

# **RESEARCH ARTICLE**

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# Are we missing the Institute of Medicine's mark? A systematic review of patient-reported outcome measures assessing quality of patient-centred cancer care

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#### **Abstract**

**Background:** The Institute of Medicine (IOM) has endorsed six dimensions of patient-centredness as crucial to providing quality healthcare. These dimensions outline that care must be: 1) respectful to patients' values, preferences, and expressed needs; 2) coordinated and integrated; 3) provide information, communication, and education; 4) ensure physical comfort; 5) provide emotional support—relieving fear and anxiety; and 6) involve family and friends. However, whether patient-reported outcome measures (PROMs) comprehensively cover these dimensions remains unexplored. This systematic review examined whether PROMs designed to assess the quality of patient-centred cancer care addressed all six IOM dimensions of patient-centred care and the psychometric properties of these measures.

**Methods:** Medline, PsycINFO, Current Contents, Embase, CINAHL and Scopus were searched to retrieve published studies describing the development and psychometric properties of PROMs assessing the quality of patient-centred cancer care. Two authors determined if eligible PROMs included the six IOM dimensions of patient-centred care and evaluated the adequacy of psychometric properties based on recommended criteria for internal consistency, test-retest reliability, face/content validity, construct validity and cross-cultural adaptation.

**Results:** Across all 21 PROMs, the most commonly included IOM dimension of patient-centred care was "information, communication and education" (19 measures). In contrast, only five measures assessed the "involvement of family and friends." Two measures included one IOM-endorsed patient-centred care dimension, two measures had two dimensions, seven measures had three dimensions, five measures had four dimensions, and four measures had five dimensions. One measure, the Indicators (Non-small Cell Lung Cancer), covered all six IOM dimensions of patient-centred care, but had adequate face/content validity only. Eighteen measures met the recommended adequacy criteria for construct validity, 15 for face/content validity, seven for internal consistency, three for cross-cultural adaptation and no measure for test-retest reliability.

**Conclusions:** There are no psychometrically rigorous PROMs developed with cancer patients that capture all six IOM dimensions of patient-centred care. Using more than one measure or expanding existing measures to cover all six patient-centred care dimensions could improve assessment and delivery of patient-centred care. Construction of new comprehensive measures with acceptable psychometric properties that can be used with the general cancer population may also be warranted.

**Keywords:** Patient-centred, Quality of care, Patient-reported outcome measures, Cancer, Reliability, Validity, Systematic review

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# **Background**

The Institute of Medicine has defined high quality health care as the provision of appropriate services in a technically competent manner, and includes good communication, shared decision-making and is consistent with patient values and preferences [1]. Optimizing the structure (e.g., hospital resources, number of staff), processes (e.g., interactions between health care providers and patients, use of effective therapies) and outcomes (e.g., survival, quality of life) of health care services are crucial to achieving high quality care [1]. In 2001, the IOM published "Crossing the Quality Chasm" a broad framework which recommended improvements to the following six areas of healthcare in order to achieve high quality care: safety; effectiveness; timeliness; efficiency; equity; and patient-centredness [1]. Within the area of patientcentredness, the IOM also endorsed Gerteis et al's six dimensions of patient-centred care [2] which state that care must be: 1) respectful to patients' values, preferences, and expressed needs; 2) coordinated and integrated; 3) provide information, communication, and education; 4) ensure physical comfort; 5) provide emotional support-relieving fear and anxiety; and 6) involve family and friends [1]. The IOM's recognition of patient-centredness as an indicator of quality acknowledges the adoption of a whole-person orientation to healthcare that goes beyond solely focusing on treatment of the disease.

A variety of sources have been used to assess the quality of care that patients receive including administrative databases, cancer registries, medical records, patient self-reported measures, physician surveys, and pharmacy and laboratory data [3]. However unlike other aspects of quality, such as efficiency, patient self-report is arguably the only way to assess constructs that relate to patientcentredness. For instance, the severity of cancer pain and levels of fatigue experienced by a patient can only reliably be assessed by the patient themselves, and self-report is widely recognised as the gold standard for such assessments [4]. The value of obtaining patient self-report data is further demonstrated by research reporting that patients' perceptions of quality of health care have been associated with important medical and psychological outcomes, including quality of life [5-8], anxiety and depression [6-9]. Patients' perceptions of quality of care have also been associated with factors that directly affect the effectiveness and efficiency of health care such as the under-utilisation of treatments [10-12] and mistrust of the medical system [13,14].

Patient-reported outcome measures (PROMs) that have been designed to assess the quality of patient-centred care include measures of: 1) *satisfaction with care*; and 2) *experiences of care*. Satisfaction with care measures investigate the extent to which an individual's

health care experiences met his/her expectations [15]. However, a range of factors unrelated to the actual health care that was delivered, such as differences among patients' expectation levels, can cause variability in satisfaction ratings, which reduce their reliability for widespread and ongoing monitoring of attempts to improve patient-centred care [15]. In contrast, experiences of care measures ask patients to indicate what actually happened during the process of care delivery, and so are less influenced by subjective patient expectations and provide more detailed information to health care providers and systems about where quality improvements are needed [16,17]. However, in order to accurately reflect the quality of care received and identify variations in patients' experiences, PROMs should meet recommended psychometric criteria for reliability (internal consistency, test re-test reliability), and validity (face, content, construct validity) [18-24].

There are few existing reviews that have assessed the psychometric properties of measures developed to identify patients' experiences of care across a range of settings and diseases [25-28]. Only one of these reviews evaluated the psychometric properties of quality of care measures designed specifically for cancer patients, but focused on satisfaction measures [27]. Further, this review [27] did not investigate the degree to which these quality of care measures assessed the six IOM-endorsed dimensions of patient-centred care [1].

This systematic review identified:

- the degree to which PROMs developed to assess the quality of patient-centred cancer care since the publication of the IOM's "Crossing the Quality Chasm" report in 2001 have addressed the IOM's six endorsed dimensions of patient-centred care [1]; and
- 2) the psychometric properties of these measures.

#### **Methods**

#### Search strategy and selection criteria

The electronic databases Medline, PsycINFO, Current Contents, Embase, CINAHL and Scopus were searched to retrieve published studies outlining the development of PROMs designed to assess the quality of patient-centred cancer care. Given the IOM's Crossing the Quality Chasm report was published in 2001 [1], databases were searched between January 2001 and December 2011 inclusive. The following combinations of keywords were used: (patient-centred or patient-centered or quality of care or satisfaction or experience\*) AND (questionnaire\* or survey\* or instrument\* or measure\* or scale\* or tool\*) AND (cancer\* or neoplasm\* or oncol\*). The use of an \* in the keywords allows words that contain that term to be captured in the literature search. For example the keyword measure\* will identify articles that contain variations of that word such

as measure, measures, measurement and measurements. The reference lists of retrieved articles were also checked to identify any additional relevant publications.

The inclusion criteria for this systematic review were studies that:

- (i) reported the development and psychometric properties (reliability and validity) of new PROMs designed to assess the quality of patient-centred cancer care, or reported the validation of an existing measure for use with a new population (e.g. patient-centred care measure translated for use with a Spanish cancer patient population). Given the IOM's recommendations were published in 2001 [1], studies describing the validation of an existing measure were eligible only if the original PROM was developed from 2001 onwards.
- (ii) described PROMs specifically developed for use with adult cancer patient populations (i.e., aged 18 years or older); and
- (iii) were published in an English language peerreviewed journal.

Publications were excluded if they:

- (i) were reviews, editorials, commentaries or protocol papers;
- (ii) reported qualitative research or used a Delphi consensus process;
- (iii) reported data from medical records, administrative databases or cancer registries (i.e., patients were not surveyed);
- (iv) focussed on cancer screening only;
- (v) predominately surveyed cancer patients under 18 years of age;
- (vi) assessed the views of health professionals such as oncologists, nurses, and general practitioners;
- (vii) examined the perceptions of relatives and/or caregivers;
- (viii) included only cancer patients with advanced cancer or those receiving end of life care; These patients were excluded because the outcome measures and care delivered to patients with advanced cancer can be unique, reflecting the specific goals of advanced disease and/or end-of-life care [29].
- (ix) reported only patient ratings of quality of care and/ or patient characteristics associated with quality of care – i.e. did not develop a measure with the aim of testing its psychometric properties; and
- (x) validation of an existing measure that was not eligible for the review (e.g. the original PROM was developed prior to 2001). PROMs developed prior to 2001 were excluded because it would have been

unreasonable to assess the degree to which such PROMs addressed the IOM's dimensions of patient-centred care given the IOM recommendations were published in 2001 [1].

#### Study and sample characteristics

The study and sample characteristics extracted from eligible publications included: the name of the measure; country of development; patient recruitment setting (e.g. hospital, cancer registry); patient eligibility criteria; sample size; consent rate; participants' socio-demographic characteristics (e.g. mean age, gender, level of education, employment status); and participants' disease and treatment characteristics (e.g. cancer type, cancer stage and/ or time since diagnosis, treatments received).

#### Items and subscales of measures

Information extracted about the characteristics of each measure included: the type of measure (i.e. satisfaction versus experiences); number of items; the type of response scale, and the names and number of subscales. Two coders (FT & SKR) independently examined each of the PROMs' items to determine whether or not the PROM contained content that related to any of the IOM's six patient-centred dimensions and how many of the six IOM-endorsed dimensions of patient-centred care were covered [1]. At least one item in the PROM needed to examine issues related to a particular IOM patient-centred care dimension (as defined below) for that area to be categorised as addressed. A conservative approach was taken when deciding whether or not a measure covered a particular dimension. For example, if a measure included an item that examined whether a patient was provided with information on long-term side effects, the measure was categorised as meeting the information and communication dimension, but not the physical comfort dimension. The physical comfort dimension was classified as present only if items assessed the provision of pain relief or the management of physical symptoms. The criteria used to classify each patientcentred care dimension, which are based on the definitions outlined in the IOM's "Crossing the Quality Chasm" report [1], are described below. Only one aspect of the dimension was needed for the PROM to be classed as covering that patient-centred care dimension.

# 1) Respect for patients' values, preferences, and expressed needs

PROMs were classified as covering this dimension if they assessed: a) whether care responded to the patient's cultural and other values, preferences and needs; b) whether patients were given the opportunity to express their views; c) whether patients were treated with respect during care; and/or d) whether patients were

informed and involved in decision making according to their preferences [1].

# 2) Coordinated and integrated care

PROMs were rated as containing this dimension if they asked: a) whether patient care was coordinated and integrated; b) whether there was timely transfer of up-to-date patient information between healthcare professionals; and/or c) whether patient transitions from one healthcare setting to another went smoothly [1].

#### 3) Provide information, communication, and education

PROMs met the criteria for this dimension if they examined whether health care professionals: a) communicated with patients in a way they could understand; and/or b) provided accurate information regarding care including diagnosis, prognosis, treatment options, follow-up care and support services, according to the patient's preferred level of information provision [1].

#### 4) Physical comfort

PROMs were classified as covering this dimension if they asked patients whether health care professionals: a) promptly provided pain relief; and/or b) attended to the patient's physical symptoms and needs [1].

# 5) Emotional support

PROMs were categorised as meeting this dimension if they assessed whether healthcare professionals: a) addressed the patients' emotional and spiritual concerns, such as anxiety, which could be experienced for a variety of reasons including uncertainty about their disease, concerns about the financial impact of treatment, or worrying about the impact of the illness on their family [1].

# 6) Involvement of family and friends

PROMs were considered to have met this dimension if they assessed whether: a) family and friends were involved in the patient's decision making and care according to the patient's preferences; and/or b) whether care was responsive to the concerns of family and friends and recognised their needs [1].

Two coders (FT & SKR) also independently examined which PROMs covered *all* aspects within each of the IOM dimensions. For instance in terms of the physical comfort dimension, PROMs that included items that addressed both of the following criteria were identified: a) promptly provided pain relief; *and* b) attended to the patient's physical symptoms and needs.

# Psychometric properties of measures

The psychometric properties of each measure were assessed against the same criteria used by Clinton-McHarg and colleagues in their review of instruments

designed to measure the psychosocial health of adolescent and young adult cancer survivors [30]. The psychometric criteria are described below.

# Internal consistency

A measure was coded as having acceptable internal consistency if correlations for the total scale and each subscale were calculated [19] and a Cronbach's alpha >0.70 (continuous or dichotomous scales) or Kuder-Richardson 20 (KR-20) >0.70 was reported for the total scale and each sub-scale [18,19].

# Test-retest reliability

Measures were recorded as having adequate test-retest reliability if the instrument had been administered twice to the same sample and: 1) the second administration occurred within 2-14 days of the first administration [20]; and 2) correlations for the total scale, subscales and items were calculated [21] and the agreement between scores achieved a Cohen's kappa co-efficient ( $\kappa$ ) > 0.60 (nominal or ordinal scales) [19] or Pearson correlation coefficient ( $\kappa$ ) > 0.70 (interval scales) [18,19] or intraclass correlation coefficient (ICC) > 0.70 (interval scales) [18,19].

#### Face validity

Measures were considered to have face validity if both those who administered it, and those who completed it, agreed it appeared to measure what it was designed to measure [22].

#### Content validity

A measure was reported to have adequate content validity if the following processes were described: 1) how the items were developed or selected [18,19]; 2) how and by whom the content was assessed [18,19]; and 3) if modifications to the content were needed that the revisions addressed the issues identified [18,19].

# **Construct validity**

Each measure was assessed as having adequate construct validity if any of the following tests were performed: 1) comparison with other existing measures [19] resulting in Pearson correlation coefficients of (r) >0.40 (convergent validity) or (r) <0.30 (divergent validity) [23]; 2) comparison of scores on the measure differ significantly between groups with known differences (discriminative validity) [18]; or 3) factor analysis [19] with Eigenvalues set at >1 [24].

# Cross-cultural adaptation

A measure was considered to have adequate crosscultural adaptation if a conceptually and linguistically equivalent version of the original form confirmed the reliability and validity reflected in the original measure [18].

#### **Coding process**

Two authors (FT & SKR) independently assessed all potentially relevant publications to determine whether they met eligibility for inclusion in the review. There was 84% agreement between the two coders' ratings. Where discrepancies emerged, inconsistent ratings were discussed between the coders until consensus was reached. Both coders also independently extracted information for the Tables from included publications to ensure accuracy. The coders then compared the information extracted and discussed any inconsistencies until agreement was reached.

#### **Results**

# Study eligibility

A total of 671 publications were identified from the electronic database searches and publication reference lists. Of these, 161 publications were reviews, editorials, commentaries or protocol papers, 40 reported qualitative research and 16 used a Delphi consensus process and were excluded. A further 108 papers reported data from medical records, administrative databases or cancer registries and 53 focussed on cancer screening only and were removed. Of the 293 remaining publications, 48 assessed the views of health professionals such as oncologists, nurses, and general practitioners, 44 focussed on the perceptions of relatives or caregivers, one related to cancer patients aged under 18 years, and 37 focused on an advanced cancer population and/or those receiving endof-life care and were excluded. Of the remaining 163 publications that surveyed adult cancer patients, 121 examined the prevalence of features of care and/or characteristics associated with patient experiences and 14 validated an existing measure that was not eligible for the review (e.g. the original PROM was developed prior to 2001). One paper that reported the development of the EORTC OUT-PATSAT35 was published in French and therefore excluded [31]. This left 27 papers that reported the development of an instrument and its psychometric properties with an adult cancer patient population, or reported the psychometric properties of a re-validated measure for use with a new population. In these papers, 21 unique PROMs were described (see Figure 1).

#### **Setting and Sample Characteristics**

Table 1 provides a detailed description of the setting and sample characteristics of the eligible studies [32-55]. Six studies were conducted in the USA [32,35,42-44,47], five in The Netherlands [37,39,40,50,52], three in England [41,49,54], two in France [53,55], and one in Australia

[33], Canada [34], Europe and Asia [38], Germany [46] and Japan [48]. Seventeen studies recruited cancer patients from hospitals or treatment centres [33,34,38-44,46-50,53-55], whereas only one study recruited patients via a population-based cancer registry [32]. The sample sizes in each study ranged from 82 to 2659 cancer patients and the consent rates varied from 43% to 85%. Thirteen studies included more than one cancer type [32-35,38,41,43,44,46-48,52,55].

#### Patient-centred care instruments

The names of the PROMs included in the review are shown in Tables 1, 2, 3, 4 and 5. As shown in Table 2, 15 measures examined patients' experiences of care [32,33,37,39-44,48-50,52,53,55] while 6 measured satisfaction [34,35,38,46,47,54]. The number of items for each measure ranged from 15 to 152, and the number of subscales ranged from 1 to 15. The type of response scales varied across the different instruments. The number of IOM-endorsed patient-centred care dimensions [1] that were included in each measure were as follows: two measures included one dimension [35,54], two measures had two dimensions [42,46], seven measures had three dimensions [34,39,41,47,48,50,55], five measures had four dimensions [32,33,37,49,53], and four measures had five dimensions [38,43,44,52]. Only one measure, the Indicators (Non-small Cell Lung Cancer) measure, covered all six dimensions of patient-centred care [40]. Table 3 summarises the PROMs that addressed each of the IOM-endorsed patient-centred care dimensions.

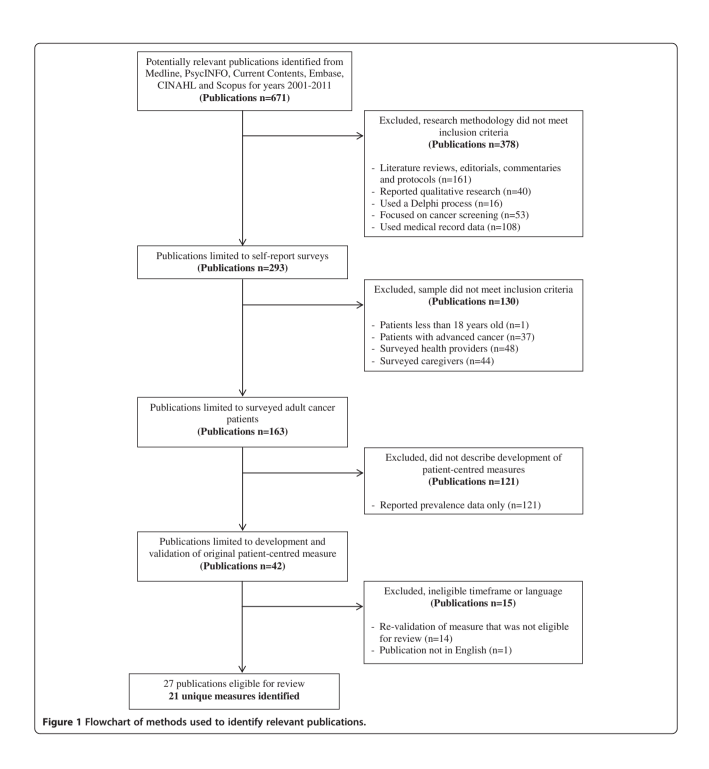
Figure 2 illustrates the frequency with which the six IOM-endorsed patient-centred dimensions were included across the 21 measures. "Information, communication and education" was the dimension most commonly included (19 measures). In contrast, only five measures assessed the "involvement and wellbeing of family and friends". Thirteen measures addressed *all* the IOM criteria for the emotional support dimension [32-34,37-41,43,48,52,53,55], 8 measures for information, communication and education [32,37,47,48,50,52,54,55] and one measure for physical comfort [44]. None of the measures addressed all the IOM criteria within the dimensions of respect for patient values, preferences and needs; coordinated and integrated care; and involvement and wellbeing of family and friends.

# Psychometric properties of instruments

A description of the psychometric properties for each PROM is reported in Table 4.

## Internal consistency

Seven of the 21 measures met the criteria considered adequate for internal consistency by reporting a Cronbach's alpha >0.70 for *both* the total scale and each sub-scale [33,42,43,46-48,55]. Of the 13 studies that reported



Cronbach's alpha only for the PROMs' subscales, six of these measures showed all subscales had a Cronbach's alpha >0.70 [34,35,44,52-54].

# Test-retest reliability

None of the five measures that examined test-retest reliability [33,35,38,49,53] met recommended adequacy criteria of a second administration within 2-14 days of

the first administration [20] and an adequate agreement between the two administrations on scores for the total scale, subscales and items [18,19].

# Face/content validity

Fifteen measures met the criteria considered adequate for face validity and content validity [33,35,37-41, 43,44,48-50,52,53,55].

Table 1 Sample characteristics of studies that have developed PROMs assessing quality of patient-centred cancer care

| Measure                             | Sample<br>size | Consent rate              | Eligibility criteria   | Setting & country  | Socio-demographics                  | Cancer type,<br>stage/ diagnosis                                 | Cancer treatments  |
|-------------------------------------|----------------|---------------------------|--|--|-------------------------------------|--|--|
| experiences of Cancer               | 623            | 69.2% participation rate. | Read English, diagnosed with leukaemia<br>or bladder or colorectal cancer between  | Cancer Prevention<br>Institute of California's   | 43.3% women                         | Colorectal cancer: 59.6%   | 38.7% surgery only   |
| Care (APECC) [32]                   |                | 49.2% response rate.      | June 1999-May 2001 (i.e., 2-5 years before<br>study enrolment), at least 20 years old at<br>diagnosis, have received cancer treatment,<br>have the cancer of interest as their first | cancer registry, USA.  | 37.7% aged<br>50-64 years           | Bladder cancer: 26.2%  | 35.1% surgery plus<br>chemotherapy or<br>radiation                   |
|                                     |                |                           | cancer diagnosis, not have any other cancer<br>between their initial diagnosis and the start   |  | 20.3% college<br>degree             | Leukaemia: 14.3%   |  |
|                                     |                |                           | of the study, have no objections from their physician of record to their participation.  |  | 71.5% married/<br>defacto           |  |  |
|                                     |                |                           |  |  | 73.8% Non-<br>Hispanic white        | 84.4% in remission.  | 12.3% surgery plus<br>chemotherapy and<br>radiation                  |
| Capear Caro Caparlination 694       |                |                           |  |  | 83.6% private<br>health insurance   | Mean of 3.5 years since diagnosis.                               | 13.9% chemotherapy<br>with or without<br>radiation but no<br>surgery |
| ancer Care Coordination             | 686            | =                         | Sample 1 (n = 245): Patients were in follow-up   | Sample 1: Six centres  | Total participants:                 | Colorectal: 82.5%  | 96% surgery  |
| uestionnaire for<br>atients [33]    |                |                           | for any cancer that had been treated between 3 to 12 months previously, had sufficient   | (2 metropolitan<br>& 4 regional).  | 46.8% women                         | Gynaecological: 7.6%   | 40.5% chemotherapy   |
|                                     |                |                           | English and were not cognitively impaired and were not receiving end of life care.   |  | Mean age: 66.1 years                | Breast: 2.6%   | 12.2% radiotherapy   |
|                                     |                |                           | and well indirectiving the of life care.   |  | 66.9% married/defacto               | Lung/mesothelioma:<br>1.3%                                       | 3.9% hormone<br>therapy  |
|                                     |                |                           |  | Sample 2: 22 public<br>and private hospitals<br>in metropolitan and<br>regional centres. | 35.3% tertiary degree<br>or diploma | Other/multiple sites: 4.7%                                       |  |
|                                     |                |                           |  |  | 23.5% employed full-time            | Primary cancer: 91.8%  |  |
|                                     |                |                           |  |  |                                     | Recurrent cancer: 3.8%   |  |
|                                     |                |                           | Sample 2 (n = 441): Patients with a newly diagnosed colorectal cancer undergoing initial surgical treatment.   | Australia  |                                     |  |  |
| ancer Patient Information           | 540            | -                         | -  | Ambulatory setting of  | 53% women                           | Breast: 19.1%  | -  |
| nportance and Satisfaction ool [34] |                |                           |  | regional cancer centre,<br>Canada.   | Mean age: 60.9 years                | Haematological: 12.5%  |  |
|                                     |                |                           |  |  |                                     | Genitourinary: 12.5%   |  |
|                                     |                |                           |  |  |                                     | Skin: 11.5%  |  |
|                                     |                |                           |  |  |                                     | Gastrointestinal: 11.5%  |  |
|                                     |                |                           |  |  |                                     | Head and neck: 11.3%   |  |
|                                     |                |                           |  |  |                                     | Gynecologic: 11.2%   |  |
|                                     |                |                           |  |  |                                     | Lung: 10.4%  |  |
|                                     |                |                           |  |  |                                     | 21.3% diagnosed in last<br>year, 48.3% between<br>2-5 years ago. |  |

Table 1 Sample characteristics of studies that have developed PROMs assessing quality of patient-centred cancer care (Continued)

| Cancer Therapy Satisfaction<br>Questionnaire (CTSQ) [35,36] | 361                                | -     | Provided written informed consent, aged 18 years or older, read and write in English,   | 14 community clinical practices, USA.                           | 63.2% women                         | Breast: 37.9%                       | First-line of therapy: 48.8%  |
|---|------------------------------------|-------|---|---|-------------------------------------|-------------------------------------|-------------------------------|
|   |                                    |       | available for follow-up evaluation, actively receiving more than one cycle of first- or |   | Mean age: 60.7 years                | Colorectal: 33.5%                   |                               |
|   |                                    |       | second-line chemo, biological or hormonal therapy for early or advanced cancer,         |   | 90.9% white                         | Lung: 25.2%                         | Second-line of therapy: 28.0% |
|   |                                    |       | mentally and physically capable of participation.                                       |   | 16.3% college/university            | Melanoma: 3.3%                      |                               |
|   |                                    |       |   |   | degree                              | Stage I: 6.1%                       | Adjuvant: 23.3%               |
|   |                                    |       |   |   |                                     | Stage II: 25.2%                     |                               |
|   |                                    |       |   |   |                                     | Stage III: 24.9%                    |                               |
|   |                                    |       |   |   |                                     | Stage IV: 43.8%                     |                               |
| Consumer Quality Index                                      | 731                                | 63%   | Older than 18 years, having received breast   | Selected from claims  | 99.7% women                         | Breast cancer: 57%                  | -                             |
| Breast Care (CQI-BC) [37]                                   |                                    |       | care in the last 24 months, not being approached in the past for CQI surveys.           | data of four health<br>insurance companies,<br>The Netherlands. | 30% aged between<br>55 and 64 years | Benign breast disorder:<br>38%      |                               |
|   |                                    |       |   |   |                                     | Breast carcinoma in situ: 5%        |                               |
|   |                                    |       |   |   |                                     | -                                   |                               |
| EORTC cancer in-patient satisfaction with care              | sfaction with care<br>asure (EORTC | 84.9% | Diagnosed with cancer, aged 18 years or older, hospitalized for at least three days,    | Surgery or medical<br>oncology wards in<br>hospitals, Belgium,  | 59% women                           | Breast: 35.1%                       | Current or planned treatment: |
| IN-PATSAT32) [38]   |                                    |       | mentally able to complete questionnaire.  | France, Germany,  | Median age: 57 years                | Gastro-intestinal: 17.2%            | 55% surgery                   |
|   |                                    |       |   | Italy, Poland, Spain,<br>Sweden, Taiwan,<br>United Kingdom.     | 21% university educated             | Gynaecologic: 10%                   |                               |
|   |                                    |       |   |   | 73.6% married/defacto               | Head and neck: 7.3%                 | 40.8% chemotherapy            |
|   |                                    |       |   |   | 40.5% full time employed            | Genito-urinary: 7.1%                | 3.2% surgery and chemotherapy |
|   |                                    |       |   |   |                                     | Haematological: 6.5%                |                               |
|   |                                    |       |   |   |                                     | Respiratory: 5.6%                   | 0.9% other                    |
|   |                                    |       |   |   |                                     | Bone: 2.8%                          |                               |
|   |                                    |       |   |   |                                     | Brain: 1.5%                         |                               |
|   |                                    |       |   |   |                                     | Melanoma: 0.8%                      |                               |
|   |                                    |       |   |   |                                     | Other: 6.2%                         |                               |
|   |                                    |       |   |   |                                     | 73.9% local/loco-<br>regional       |                               |
|   |                                    |       |   |   |                                     | 26% metastatic                      |                               |
|   |                                    |       |   |   |                                     | Median of 15 weeks since diagnosis. |                               |
| Indicators (Head & Neck                                     | 158                                | 84%   | Patients with head and neck cancer newly  | Selected from clinic  | 27% women                           | Head & neck cancer.                 | First treatment:              |
| Cancer) [39]  |                                    |       | diagnosed between May to December 2003  | lists at a university<br>hospital, The<br>Netherlands           | Mean age: 62 years                  | Larynx and hypharynx:<br>38%        | Operation: 56%                |
|   |                                    |       |   | . vea renarios  | 23% highly educated                 |                                     | Radiotherapy: 37%             |
|   |                                    |       |   |   |                                     |                                     |                               |

Table 1 Sample characteristics of studies that have developed PROMs assessing quality of patient-centred cancer care (Continued)

|  |              |                |   |  |                                   | Cavity of the mouth: 36%    |                     |
|--|--------------|----------------|---|--|-----------------------------------|-----------------------------|---------------------|
|  |              |                |   |  |                                   | Other: 26%                  | Chemotherapy: 7%    |
|  |              |                |   |  |                                   | =                           |                     |
| Indicators (Non-small<br>Cell Lung Cancer) [40]      | 100          | 76%            | Patients newly diagnosed with non-small cell lung cancer between September 2004                 | 6 hospitals, The Netherlands.                | 34% women                         | Non-small cell lung cancer. | -                   |
|  |              |                | and February 2005.  |  | Mean age: 66 years                | Stage IV: 24%               |                     |
| Medical Care Questionnaire                           | Phase        | Phase 3:       | Adult patients from all tumor groups attending  | A regional hospital,                         | Phase 3:                          | Phase 3:                    | =                   |
| (MCQ) [41]   | 3: 200       | 70%            | the Medical Oncology Unit, could read and<br>understand English, were not exhibiting overt      | England.                                     | 81% women                         | Gynecological: 38%          |                     |
|  | Phase 4: 477 | Phase 4: 79.6% | cognitive dysfunction or signs of distress.   |  | 42% aged 45-59 years              | Breast: 26.5%               |                     |
|  |              |                |   |  | 74% married/de facto              | Genitourinary: 16.5%        |                     |
|  |              |                |   |  | 13% employed full time            | Sarcoma: 5.5%               |                     |
|  |              |                |   |  |                                   | Gastrointestinal: 4.5%      |                     |
|  |              |                |   |  | Phase 4:                          | Melanoma: 1%                |                     |
|  |              |                |   |  | 74.2% women                       | Other: 8%                   |                     |
|  |              |                |   |  | 40.9% aged 45-59 years            |                             |                     |
|  |              |                |   |  | 75.1% married/de facto            | Phase 4:                    |                     |
|  |              |                |   |  | 44.2% employed full time          | Gynecological: 33.8%        |                     |
|  |              |                |   |  |                                   | Breast: 23.5%               |                     |
|  |              |                |   |  |                                   | Genitourinary: 21.4%        |                     |
|  |              |                |   |  |                                   | Melanoma: 8.4%              |                     |
|  |              |                |   |  |                                   | Sarcoma: 7.5%               |                     |
|  |              |                |   |  |                                   | Other: 5.5%                 |                     |
|  |              |                |   |  |                                   | -                           |                     |
| Modified Version of the                              | 87           | 74%            | Females aged 18 years or older, confirmed   | Four hospital-based                          | 100% women                        | Breast cancer.              | 89.7% chemotherapy  |
| Perceived Involvement in<br>Care Scale (M-PICS) [42] |              |                | diagnosis of breast cancer, reported pain of<br>at least moderate intensity (score ≥4 on the    | outpatient oncology<br>clinics and a private | Mean age: 50.4 years              | Stage I: 12.6%              | 43.7% radiation     |
|  |              |                | Brief Pain Inventory's Worst Pain Intensity item) over prior two weeks, absence of any          | hospital-affiliated oncology practice,       | 31% Caucasian                     | Stage II: 24.1%             | therapy             |
|  |              |                | gross cognitive impairment, literate in English   | USA.   | 50.6% married/partnered           | Stage III: 13.8%            |                     |
|  |              |                | or Spanish.   |  | 64.5% at least 13 years education | Stage IV: 49.4%             |                     |
|  |              |                |   |  | 26.2% employed                    |                             |                     |
| Oncology Patients'                                   | 436          | -              | 18 years or older, registered with the receptionist   | A haematology-                               | 66% women                         | Breast: 40%                 | In active treatment |
| Perceptions of the<br>Quality of Nursing Care        |              |                | on the days of data collection, had received cancer nursing care in the clinic or hospital, not | oncology clinic,<br>USA.                     | Mean age: 54.8 years              | Melanoma: 9%                |                     |
| Scale (OPPQNCS) [43]                                 |              |                | extremely ill or confused.  |  | 93% white                         | Lung: 6%                    |                     |
|  |              |                |   |  | 81% more than high                | Renal cell: 4%              |                     |
|  |              |                |   |  | school education                  | Squamous cell: 4%           |                     |

Table 1 Sample characteristics of studies that have developed PROMs assessing quality of patient-centred cancer care (Continued)

|  |      |        |   | _  | _                                   | Prostate: 3%                        | _  |
|--|------|--------|---|--|-------------------------------------|-------------------------------------|--|
|  |      |        |   |  |                                     | Other: <3%                          |  |
|  |      |        |   |  |                                     | -                                   |  |
| Pain Care Quality Survey<br>(PainCQ) [44,45] | 109  | -      | 18+ years of age, inpatients on one of the designated units with an expected stay of  | Hospitals in three geographically  | 58.7% women<br>Mean age: 53.1 years | Leukaemia/lymphoma:<br>15.6%        | Reason for hospitalization:                            |
|  |      |        | more than 24 hours, diagnosis of cancer,<br>surgery for cancer, a suspected cancer<br>diagnosis or a hematological disorder and   | diverse settings<br>with medical or<br>surgical oncology   | 88.1% non-Hispanic white            | Uterine/Cervical/Ovarian:           | Surgery: 40.4%   |
|  |      |        | a positive response to screening regarding<br>the presence of pain, cognitively and<br>physically able to complete survey.  | units, USA.  | 66.1% married/partnered             | Prostate and genitourinary: 11.9%   | Supportive care and management of complications: 45.9% |
|  |      |        | h   |  | 25.7% college graduate              | Colorectal: 10.1%                   | Treatment of cancer:                                   |
|  |      |        |   |  |                                     | Lung: 7.3%                          | 8.3%   |
|  |      |        |   |  |                                     | Gastrointestinal: 7.3%              | Other: 5.5%  |
|  |      |        |   |  |                                     | Breast: 6.4%                        |  |
|  |      |        |   |  |                                     | Other cancers: 22.0%                |  |
|  |      |        |   |  |                                     | Non-cancer diagnosis: 7.3%          |  |
|  |      |        |   |  |                                     | 34.9% local or regional             |  |
|  |      |        |   |  |                                     | 41.3% advanced cancer               |  |
| PASQOC questionnaire [46]                    | 2659 | 78.6%  | Aged 18 years or older, the presence of any cancer suitable for outpatient treatment, read and write in German, mentally and physically able to complete questionnaire. | 24 institutions including<br>15 private group<br>practices, 6 single<br>oncologist-led practices,<br>3 hospital day clinics,<br>Germany. | 56% women                           | Breast: 22.9%                       | 80.9% chemotherapy                                     |
|  |      |        |   |  | Mean age: 61.7 years                | Intestine: 19.8%                    | 58.0% surgery  |
|  |      |        |   |  | s, 8.7% employed full-time          | Lymphoma: 15.2%                     |  |
|  |      |        |   |  |                                     | Haematological: 12.3%               |  |
|  |      |        |   |  |                                     | Other: 29.8%                        |  |
|  |      |        |   |  |                                     | 58.2% distant metastases            |  |
|  |      |        |   |  |                                     | 61.9% diagnosed within last 3 years |  |
| Patient Satisfaction with                    | 891  | =      | Abnormal breast, cervical, colorectal and prostate  | Multiple patient   | 81.3% women                         | Breast: 64.2%                       | =  |
| Cancer Care [47]                             |      |        | cancer test finding or a new diagnosis of these cancers without any prior history of cancer   | Navigation Research<br>Program recruitment   | Mean age: 51.4 years                | Cervix: 10.8%                       |  |
|  |      |        | treatment other than non-melanoma skin  | sites (eg, clinics or  | 43.2% white                         | Colorectal: 12.0%                   |  |
|  |      |        | cancer, fluent in English.  | hospitals), within nine<br>largely racial/ethnic   | 40.4% married/defacto               | Prostate: 12.6%                     |  |
|  |      |        |   | minority and low-  | 12.9% college graduate              | Multiple sites: 0.5%                |  |
|  |      |        |   | income communities,<br>USA.  | 29.9% full-time employed            |                                     |  |
| Perceived Physician's                        | 147  | 47 74% | Patients with cancer who were aware of their cancer diagnosis, met their doctor more than once and did not have a debilitating condition.                               | A hospital, Japan.   | 66.7% women                         | Breast cancer: 50.3%                | -  |
| Communication Style<br>Scale [48]            |      |        |   |  | Mean age: 57.6 years                | Gastric cancer: 30.6%               |  |
| cale [48]                                    |      |        |   |  | 20.4% university educated           | Lung cancer: 15.0%                  |  |

Table 1 Sample characteristics of studies that have developed PROMs assessing quality of patient-centred cancer care (Continued)

|                                  |     |       |   | <u> </u>                                  |  |  |  |
|----------------------------------|-----|-------|---|---|--|--|--|
|                                  |     |       |   |   |  | Other cancer: 4.1%   |  |
|                                  |     |       |   |   |  | 55.1% disease free   |  |
|                                  |     |       |   |   |  | 44.9% recurrent or metastatic disease  |  |
| Prostate Care Questionnaire      | 865 | 69.2% | Patients diagnosed with, or treated for prostate  | 5 hospitals, England                      | 100% male                                | Prostate cancer  | =  |
| or Patients (PCQ-P) [49]         |     |       | cancer within the past two years, who were<br>not too ill to participate.   |   | 40.5% aged 65-74 years                   |  |  |
|                                  |     |       |   |   | 92.8% white                              | -  |  |
|                                  |     |       |   |   | 21.3% employed                           |  |  |
| QUOTE Breast                     | 276 | 43%   | Experience with any type of surgery for breast  | 5 hospitals, The                          | 100% women                               | Breast cancer  | 54% lumpectomy   |
| Cancer [50,51]                   |     |       | cancer 3 -15 months before the start of the study, age older than 17 years and mental competence as judged by the breast nurse. | Netherlands.                              | Mean age: 57 years                       |  | 52% (modified)<br>radical mastectomy                   |
|                                  |     |       |   |   | 32% college/university                   | 16% diagnosed 3-6 months   | 50% radiotherapy                                       |
|                                  |     |       |   |   | educated                                 | ago, 47% 7-12 months<br>ago, 34% 13-18 months<br>ago, 3% more than<br>18 months ago. | 38% chemotherapy                                       |
| QUOTE <sup>chemo</sup> [52] 345  | 345 | 59.3% | 60 most recent patients from each hospital who  | Hospital records                          | 67% women                                | Breast: 47.2%  | 16.2% chemotherapy                                     |
|                                  |     |       | were new to chemotherapy, aged 18 years or older, able to read Dutch.   | from 10 hospitals,<br>The Netherlands.    | Mean age: 55.7 years                     | Digestive-gastrointestinal:  | only   |
|                                  |     |       |   |   | 79% lived with partner                   | 21.5%  | 74.5% chemotherapy                                     |
|                                  |     |       |   |   | 28% highly educated                      | Haematologic: 10.6%<br>Lung: 9.7%  | & surgery  |
|                                  |     |       |   |   | 52.8% employed                           | Gynaecological: 6.2%<br>Urologic: 2.9%   | 45.5% chemotherapy<br>& radiotherapy                   |
|                                  |     |       |   |   |  | Other: 1.8%  | 21.1% chemotherapy<br>& hormone replacement<br>therapy |
|                                  |     |       |   |   |  | Mean of 11.5 months since diagnosis.   | 6.2% chemotherapy<br>& immunotherapy                   |
|                                  |     |       |   |   |  |  | 74.8% curative intent                                  |
|                                  |     |       |   |   |  |  | 25.2% palliative intent                                |
| EPERES-60 [53]                   | 820 | 84%   | A first diagnosis of invasive non-metastatic breast   | Public and private                        | 100% women                               | Breast cancer.   | -  |
|                                  |     |       | cancer, at least two contacts for cancer with one of the health professionals in one of the two                                 | cancer centres in<br>two regions, France. | Mean age: 58 years                       |  |  |
|                                  |     |       | regions between diagnosis and the first year of follow-up, signed informed consent.   | two regions, marice.                      | 55.1% living with spouse/partner         | -  |  |
|                                  |     |       |   |   | 15.7% higher education<br>43.7% employed |  |  |
| atisfaction with Cancer          | 82  | 76%   | Newly diagnosed patients with head and neck   | 4 hospitals, England.                     | 34% women                                | Head and neck cancer.  | Planned treatments:                                    |
| nformation Profile<br>SCIP) [54] |     | . 2/0 | cancér.   | -   |  | Most common sites<br>tongue and laryngeal/<br>glottis                                | 27% surgery only                                       |
| CIP) [54]                        |     |       |   |   |  |  | 26% radiotherapy<br>only                               |
|                                  |     |       |   |   |  |  |  |

Table 1 Sample characteristics of studies that have developed PROMs assessing quality of patient-centred cancer care (Continued)

|              |     |       |  |                     |   |   | 31% surgery and radiotherapy   |
|--------------|-----|-------|--|---------------------|---|---|--|
|              |     |       |  |                     | Mean age: 60 years  | Stages I and II: approximately 50%;     | 11% radiotherapy<br>and chemotherapy   |
| CAT DAD (F)  |     |       |  |                     | 92% white<br>61% married/de facto   | Stages III and IV:<br>approximately 50% | 5% surgery,<br>radiotherapy<br>and chemotherapy  |
| SAT-RAR [55] | 297 | 55.8% | Aged > 18 years, curative irradiation and satisfactory 1 general status (<3 on WHO performance scale). | 16 centers, France. | Breast cancer ( $n = 98$ ):   | Non-small cell lung                     | 100% radiotherapy  |
|              |     |       |  |                     | Mean age: 56 years  | cancer.                                 | Respiratory gating:<br>44% (breast cancer)<br>& 67% (lung cancer)                              |
|              |     |       |  |                     | 100% women  | Breast cancer.                          |  |
|              |     |       |  |                     | Mean days hospitalized:<br>4.7 days   | -                                       |  |
|              |     |       |  |                     | Non-small cell lung cancer (n = 199):                                       |   | 1 or more acute<br>toxicity during<br>treatment: 96%<br>(breast cancer) &<br>86% (lung cancer) |
|              |     |       |  |                     | Mean age: 65 years<br>15% women   |   |  |
|              |     |       |  |                     | Mean days hospitalized:<br>3.8 days Mean number<br>of hospitalizations: 7.8 |   | (- 3/  |

Table 2 Measurement features of PROMs and included IOM-endorsed patient-centred care dimensions

| Measure  | Satisfaction/<br>experience | Number of items           | Response scale   | Subscales                              | IOM patient-centred dimensions |
|--|-----------------------------|---------------------------|--|--|--------------------------------|
| Assessment of Patient  | Experiences                 | 33                        | Not a problem, A small problem,  | Getting needed care                    | Emotional support              |
| Experiences of Cancer<br>Care (APECC) [32]                     |                             |                           | A big problem.   | Timeliness of care                     | Information & communication    |
|  |                             |                           | Never, Sometimes, Usually,   | Waiting time in physician's office     | Integrated & coordinated care  |
|  |                             |                           | Always.  | Information exchange                   | Respectful to patients' values |
|  |                             |                           |  | Physicians' affective behavior         |                                |
|  |                             |                           |  | Physicians' knowledge                  |                                |
|  |                             |                           | On time, < 15 minutes, 16-30 minutes,                                    | Interaction with nurses                |                                |
|  |                             |                           | 31-45 minutes, > 45 minutes.   | Interaction with office staff          |                                |
|  |                             |                           |  | Health promotion                       |                                |
|  |                             |                           |  | Coordination of care                   |                                |
|  |                             |                           | Poor, Fair, Good, Very good, Excellent.                                  | Overall rating of care                 |                                |
|  |                             |                           | Yes definitely, Yes somewhat, No.  |  |                                |
|  |                             |                           | 0 (worst doctor possible) to 10 (best doctor possible)                   |  |                                |
|  |                             |                           | Definitely yes, Probably yes, Not sure,<br>Probably not, Definitely not. |  |                                |
| Cancer Care Coordination<br>Questionnaire for Patients<br>[33] | Experiences                 | 20                        | Strongly disagree, Disagree, Neutral,<br>Agree, Strongly agree.          | Communication                          | Emotional support              |
|  |                             |                           |  | Navigation                             | Family & friends               |
| [55]   |                             |                           | Never, Rarely, Sometimes, Frequently, Always.                            |  | Information & communication    |
|  |                             |                           |  |  | Integrated & coordinated care  |
| Cancer Patient Information                                     | Satisfaction                | 24                        | 5 point scale from Not important (0) to                                  | Information importance                 | Emotional support              |
| Importance and Satisfaction<br>Tool [34]                       |                             |                           | Very important (4).  | Information satisfaction               | Information & communication    |
|  |                             |                           | 5 point scale from Not satisfied (0) to<br>Very satisfied (4).           |  | Physical comfort               |
| Cancer Therapy Satisfaction                                    | Satisfaction                | 16                        | 5 point scale with 1 representing the                                    | Expectation of therapy                 | Respectful to patients' values |
| Questionnaire (CTSQ) [35,36]                                   |                             |                           | worst response and 5 representing the best response.                     | Feelings about side effects            |                                |
|  |                             |                           | and best response.   | Satisfaction with therapy              |                                |
| Consumer Quality Index   | Experiences                 | 152 (118 items related    | Never, Sometimes, Usually, Always.                                       | Conduct of professionals during        | Emotional support              |
| Breast Care (CQI-BC) [37]                                      |                             | to patients' experiences) |  | breast examination                     | Information & communication    |
|  |                             |                           | Yes, No.   | Conduct of general practitioner        | Integrated & coordinated care  |
|  |                             |                           |  | Conduct of nurses                      | Respectful to patients' values |
|  |                             |                           | A big problem, A small problem,  | Conduct of surgeon                     |                                |
|  |                             |                           | No problem.  | Autonomy regarding treatment           |                                |
|  |                             |                           |  | Autonomy regarding follow-up treatment |                                |

Table 2 Measurement features of PROMs and included IOM-endorsed patient-centred care dimensions (Continued)

|  |              |                        |   | Conduct of professionals during radiotherapy                        |                                |
|--|--------------|------------------------|---|---|--------------------------------|
|  |              |                        |   | Information on radiotherapy   |                                |
|  |              |                        |   | Conduct of professionals during chemotherapy                        |                                |
|  |              |                        |   | Information on chemotherapy   |                                |
|  |              |                        |   | Cooperation   |                                |
|  |              |                        |   | Accessibility of care   |                                |
|  |              |                        |   | Continuity psychosocial care  |                                |
|  |              |                        |   | Continuity physiotherapy  |                                |
|  |              |                        |   | Continuity rehabilitation   |                                |
| EORTC cancer in-patient                  | Satisfaction | 32                     | Poor, Fair, Good, Very good,                                | Doctors' technical skills   | Emotional support              |
| satisfaction with care<br>measure (EORTC |              |                        | Excellent.  | Doctors' interpersonal skills                                       | Information & communication    |
| N-PATSAT32) [38]                         |              |                        |   | Doctors' information provision                                      | Integrated & coordinated care  |
|  |              |                        |   | Doctors' availability   | Physical comfort               |
|  |              |                        |   | Nurses' technical skills  | Respectful to patients' values |
|  |              |                        |   | Nurses' interpersonal skills  |                                |
|  |              |                        |   | Nurses' information provision                                       |                                |
|  |              |                        |   | Nurses' availability  |                                |
|  |              |                        |   | Exchange of information   |                                |
|  |              |                        |   | Other hospital staff interpersonal skills and information provision |                                |
|  |              |                        |   | Waiting time  |                                |
|  |              |                        |   | Hospital access   |                                |
|  |              |                        |   | Comfort   |                                |
|  |              |                        |   | General   |                                |
|  |              |                        |   | satisfaction  |                                |
| ndicators (Head & Neck                   | Experiences  | 23 specific indicators | -   | Patient-oriented quality of care                                    | Emotional support              |
| Cancer) [39]                             |              | for patients           |   | Organisational quality of care                                      | Information & communication    |
|  |              |                        |   | Medical/technical quality of care                                   | Integrated & coordinated care  |
| ndicators (Non-small Cell                | Experiences  | 56                     | 1 = Not done, 2 = Done, but                                 | Access  | Emotional support              |
| ung Cancer) [40]                         |              |                        | inadequately, 3 = Done adequately,<br>4 = Done excellently. | Follow up   | Family & friends               |
|  |              |                        | . Done executivity.   | Communication and respect   | Information & communication    |
|  |              |                        |   | Patient & family involvement  | Integrated & coordinated care  |
|  |              |                        | Yes, No   | Information   | Physical comfort               |
|  |              |                        |   | Coordination  | Respectful to patients' values |

Table 2 Measurement features of PROMs and included IOM-endorsed patient-centred care dimensions (Continued)

|  |              |                       |  | Physical support                      |                                |
|--|--------------|-----------------------|--|---------------------------------------|--------------------------------|
|  |              |                       |  | Emotional & psychosocial support      |                                |
| Medical Care Questionnaire                           | Experiences  | 15                    | -  | Communication                         | Emotional support              |
| (MCQ) [41]   |              |                       |  | Preferences                           | Integrated & coordinated care  |
|  |              |                       |  | Coordination                          | Respectful to patients' values |
| Modified Version of the                              | Experiences  | 20                    | 1 = AII the time to $5 = Never$ .          | Health care provider information      | Information & communication    |
| Perceived Involvement<br>in Care Scale (M-PICS) [42] |              |                       |  | Patient information                   | Respectful to patients' values |
| (, []  |              |                       |  | Patient decision making               |                                |
|  |              |                       |  | Health care provider facilitation     |                                |
| Oncology Patients'                                   | Experiences  | 40 (and 18-item short | 1 = Never to $6 = $ Always, Didn't matter, | Responsiveness                        | Emotional support              |
| Perceptions of the<br>Quality of Nursing Care        |              | form created)         | Don't know.                                | Individualization                     | Family & friends               |
| Scale (OPPQNCS) [43]                                 |              |                       |  | Coordination                          | Information & communication    |
|  |              |                       |  | Proficiency                           | Integrated & coordinated care  |
|  |              |                       |  |                                       | Respectful to patients' values |
| Pain Care Quality Survey                             | Experiences  | 33                    | 1 = Strongly disagree to 6 = Strongly      | PainCQ-Interdisciplinary scale:       | Family & friends               |
| (PainCQ) [44,45]                                     |              |                       | agree.                                     | Partnership with healthcare team      | Information & communication    |
|  |              |                       |  | Comprehensive interdisciplinary       | Integrated & coordinated care  |
|  |              |                       |  | pain care                             | Physical comfort               |
|  |              |                       |  | PainCQ-Nursing scale:                 | Respectful to patients' values |
|  |              |                       |  | Being treated right                   |                                |
|  |              |                       |  | Comprehensive nursing pain care       |                                |
|  |              |                       |  | Efficacy of pain management           |                                |
| PASQOC questionnaire [46]                            | Satisfaction | 120                   | Nominal or ordinal scales and some         | 5 dimensions reported                 | Information & communication    |
|  |              |                       | interval scales                            | Patient-provider relationship         | Respectful to patients' values |
|  |              |                       |  | Premises                              |                                |
|  |              |                       |  | Information on diagnosis & treatment  |                                |
|  |              |                       |  | Information on treatment consequences |                                |
|  |              |                       |  | Relationship between patient & nurse  |                                |
| Patient Satisfaction with                            | Satisfaction | 18                    | 1 = Strongly Agree to 5 = Strongly         | 1 component structure – satisfaction  | Information & communication    |
| Cancer Care [47]                                     |              |                       | Disagree                                   | with cancer care                      | Integrated & coordinated care  |
|  |              |                       |  |                                       | Respectful to patients' values |
| Perceived Physician's                                | Experiences  | 27                    | 1 = Strongly Disagree, 3 = Neutral,        | Acceptive                             | Emotional support              |
| Communication Style<br>Scale [48]                    |              |                       | 5 = Strongly Agree.                        | Patient-centered                      | Information & communication    |
| 111.0 [10]   |              |                       |  | Attentive                             | Respectful to patients' values |
|  |              |                       |  | Facilitative                          |                                |
|  |              |                       |  |                                       |                                |

Table 2 Measurement features of PROMs and included IOM-endorsed patient-centred care dimensions (Continued)

| Prostate Care Questionnaire | Experiences | 106 (Sections A-E). | Various scales – please see                                      | Information from additional file 1                                    | Information & communication    |  |  |  |             |  |
|-----------------------------|-------------|---------------------|--|---|--------------------------------|--|--|--|-------------|--|
| for Patients (PCQ-P) [49]   |             |                     | reference [63]   | Section A: GP visits and referral                                     | Integrated & coordinated care  |  |  |  |             |  |
|                             |             |                     |  | Explanation   | Physical comfort               |  |  |  |             |  |
|                             |             |                     |  | Experience of referral  | Respectful to patients' values |  |  |  |             |  |
|                             |             |                     |  | Taking the problem seriously  |                                |  |  |  |             |  |
|                             |             |                     |  | Section B: Tests at the hospital                                      |                                |  |  |  |             |  |
|                             |             |                     |  | Explanation & support   |                                |  |  |  |             |  |
|                             |             |                     |  | Quality of care   |                                |  |  |  |             |  |
|                             |             |                     |  | Appointment   |                                |  |  |  |             |  |
|                             |             |                     |  | Section C: Diagnosis and treatment decision                           |                                |  |  |  |             |  |
|                             |             |                     |  | Explanation & support   |                                |  |  |  |             |  |
|                             |             |                     |  | Making treatment decision   |                                |  |  |  |             |  |
|                             |             |                     |  | Getting the diagnosis   |                                |  |  |  |             |  |
|                             |             |                     |  | Length of wait  |                                |  |  |  |             |  |
|                             |             |                     |  | Section D: Treatment and discharge                                    |                                |  |  |  |             |  |
|                             |             |                     |  | Preparation for discharge   |                                |  |  |  |             |  |
|                             |             |                     |  | Treatment   |                                |  |  |  |             |  |
|                             |             |                     |  |   |                                |  |  |  | Information |  |
|                             |             |                     |  | Section E: Monitoring   |                                |  |  |  |             |  |
|                             |             |                     |  | Explanation & reassurance   |                                |  |  |  |             |  |
|                             |             |                     |  | Advice  |                                |  |  |  |             |  |
|                             |             |                     |  | Choice  |                                |  |  |  |             |  |
| QUOTE Breast Cancer [50,51] | Experiences | 33                  | Performance:   | Patient education regarding aspects                                   | Information & communication    |  |  |  |             |  |
|                             |             |                     | Never, Sometimes, Usually, Always.                               | related to postoperative treatment                                    | Integrated & coordinated care  |  |  |  |             |  |
|                             |             |                     |  | Services by the breast nurse  | Respectful to patients' values |  |  |  |             |  |
|                             |             |                     | Yes, No.   | Services by the surgeon   |                                |  |  |  |             |  |
|                             |             |                     |  | Patient education regarding activities at home                        |                                |  |  |  |             |  |
|                             |             |                     | Not applicable/I do not know added to a subset of items.         | Patient education regarding aspects related to preoperative treatment |                                |  |  |  |             |  |
|                             |             |                     | Importance:  |   |                                |  |  |  |             |  |
|                             |             |                     | Not important, Fairly important, Important, Extremely important. |   |                                |  |  |  |             |  |
| QUOTE <sup>chemo</sup> [52] | Experiences | 67                  | Performance:   | Treatment-related information   | Emotional support              |  |  |  |             |  |
|                             |             |                     | Yes, No.   | Prognosis information   | Family & friends               |  |  |  |             |  |

Table 2 Measurement features of PROMs and included IOM-endorsed patient-centred care dimensions (Continued)

|  |              |    | Importance:   | Rehabilitation information  | Information & communication                   |
|--|--------------|----|---|---|---|
|  |              |    |   | Coping information  | Physical comfort                              |
|  |              |    | Not important, Fairly important,  | Interpersonal communication                                       | Respectful to patients' values                |
|  |              |    | Important, Very important.  | Tailored communication  |   |
|  |              |    |   | Affective communication   |   |
| REPERES-60 [53]  | Experiences  | 60 | Bad, Fair, Good, Very good, Excellent.                                  | Access to primary care  | Emotional support                             |
|  |              |    |   | Access to secondary care  | Information & communication                   |
|  |              |    | Completely agree, Agree generally,<br>No marked opinion, Do not really  | Competence and communication skills of primary care doctors       | Integrated & coordinated care                 |
|  |              |    | agree, Do not agree at all.   | Competence of secondary care doctors                              | Respectful to patients' values                |
|  |              |    |   | Communication skills of secondary care doctors                    |   |
|  |              |    | Bad, Fair, Good, Very good, Excellent,                                  |   |   |
|  |              |    | Not concerned.  | Choice among different doctors                                    |   |
|  |              |    |   | Human qualities of doctors  |   |
|  |              |    |   | Global satisfaction   |   |
|  |              |    |   | Cover for medical expenses  |   |
|  |              |    |   | Listening abilities and information provided by doctors           |   |
|  |              |    |   | Organisation and follow-up of medical care provision              |   |
|  |              |    |   | Psychological support   |   |
|  |              |    |   | Material environment  |   |
| Satisfaction with Cancer<br>Information Profile<br>(SCIP) [54] | Satisfaction | 21 | Too much, About right, Too little,<br>None wanted.                      | Satisfaction with the amount and content of information           | Information & communication                   |
|  |              |    | Very satisfied, Satisfied, Neither,<br>Dissatisfied, Very dissatisfied. | Satisfaction with the form and timing of the information received |   |
| SAT-RAR [55]   | Experiences  | 23 | Poor, Fair, Good, Very good,<br>Excellent.                              | Perception of the radiotherapist or radiotherapy technicians      | Emotional support Information & communication |
|  |              |    | Not at all, A little, Quite a bit, Very                                 | Global satisfaction   | Physical comfort                              |
|  |              |    | much.   | Treatment experience  | •   |
|  |              |    | Poor, Moderate, Good, Very good, Excellent.                             | •   |   |
|  |              |    | Disagree, Unsure, Tend to agree,<br>Agree, Strongly agree.              |   |   |

Table 3 IOM patient-centred care dimensions captured by PROMs

| Measure   |                   |                  | IOM patient-co              | entred care dimensions        | <b>i</b>         |                                |
|---|-------------------|------------------|-----------------------------|-------------------------------|------------------|--------------------------------|
|   | Emotional support | Family & friends | Information & communication | Integrated & coordinated care | Physical comfort | Respectful to patients' values |
| APECC [32]  | √                 |                  | $\sqrt{}$                   | $\sqrt{}$                     |                  | √                              |
| Cancer Care Coordination Questionnaire for Patients [33]            | $\sqrt{}$         | $\checkmark$     | $\sqrt{}$                   | $\checkmark$                  |                  |                                |
| Cancer Patient Information Importance and<br>Satisfaction Tool [34] | $\sqrt{}$         |                  | $\sqrt{}$                   |                               | $\sqrt{}$        |                                |
| CTSQ [35,36]  |                   |                  |                             |                               |                  | $\checkmark$                   |
| CQI-BC [37]   | $\sqrt{}$         |                  | $\sqrt{}$                   | $\checkmark$                  |                  | $\checkmark$                   |
| EORTC IN-PATSAT32 [38]  | $\sqrt{}$         |                  | $\sqrt{}$                   | $\checkmark$                  | $\sqrt{}$        | $\checkmark$                   |
| Indicators (Head & Neck Cancer) [39]                                | $\sqrt{}$         |                  | $\sqrt{}$                   | $\checkmark$                  |                  |                                |
| Indicators (Non-small Cell Lung Cancer) [40]                        | $\sqrt{}$         | $\sqrt{}$        | $\sqrt{}$                   | $\checkmark$                  | $\sqrt{}$        | $\checkmark$                   |
| MCQ [41]  | $\sqrt{}$         |                  |                             | $\checkmark$                  |                  | $\checkmark$                   |
| M-PICS [42]   |                   |                  | $\sqrt{}$                   |                               |                  | $\checkmark$                   |
| OPPQNCS [43]  | $\sqrt{}$         | $\sqrt{}$        | $\sqrt{}$                   | $\checkmark$                  |                  | $\checkmark$                   |
| PainCQ [44,45]  |                   | $\sqrt{}$        | $\sqrt{}$                   | $\checkmark$                  | $\sqrt{}$        | $\checkmark$                   |
| PASQOC questionnaire [46]   |                   |                  | $\sqrt{}$                   |                               |                  | $\checkmark$                   |
| Patient Satisfaction with Cancer Care [47]                          |                   |                  | $\sqrt{}$                   | $\checkmark$                  |                  | $\checkmark$                   |
| Perceived Physician's Communication Style<br>Scale [48]             | $\checkmark$      |                  | $\checkmark$                |                               |                  | $\checkmark$                   |
| PCQ-P [49]  |                   |                  | $\sqrt{}$                   | $\checkmark$                  | $\sqrt{}$        | $\checkmark$                   |
| QUOTE Breast Cancer [50,51]   |                   |                  | $\sqrt{}$                   | $\checkmark$                  |                  | $\sqrt{}$                      |
| QUOTE <sup>chemo</sup> [52]   | $\sqrt{}$         | $\sqrt{}$        | $\checkmark$                |                               | $\sqrt{}$        | $\checkmark$                   |
| REPERES-60 [53]   | $\sqrt{}$         |                  | $\checkmark$                | $\checkmark$                  |                  | $\sqrt{}$                      |
| SCIP [54]   |                   |                  | $\checkmark$                |                               |                  |                                |
| SAT-RAR [55]  | $\sqrt{}$         |                  | $\sqrt{}$                   |                               | $\sqrt{}$        |                                |

# **Construct validity**

Eighteen measures met the criteria for adequate construct validity [32,33,35,37,38,41-44,46-50,52-55]. Sixteen measures conducted factor analyses [32,33,35,37,41-44,46-50,52,53,55] (although only seven reported eigenvalues) [33,41,44,47,48,50,53], nine measures examined convergent validity (r > 0.40) or divergent validity (r < 0.30) with existing instruments [35,38,42,46,47,49,52-54] and six measures demonstrated significant differences on scores between known groups [35,38,41,42,53,55].

## Cross-cultural adaptation

Three measures were re-validated with non-English speaking populations. The EORTC IN-PATSAT32 was validated with Sri Lankan cancer patients [56]; the Modified version of the Perceived Involvement in Care Scale (M-PICS) was validated with Lithuanian cancer patients [57]; and the Oncology Patients' Perceptions of the Quality Nursing Care Scale (OPPQNCS) was validated with Turkish cancer patients [58].

Table 5 summarises which PROMs met the psychometric criteria considered adequate, as described above.

# Psychometric properties of PROMs containing all six IOM patient-centred care dimensions

The Indicators (Non-small Cell Lung Cancer) measure [40] was the only PROM that contained items covering all six IOM dimensions of patient-centred care. This measure met the criteria considered adequate for face/content validity, but not for any other psychometric criteria evaluated in this review.

#### **Discussion**

This is the first review to identify how many of the six IOM-endorsed dimensions of patient-centred care [1] are covered in existing PROMs assessing the quality of cancer care. Our findings demonstrate that since the publication of the IOM's *Crossing the Quality Chasm* report in 2001 [1], only one of 21 patient-centred cancer care instruments, the Indicators (Non-small Cell Lung Cancer) measure, included questions relating to the six

Table 4 Psychometric properties of PROMs assessing quality of patient-centred cancer care

| Measure  | Face validity/content validity  |   | <i>'</i>     | Internal<br>consistency | Test-retest reliability                                    | Cross-<br>cultural<br>adaptation   |   |  |                                    |  |  |  |  |  |  |  |  |  |                                       |
|--|---|---|--------------|-------------------------|--|--|---|--|------------------------------------|--|--|--|--|--|--|--|--|--|---------------------------------------|
|  |   | Factor analysis   | Known groups | Existing measure        |  |  | • |  |                                    |  |  |  |  |  |  |  |  |  |                                       |
| Assessment of Patient<br>Experiences of Cancer                         | All items underwent cognitive testing with nine cancer  | Confirmatory factor analysis indicated a reasonably good fit                              | -            | -                       | Getting needed care: α =.76                                | -  | - |  |                                    |  |  |  |  |  |  |  |  |  |                                       |
| Care (APECC) [32]  | survivors to ensure that the<br>questions and response options<br>were understandable and   | for the 10-factor model (comparative fit index = $0.93$ ).                                |              |                         |  |  |   |  | Timeliness of care: $\alpha = .62$ |  |  |  |  |  |  |  |  |  |                                       |
|  | related to the concept being measured.  |   |              |                         | Waiting time in physician's office: $\alpha = .65$         |  |   |  |                                    |  |  |  |  |  |  |  |  |  |                                       |
|  |   |   |              |                         | Information exchange: $\alpha = .92$                       |  |   |  |                                    |  |  |  |  |  |  |  |  |  |                                       |
|  |   |   |              |                         |  |  |   | Physicians'<br>affective<br>behavior: $\alpha = .92$ |                                    |  |  |  |  |  |  |  |  |  |                                       |
|  |   |   |              |                         | Physicians'<br>knowledge: α<br>=.86                        |  |   |  |                                    |  |  |  |  |  |  |  |  |  |                                       |
|  |   |   |              |                         | Interaction with nurses: $\alpha = .82$                    |  |   |  |                                    |  |  |  |  |  |  |  |  |  |                                       |
|  |   |   |              |                         |  |  |   |  |                                    |  |  |  |  |  |  |  |  |  | Interaction with office staff: α =.90 |
|  |   |   |              |                         | Health<br>promotion: α<br>=.88                             |  |   |  |                                    |  |  |  |  |  |  |  |  |  |                                       |
|  |   |   |              |                         |  |  |   | Coordination of care: N/A                            |                                    |  |  |  |  |  |  |  |  |  |                                       |
|  |   |   |              |                         | Overall rating of care: α=.87                              |  |   |  |                                    |  |  |  |  |  |  |  |  |  |                                       |
| Cancer Care Coordination<br>Questionnaire for Patients [33]            | Literature review undertaken to identify relevant issues and existing instruments and focus groups and semi-structured interviews with 24 cancer patients and carers and 29 clinicians. | Exploratory factor analysis.<br>Principal factor method followed<br>by a promax rotation. | -            | -                       | Total scale: $\alpha$ =.88<br>Communication: $\alpha$ =.87 | patients completed<br>the survey twice -<br>mailed 2 weeks<br>after receipt of first                                 | - |  |                                    |  |  |  |  |  |  |  |  |  |                                       |
|  | and carers and 29 clinicians.  Draft questionnaire was reviewed by clinicians and re- searchers to assess face validity and clarity of wording.   | Factor loadings >.40 with the exception of one item (0.37). Eigenvalues>1                 |              |                         | Navigation:<br>α=.73                                       | survey.  Kappa for individual items ranged from 0.29 to 0.69. Four items with values less than 0.40 were eliminated. |   |  |                                    |  |  |  |  |  |  |  |  |  |                                       |
| Cancer Patient Information<br>Importance and Satisfaction<br>Tool [34] | Literature review and extensive qualitative interviews with cancer patients. Tool was field-  | -   | -            | -                       | Information importance: α=.89                              | -  | - |  |                                    |  |  |  |  |  |  |  |  |  |                                       |

Table 4 Psychometric properties of PROMs assessing quality of patient-centred cancer care (Continued)

|   | tested with 10 cancer patients<br>who completed tool and pro-<br>vided feedback about its clarity<br>and ease of completion.  |  |  |  | Information satisfaction: α=.92                                   |   |
|---|---|--|--|--|---|---|
| Cancer Therapy Satisfaction<br>Questionnaire (CTSQ) [35,36] | Interviews with 70 oncology patients, 4 oncology nurses and 7 physicians. Focus groups with 14 oncology nurses. Content validity tested with 30 patients who completed the survey and | Exploratory factor analysis using oblique promax rotation. | Cancer stage (I, II, III, IV) P <0.001 for 1 subscale.                 | Treatment Satisfaction<br>Questionnaire for Medication<br>6 correlations > .40 | Expectations of<br>therapy: α=.87                                 | 85 patients - completed follow- up questionnaires one week after baseline assessment. |
|   | were interviewed, followed by<br>retesting in an additional 10<br>patients.   |  |  |  | Feelings about side effects: $\alpha$ =.77                        | Intraclass correlation:   |
|   |   |  |  |  | Satisfaction with therapy: α=.82                                  | Expectations of therapy: .56  |
|   |   |  | Side effects (with, without) <i>P</i> <0.05 for 2 subscales.           | EORTC QLQ-C30 1 correlation >.40   |   | Feelings about side effects: .77  |
|   |   |  | performance<br>status (Grade 0,<br>1, 2, 3) P<0.005<br>for 1 subscale. |  |   | Satisfaction with therapy: .75  |
| Consumer Quality Index Breast<br>Care (CQI-BC) [37]         | Three focus groups with 27 breast cancer patients, existing Dutch questionnaires on breast care and key stakeholders' input used for questionnaire                                    | Explorative factor analysis.                               | -  | -  | Conduct of professionals during breast examination: $\alpha$ =.91 | _   |
|   | development.  |  |  |  | Conduct of general practitioner: a=.89                            |   |
|   |   |  |  |  | Conduct of nurses: α=.88  |   |
|   |   |  |  |  | Conduct of surgeon: $\alpha$ =.91                                 |   |
|   |   |  |  |  | Autonomy<br>regarding<br>treatment: α=.84                         |   |
|   |   |  |  |  | Autonomy regarding follow-<br>up treatment:<br>α=.93              |   |

| Table 4 Psychometric properties of PROMs assessing quality of patient-centred cancer care (Continued) |
|---|
|   |

tion with care measure (EORTC

EORTC QLQ-SAT32) [64,65], as

specialists and cancer patients.

well as interviews with oncology

IN-PATSAT32) [38]

EORTC cancer in-patient satisfac- Adapted from existing patient Age (less than 57 Oberst Patients' Perception of satisfaction questionnaires years, 57 years or Care Quality and Satisfaction (Comprehensive Assessment of more) P < 0.05 Scale correlations (-0.21 to -0.61).  $\alpha$ =.85-.87 for 4 subscales. Satisfaction with Care and

Education (less

P < 0.05 for 2

subscales.

than compulsory,

post-compulsory)

technical skills: Doctors' interpersonal skills: α=.91-.94 Doctors' information provision:  $\alpha$ =.90 Doctors' availability: α=.86 Nurses' technical skills:  $\alpha$ =.90-.94

Doctors'

-.94

-.91

Conduct of professionals during radiotherapy: α=.89 Information on radiotherapy: a=.89 Conduct of professionals during chemotherapy:  $\alpha = .90$ Information on chemotherapy: a=.85 Cooperation: a=.88 Accessibility of care: α=.68 Continuity psychosocial care:  $\alpha$ =.83 Continuity physiotherapy: α=.82 Continuity rehabilitation:  $\alpha = .80$ 

> 113 patients Validated recruited from one with Sri centre for follow-up Lankan approximately 2 cancer weeks after first patients assessment. (n=343)[56]

Nurses'

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|   |   |   |                                       |                      | Nurses'<br>interpersonal<br>skills: a=.9093<br>Nurses'<br>information<br>provision: a=.94<br>96<br>Nurses'<br>availability: a=.83<br>92 | Intra-class<br>correlations<br>coefficients for the<br>scales ranged from<br>0.70-0.85 and was<br>0.66 for the general<br>satisfaction item. |  |
|---|---|---|---------------------------------------|----------------------|---|--|--|
|   |   |   | Treatment related toxicity            | EORTC QLQ-C30 r<.30. | Exchange of information: N/A  |  |  |
|   |   |   | (yes, no) P <0.05<br>for 7 subscales. |                      | Other hospital staff interpersonal skills and information provision: α=.86 –.90   |  |  |
|   |   |   |                                       |                      | Waiting time:<br>α=.80–.84  |  |  |
|   |   |   |                                       |                      | Hospital access: $\alpha$ =.56–.71  |  |  |
|   |   |   |                                       |                      | Comfort: N/A  |  |  |
|   |   |   |                                       |                      | General satisfaction: N/A   |  |  |
| Indicators (Head & Neck Cancer)<br>[39]         | Systematically searched for recommendations in literature, performed a systematic consensus procedure based on evidence-based guidelines and sought opinions of 15 professionals and 30 patients with head and neck cancer. | - | -                                     | _                    | -   | _  |  |
| Indicators (Non-small Cell Lung<br>Cancer) [40] | Recommendations for patient-<br>centred care extracted from clin-   | - | -                                     | -                    | Access: $\alpha$ =.87<br>Follow up: $\alpha$ =.78   | =  |  |
|   | ical guidelines and conducted<br>semi-structured interviews with<br>30 head and neck cancer pa-<br>tients and 7 patient representa-   |   |                                       |                      | Communication and respect: N/A  |  |  |
|   | tives from the Dutch national association of patients with lung   |   |                                       |                      | Patient & family involvement: α=.85   |  |  |
|   | cancer. Two researchers trans-<br>lated recommendations into in-<br>dicators which were considered<br>by a panel of four researchers.   |   |                                       |                      | Information: $\alpha$ =.78  |  |  |
|   | by a paner of four researchers.   |   |                                       |                      | Coordination:<br>α=.22<br>(specialists) &   |  |  |
|   |   |   |                                       |                      | (Specialists) &   |  |  |

Table 4 Psychometric properties of PROMs assessing quality of patient-centred cancer care (Continued)

Table 4 Psychometric properties of PROMs assessing quality of patient-centred cancer care (Continued)

|  |  |  |   |   | α=.68 (oncology<br>nurses)<br>Physical support:<br>N/A   |   |
|--|--|--|---|---|--|---|
|  |  |  |   |   | Emotional & psychosocial support: a=.67                  |   |
| Medical Care Questionnaire<br>(MCQ) [41]   | Literature review of existing<br>instruments, modification (items<br>removed, reworded, generated)<br>of an existing instrument by an  | Exploratory factor analysis using oblique rotation (Phase 3) and confirmatory factor analysis (Phase 4). | Cancer type<br>(Breast,<br>Genitourinary,<br>Gynecological, | -   | Communication: -<br>α=.69                                | -   |
|  | expert panel (3 medical oncologists and one oncologist   | Factor loadings >.40   | Melanoma,<br>Sarcoma, Other)                                |   | Preferences:<br>α=.84                                    |   |
|  | in training) using a consensus<br>procedure, instrument<br>administered to 200 oncology<br>outpatients and then refined.   | Eigenvalues > 1  | P < 0.05 for 3 subscales.                                   |   | Coordination:<br>α=.75                                   |   |
| Modified Version of the<br>Perceived Involvement in Care<br>Scale (M-PICS) [42]          | Literature review and consultation with pain clinicians guided augmentation and  | Exploratory factor analysis – principal components analysis with oblique rotation.                       | Age <i>P</i> <0.01 for 1 subscale.                          | Barriers Questionnaire-II   | Total scale: α= -<br>.87                                 | Validated<br>with<br>Lithuanian                                   |
|  | addition of items on the original<br>Perceived Involvement in Care<br>Scale (PICS).  |  | Ethnicity (Latina;<br>Caucasian/<br>African-<br>American)   | 2 correlations >.40   | Health care provider information: α= .90                 | cancer<br>patients<br>(n=30) [57]                                 |
|  |  |  | P<0.01 for 2 subscales.                                     | Mental Health Inventory   | Patient<br>information: α=<br>.82                        |   |
|  |  |  |   | 3 correlations <.30   | Patient decision<br>making: α= .80                       |   |
|  |  |  |   | Medical Outcomes Study Short-<br>Form 12: Mental Component<br>Scale   | Health care<br>provider<br>facilitation: α=              |   |
|  |  |  |   | 3 correlations <.30   | .80  |   |
|  |  |  |   | Medical Outcomes Study Short-<br>Form 12: Physical Component<br>Scale |  |   |
|  |  |  |   | 4 correlations <.30   |  |   |
|  |  |  |   | Patient Satisfaction<br>Questionnaire                                 |  |   |
|  |  |  |   | 3 correlations >.40   |  |   |
| Oncology Patients' Perceptions<br>of the Quality of Nursing Care<br>Scale (OPPQNCS) [43] | Interviews with cancer patients<br>about their perceptions of<br>nursing care, items and<br>subscales generated from this<br>qualitative work, an expert<br>methods consultant evaluated | Exploratory factor analysis – principal components analyses with promax (oblique) rotation.              | -   | -   | Total scale: $\alpha$ =99 Responsiveness: $\alpha$ = .99 | Validated<br>with<br>Turkish<br>cancer<br>patients<br>(n=54) [58] |

Table 4 Psychometric properties of PROMs assessing quality of patient-centred cancer care (Continued)

|  | items for clarity and relevance<br>and a nine-member expert rater<br>panel consisting of 5 cancer pa-                               |   |   | Individualization: $\alpha$ = .97                      |
|--|---|---|---|--|
|  | tients, a nurse, a nurse re-<br>searcher, the executive director  |   |   | Coordination: α=<br>.87                                |
|  | of a patient advocacy group and<br>a survey scientist, reviewed and<br>reduced items.   | Factor loadings >.40  |   | Proficiency: a=<br>.95                                 |
| Pain Care Quality Survey<br>(PainCQ) [44,45] | Conducted 33 qualitative interviews with cancer patients in pain. Items were constructed  | Exploratory factor analysis – - principal axis factoring using an oblimin rotation. | -   | PainCQ<br>Interdisciplinary<br>scale:                  |
|  | using this qualitative work,<br>existing tools, recommendations<br>for item development in the<br>literature, and consultation with |   |   | Partnership with healthcare team: $\alpha$ =.85        |
|  | a national expert in tool<br>development. Two panels of<br>pain and quality experts<br>reviewed items (removed,                     |   |   | Comprehensive<br>interdisciplinary<br>pain care: a=.76 |
|  | reworded or added items).  Cognitive interviews were then   |   |   | PainCQ-Nursing<br>scale:                               |
|  | undertaken with 39 hospitalized cancer patients reporting pain.   |   |   | Being treated right: α=.95                             |
|  |   | Factor loadings >.40  |   | Comprehensive nursing pain care: $\alpha$ =.77         |
|  |   | Eigenvalues > 1   |   | Efficacy of pain management: $\alpha$ =.87             |
| PASQOC questionnaire [46]                    | Existing survey re-designed and content similar to other surveys. Focus group discussions with 29 patients in four centres.         | Factor analysis -   | Short Form-36 (SF-36) correlations (.042 to161) | Total scale: α=93                                      |
|  | Pre-testing the questionnaire for appropriateness of the questions and length with 280 patients                                     |   |   | Patient-provider<br>relationship:<br>α=.81             |
|  | from 14 centres.  |   |   | Premises: α=.76  |
|  |   |   |   | Information on diagnosis & treatment: $\alpha$ =.71    |
|  |   |   |   | Information on treatment consequences: $\alpha$ =.87   |
|  |   |   |   | Relationship<br>between patient<br>& nurse: α=.72      |
|  |   |   |   |  |

Table 4 Psychometric properties of PROMs assessing quality of patient-centred cancer care (Continued)

| Patient Satisfaction with Cancer<br>Care [47]            | Item pool based on literature review and existing measures,  | Factor analysis – principal components analysis                                  | -  | Communication and Attitudinal Self-Efficacy – Cancer 1 correl-    | Total scale: α=<br>.95 and .96 (for  | -   | - |
|--|--|--|--|---|--|---|---|
|  | expert feedback, group discussion and consensus.   | Factor loadings >.40   |  | ation >.40 (sample 2).  | two samples).  |   |   |
|  | alseassion and consensus.  | Eigenvalue >1  |  |   |  |   |   |
| Perceived Physician's<br>Communication Style Scale [48]  | Initial item pool created from<br>literature review and 25 cancer  | Factor analysis using varimax procedure.   | -  | -   | Total scale: α=<br>.95   | -   | = |
|  | patients' opinions about the physician's communication   | Factor loadings >.40   |  |   | Acceptive: α=.90   |   |   |
|  | style.114 nurses assessed the content validity of items. Items   | Eigenvalues > 1  |  |   | Patient-centered: $\alpha$ =.90  |   |   |
|  | pre-tested with 70 breast cancer patients.   |  |  |   | Attentive: α=.73   |   |   |
|  | •  |  |  |   | Facilitative: α=.76  |   |   |
| Prostate Care Questionnaire for<br>Patients (PCQ-P) [49] | Initial items developed through a literature review and interviews with patients and service providers.  | Exploratory principal components analysis with varimax rotation.                 | -  | National Centre for Social<br>Research Shortened<br>Questionnaire | Section A: α=.80   | 148 (50%) patients<br>from two hospitals<br>completed retest        | - |
|  | Semi-structured interviews with 20   | Factor loadings for each section   |  | Sections B & C  | Section B: α=.63   | survey mailed 3<br>weeks later.                                     |   |
|  | prostate cancer patients who com-<br>pleted survey.  | of 0.3 and higher presented in an additional file.                               |  | r <.30  | Section C: α=.77   | weeks later.  |   |
|  |  | an additional file.  |  | Sections D & E r >.40   | Section D: α=.80   |   |   |
|  |  |  |  |   | Section E: α=.68   | Intraclass<br>correlation<br>coefficient:                           |   |
|  |  |  |  |   |  | Section A: .68  |   |
|  |  |  |  |   |  | Section B: .57  |   |
|  |  |  |  |   |  | Section C: .61  |   |
|  |  |  |  |   |  | Section D: .73  |   |
|  |  |  |  |   |  | Section E: .70  |   |
|  |  |  |  |   |  | Identical responses<br>to individual<br>questions: 52.6% to<br>100% |   |
| QUOTE Breast Cancer [50,51]                              | Based on eight focus groups<br>with 72 breast cancer patients<br>and concept mapping sessions<br>with 67 breast cancer patients a<br>pilot questionnaire was<br>developed. Two researchers | Exploratory factor analysis (i.e., principal axis factoring) – oblique rotation. | Age (18-49, 50-<br>65, >65 years) no<br>difference.            |   | Patient<br>education<br>regarding<br>aspects related<br>to postoperative<br>treatment: α=.83 | -   | - |
|  | categorised and reduced the aspects of care identified.  | Factor loadings >.40 except for 0.35 loading and 6 separate items.               |  |   | Services by the breast nurse: α=.89  |   |   |
|  |  | Eigenvalues >1   | Education<br>(primary school,<br>secondary<br>school, college/ |   | Services by the surgeon: α=.85  Patient education  |   |   |

Table 4 Psychometric properties of PROMs assessing quality of patient-centred cancer care (Continued)

|                             |  | · · · · · · · · · · · · · · · · · · · |   |  |  |
|-----------------------------|--|---------------------------------------|---|--|--|
|                             |  |                                       | university) no<br>difference.                                       |  | regarding<br>activities at<br>home: α=.70                                    |
|                             |  |                                       | Time since<br>surgery (0-6, 6-<br>12, >12 months)<br>no difference. |  | Patient education regarding aspects related to preoperative treatment: a=.81 |
| QUOTE <sup>chemo</sup> [52] | Items developed via existing   | Confirmatory factor analysis.         | -   | QUOTE <sup>chemo</sup> Importance with:    | Performance:   |
|                             | measure, literature review and 5 focus groups (n=33) as well as individual interviews with 5 cancer patients. 10 coders categorised the items into the | Factor loadings >.40                  |   |  | Treatment-related information: $\alpha$ =.92                                 |
|                             | seven dimensions.  |                                       |   |  | Prognosis<br>information: α=<br>.72  |
|                             |  |                                       |   |  | Rehabilitation<br>information: α=<br>.87                                     |
|                             |  |                                       |   | Information Satisfaction<br>Questionnaire  | Coping information: a= .78   |
|                             |  |                                       |   | 7 correlations r <.30                      | Interpersonal communication: $\alpha$ = .89                                  |
|                             |  |                                       |   |  | Tailored communication: $\alpha$ = .86                                       |
|                             |  |                                       |   |  | Affective communication: $\alpha$ = .91                                      |
|                             |  |                                       |   | Threatening Medical Situation<br>Inventory | Importance:  |
|                             |  |                                       |   | 7 correlations r <.30                      | Treatment-related information: $\alpha$ =.90                                 |
|                             |  |                                       |   | Impact of Event Scale:                     | Prognosis<br>information: α=<br>.76  |
|                             |  |                                       |   | - Intrusion 7 correlations r <.30          | Rehabilitation information: α= .86   |
|                             |  |                                       |   | - Avoidance 7 correlations r <.30          | Coping information: a= .81   |
|                             |  |                                       |   |  |  |

Table 4 Psychometric properties of PROMs assessing quality of patient-centred cancer care (Continued)

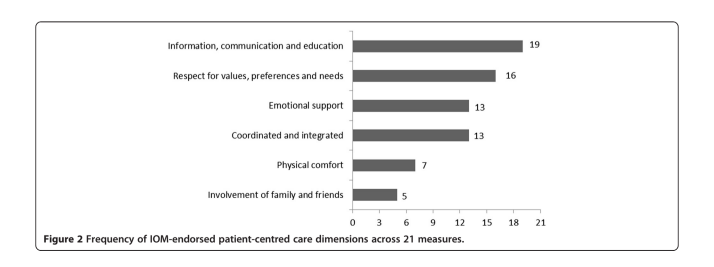
|                 |   |  |   |                      | Interpersonal communication: α= .90                                     |  |
|-----------------|---|--|---|----------------------|---|--|
|                 |   |  |   |                      | Tailored communication: $\alpha$ = .81                                  |  |
|                 |   |  |   |                      | Affective communication: $\alpha$ = .88                                 |  |
| REPERES-60 [53] | Literature review and two focus<br>groups with 30 breast cancer<br>patients used to generate items  | Principal components factor analysis (varimax rotation). | Age (less than 58<br>years, 58 years or<br>older)               | EORTC QLQ-C30 r <.30 | Access to primary care: α=.88   | 166 (91%) patients -<br>sent retest 1 week<br>later. |
|                 | and identify domains. Adapted<br>existing Consumer Satisfaction<br>Survey and developed new<br>items based on patient focus   | Factor loadings >.40                                     | P <0.05 for 11 subscales.                                       |                      | Access to secondary care: α=.82   |  |
|                 | groups and experts' input. Test version of questionnaire tested with breast cancer patients to assess understanding and comprehensiveness of items, which led to minor alterations. | Eigenvalues >1   | Education<br>(primary and<br>secondary,<br>higher<br>education) |                      | Competence and communication skills of primary care doctors: a=.93      | ltem-to-item<br>agreement:                           |
|                 |   |  | P <0.05 for 8 subscales.  |                      | Competence of secondary care doctors: $\alpha$ =.94                     | Kappa ranged from<br>0.44 to ≥ 0.70                  |
|                 |   |  |   |                      | Communication skills of secondary care doctors: $\alpha$ =.92           |  |
|                 |   |  |   |                      | Choice among<br>different doctors:<br>α=.87                             |  |
|                 |   |  |   |                      | Human qualities of doctors: α=.94                                       |  |
|                 |   |  |   |                      | Global<br>satisfaction:<br>α=.74  |  |
|                 |   |  |   |                      | Cover for<br>medical<br>expenses: α=.90                                 |  |
|                 |   |  |   |                      | Listening abilities<br>and information<br>provided by<br>doctors: a=.93 |  |
|                 |   |  |   |                      | Organisation and follow-up of medical care provision: α=.87             |  |

Table 4 Psychometric properties of PROMs assessing quality of patient-centred cancer care (Continued)

|   |   |   |   |  | Psychological<br>support: α=.88  |
|---|---|---|---|--|--|
|   |   |   |   |  | Material<br>environment:<br>α=.89  |
| Satisfaction with Cancer<br>Information Profile (SCIP) [54] | Four items derived from Satisfaction with Information About Medicines Scale (SIMS) and additional items from patient interviews.  | -   | -   | Illness Perception Questionnaire-<br>Revised | Satisfaction with the amount and content of information: $\alpha$ =.89             |
|   |   |   |   | 2 correlations r <.30                        | Satisfaction with the form and timing of information: $\alpha$ =.87                |
| SAT-RAR [55]  | Review of the literature,<br>selection and formation of items<br>based on relevant themes and a<br>cancer care satisfaction<br>questionnaire, pilot-testing of                    | Exploratory factorial analysis with varimax rotation followed by Confirmatory factorial analysis. | Education (at<br>least high school<br>completed, high<br>school not<br>completed) | -  | Total scale: α=.86   |
|   | preliminary questionnaire with<br>10 patients to evaluate rele-<br>vance, content validity and com-<br>prehensibility of items, survey<br>reviewed by clinicians in the<br>study. |   | P <0.01 in 1<br>subscale.   |  | Perception of<br>the<br>radiotherapist or<br>radiotherapy<br>technicians:<br>a=.94 |
|   |   | Most factor loadings >.40,<br>except 3 factor loadings >.30                                       | Marital status<br>(married, not<br>married)                                       |  | Global<br>satisfaction:<br>a=.80   |
|   |   |   | P < 0.01 in 1 subscale.   |  | Treatment experience:  |
|   |   |   | Work status<br>(employed, not<br>employed)  |  | α=.75  |
|   |   |   | P <0.01 in 2 subscales.   |  |  |
|   |   |   | Type of<br>radiotherapy<br>(respiratory<br>gating, no<br>respiratory<br>gating)   |  |  |
|   |   |   | P <0.01 in 2 subscales.   |  |  |

Table 5 PROMs demonstrating adequate psychometric properties based on recommended criteria

| Measure  | Face/content validity | Co              | onstruct vali   | dity             | Internal consistency |  | Cross-cultural adaptation |
|--|-----------------------|-----------------|-----------------|------------------|----------------------|--|---------------------------|
|  |                       | Factor analysis | Known<br>groups | Existing measure |                      |  |                           |
| APECC [32]   |                       | √               |                 |                  |                      |  |                           |
| Cancer Care Coordination Questionnaire for Patients [33]         | $\sqrt{}$             | $\checkmark$    |                 |                  | $\checkmark$         |  |                           |
| Cancer Patient Information Importance and Satisfaction Tool [34] |                       |                 |                 |                  |                      |  |                           |
| CTSQ [35,36]   | $\checkmark$          | $\checkmark$    | $\sqrt{}$       | $\sqrt{}$        |                      |  |                           |
| CQI-BC [37]  | $\checkmark$          | $\checkmark$    |                 |                  |                      |  |                           |
| EORTC IN-PATSAT32 [38]   | $\checkmark$          |                 | $\sqrt{}$       | $\sqrt{}$        |                      |  | $\checkmark$              |
| Indicators (Head & Neck Cancer) [39]                             | $\checkmark$          |                 |                 |                  |                      |  |                           |
| Indicators (Non-small Cell Lung Cancer) [40]                     | $\checkmark$          |                 |                 |                  |                      |  |                           |
| MCQ [41]   | $\checkmark$          | $\checkmark$    | $\sqrt{}$       |                  |                      |  |                           |
| M-PICS [42]  |                       | $\checkmark$    | $\sqrt{}$       | $\sqrt{}$        | $\sqrt{}$            |  | $\checkmark$              |
| OPPQNCS [43]   | $\checkmark$          | $\checkmark$    |                 |                  | $\sqrt{}$            |  | $\checkmark$              |
| PainCQ [44,45]   | $\checkmark$          | $\checkmark$    |                 |                  |                      |  |                           |
| PASQOC questionnaire [46]  |                       | $\checkmark$    |                 | $\sqrt{}$        | $\sqrt{}$            |  |                           |
| Patient Satisfaction with Cancer Care [47]                       |                       | $\checkmark$    |                 | $\sqrt{}$        | $\sqrt{}$            |  |                           |
| Perceived Physician's Communication Style<br>Scale [48]          | $\checkmark$          | $\sqrt{}$       |                 |                  | $\checkmark$         |  |                           |
| PCQ-P [49]   | $\checkmark$          | $\checkmark$    |                 | $\sqrt{}$        |                      |  |                           |
| QUOTE Breast Cancer [50,51]                                      | $\sqrt{}$             | $\sqrt{}$       |                 |                  |                      |  |                           |
| QUOTE <sup>chemo</sup> [52]                                      | $\sqrt{}$             | $\sqrt{}$       |                 | $\checkmark$     |                      |  |                           |
| REPERES-60 [53]  | $\sqrt{}$             | $\sqrt{}$       | $\sqrt{}$       | $\checkmark$     |                      |  |                           |
| SCIP [54]  |                       |                 |                 | $\checkmark$     |                      |  |                           |
| SAT-RAR [55]   | $\checkmark$          | $\sqrt{}$       | $\sqrt{}$       |                  | $\checkmark$         |  |                           |



IOM dimensions of patient-centred care [40]. However this measure only met the criteria considered acceptable for face/content validity. Further psychometric testing of the Indicators (Non-small Cell Lung Cancer) measure is required before more definitive conclusions can be drawn about its reliability and validity.

Across measures, the most commonly included patientcentred care dimensions were "information, communication and education" (19 of 21 measures) followed by "respectful to patients' values, preferences, and expressed needs" (16 of 21 measures). In contrast, only seven measures examined patient's perceptions of "physical comfort" and five assessed the "involvement and wellbeing of family and friends." Possible explanations for the lesser focus on issues related to family and friends could include: 1) researchers/health professionals perceiving issues related to information and communication as the most important features of patient-centredness; 2) that the patients and survey developers involved in item selection only wished to focus on specific aspects of care; and 3) issues related to family and friends are considered a less crucial feature of cancer care. Furthermore, the measures may not have adequately captured the IOM's six dimensions of patient-centred care because they were not developed for that purpose. For example, a measure's objective may have been to focus solely or primarily on physical comfort, rather than to address the IOM's six dimensions of patient-centred care. Nevertheless, the lack of PROMs that included all six IOM dimensions of patient-centred care [1] limits the potential of these existing measures to capture the whole-person orientation of health care and is likely to result in an incomplete representation of the quality of care provided to cancer patients.

Improvements to the reliability of existing patientcentred care PROMs and better reporting of their internal consistency, are needed. Only seven of the 21 measures met the criteria considered adequate for internal consistency by reporting a Cronbach's alpha >0.70 for the total scale and each sub-scale [33,42,43,46-48,55]. A further six measures showed that all subscales had a Cronbach's alpha >0.70 [34,35,44,52-54], but failed to report the internal consistency for the total scale. However, interpretation of internal consistency findings should always consider that when a subscale has a large number of items, Cronbach's alpha can be artificially high [59,60]. Test-retest validity was very rarely considered during the development of PROMs assessing patient-centred cancer care. Although four of the five measures that examined test-retest reliability administered a second survey within 2-14 days [33,35,38,53], none of the measures demonstrated acceptable agreement between scores for the total scale, subscales and items across the two administrations [18,19]. However possible explanations for the lack of adequate test-retest reliability among PROMs assessing patient-centred cancer care may include that: 1) patients' experiences of care, particularly for those receiving active treatment, actually changed between the initial and second administration of the measure; and 2) completing the initial measure altered patients' expectations of patient-centred care and as a result patients rated their care differently during the second administration of the measure. Nonetheless, future research that develops PROMs of patient-centred cancer care, or validates existing measures should examine test-retest reliability, with the aim of achieving high item-to-item agreement. Item-to-item agreement is necessary [21], as high agreement between overall subscale scores can be obtained even when corresponding items within the subscale are answered differently across the two administrations.

In terms of the validity of the PROMs developed to assess patient-centred care, most of the measures met the criteria considered adequate for face/content validity (15 of 21 measures) and construct validity (18 of 21 measures). Factor analysis was the most common strategy adopted to measure construct validity (16 measures), however, few studies indicated whether eigenvalues >1 [24] were achieved [33,41,44,47,48,50,53]. Eigenvalues are used to determine the number of subscales within the measure by applying the eigenvalues >1 rule which produces psychometrically reliable and psychologically meaningful results [24]. Thus improvements to reporting whether eigenvalues were >1 appears necessary for PROMs that examine patient-centred cancer care.

The context in which these PROMs assessed patientcentred cancer care should be considered. Most measures were developed with cancer patients recruited from hospitals or treatment centres [33,34,38-44,46-50,53-55]. Only one measure was developed with patients recruited via a population-based cancer registry [32], despite benefits of such recruitment including the ability to sample a representative group of patients at different stages of the disease and with varied experiences of cancer care [61]. Although measuring the quality of patient-centred cancer care during initial treatment and hospital visits is crucial, undertaking such assessments with cancer survivors who no longer visit the hospital regularly is also important. For instance, women diagnosed with breast cancer have reported that the quality and duration of their follow-up consultations with clinicians had declined compared to the quality and duration of their initial treatment experiences [62].

The limitations of this review include that studies available in a non-English language peer-reviewed journal and the grey literature were excluded which could have led to some bias in the findings. Furthermore, the survey developers' reasons for constructing the PROM's should be considered. It is possible that the PROM's objective may have been to focus on specific features of patient-centred care rather than to include items that covered the IOM's six dimensions of patient-centred care. This may explain why most PROMs did not adequately address

the IOM's six dimensions of patient-centred care. Additionally, insufficient or unavailable reporting of the 21 PROMs' psychometric properties may have influenced the ratings regarding the adequacy of the measure's psychometric properties. We did not contact the authors of each PROM to enquire if additional unpublished psychometric information was available for that measure.

# **Conclusions**

Quality improvements to the health care system can be guided by PROMs assessing the quality of patient-centred cancer care. The Indicators (Non-small Cell Lung Cancer) measure [40] was the only identified PROM that included questions relating to the six IOM endorsed dimensions of patient-centred care [1], however psychometric inadequacies and/or incomplete reporting indicates that further psychometric testing of this measure is required. Using more than one measure or further developing existing measures to include all six patient-centred care dimensions could improve the assessment and the delivery of patient-centred care. Additionally, given the lack of psychometrically rigorous PROMs developed to assess patient-centred cancer care that capture the six IOM dimensions, the construction of new comprehensive measures whose psychometric properties are adequate may also be warranted.

#### Abbreviations

 $\hbox{IOM: Institute of Medicine; PROMs: Patient-reported outcome measures.} \\$ 

#### Competing interests

The authors declare that they have no competing interests.

#### Authors' contributions

FT, SKR, RWSF, TCM, MLC and CLP were involved in study conception and design of the systematic review. FT and SKR undertook literature searches, coded the studies for eligibility and evaluated and extracted information from eligible studies. FT drafted the manuscript. All the authors revised the article critically and approved the final version of the manuscript.

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#### References

- Institute of Medicine: Crossing The Quality Chasm: A New Health System for the 21st Century. Washington DC: National Academy Press; 2001.
- Gerteis M, Edgman-Levitan S, Daley J: Through the Patient's Eyes. Understanding and Promoting Patient-centered Care. San Francisco, CA: Jossey-Bass; 1993.
- 3. Hayman JA: Measuring the quality of care in radiation oncology. Semin Radiat Oncol 2008, 18:201–206.
- 4. Fink R: Pain assessment: the cornerstone to optimal pain management. BUMC Proceedings 2000, 13:236–239.

- Wong WS, Fielding R: The association between patient satisfaction and quality of life in Chinese lung and liver cancer patients. Med Care 2008, 46:293–302.
- Von Essen L, Larsson G, Oberg K, Sjoden PO: 'Satisfaction with care': associations with health-related quality of life and psychosocial function among Swedish patients with endocrine gastrointestinal tumours. Eur J Cancer Care 2002, 11:91–99.
- Kim S, Bae J-M, Kim Y-W, Ryu KW, Lee JH, Noh J-H, Sohn T-S, Homg S-K, Lee MK, Park SM, Yun YH: Self-reported experience and outcomes of care among stomach cancer patients at a median follow-up time of 27 months from diagnosis. Support Care Cancer 2008, 16:831–839.
- Frojd C, Lampic C, Larsson G, von Essen L: Is satisfaction with doctors' care related to health-related quality of life, anxiety and depression among patients with carcinoid tumours? A longitudinal report. Scand J Caring Sci 2009. 23:107–116.
- Mager WM, Andrykowski MA: Communication in the cancer 'bad news' consultation: patient perceptions and psychological adjustment. Psychoencology 2002, 11:35–46.
- Bickell NA, Weidmann J, Fei K, Lin JJ, Leventhal H: Underuse of breast cancer adjuvant treatment: patient knowledge, beliefs, and medical mistrust. J Clin Oncol 2009, 27:5160–5167.
- Kahn KL, Schneider EC, Malin JL, Adams JL, Epstein AM: Patient centered experiences in breast cancer: predicting long-term adherence to tamoxifen use. Med Care 2007, 45:431–439.
- Mandelblatt JS, Sheppard VB, Hurria A, Kimmick G, Isaacs C, Taylor KL, et al: Breast cancer adjuvant chemotherapy decisions in older women: the role of patient preference and interactions with physicians. J Clin Oncol 2010, 28:3146–3153.
- Shin DW, Park JH, Shim EJ, Hahm MJ, Park EC: Predictors and outcomes of feeling of insufficient consultation time in cancer care in Korea: results of a nationwide multicenter survey. Support Care Cancer 2012, 20:1965–1973.
- Kowalski C, Nitzsche A, Scheibler F, Steffen P, Albert U-S, Pfaff H: Breast cancer patients' trust in physicians: the impact of patients' perception of physicians' communication behaviors and hospital organizational climate. Patient Educ Couns 2009, 77:344–348.
- Crow R, Gage H, Hampson S, Hart J, Kimber A, Storey L, Thomas H: The measurement of satisfaction with healthcare: implications for practice from a systematic review of the literature. *Health Technol Assess* 2002, 6:1–244.
- Cleary PD, Edgman-Levitan S: Health care quality. Incorporating consumer perspectives. JAMA 1997, 278:1608–1612.
- Mitchell PH, Heinrich J, Moritz P, Hinshaw AS: Measurement into practice. Summary and recommendations. Med Care 1997, 35(Suppl 11):NS124–127.
- Lohr KN, Aaronson NK, Alonso J, Burnam MA, Patrick DL, Perrin EB, Roberts JS: Evaluating quality-of-life and health status instruments: development of scientific review criteria. Clin Ther 1996, 18:979–992.
- McDowell I: Measuring Health: A Guide to Rating Scales and Questionnaires. New York: Oxford University Press; 2006.
- 20. Marx RG, Menezes A, Horovitz L, Jones EC, Warren RF: A comparison of two time intervals for test-retest reliability of health status instruments. *J Clin Epidemiol* 2003, **56**:730–735.
- Viswanathan M: Measurement Error and Research Design. CA: Sage Publications; 2005.
- Anastasi A, Urbina S: Psychological Testing. Prentice Hall: Upper Saddle River, NJ: 1997.
- Cohen J: Statistical Power Analysis for the Behavioural Sciences. Hillsdale, NJ: Erlbaum; 1988.
- 24. Kaiser HF: The application of electronic computers to factor analysis. *Educ Psychol Meas* 1960, **20**:141–151.
- Castle NG, Brown J, Hepner KA, Hays RD: Review of the literature on survey instruments used to collect data on hospital patients' perceptions of care. Health Serv Res 2005, 40:1996–2017.
- Rubin HR: Patient evaluations of hospital care. A review of the literature. Med Care 1990, 28(Suppl 9):S3–9.
- Bredart A, Sultan S, Regnault A: Patient satisfaction instruments for cancer clinical research or practice. Expert Rev 2010, 10:129–141.
- Hudon C, Fortin M, Haggerty JL, Lambert M, Poitras ME: Measuring patients' perceptions of patient-centered care: a systematic review of tools for family medicine. Ann Fam Med 2011, 9:155–164.
- Hearn J, Higginson IJ: Outcome measures in palliative care for advanced cancer patients: a review. J Public Health Med 1997, 19:193–199.

- Clinton-McHarg T, Carey M, Sanson-Fisher R, Shakeshaft A, Rainbird K: Measuring the psychosocial health of adolescent and young adult (AYA) cancer survivors: a critical review. Health Qual Life Outcomes 2010, 8:25.
- Poinsot R, Altmeyer A, Conroy T, Savignoni A, Asselain B, Léonard I, Marx E, Cosquer M, Sévellec M, Gledhill J: Multisite validation study of questionnaire assessing out-patient satisfaction with care questionnaire in ambulatory chemotherapy or radiotherapy treatment. *Bull Cancer* 2006, 93:315–327.
- Arora NK, Reeve BB, Hays RD, Clauser SB, Oakley-Girvan I: Assessment of quality of cancer-related follow-up care from the cancer survivor's perspective. J Clin Oncol 2011, 29:1280–1289.
- Young JM, Walsh J, Butow PN, Solomon MJ, Shaw J: Measuring cancer care coordination: development and validation of a questionnaire for patients. BMC Cancer 2011, 11:298.
- Fitch F, McAndrew A: A performance measurement tool for cancer patient information and satisfaction. J Cancer Educ 2011, 26:612–618.
- Trask P, Tellefsen C, Epspindle D, Getter C, Hsu M: Psychometric validation of the Cancer Therapy Satisfaction Questionnaire. Value Health 2008, 11:669–679.
- Abetz L, Coombs JH, Keininger DL, Earle CC, Wade C, Bury-Maynard D, Copley-Merriman K, Hsu M-A: Development of the cancer therapy satisfaction questionnaire: item generation and content validity testing. Value Health 2005, 8(Suppl 1):S41–53.
- Damman OC, Hendriks M, Sixma HJ: Towards more patient centred healthcare: a new Consumer Quality Index instrument to assess patients' experiences with breast care. Eur J Cancer 2009, 45:1569–1577.
- Bredart A, Bottomley A, Blazeby JM, Conroy T, Coens C, D'Haese S, et al: An
  international prospective study of the EORTC cancer in-patient satisfaction
  with care measure (EORTC IN-PATSAT32). Eur J Cancer 2005, 41:2120–2131.
- Ouwens MM, Marres HA, Hermens RR, Hulscher MM, van den Hoogen FJ, Grol RP, Wollersheim HC: Quality of integrated care for patients with head and neck cancer: development and measurement of clinical indicators. Head Neck 2007, 29:378–386.
- Ouwens M, Hermens R, Hulscher M, Vonk-Okhuijsen S, Tjan-Heijnen V, Termeer R, Marres H, Wollersheim H, Grol R: Development of indicators for patient-centred cancer care. Support Care Cancer 2010, 18:121–130.
- Harley C, Adams J, Booth L, Selby P, Brown J, Velikova G: Patient experiences of continuity of cancer care: development of a new Medical Care Questionnaire (MCQ) for oncology outpatients. Value Health 2009, 12:1180–1186.
- 42. Smith MY, Winkel G, Egert J, Diaz-Wionczek M, DuHamel KN: Patient-physician communication in the context of persistent pain: validation of a Modified Version of the Patients' Perceived Involvement in Care Scale. *J Pain Symptom Manage* 2006, 32:71–81.
- Radwin L, Alster K, Rubin KM: Development and testing of the Oncology Patients' Perceptions of the Quality of Nursing Care Scale. Oncol Nurs Forum 2003, 30:283–290.
- 44. Beck SL, Towsley GL, Pett MA, Berry PH, Smith EL, Brant JM, Guo J-W: Initial psychometric properties of the Pain Care Quality Survey (PainCQ). *J Pain* 2010, 11:1311–1319.
- Beck SL, Towsley GL, Berry PH, Brant JM, Smith EM: Measuring the quality of care related to pain management: a multiple-method approach to instrument development. Nurs Res 2010, 59:85–92.
- Kleeberg UR, Tews JT, Ruprecht T, Hoing M, Kuhlmann A, Runge C: Patient satisfaction and quality of life in cancer outpatients: results of the PASQOC study. Support Care Cancer 2005, 13:303–10.
- Jean-Pierre P, Fiscella K, Freund KM, Clark J, Darnell J, Holden A, Post D, Patierno SR, Winters PC, Patient Navigation Research Program Group: Structural and reliability analysis of a patient satisfaction with cancer-related care measure: a multisite patient navigation research program study. Cancer 2011, 117:854–861.
- Takayama T, Yamazaki Y, Katsumata N: Relationship between outpatients' perceptions of physicians' communication styles and patients' anxiety levels in a Japanese oncology setting. Soc Sci Med 2001, 53:1335–1350.
- Tarrant C, Baker R, Colman AM, Sinfield P, Agarwal S, Mellon JK, Steward W, Kockelbergh R: The prostate care questionnaire for patients (PCQ-P): reliability, validity and acceptability. BMC Health Serv Res 2009, 9:199.
- de Kok M, Sixma HJM, van der Weijden T, Kessels AGH, Dirksen CD, Spijkers KFJ, et al: A patient-centred instrument for assessment of quality of breast cancer care: results of a pilot questionnaire. Qual Saf Health Care 2010, 19:e40.

- 51. de Kok M, Scholte RW, Sixma HJ, van der Weijden T, Spijkers KF, van de Velde CJ, Roukema JA, ven der Ent FW, Bell AV, von Meyenfeldt MF: The patient's perspective of the quality of breast cancer care. The development of an instrument to measure quality of care through focus groups and concept mapping with breast cancer patients. Eur J Cancer 2007. 43:1257–1264.
- van Weert JCM, Jansen J, de Bruijn GJ, Noordman J, van Dulmen S, Bensing JM: QUOTEchemo: a patient-centred instrument to measure quality of communication preceding chemotherapy treatment through the patient's eyes. Eur J Cancer 2009, 45:2967–2976.
- Defossez G, Mathoulin-Pelissier S, Ingrand I, Gasquet I, Sifer-Riviere L, Ingrand P, Salamon R, Migeot V, the REPERES research network: Satisfaction with care among patients with non-metastatic breast cancer: development and first steps of validation of the REPERES-60 questionnaire. BMC Cancer 2007. 7:129.
- Llewellyn C, Horne R, McGurk M, Weinman J: Development and preliminary validation of a new measure to assess satisfaction with information among head and neck cancer patients: the Satisfaction with Cancer Information Profile (SCIP). Head Neck 2005, 28:540–548.
- Bredart A, Morvan E, Savignoni A, Giraud P, Respiratory Gated Radiotherapy Study Group S-R: Patient's perception of care quality during radiotherapy sessions using respiratory gating techniques: validation of a specific questionnaire. Cancer Invest 2011, 29:145–152.
- Jayasekara H, Rajapaksa L, Bredart A: Psychometric evaluation of the European Organization for Research and Treatment of Cancer in-patient satisfaction with care questionnaire ('Sinhala' version) for use in a South-Asian setting. Int J Qual Health Care 2008, 20:221–226.
- 57. Jacobsen R, Samsanaviciene J, Liuabarskiene Z, Sciupokas A: Barriers to pain management among Lithuanian cancer patients. *Pain Pract* 2010, **10**:145–157.
- Can G, Akin S, Aydiner A, Ozdilli K, Durna Z: Evaluation of the effect of care given by nursing students on oncology patients' satisfaction. Eur J Oncol Nurs 2008, 12:387–392.
- Streiner D, Norman G: Health Measurement Scales: A Practical Guide to their Development and Use. New York: Oxford University Press, Fourth; 2008.
- Terwee CB, Bot SD, de Boer MR, van der Windt DA, Knol DL, Dekker J, Bouter LM, de Vet HC: Quality criteria were proposed for measurement properties of health status questionnaires. J Clin Epidemiol 2007, 60:34–42.
- Sanson-Fisher R, Carey M, Mackenzie L, Hill D, Campbell S, Turner D: Reducing inequities in cancer care: the role of cancer registries. Cancer 2009. 115:3597–3605.
- Lawler S, Spathonis K, Masters J, Adams J, Eakin E: Follow-up care after breast cancer treatment: experiences and perceptions of service provision and provider interactions in rural Australian women. Support Care Cancer 2011, 19:1975–1982.
- The Prostate Care Questionnaire for Patients (PCQ-P). http://winden.co.uk/ Surveys/Prostate%20Disease/Appendix%2023%20Patient%20v5.1%2008-02-08%2020-02-08.pdf. Accessed 16 November 2012. September 2007.
- Bredart A, Mignot V, Rousseau A, Dolbeault S, Beauloye N, Adam V, Elie C, Leonard I, Asselain B, Conroy T: Validation of the EORTC QLQ-SAT32 cancer inpatient satisfaction questionnaire by self- versus interviewassessment comparison. Patient Educ Couns 2004, 54:207–212.
- Bredart A, Razavi D, Delvaux N, Goodman V, Farvacques C, Van Heer C: A comprehensive assessment of satisfaction with care for cancer patients. Support Care Cancer 1998, 6:518–523.

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