Information and communication technology for managing pain in palliative care: a review of the literature

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ABSTRACT

Background Information and communication technology (ICT) systems are being developed for electronic symptom reporting across different stages of the cancer trajectory with research in palliative care at an early stage.

Aim/design This paper presents the first systematic search of the literature to review existing ICT systems intended to support management of pain in palliative care patients with cancer. The review was conducted according to Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines for systematic reviews and meta-analyses.

Data sources Four databases (Embase, MEDLINE, PsycINFO and Healthcare Management Information Consortium) from 1990 to December 2012 were searched, with exclusion of papers based on their description of ICT systems and language used.

Results 24 articles met the inclusion criteria, many of which reported the use of non-experimental research designs. Studies were identified at different stages of development with no systems having reached implementation. Most systems captured pain as part of quality-of-life measurement with wide variation in approaches to pain assessment.

Conclusions ICT systems for symptom reporting are emerging in the palliative care context. Future development of ICT systems need to increase the quality and scale of development work, consider how recommendations for pain measurement can be integrated and explore how to effectively use system feedback with patients.

INTRODUCTION

Information and communication technology (ICT) includes all digital technologies that facilitate the electronic capture, processing, storage and exchange of information.1 The application of ICT in healthcare settings in the UK has been highlighted as a means of improving patient outcomes2 and ensuring that patients receive high-quality care.3 Electronic systems have been developed that use ICT to facilitate the capture of clinical data directly from patients, with early indications of patient acceptance of this approach.4 While systems are being developed to support different stages of the illness trajectory in cancer, the use of ICT to capture clinical data specifically in palliative care is at an early stage.5 Patients engaged with palliative care experience multiple physical symptoms,6 which affect their quality-of-life and psychological well-being.7 Pain is reported by 64% of patients with advanced cancer,68 but undertreatment is common.9 Frequently cited barriers to adequate pain management include knowledge deficits, inadequate pain assessment and misconceptions regarding pain.10 The use of ICT could be seen as an approach to address pain management barriers by facilitating greater communication between patients and health professionals contributing to the delivery of palliative care.11 There is scope for the development and implementation of online palliative care symptom reporting systems in which such timeliness and methods of communication can be explored, with exciting opportunities for future development.12
of pain in palliative care patients with cancer. The identification of implemented systems or those under development to support pain reporting by palliative care patients will indicate the status of system development at present and inform future work to improve palliative pain management using ICT systems. Rather than seeking to review the benefits of ICT systems and the quality of their effectiveness on implementation, this paper seeks to outline their structure. Palliative care has been acknowledged as posing its own specific challenges to systematic review methods, with an evidence base largely unsuitable for traditional forms of review and synthesis. The approach taken by the current review is to better understand the types of available systems through a descriptive analysis, rather than providing an assessment of the levels of quality or bias of available evidence.

METHOD
A systematic review of qualitative and quantitative literature was undertaken to identify the function and structure of existing ICT systems used in the management of pain in palliative care patients. This review takes a broad search of ICT systems used in patients with cancer with interpretation of findings orientated towards understanding the use of ICT systems in palliative care. The review was undertaken in five stages: (1) development of search strategy; (2) generation of inclusion and exclusion criteria; (3) assessment of relevance; (4) data extraction and tabulation; and (5) overview of system characteristics and descriptive analysis.

Search strategy
The search strategy was compiled to identify published reports of ICT systems for pain management. MeSH headings and keywords were identified (as outlined in online supplementary appendix A) and relevant databases were selected and searched in consultation with a healthcare information specialist based at the University of Leeds. The databases Embase, MEDLINE, PsycINFO and Healthcare Management Information Consortium were searched for literature published between 1990 and December 2012. The following journals were hand searched for relevant articles: Journal of Pain and Symptom Management, Supportive Care in Cancer, European Journal of Pain and BMJ Supportive & Palliative Care. Potentially relevant references from bibliographies and citation indices were identified and abstracts assessed against selection criteria. The literature search was carried out during December 2012.

Inclusion and exclusion criteria
Inclusion criteria were developed through consensus by the research team. To be included articles had to describe the use of ICT in the management of pain in patients with cancer. The term ICT was chosen intentionally as a broad term applying to a wide range of technology use in healthcare to allow capture of existing systems documented in published literature. Management in the current search refers to the identification, assessment or monitoring of pain in patients with cancer. To be considered a management tool, the ICT system must capture information generated by patients, which is shared, reviewed or assessed by a health professional.

Papers were excluded that (i) provide no description of system structure and content; (ii) describe systems that were not designed for, or used by, patients with cancer pain; and (iii) discuss cancer pain, but not involving the use of ICT in pain management. Due to resource limits, it was intended that non-English articles be removed during the assessment of article relevance.

Study selection
Titles and abstracts were screened for relevant articles, for which full-text reports were sought. Full-text reports were assessed against the inclusion/exclusion criteria to identify eligible reports for the descriptive analysis. Details of the study identification and selection process are shown in the flowchart in figure 1, developed using Preferred Reporting Items for Systematic Reviews and Meta-Analyses guidelines.

Data extraction, appraisal and synthesis
Data were extracted by MJA (lead author) into a tabulated form (see online supplementary appendix B) and checked by a second reviewer. Data were extracted from full-text reports meeting the inclusion criteria. To describe the stage of system development, the human-centred systems development methodology was used for categorising identified studies. This methodology comprises (i) planning and project selection, (ii) analysis, (iii) design and (iv) implementation. Planning and project selection refers to defining the problem and considering possible solutions. Analysis relates to examination of patient expectations and requirements for a proposed system, including early usability work. The design stage involves testing systems with patients and evaluating ease of use and preferences for layout. The implementation stage refers to an evaluative component of development and as such is referred to as the evaluation stage in the article, which includes pilot studies using a system in clinical practice (eg, in cancer centres) and summative evaluations of patient experience of using systems as part of their care.

RESULTS
The search identified 1898 citations initially, of which 24 articles met the inclusion criteria (see figure 1). A large proportion of studies were rejected following title and abstract screening (n=1456). As shown in figure 1, several full-text sources (n=76) were
examined, from which n=53 studies were excluded. Reasons for exclusion of full-text sources included not focusing on cancer pain, discussion of systems that were not designed for communication of pain data (eg, virtual reality simulators) and systems that did not involve any interaction with a patient. Our assessment of titles and abstracts identified no non-English papers that met the inclusion criteria. Online supplementary appendix B contains an overview of all included papers and outlines the various ICT systems described. The descriptive analysis of included articles is organised into three categories: (i) study design, location and population; (ii) system design and development; and (iii) communication facilitation.

Study design, location and population

Study design

Of the 24 included studies, 7 unique study designs were described: randomised experiment, non-randomised experiment, observational, survey design, system description, expert opinion and a non-specified non-experimental study design. The largest proportion of included papers used a non-specified non-experimental study design, including reports of patients completing usability evaluations or providing feedback on specific system features using a range of methods including qualitative interviews and descriptive evaluation.

Location of system

Of the 24 included studies, 17 independent systems were identified. As shown in table 1, the majority of systems (n=10) were identified from research that was conducted in the USA. Other countries in which systems were identified include Sweden, the UK, Austria, Germany, Japan and Norway.

Target population for system

Of the 24 included studies, the target population or intended users of systems were oncology patients, palliative home care patients, oncology outpatients, hospice and palliative care patients, patients undergoing palliative radiotherapy, a mix of palliative care patients, carers and health professionals, brain tumour outpatients, patients requiring surgery for cancer, patients with cancer with incurable metastatic or locally advanced disease, patients receiving chemotherapy for cancer, breast cancer survivors, patients with cancer with depression and/or pain, and a mix of members of the public with pain and people with cancer pain.
Sample population

Fifteen of the twenty-four included studies recruited patients with cancer only, the remaining articles recruited mixed groups of health professionals and patients, patients with cancer and the general public, and clinicians. In the two studies that provided overviews of new and emerging systems, no research participants were involved.

System design and development

Figure 2 outlines the classification of included articles in the stages of system development. Of the 24 included studies, no papers discussed the planning stages of systems, 11 studies detail systems in the analysis stage, 4 studies described the design stage, and 9 studies described the evaluation stage. Only two systems, from across five articles, are represented at more than one stage of development, classified in both analysis and evaluation stages. There are a number of systems (n=4) in the evaluation category that do not have representation in earlier categories.

System focus and the measurement of pain
The focus of systems differed across the 24 included studies. Six studies documented systems that focused on the capture of pain information only, while three systems considered the management of pain and depression together as part of a collaborative care approach. The remaining 15 studies outlined systems that considered pain amid a wider range of symptoms in the context of quality-of-life measurement.

A variety of measures to capture pain information were incorporated into systems described in the included articles as outlined in table 2. Systems only considering pain incorporated the McGill Pain Questionnaire, bespoke visual analogue scales...
asking patients to measure pain intensity and presentation of visual images of an affected region of the body for patients to indicate the location of pain. Papers outlining a system for capturing pain and depression comprised a mix of items taken from the PHQ-9 depression scale, BPI, single items about global improvement, medication adherence, side effects and whether a call from a nurse was required. Additional BPI questions were included for patients with pain, and additional PHQ-9 questions for those with depression.

Systems measuring pain as part of a wider collection of symptoms used a variety of existing and bespoke symptom and quality-of-life measurement tools. The C-SAS, EORTC QLQ-C30, EORTC BN20, MDASI, M.D. Anderson Symptom Inventory, MPQ, McGill Pain Questionnaire, PHQ9, Brief Patient Health Questionnaire for depression, PRO, patient-reported outcome; SCFS-6, Schwartz cancer fatigue scale; SGA, Subjective Global Assessment of Nutritional Intake; VAS, Visual analogue scale.

Communication facilitation As shown in table 3, of the 24 included studies, the intended location of use of systems for patients varied between waiting room (n=12), home setting (n=8) and remote use in the community (n=4). Systems intended for use in the waiting room capture and collate information for review by a health professional. No consistency in the use of measures for capturing pain reports from patients was evident, and there was variation in the compilation and presentation of measures of pain across all systems independent of the system focus.

Table 2 Overview of tools used for pain measurement in the identified systems

<table>
<thead>
<tr>
<th>System focus</th>
<th>Reference</th>
<th>Measure(s) used to capture pain reports from a patient</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pain</td>
<td>30</td>
<td>Computerised extension of the MPQ</td>
</tr>
<tr>
<td></td>
<td>21</td>
<td>Areas of pain and no pain were digitised into a summation image of a female torso, with blackened areas indicating pain areas and white indicating no pain</td>
</tr>
<tr>
<td></td>
<td>33</td>
<td>A pain diary form consisting of a VAS (0–100 mm) for pain intensity and boxes for the number of consumed extra doses of analgesics</td>
</tr>
<tr>
<td></td>
<td>34</td>
<td>Not described in paper</td>
</tr>
<tr>
<td></td>
<td>35</td>
<td>The pain diary included a unidimensional tool for assessment of pain intensity using a 100 mm VAS where the scale was anchored by the words ‘No pain’ (VAS 0 mm) and ‘Severe pain’ (VAS 100 mm). It also included a question about consumed extra doses of analgesics</td>
</tr>
<tr>
<td></td>
<td>38</td>
<td>Computerised extension of the MPQ and 16 additional items: two items related to the body outline, three 0–10 items related to current pain, least pain and worst pain intensity, three items measuring goals for pain levels and amount of time pain greater than tolerable level, one item measuring patient satisfaction with pain level, and one item addressing a range of concepts (expectations about the pain, effectiveness of previous pain treatments, pain medication treatment pattern, non-drug treatments used for pain, tendency to tell or not tell others about the pain, onset of pain and beliefs about the cause of the pain)</td>
</tr>
<tr>
<td>Pain and depression</td>
<td>18</td>
<td>7 core items, 2 items from PHQ-9, average pain from BPI, single items about global improvement, medication adherence, side effects and whether a call from a nurse was required. Additional BPI questions were included for patients with pain, and additional PHQ-9 questions for those with depression</td>
</tr>
<tr>
<td>Quality-of-life</td>
<td>26</td>
<td>86 items, reflecting common cancer-related and treatment-related symptoms, including pain</td>
</tr>
<tr>
<td></td>
<td>27</td>
<td>An 11-point (0–10) pain-intensity numerical scale as an additional item</td>
</tr>
<tr>
<td></td>
<td>23</td>
<td>Not described in paper</td>
</tr>
<tr>
<td></td>
<td>16</td>
<td>MDASI, a brief, validated measure of 13 common cancer-related symptoms over the previous 24 h. Each symptom is rated on an 11-point scale, with 0 being ‘not present’ and 10 being ‘as bad as you can imagine’</td>
</tr>
<tr>
<td></td>
<td>25</td>
<td>ESAS to be completed daily and the EQ-5D weekly</td>
</tr>
<tr>
<td></td>
<td>24</td>
<td>Symptom-specific questions, including an 11-point scale for reporting pain level</td>
</tr>
<tr>
<td></td>
<td>20</td>
<td>QOL assessment EORTC QLQ-C30 and the EORTC BN20</td>
</tr>
<tr>
<td></td>
<td>28</td>
<td>EORTC QLQ-C30</td>
</tr>
<tr>
<td></td>
<td>29</td>
<td>Intensity of symptoms (pain, fatigue, nausea, anxiety, depression and drowsiness) rated using a 0 to 100 VAS</td>
</tr>
<tr>
<td></td>
<td>27</td>
<td>ESAS, EORTC QLQ-C30, screening item about pain intensity at its worst in the last 24 h (taken from BPI), 15 items on physical function, PHQ-9, two questions from the SGA, five questions related to need for assistance, patients’ opinions on time expenditure and preferences for computerised vs paper and pencil-based assessment, and a computerised pain body map</td>
</tr>
<tr>
<td></td>
<td>31</td>
<td>Visual display of relevant information from multiple sources can be captured including PRO assessments, patient history and notes</td>
</tr>
<tr>
<td></td>
<td>32</td>
<td>Aches and pain captured as part of C-SAS</td>
</tr>
<tr>
<td></td>
<td>36</td>
<td>A patient-reported symptom severity screening scale that generates a real-time, point-of-care report. Contains 38 items that produce scores for individual patient-reported problems</td>
</tr>
<tr>
<td></td>
<td>22</td>
<td>As above</td>
</tr>
<tr>
<td></td>
<td>37</td>
<td>An electronic version of the 1970 edition of the MPQ, a pain barriers questionnaire and the SCFS-6</td>
</tr>
</tbody>
</table>

BPI, Brief Pain Inventory; C-SAS, Chemotherapy Symptom Assessment Scale; EORTC BN20, European Organisation for Research and Treatment of Cancer brain cancer module; EORTC QLQ-C30, European Organisation for Research and Treatment of Cancer Quality-of-Life Questionnaire; EQ-5D, EuroQol European Quality-of-Life-5 Dimensions; ESAS, Edmonton Symptom Assessment Scale; MDASI, M.D. Anderson Symptom Inventory; MPQ, McGill Pain Questionnaire; PHQ9, Brief Patient Health Questionnaire for depression; PRO, patient-reported outcome; SCFS-6, Schwartz cancer fatigue scale; SGA, Subjective Global Assessment of Nutritional Intake; VAS, Visual analogue scale.
professional, typically prior to consultation. Where the intended location for patients is the home setting,16 23 25 29 32–35 ICT systems add to an existing dialogue about pain or its changing nature. Systems in this category typically involve the monitoring of pain symptoms, either actively by nursing staff or through thresholds that generate alerts to health professionals when patient scores are reported outside of a predetermined acceptable range. Systems intended for remote use by patients17–19 24 provide flexibility in ways to report pain using web-based systems alongside automated or patient-initiated symptom reporting via landline or computer. One system24 was specifically designed for use with hospice and palliative care patients to accommodate a situation where patient status may change rapidly and/or need regular monitoring.

System feedback

Systems described in the included studies facilitated two forms of communication: patient to health professional (n=15) (where a patient enters data that are sent directly to a health professional without receiving feedback),5 16 18–23 26 27 29 31 33 35 36 and feedback of information presented to patient and health professional following patient entry of information (n=8).17 24 25 28 30 32 34 37 One paper38 did not report on the feedback provided by the system.

Systems facilitating patient to health professional communication5 16 18–23 26 27 29 31 33 35 36 did not create any additional dialogue between these two groups. Systems informed or alert ed health professionals to the experience of pain in a patient, which may be used to impact on decision making regarding pain management. This feedback provided only to the health professional was in the form of patient score summaries that could be printed, quality-of-life profiles showing longitudinal charts or graphical summaries for patients, and summation pictures of areas where pain is reported. The modes of technology that were used to capture data from patients for this purpose included landline and mobile telephone, and handheld devices (such as tablet computers).

Systems providing feedback to patients and health professionals (n=8) provided different forms of output for each group. For patients, these systems reported one or more of the following types of feedback:
1. specific self-care advice when symptoms are reported out of a predefined range
2. graphical display of scores over time
3. tailored educational materials relating to symptoms
4. personalised feedback from a nurse (via phone or online)

A combination of these types of feedback was present in one system,24 which provided presentations of scores over time and tailored educational materials. For health professionals, the types of information provided through this category of system are similar to those providing information from a patient to a health professional only. For example, patient scores can be printed out and reviewed by a health professional or used during a consultation with a patient. Additional features were also identified in this category of systems, with the use of predefined ranges set to prompt email alerts. One system20 used the information reported by a patient to produce summaries of pain data and an evidence-based treatment plan to support clinical management decisions.

**DISCUSSION**

This review has found that few ICT systems for pain reporting in palliative care patients are evaluated in the context of experimental research and no systems are reported to be implemented in clinical practice. Over half of the articles reviewed describe systems that were in the early stages of design and provided little detail on the overall development approach being taken. When identifying how ICT systems facilitate pain management, it was found that most ICT systems conduct pain assessment in the context of wider quality-of-life measurement in patients with cancer. A focus only on pain symptoms was seen in a small proportion of the included ICT systems, where reports were collected directly from the patient for use by a health professional with no feedback provided to the patient. The possibilities of feedback that can be generated by ICT systems have only begun to be explored with patients and health professionals with a small number of systems.

ICT system development is an emerging area of research in healthcare that may reflect the low numbers of systems identified across the stages of

<table>
<thead>
<tr>
<th>Location for Health Professional</th>
<th>Location for patient</th>
<th>Home setting</th>
<th>Remote use in community</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clinical environment</td>
<td>(n=12) 20–22 25 26 28 30 31 37 38 48</td>
<td>(n=5) 25 32–35</td>
<td>(n=4) 7–19 24</td>
</tr>
<tr>
<td>Remotely (eg, web-based systems)</td>
<td>No articles</td>
<td>(n=2) 16 23</td>
<td>No articles</td>
</tr>
<tr>
<td>No setting defined</td>
<td>No articles</td>
<td>(n=1) 29</td>
<td>No articles</td>
</tr>
</tbody>
</table>

Table 3 The location for systems to be used by patients and health professionals in the identified articles (numbers correspond to study number in online supplementary appendix B)
system development. This is highlighted through the generally low quality of development research found in symptom-reporting systems.49 Concerns have been raised about the use of technologies in cancer care when implementation occurs prior to established clinical efficacy or use.50 Encouraging structured development of ICT systems with continued dissemination of findings is a strategy that can support high-quality research generation across all areas of application in healthcare.

Many systems identified in this review were designed to capture information from a patient for use by a healthcare professional in a clinical setting, with a patient relaying symptom reports without engaging in active forms of communication. Although cancer symptom reporting tends to occur inside healthcare institutions,51 the unidirectional provision of pain reports to a health professional (via ICT systems) ignores the benefits to both parties (particularly the patient) that could derive from feedback or tailored support in response to data entered into an ICT system. Although feedback was provided by some of the systems identified here, there is now a clear need for future research to explore how ICT system feedback can be used to improve pain management in palliative care.

The lack of consistency in pain measures identified in ICT systems may impact on the assessment and management of patients in palliative care. When used in palliative care, pain assessment tools can include dimensions and items of limited relevance for patients with advanced cancer.52 This can negatively influence patients, such as reducing compliance to assessment, and should be considered to ensure clinical relevance of selected tools and measures. Well-validated, multidimensional assessment tools such as the BPI43 and McGill Short Form Questionnaire53 have been suggested for use in capturing the experience of pain.54

A descriptive analysis of the literature was chosen as a tool for understanding the current state of ICT system development in the literature to inform thinking for future areas of research and development. The scope of systems included in the review was limited to systems reported in research literature. Inclusion of systems was reliant on developers and researchers disseminating the design processes of an ICT system in the research literature. In order to counter this limitation, the search strategy was broadened to include all cancer not just advanced cancer (ie, palliative care). The strength of this is that we are likely to have broadened the search and captured a greater number of ICT systems, but not all included systems have been developed and validated in palliative sample populations that may not accurately reflected our target population (palliative care patients with advanced cancer and pain) in terms of their needs, experiences and symptom profiles.

The use of ICT systems presents an approach to improving the management of pain in palliative care patients with cancer. This is the first review of ICT systems used to manage pain in palliative care. A number of systems were identified in this systematic review of the literature, with systems focusing specifically on pain alongside those capturing more varied measures of quality-of-life. Future ICT system development needs to consider the palliative care context closely to assess how existing and emerging systems can be integrated effectively. ICT systems in palliative care need to increase the quality and scale of development work, adhere to recommended pain measure use and consider how to effectively use feedback to patients. If achieved, ICT system development in palliative care promises a platform on which patients and health professionals can engage in an efficient and meaningful dialogue to improve the management of pain.

Contributors MJA and BMB conceived the paper and generated early drafts of the paper. MJA led the reviewing process. ST, MRM and MIB contributed to the reviewing process and revised later drafts of the paper.

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REFERENCES


Review

42 Aaronson NK, Cull A, Kaasa S, et al. The European Organization of Research and Treatment of Cancer (EORTC)


