Nursing documentation in palliative care: An integrative review

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Abstract

Purpose: Nursing documentation is essential in ensuring communication between team members, continuity of care and evidence based practice. The purpose of this paper is to review and synthesise research pertaining to nursing documentation in specialist palliative care.

Method: An Integrative review (IR) utilising Whittemore and Knafi’s five stage process was employed. Electronic searches of: Scopus, Medline, Cinahl, Web of Science, Academic Search Complete databases (2010-2017) were conducted. 10,842 articles were retrieved which were reduced to five articles for review. Data extraction, quality assessment (Critical Appraisal Skills Programmes – CASP) and thematic analysis were conducted on the included articles.

Results: Two overarching themes emerged in this review, symptom management and engagement. Symptom management focused on documentation of symptoms while engagement highlighted emotional care, information giving, providing support, coordinating care, supporting families and education.

Conclusions: Clear, accurate, and complete documentation is crucial to the delivery of quality health care and pivotal to effective communication within the team. Although this is important in all aspects of care it is arguably even more so in end of life care. In order to audit the care provided, such care must be clearly identified and documented.

Keywords: Palliative care, documentation, care provision, continuous professional development

Introduction

The World Health Organizations [1] position on palliative care describes the total care of patients in order to achieve the best possible quality of life for patients and their families, encompassing care of body, mind, and spirit. Inherent to the delivery of safe, ethical and effective nursing practice is clear accurate and comprehensive documentation [2,3]. Documentation refers to any written information regarding a patient that describes their status or the care/services provided to the patient by a nurse [4]. In addition nursing documentation serves as a legal document, contributes credibility to nursing practices and enhances the professional image and presence of nurses [5]. The nursing process of; assessment, planning, implementation and evaluation has frequently been used as a framework for documentation across various care settings [6]. When caring for a patient, nurses’ documentation provides a clear picture of; the status of the patient, the actions of the nurse and care outcomes [4]. Clear and coherent documentation can prevent negative outcomes caused by miscommunication [7]. Nursing documentation is a communication tool demonstrating not only evidence of what the nurse actually does for the patient, but also provides a tool to audit the written record of the patient's journey [8]. While having clear documentation is an essential requirement for nursing practice, there is also a need for the profession to make their contribution to healthcare visible [9-11].

While nurses recognise that documentation is important, their first priority and focus has been on patient care with documentation a lower priority [12]. Over the last few decades nursing has developed towards independent practice with explicit knowledge of nursing care. However, these developments require nurses’ not only to perform interventions but also document their decision making processes and identify why
a nursing action has been prompted [13]. Nursing practice requires documentation to ensure continuity of care, planning, and accountability, as well as the promotion and uptake of evidence-based practice [14]. Nursing documentation provides an efficient way to communicate crucial patient information with members of the healthcare team [3]. Although this is important in all aspects of care it is arguably even more so in end of life care [15] as this may be the last opportunity to provide patients and their families with ‘impeccable assessment and management of symptoms’ [1] and thus positively influencing the grieving processes following a death. In nursing practice physical care needs are generally easy to identify and document. However, other aspects of care may be less easily identified and communicated to others for example; psychological care (empathy, communication, comfort, and support), social care (isolation/connections, feeling safe, effects of role changes on individuals and patient being involved in his/her own care) and spiritual care (being present, applying meaning to their journey and working with changing hopes).

This paper focuses on palliative care nursing documentation through an integrative review and aims to identify what nurses in this area of care document. Such a review is necessary in order to identify whether or not nursing documentation reflects the provision of palliative care as positioned by the World Health Organization [1].

Methods
The review utilised Whittemore and Knafi [16] framework and was conducted and reported in five stages (problem identification, literature search, data evaluation and extraction, data analysis, and presentation of results). Particular attention was paid to the development of the review question, search strategy, appraisal of study quality and method, identification of common aspects, and formation of themes for presentation.

Stage 1: Problem identification
A clear problem identification and review purpose provided focus, boundaries and facilitated all stages of the review. To assist with the review a clear identification of the problem variables of interest (concepts, target population, health care problem) and appropriate sampling frame (type of empirical studies, inclusion of theoretical literature) was developed. Using PEO framework (Population, Exposure and Outcome [17] Table 1).

Explanatory investigations highlighted a number of different terms encompassing palliative care including: end of life care, terminal care, hospice, life limiting condition, life threatening illness, cancer care, and specialist palliative care. Reviewing a broad range of terms is central in providing a comprehensive understanding of the topic area [18]. Search parameters which guided the inclusion of papers are presented in Table 1. The time period (01 January 2010 to 10 January 2017) was identified as appropriate to ensure currency of relevant literature and reflect developments in palliative care and nursing practice. This seven year search period was chosen to reflect current evidence based knowledge regarding nursing documentation of palliative care and deemed as appropriate given the scope of integrative reviews to include a range of studies for different designs.

Stage 2: Literature search
The data-bases Scopus, Medline, Cinahl, Web of Science and Academic Search Complete were searched within the review parameters (Table 2). Boolean and Truncation were used to capture relevant variations of terms (Table 3). Finally an ancestry search of the reference lists of the identified studies was conducted.

Table 1. PEO Framework formulating the literature review question.

| P - Population  | Palliative care nurses
|-----------------|--------------------------|
| E - Exposure    | ‘care’
| O - Outcome     | ‘documentation’

Table 2. Search parameters.

| Inclusion criteria | Exclusion criteria |
|--------------------|-------------------|
| Nursing documentation in palliative care environments | Studies that reported on advanced care planning, survivorship care planning or cancer care planning |
| Papers from peer reviewed journals published from 01 January 2010- 10 January 2017 | Non-peer reviewed studies |
| Articles written in English | Non-English articles |
| Original quantitative, qualitative research, integrative literature and systematic reviews | Review articles that did not use a systematic process to identify the literature |

Table 3. Search strategy.

Search 1 - nurs* documentation OR nurs* record OR nurs* note* OR care plan* OR nurs* kardex
Search 2 - palliative care OR end of life care OR terminal care OR hospice OR life limiting condition* OR life threatening illness OR cancer care OR specialist palliative care
Search 3 - search 1 + search 2

The results of the computerised search process are presented in PRISMA format in Figure 1 where 5 articles, 4 quantitative studies and 1 qualitative study were identified for inclusion in this review. The articles country of origin included Norway (n=1), Canada (n=1), USA (n=2) and Sweden (n=1). The initial search identified 10842 results from which 2629 duplicates were removed. The remaining 8213 papers were screened broadly by title review and 8013 papers were omitted following this process. Of the remaining 200 papers an abstract review was conducted with 187 papers excluded as not meeting the criteria outlined in Table 2. A full text review of the remaining...
13 articles was completed by two independent assessors to identify if they met the inclusion/exclusion criteria (Table 2). The independent reviewers met to discuss the articles and based on their independent review an agreement was reached that; 8 further papers not meeting the criteria were excluded. The reference lists of the identified studies did not yield any additional papers. The search and application of the inclusion/exclusion criteria resulted in 5 articles identified for inclusion in this review (Figure 1).

**Stage 3: Data evaluation and extraction**

Studies were appraised using the Critical Appraisal Skills Programmes Checklists [20] and graded low, medium or high. However as the review sought to extract key principles from the studies no disqualifications were made on the grounds of quality rather the quality assessment process assisted in building a picture of the underlying assumptions and methods that currently characterise the field. Initial data extraction captured the study characteristics including, setting, design of study, sample strategy, data collection and key issues identified in the research (Table 4) and subsequent data extraction collating findings into themes.

**Stage 4: Data analysis**

Within data analysis, data were ordered, coded, categorised and summarised into a unified and integrated conclusion. Themes were identified from each study and synthesised to form final themes. This was an iterative process of engagement and re-engagement with the studies where the findings of all included studies were carefully read line by line. Extracted information was compared and patterns recorded as they became apparent. This comparative analysis process was further scrutinised and discussed by the reviewers and it was possible to discern groupings of similar information leading to the identification of two key themes. Given the heterogeneity of the included literature meta-analysis was not undertaken and given that only one qualitative study was included meta-synthesis or discourse analysis was not possible. Therefore, a narrative synthesis of extracted data was undertaken (Table 4) with emerging thematic headings. This allowed for coding according to similarities and differences and verification for accuracy and relevancy by all reviewers [21] and data in each

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**Figure 1. PRISMA 2009 Flow Diagram.**

Records identified through database searching (n = 10845)

Records after duplicates removed (n = 8183)

Records screened (n = 200)

Full-text articles assessed for eligibility (n = 13)

Studies included in qualitative synthesis (n = 1)

Studies included in qualitative synthesis (meta-analysis) (n = 6)

Records excluded (n = 187)

- Related to social worker documentation n=1
- Related to end of life care tool n=1
- Related to prevalence and use of electronic documentation n=1
- Related to promoting health at end of life n=1
- Related to surwovice care planning n=1
- Referred to advanced care planning n=1
- Did not refer to nursing documentation n=2
Table 4. Data extraction table.

| Author, Publication year, Title, and Country | Study aim | Methodology/design | Sample strategy and sample size | Data collection method | Analytical approach | Summary of main findings and quality assurance |
|--------------------------------------------|-----------|--------------------|--------------------------------|-----------------------|--------------------|-----------------------------------------------|
| Doody et al. 2018, “Improving documentation of quality measures in the electronic health record” United States | Name Practitioners documentation quality oncology practice incentive (QOPI) measures in the electronic health record (EHR) is evaluated in this study. | Retrospective review of medical charts completed in August 2012. 100 medical charts of Cancer Center patients. | 100 medical charts reviewed for evidence of accurate supportive care QOPI measures. Areas of deficiency in documentation were identified and used to develop educational intervention (EI) for the NP staff. | Data analyzed using SPSS software. The accuracy used in the EI were based on a selected 80% compliance level. | Main Findings: Pre EI chart audit of 100 records was found to have failed below 80% compliance level and identified. Documentation and appropraiteness of management plan for drug resistant moderate to severe pain. Documentation of pain, management and review of intervention, for pain, bowel management, and assessing emotional well-being and oral chemotherapy management. Quality assurance: This paper uses a quality measure that we included in QOPI that requires to obtain a rapid and objective measurement process quality. Limitations were noted by the authors. |
| Fromm, S., Blake, J.P., Stokes, R., Gauthier, F., Hriso Smith, T., Steel, K., Monia, J.N. 2014, “Care planning needs of palliative home care clients” Development of the interRAI palliative care assessment clinical assessment protocols (CAPs) Canada | To describe the interRAI PC CAP development process and provide an overview of the distributional properties of the eight interRAI PC CAPs among community-dwelling adults receiving palliative home care. | Secondary data analysis used interRAI PC assessments (n = 6,799) collected as part of regular clinical practice at baseline (n = 6,799) and follow-up (n = 1,000). Clients across six regional jurisdictions in Ontario, Canada assessed to receive palliative home care services between 2006 and 2011 were included (mean age 70.7 years; 11.3% males). Descriptive analyses focused on the right interRAI PC CAPs: Fatigue, Distress Tolerance, Nutrition, Pressure Ulcers, Pain, Dyspnea, Mood Disturbance and Delirium. | Data were collected from six Community Care Access Centre’s (CCAC) sites located across geographical regions and ranged from primarily metropolitan urban to more rural and geographically dispersed districts. Each CCAC is ranked of syndromes, specialized supports, including palliative care, for persons under its jurisdiction. | De-identified cross-sectional data from 6,799 interRAI PC assessments gathered between 2006 and 2011 from palliative home care clients in Ontario, Canada were included for analysis. | Main Findings: Nearly 9 in 10 persons triggered at least one CAP (87.9% in n = 5,950) and approximately two thirds triggered more than two CAPs. The interRAI PC Clinical Assessment Protocols (CAPs) focus on specific clinical, functional, and life quality issues. Using algorithms embedded within the interRAI PC, the CAPs alert the user to specific problems and indicate whether it is an appearance or potential for improvement. Both can be addressed in the care plan. Each CAP contains a summary statement, goals of care, triggers, and guidelines. Triggering rates ranged from 74% for the Fatigue to less than 12% for the Delirium and Pressure Ulcers. The hierarchical CAP triggering structure suggested Fatigue and Dyspnea CAPs were present issues prevalent among the majority of clients while Delirium and Pressure Ulcers CAPs rarely triggered in isolation. More often trigger within the symptom trajectory. Fatigue was the most commonly triggered CAP, triggered by 98.8% of persons who triggered only one CAP to over 90% of persons who triggered three or more CAPs. Captured in the percentage of persons who triggered only one CAP, the Fatigue CAP, Pain CAP, Nourishment CAP, and Dyspnea CAP emerged as early triggered CAPs. In contrast, the Delirium and Pressure Ulcers CAPs emerged as late triggered CAPs. Consequentially, persons who triggered only one CAP, were more likely to trigger the Fatigue, Pain, Nourishment, or Dyspnea CAPs and were least likely to trigger the Delirium and Pressure Ulcers CAP. Quality assurance: Three phase development process – review, consultation with experts, creation of triggering algorithms. Spearman rank correlations suggest majority of CAPs are reasonably independent from each other (spiceur correlations between Fatigue and...
Continuation of Table 4

| Author(s) | Title and Publication Details | Description | Methodology | Main Findings |
|-----------|-------------------------------|-------------|-------------|--------------|
| Olsen, C.A., Forsberg, C., Bringner, E. (2013) | 'Documentation of Nursing Care in advanced Home Care' Sweden | The aim of this study was to describe documentation of nursing care within AHC. | Sixty nursing records from two AHC-units in Stockholm Sweden were collected. Thirteen patient records were estimated to be two-thirds of the records from patients admitted to the AMC unit at the time of data collection were randomly collected from each unit. The patients had been admitted to AHC between 25 to 208 days, median 124 days, and mean 122 days. | 30 computerized records randomly selected from currently admitted patients. The records were anonymous for the researchers and data processed in a way so that it could not be traced to a specific patient, nurse, or the patient’s family. Document analysis with inspiration from content analysis according to Elo and Kyngäs. The analysis process consisted of three phases: preparing, organizing, and reporting data. Only the manifest content was analyzed using an inductive approach. The analysis was executed manually. |
| Almassalka, F., Xua, D., Keenan, G.M., Khokhar, A., Yao, T., Chen, Y.C., Johnson, A., Assael, R., Wilke, D.J. (2012) | 'Data Mining Nursing Care Plans of End-of-Life (EOL) Hospitalized Patients: A Study to Improve Healthcare Decision Making' USA | Revealing hidden patterns and knowledge present in nursing care information documented with standardized nursing terminologies on end-of-life (EOL) hospitalized patients. | Quantitative analysis of care plans and nursing documentation using association mining, clustering, and classification techniques to glean hidden information. Confidence sampling to select units. Four agreed to participate, two large community hospitals, one university hospital, and one small community hospital. Eight units recruited four times hospitals to participate for either a 12-month (four units) or 24-month (four units) study period. Data derived from a primary dataset gathered on 23,451 medical surgical patients (40,747 unique episodes). With analysts conducted on a subset of 1,425 EOL episodes derived from the primary dataset. The data were extracted from the Hands-On Automated Nursing Data System database of nursing care plans episodes coded with NANDA-I, Nursing Outcomes Classification, and Nursing Intervention Classification (NIC) terminologies. Statistical analysis using SPSS with t-tests or chi-square tests. Two-tailed alpha level of 0.05. | Main findings: Only 41.3% of care plans included pain, 35.5% of patients were discharged to a hospice with severe pain than expected. The most effective nursing intervention classified pain intervention was only used in 81/545 cases. Standardized terminology can more effectively demonstrate nursing knowledge within documentation. Standardized documentation terminology can reduce bias and confounding factors in data mining which can in turn inform clinical decision making in end of life care. Quality assurance: Data mining techniques need to be applied carefully to the underlying datasets. Although generic at some abstract level, these techniques involve many technical steps that need to be tailored to the underlying data, and their complexity varies with the characteristics of the data. |
| Stoeldal, S.A., Wørnøe, L., Sortbye, L., Schou, Bredal, L. | To measure the level of agreement between data from Crossectional study collecting data retrospectively from Sample strategy involved interviewing nurses | The resident assessment instrument for palliative care interviewed data compared with electronic patient | Main Findings: The best agreement between nurse interview data and EPE data were |
Continuation of Table 4

| Larale, A. (2011) | interviews with nurses and data from EPR on the prevalence of physical symptoms, clinical signs and treatment during hospitalised patients’ last three days of life. | August to December 2007 from electronic patient records on 9 wards (four medical wards, one oncology ward, three surgical wards and one intensive care unit). Records of patients (n=112) were reviewed on patients that had died in the hospital wards over the five month period, 57 female and 55 male patients. Only nurses who knew the patient best or were at the bedside when patients died were interviewed. | delivering care to the patient in the last three days of life, a total of 115 patients within the study. However unclear as to how many nurses were interviewed regarding the 112 patients to make the comparison between nurses and electronic patient record. | (RAI-PAC), version 9.0 was used to collect data regarding patients’ physical symptoms, signs and treatment during the last three days of life in the hospital. The RAI-PAC is a 72 items instrument collecting data on (social demographic status, health condition (including symptoms and clinical signs), oral and nutritional status, skin condition, cognition, communication, mood, psychosocial well-being, functional status, continence, medications, treatment and procedures, responsibility and directives, social support and discharge). Structured interviews conducted with nurses using the RAI-PAC if a nurse did not know whether a symptom or clinical sign was present or whether treatment was given, the item was coded ‘missing data’. An additional RAI-PAC assessment was filled out for each patient based on their electronic record. | records data using RAI-PAC. Data analysed using SPSS, version 17. Cohen’s Kappa coefficient used to assess agreement between interview and EPR data. Symptoms and signs were defined as absent, present, but not exhibited in last three days; exhibited on one of the last three days; exhibited daily in last three days. The variables were dichotomised as not present during the last three days of life. The treatment variables were defined as: no order and no change; ordered, but not implemented; given in error for 1-2 days of the last three days; given daily for last three days; not given, because declined. The treatment variables were dichotomised as: not given in the last three days of life or given one or two days after. observed for dyspnoea, pain and nausea, and present an agreement on patient fatigue. There was good agreement between nurse interview data and EPR data for the clinical signs falls. There was moderate agreement for vomiting, peripheral oedema and fever. There was a fair level of agreement for constipation, excessive sweating, difficulty in falling or staying awake, difficulty in cleaning airway secretion and diarrhea. A poor level of agreement was suggested for dry mouth, blushing and sleep interfering with normal functioning. The agreement was good between nurses assessments and EPR data for monitoring and oxygenator related positive airway pressure (PAP)/continuous positive airway pressure (CPAP). There was moderate agreement for the administration of intravenous medication, oxygen therapy and wound care. There was only fair agreement on whether a nursing or repositioning programme was applied. Quality assurance: only focused on clinical signs and symptoms or aspects of psychological, social and emotional care. Does not capture nurse patient interaction. Only one hospital. Retrospective data collected and recall bias may have affected the complement of the documentation and the reliability. |

Theme One: Symptom management

Within the review of the studies identified it was evident that symptom management was the main focus of nurses’ documentation. While a broad range of symptoms was identified across the studies reviewed (Table 5) only pain was identified as being specifically documented across all studies [23-27].

Pressure ulcer/wound care [23,24,26] was identified in three studies, and fatigue [24,26], intravenous medication [23,24], nutrition [23,26], oedema [23,24], and oxygen therapy [23,24] identified as documented within two studies. The remaining symptoms were identified as documented individually within a single study (Table 5). While pain was identified across all studies Almasalha et al [25] highlighted within their study that only 41.3% of care plans reviewed included pain documentation and 55% of patients were discharged to a hospice with worse pain than expected. While pain was rated on the Nursing Outcomes Classification System and recorded on the plan of care, when compared at discharge the most effective nursing intervention classification of pain intervention was only used in 13.6% of cases [25].

Documentation of assessment was evident in Esper and Walker’s [27] audit, where assessment of bowel function at the time of narcotic prescription, assessment of narcotic efficacy on the return visit following initial or prescription change and assessment of bowel function post-narcotic prescription were highlighted. However, within Esper and Walker’s [27] audit these assessments fell below the 80% compliance level set. Furthermore, documentation of the plan and appropriateness of the plan for addressing moderate to severe pain, and measures for addressing oral chemotherapy management also fell below the 80% level [27]. Furthermore, Ohlen et al [23] highlight that the documentation often began with a description of patient status followed by an assessment from the nurse, typically concerning medication, pain management, medical technical interventions and nutrition.
Theme Two: Engagement

The theme of engagement was evident within two studies [23,27] and encompassed the aspects of advice, coordination, consultation and personal support. Within Ohlen et al [23] study it was evident within the nursing documentation that nurses’ advised patients and also gave the patient and/or family support in different matters. These dialogues were concerned with providing information of different activities aimed at achieving a higher degree of well-being for the patient and family for example, aquatic training or creative work like painting [23]. Documentation highlighted that nurses comforted and supported patients to better cope with their present situation by being present and listening [23]. However, assessment of emotional well-being and the nursing plan for addressing emotional well-being fell below the 80% compliance level set [27].

Within nurses’ documentation coordination and consultation consisted of nurses engaging with multidisciplinary team members regarding concerns about the patient or patient care [23]. This coordination and consultation was mainly performed from the office by telephone [23]. Within coordination and consultation care nurses provided personal assistance to the patient by assisting patients to book or check appointments, order mobility equipment, medical or personal aids and submit applications for travel services. These activities required a nursing assessment and the involvement of other professionals [23].

Discussion

Palliative care also strives to improve the sense of wellbeing of the person’s informal support network, including family members and other caregivers [28]. However, from this review it is apparent that the gap between nurses recognising the significance of non-physical aspects of care and recognising the importance of documenting these persists. In the studies reviewed psychosocial, spiritual and cultural issues were rarely mentioned and documentation focused on physical symptoms and their management. This raises the question of what is considered by nurses as important to document and may reflect the apparent difficulty in writing about non-physical aspects of care, compared to speaking about them and raises the question does this then become lost information? Findings from this review suggests that an inconsistency exists between nursing documentation and actual nursing activities performed and this is consistent with De Marinis et al [29] who identified this inconsistency at 47% (n=742 nursing activities). Several researchers note that too often nurses perform the necessary direct nursing care and forget to document due to distractions, or that entries made in nursing documentation don’t truly reflect the comprehensive care provided [7,29-31]. It may also be the case that nurses are concerned with direct nursing care and documentation can be sacrificed in times of heavy workloads and interruptions [32]. This has been further compounded in recent stringent economic times where under staffing has occurred [33] thus generating a further increase in direct nursing care workload and reduced focus on documentation [34]. However, as nurses have frequent and continual contact with patients throughout the day, their role in documenting any changes or deteriora-

| Symptom                        | Esper and Walker (2015) | Freeman et al. (2014) | Ohlen et al. (2013) | Almasalha et al. (2012) | Steindal et al. (2011) |
|-------------------------------|-------------------------|-----------------------|---------------------|-------------------------|-----------------------|
| Dyspnea                       | --                      | √                     | --                  | --                      | --                    |
| Fever                         | --                      | --                    | √                   | --                      | --                    |
| Constipation                  | --                      | --                    | --                  | --                      | √                     |
| Delirium                      | --                      | √                     | --                  | --                      | --                    |
| Excessive sweating            | --                      | --                    | --                  | --                      | √                     |
| Unexpected bleeding           | --                      | --                    | √                   | --                      | --                    |
| Vomiting                      | --                      | --                    | --                  | --                      | √                     |
| Clinical sign falls           | --                      | --                    | --                  | --                      | √                     |
| CPAP                          | --                      | --                    | --                  | --                      | √                     |
| BiPAP                         | --                      | --                    | --                  | --                      | √                     |
| Diarrhoea                     | --                      | --                    | --                  | --                      | √                     |
| Dry mouth                     | --                      | --                    | --                  | --                      | √                     |
| Bloating                      | --                      | --                    | --                  | --                      | √                     |
| Turning or repositioning      | --                      | --                    | --                  | --                      | √                     |
| Clearing airway secretion / suctioning | --            | --                    | --                  | --                      | √                     |
| Oral chemotherapy management  | √                      | --                    | --                  | --                      | --                    |
| Catherisation                | --                      | √                     | --                  | --                      | --                    |
| Appetite problems             | --                      | --                    | √                   | --                      | --                    |
| Sleep                         | --                      | --                    | --                  | --                      | √                     |

Table 5. Symptom management identified across studies.
tion in health status is critical [3].

Within this review it is evident that nurses don’t clearly document what they do beyond physical care/symptom management and given the philosophy of palliative care there is a need to record physical, psychological, social and spiritual aspect of care. Nursing documentation can be seen as an opportunity for nurses to showcase what they actually do for the patient [8,35]. The purpose of nursing documentation is to provide a permanent record of nursing’s contribution to patient outcomes [36]. This is attained if nursing documentation is a chronological verification of events showing how nursing interventions assimilate with the overall health care plan [32,37]. By presenting a record of the nurse’s contribution to patient care, nursing documentation fulfills many functions, such as: supplying a framework for the continuity of patient care by communicating the patient’s condition and care to all members of the health care team; providing evidence of care in a legal context; supporting the evaluation of the effectiveness of care; and acting as a database of nursing knowledge for information about future planning in health care and, can be used for risk planning, learning experiences for students, and the protection of patients’ rights [32]. Given the long tradition of holistic care, nursing as a profession must create an environment of documentation that captures the essential and valued aspect of care.

This review reveals several inadequacies in the documentation of the nursing interventions afforded to patients in receipt of specialist palliative care. One of the main issues is the lack of documentation regarding the broad range of nursing interventions. Studies indicated the predominance of documentation of a biomedical nature and insufficient recording of psychological, social, cultural and spiritual aspects of care [36,38]. What may be helpful, is the use of standardised terminology to effectively demonstrate nursing knowledge within documentation and reduce bias and confounding factors in data mining which can in turn inform clinical decision making. Inadequate documentation of the steps of the nursing process is evident in nursing documents with insufficient documentation of assessment and the use of assessment tools [36,39]. The difficulty for nurses is the need to balance the delivery of care with the need to document the interventions. While documentation is reported as taking up to 50% of nurses’ time per shift [40] this needs to be considered in light of the essential functions it serves. Poor communication is known to contribute significantly to the incidence of adverse events in healthcare [41] and is therefore an important objective of initiatives to improve patient safety. Adverse events can be reduced if a complete and accurate account of the patient’s condition, care and response to care is available to all healthcare team members [42].

Documentation is essential for education, research and quality assurance [43,44] and can be used to predict mortality [45]. According to Jefferies et al [46], quality nursing documentation must meet seven criteria: (1) patient-centred, (2) contains the actual work of nursing, (3) reflects the nurses’ clinical judgment, (4) is presented in a logical sequence, (5) is written in real time, (6) records variances in care and (7) fulfills legal requirements. However, even with this knowledge documentation remains poor [42], with insufficient documentation of; assessment and nursing care and inaccuracy of documentation evident [47]. This is compounded by incongruence between what is documented and the actual physical status of the patient [48] and limited documentation of the work of nurses [35]. These inadequacies could be associated with challenges in two areas: the nurses’ individual characteristics and work environments [49]. Nurses’ perceptions and attitudes towards documentation impact the quality of how and what they document. While nurses consider documentation important they also consider it to be a burdensome task that takes them away from direct patient care [7]. Furthermore, knowledge of the nursing process, which forms the basis for nursing documentation, has been found to be insufficient [5]. In tandem with this is nurses awareness of increasing liability for their practise and as a result their documentation may be negatively affected [50,51].

The workplace environment can also contribute to poor documentation where heavy workloads, arduous documentation forms, fragmented language (i.e. use of colloquial language), inadequate resources and hospital culture all impact upon the quality of nurses’ documentation and time to document [35,42]. Given the significance of nursing documentation and poor documentation practises, there have been calls and efforts to improve documentation quality [52]. Varying education strategies have been utilised to improve knowledge, skills and documentation practises these include: the introduction of a writing coach programme to improve documentation quality [32], augmenting training using written practise standards [53] and standardising nursing records, or a nursing model [54].

This review highlights a need for documentation to be developed in order to fully reflect how nurses integrate the philosophy of palliative care into their practice and for priority to be afforded to the documentation of nurses’ work. In recent years the emphasis on quality has resulted in documentation being considered an important mechanism to evaluate care performance given by nurses [55]. It follows therefore that attention should be focused towards educating nurses in the area of documenting care so that nurses have the ability to describe on paper all elements of the holistic care they provide [29] and demonstrate how nursing interventions affects client outcomes [56]. Such educational programmes should provide nurses with the competence to document adequately but also encouraged nurses to view their documentation as a crucial aspect of care and recognise its value to other disciplines outside the profession [35]. However, while Jefferies et al [32] pre and post programme documentation audit, illustrated vast improvements in nursing documentation, Dehghan et al [57] found that nursing documentation did not improve after
a two year clinical governance programme, which included education. In addition, it is recognised that regular auditing of clinical records can improve standards of record keeping and hence patient care [42]. However, from our review what is evident is that nurses need to move beyond policy aspiration to a regular daily practice within palliative care documentation. This commitment requires a willingness to change attitudes and practice, not merely by the introduction of policy or best practice imperatives, but by services, nurse managers and nurses adopting such a change in order for the potential of nursing documentation to be fully realised.

Conclusion
Nurses often state that "I did it; I just did not document it," but regulatory and legal views on documentation counter argue with the adage "if it was not documented; it was not done" [58]. Having a quality nursing document is important as it improves communication, avoids duplication, provides clear information, avoids irrelevant data been entered and inaccuracies occurring. Within palliative care nursing documentation needs to reflect the specialist and expert nursing afforded to patient and their families, moving beyond the recording of physical care and symptom management. Nurses need to demonstrate the value they place on all aspects of their care by documenting the total care provided. To support nurses’ documentation of care, education and support is required, but this education has to occur within a culture of change within the whole organisation/service in order to achieve sustainable changes in attitudes and practice.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions

| Authors’ contributions              | OD | MEB | SM | KS |
|------------------------------------|----|-----|----|----|
| Research concept and design        | ✓  |     | ✓  | ✓  |
| Collection and/or assembly of data | ✓  | ✓   | ✓  | ✓  |
| Data analysis and interpretation   | ✓  | ✓   | ✓  |   |
| Writing the article                | ✓  |     |   |   |
| Critical revision of the article   | ✓  | ✓   |   |   |
| Final approval of article          | ✓  | ✓   |   |   |
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