.
- 194. Brown DJ, McMillan DC and Milroy R. The correlation between fatigue, physical function, the systemic inflammatory response, and psychological distress in patients with advanced lung cancer. *Cancer* 2005; 103: 377–382.
- 195. Eriksson LE, Bratt GA, Sandstrom E, et al. The two-year impact of first generation protease inhibitor based antiret-roviral therapy (PI-ART) on health-related quality of life. *Health Qual Life Outcomes* 2005; 3: 32.
- 196. Jönsson H, Bergh C, Nielsen M, et al. Good quality of life after heart surgery. Comparable ratings by patients and their relatives. *Lakartidningen* 1999; 96: 5233–5236.

- 197. Bedard M, Molloy DW, Squire L, et al. The Zarit Burden Interview: a new short version and screening version. *Gerontologist* 2001; 41: 652–657.
- 198. O'Rourke N and Tuokko HA. Psychometric properties of an abridged version of The Zarit Burden Interview within a representative Canadian caregiver sample. *Gerontologist* 2003; 43: 121–127.
- 199. Gort AM, Mazarico S, Balleste J, et al. Use of Zarit scale for assessment of caregiver burden in palliative care. *Med Clin* 2003; 121: 132–133.
- 200. Arai Y, Tamiya N and Yano E. The short version of the Japanese version of the Zarit Caregiver Burden Interview (J-ZBI_8): its reliability and validity. *Nihon Ronen Igakkai Zasshi* 2003; 40: 497–503.
- 201. Guyatt GH, Feeny DH and Patrick DL. Measuring health-related quality of life. *Ann Intern Med* 1993; 118: 622–629.
- 202. McHorney CA and Tarlov AR. Individual-patient monitoring in clinical practice: are available health status surveys adequate? *Qual Life Res* 1995; 4: 293–307.
- 203. Kaasa S, Hjermstad MJ and Loge JH. Methodological and structural challenges in palliative care research: how

have we fared in the last decades? *Palliat Med* 2006; 20: 727–734.

- 204. US Department of Health and Human Sciences Food and Drug Administration (FDA). *Guidance to industry patient reported outcome measures: use in medical product development to support labeling claims*. Silver Spring, MD: Food and Drug Administration, 2009.
- 205. Dawson J, Doll H, Fitzpatrick R, et al. The routine use of patient reported outcome measures in healthcare settings. *BMJ* 2010; 340: c186.
- 206. Mapi Research Trust. Patient reported outcome and quality of life instruments database. 2009.
- 207. Whalen KJ and Buchholz SW. The reliability, validity and feasibility of tools used to screen for caregiver burden: a systematic review. *Joanna Briggs Inst Libr Syst Rev* 2009; 7: 1372–1429.
- 208. Deeken JF, Taylor KL, Mangan P, et al. Care for the caregivers: a review of self-report instruments developed to measure the burden, needs, and quality of life of informal caregivers. J Pain Symptom Manage 2003; 26: 922–953.
- 209. World Health Organization (WHO). *National cancer control programmes: policies and managerial guidelines*. 2nd ed. Geneva: WHO, 2002.