Variations in Clinical Practice: Assessing Clinical Care Processes According to Clinical Guidelines in a National Cohort of Hospice Patients

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Abstract

Background: National clinical guidelines have been developed internationally to reduce variations in clinical practices and promote the quality of palliative care. In The Netherlands, there is considerable variability in the organisation and care processes of inpatient palliative care, with three types of hospices – Volunteer-Driven Hospices (VDH), Stand-Alone Hospices (SAH), and nursing home Hospice Units (HU). Aim: This study aims to examine clinical practices in palliative care through different hospice types and identify variations in care. Methods: Retrospective cohort study utilising clinical documentation review, including patients who received inpatient palliative care at 51 different hospices and died in 2017 or 2018. Care provision for each patient for the management of pain, delirium, and palliative sedation were analysed according to the Dutch national guidelines. Results: 412 patients were included: 112 patients who received treatment for pain, 53 for delirium, and 116 patients underwent palliative sedation therapy. Care was provided in accordance with guidelines for pain in 32%, 61% and 47% (P = .047), delirium in 29%, 78% and 79% (P = .0016), and palliative sedation in 35%, 63% and 42% (P = .067) of patients who received care in VDHs, SAHs, and HUs respectively. When all clinical practices were considered, patient care was conducted according to the guidelines for 33% of patients in VDHs, 65% in SAHs, and 50% in HUs (P < .001). Conclusions: The data demonstrate that care practices are not standardised throughout Dutch hospices and exhibit significant variations between type of hospice.

Keywords

hospice, palliative care, delirium, pain, palliative sedation, guidelines, quality care

Introduction

Clinical guidelines have been developed internationally to reduce variations in care provision, enable effective and efficient care, encourage prudence and improve quality of care.¹⁻⁴⁻¹⁻⁴ There is considerable evidence from a wide range of health settings that variations in clinical practices are associated with poor health outcomes and low-quality care.¹⁻⁷ Clinical guidelines aim to improve structures and processes of care through standardisation of clinical practice, to enable appropriate care for every patient irrespective of their circumstances. As a result, guideline adherence has demonstrated improvements in patient and clinical outcomes, such as improved survival in cancer treatment and symptom management.²⁻⁸⁻⁹

Quality of care is a broad concept that incorporates notions of appropriateness, effectiveness, acceptability, equity, accessibility, efficacy and humanity.⁵⁻¹⁰⁻¹¹ Patients have described quality palliative care as consistent care; through assessment, communication, clinical decision-making, treatment, and multidisciplinary involvement.¹²⁻¹⁴ Every patient

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should ideally have access to the same care which is appropriate to their needs, independent of their background, underlying illnesses, and site of care. The Donabedian model proposes a framework for evaluating health services and quality of care. According to this framework, health care (and the quality of that care) consists of three domains: structure, process and outcomes. This study measures care processes, as the actions that deliver care, yet these processes reflect care structures and influence outcomes. As a result, guideline adherence has thus been employed widely as a quality measure associated with all three domains of quality of care.

Dutch national guidelines for palliative care were originally developed in the 1990s, a central impetus being considerable variations in clinical practices evident at the time. Palliative sedation was a notable clinical practice where there were inconsistencies in care provision identified, and thus concrete guidance was needed to direct best practice care that was evidence-based, collaborative and promoted transparent clinical decision-making. There are now over forty national palliative care guidelines focusing on differing clinical practices, in order to promote consistent and quality care provision. They consist of evidence-based guidance regarding the aetiology, prevention, diagnosis, management, evaluation and engagement of multidisciplinary care for specific clinical scenarios and populations.

The Netherlands contains approximately 300 hospices, which provide inpatient palliative care accessible to patients with an estimated life expectancy of less than three months. They can be divided into three types, including hospices where care is primarily provided by volunteers (VDH), palliative units that operate as part of nursing homes (HU), and independent hospices with specialist-trained palliative care staff (SAH), as described in Figure 1. There are distinct organisations responsible for these types of hospices, which may vary in structures of governance, frameworks of care provision, reporting and quality standards. As a result, hospice care in the Netherlands is subject to considerable structural variability, that may impact upon the processes of care provision, and outcomes.

This study aims to examine clinical practices in palliative care through different hospice types, to identify variations in care processes and settings in The Netherlands. To study these variations in care we focused on three clinical practices with associated national guidelines that have specific processes of care that could be measured through documentation review:

- Pain (a symptom),
- Delirium (a syndrome), and
- Palliative Sedation (an intervention).

These practices were identified due to their importance and prevalence in hospice care, as three practices that are fundamental to quality palliative care provision. In particular, palliative sedation is a practice that has received increasing attention in the Netherlands over the preceding decade, in part due to increasing incidence, which in 2017 was reported in 23% of deaths nationwide with higher levels (28%) in the hospice setting. Pain is a very common symptom in hospice populations, the treatment of which is integral to quality palliative care. Delirium is a syndrome affecting 25-45% of patients admitted to hospices, and for approximately 20% of patients can take an agitated form, that can be very distressing for patients, family members and staff.

### Methods

#### Study Design

This retrospective observational cohort study was performed using patient records of patients admitted to hospices in the Netherlands and died in 2017 or 2018. The Strengthening the Reporting of Observational Studies in Epidemiology (STROBE) statement was used for reporting.

| Hospice type | Volunteer-driven hospice (VDH) | Stand-alone hospice (SAH) | Hospice unit nursing home (HU) |
|--------------|---------------------------------|---------------------------|---------------------------------|
| Organisational structure | Volunteer Driven | Professional-driven |                      |
| Governing association | Volunteers palliative terminal care | Dutch association of Hospice Care | Actiz |
| Access | Patients with estimated life expectancy <3 months |                      |                               |
| Staff | Trained volunteers 24/7, patient’s GP, district nurses | Specialised nurses 24/7, doctors (GP, physician, elderly care specialists), chaplain, psychologist, supported by trained volunteers |                      |

**Figure 1.** Organisation of hospice care in the Dutch healthcare system.
Setting and Participants

Hospices were randomly selected using a random number generator, with equal numbers of VDH, SAH and HUs invited to participate. Hospices for specific patient populations, such as paediatric hospices, were not included. 17 hospices of each hospice type were selected (51 in total), representing geographical locations of the Netherlands.

From each participating hospice an overview was made of all patients who were admitted to the hospice and died in 2017 or 2018. 16 patients from each hospice were randomly selected for data collection. Of these, four patients from each hospice were randomly selected for an in-depth exploration, for which data on clinical practices was collected. Sample size was determine by initial calculations, predicting approximately 200 patients would be required to demonstrate differences in clinical practices between hospice types. It was decided to include all patients (from the 16 selected at each hospice site) that had palliative sedation reported from the mid-point of data collection, not only those who had in-depth analysis.

Data Collection

Patient clinical records were reviewed by the researchers, who were experienced clinicians (nurses, doctors and allied health) and entered manually using electronic case report forms in Castor Electronic Data Capture System. All data collected was pseudo-anonymised at the level of the patient and hospice. Data collection was developed and tested in an initial pilot in 2017 which has been described elsewhere. Data was collected by the researchers from December 2018 to June 2021.

Distinct episodes of patient care provision – termed ‘clinical practices’ were identified in the records for these patients, included pain, delirium and palliative sedation. The researchers assessed all available clinical documentation in the first 72 hours after admission, the middle 72 hours, and the last 72 hours, to identify if these clinical practices of pain, delirium, or palliative sedation occurred. Every episode of care related to that clinical practice was recorded, describing the assessment, problem classification, pharmacological and non-pharmacological management, evaluation of treatment and staff member providing care. For pharmacological treatment, type, dosing, and alterations to medication were collected. The clinical care was assessed longitudionally to identify if patient care was escalated appropriately over the individual course of treatment.

The following criteria described in Figure 2 were assessed for each individual patient in accordance with the national guidelines, which could be reliably measured through the data.

Ethics

This research was reviewed by the institutional review board of the UMC Utrecht (18-373/C, 18/05/2018) and not considered subject to the Medical Research Involving Human

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| Pain | 22 |
|------|----|
| 1. Was medication treatment according to the guidelines – regarding types of medications used? |
| 2. After interventions were performed, was this evaluated and documented? |

Given variability in analgesia dosing dependent on previous use of opioids which could not be fully assessed through retrospective clinical dossier review, the doses of medication were not assessed according to the guidelines.

| Delirium | 31 |
|---------|----|
| 1. Was the existence of possible features of delirium described, such as: restlessness, disorientation, concentration problems or diurnal unrest. |
| 2. Was medication treatment according to the guidelines – both in type and doses of medication used? |

| Palliative Sedation | 30 |
|---------------------|----|
| 1. Was an indication for palliative sedation described? |
| 2. Was medication treatment according to the guidelines – both in type and doses of medication used? |
| 3. After interventions were performed, was this evaluated and documented? |

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Figure 2. Criteria used to analyse each clinical practice in accordance with national guidelines.
Subjects (WMO) Act of the Netherlands. In line with the principles of Good Clinical Practice, local consent from hospices was obtained.

**Data Analyses**

A core team of clinician researchers (EG, ST, MG) analysed the data who had experience as nurses or physicians in palliative medicine. Where there were uncertainties, these were discussed as a group. If the data collected was inconclusive in describing care that did not follow the guidelines, then this was adjudged ‘according to the guidelines’. Data analysis employed SPSS v.26 and comprised primarily of descriptive statistics (proportions, means, and standard deviations where appropriate) and chi-squared test for between-group differences.

**Results**

192 patients were screened to identify episodes of pain and delirium, and 412 patients screened for palliative sedation. 112 patients (of 192, 58%) were included who received treatment for pain, 53 (of 192, 28%) for delirium, and 116 patients (of 412, 28%) underwent palliative sedation therapy. Mean age was between 72.7 and 75.7 years. Table 1 describes their demographic and clinical characteristics according to hospice type.

**Pain**

The majority (91%) of patients received medication treatment in accordance with guidelines, yet only 49% had routine evaluation after treatment. The proportion of patients for who all criteria were fulfilled was highest in SAH units (61%) and lowest in VDHs (32%). Table 2 describes these results per hospice type.

### Table 1. Patient demographics and clinical characteristics.

|                | VDH | SAH | HU  |
|----------------|-----|-----|-----|
| **Pain**       |     |     |     |
| Number patients| 34  | 41  | 37  |
| Sex–female (%) | 23 (68) | 20 (49) | 19 (51) |
| Age–mean (range)| 75.5 (47-99) | 73.4 (51-94) | 72.7 (39-91) |
| Main diagnosis |
| -Cancer (%)    | 26 (76) | 36 (88) | 30 (81) |
| -Organ failure (%) | 4 (12) | 3 (7) | 4 (11) |
| -Neurological (%) | 1 (3) | 1 (2) | 0 |
| -Other (%)     | 3 (9) | 1 (2) | 3 (8) |
| Major comorbidities–mean | .94 | .76 | 1.0 |
| **Delirium**   |     |     |     |
| Number patients| 21  | 18  | 14  |
| Sex–female (%) | 11 (52) | 5 (28) | 5 (36) |
| Age–mean (range)| 79.8 (44-99) | 75.4 (56-92) | 76.6 (54-90) |
| Main diagnosis |
| -Cancer (%)    | 13 (62) | 15 (83) | 8 (57) |
| -Organ failure (%) | 5 (24) | 3 (17) | 3 (21) |
| -Neurological (%) | 0 | 0 | 1 (7) |
| -Other (%)     | 3 (14) | 0 | 2 (14) |
| Major comorbidities–mean | 1.23 | .83 | 1.07 |
| **Palliative sedation** |     |     |     |
| Number patients| 34  | 32  | 50  |
| Sex–female (%) | 18 (53) | 17 (53) | 28 (56) |
| Age – mean (range)| 77.9 (44-99) | 71.8 (38-91) | 74.6 (42-97) |
| Main diagnosis |
| -Cancer (%)    | 26 (76) | 27 (84) | 38 (76) |
| -Organ failure (%) | 2 (6) | 2 (6) | 5 (10) |
| -Neurological (%) | 1 (3) | 2 (6) | 1 (2) |
| -Other (%)     | 2 (6) | 1 (3) | 3 (6) |
| Major comorbidities-mean | 1.21 | .72 | 1.22 |
| Days prior to death–palliative sedation initiated |
| Median         | 1.0  | 1.0  | 1.0  |
| Range          | 0-7  | 0-9  | 0-7  |
**Reasons for Medication Treatment Diverging From Guidelines.** Ten patients (9%) received medication treatment that did not conform to the guidelines, all of which involved the use of midazolam documented for pain. Three patients were given midazolam as a single agent for pain. Seven patients were treated with midazolam in conjunction with an opioid for pain. None of these patients were documented as receiving palliative sedation at the time of these treatments.

**Delirium**

Most patients who experienced a delirium were assessed for anticipation of a prodrome (75%) and had medication treatment in accordance with guidelines (77%). The proportion of patients in which all criteria were met was highest in HU (79%) and SAH (78%) units, and lowest in VDHs (29%). Table 3 describes these results per hospice type.

**Reasons for Treatment Diverging From Guidelines.** 12 patients received medication treatment that did not conform to the guidelines. Five patients were given benzodiazepines as a single agent and five treated with benzodiazepines prior to antipsychotics. Two patients were treated with other forms of antipsychotics – quetiapine (step 2 in the guidelines) and dipiperon (not in guidelines) - prior to the use of haloperidol (step 1).

**Palliative Sedation**

Patients who underwent palliative sedation therapy had, in most cases, a known clinical indication for therapy.

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### Table 2. Patient treatment for pain in accordance with guidelines (2 criteria) per hospice type.

| Type        | Hospice | VDH | SAH | HU  |
|-------------|---------|-----|-----|-----|
| Medication use according to guidelines | Yes | 31 (91.2%) | 40 (97.6%) | 32 (84.2%) |
|             | No     | 3 (8.8%) | 1 (2.4%)  | 6 (15.8%)  |
| Evaluation of treatment     | Yes | 12 (35.3%) | 26 (68.4%) | 17 (42.4%) |
|             | No     | 26 (64.7%) | 15 (36.6%) | 21 (55.3%) |
| Number of criteria (n = 2) met | 0 | 2 (5.9%) | 0 | 3 (7.9%) |
|             | 1 | 21 (58.1%) | 16 (39.0%) | 17 (42.4%) |
|             | 2 | 11 (32.4%) | 25 (61.0%) | 18 (47.4%) |

### Table 3. Patient treatment according to delirium guidelines (2 criteria) per hospice type.

| Type          | Hospice | VDH | SAH  | HU  |
|---------------|---------|-----|------|-----|
| Anticipation of prodrome | Yes | 12 (57.1%) | 14 (77.8%) | 14 (100%) |
|               | No     | 9 (42.9%) | 3 (16.7%) | 0 |
|               | Missing data | 0 | 1 (5.6%) | 0 |
| Medication treatment according to guidelines | Yes | 13 (61.9%) | 17 (94.4%) | 11 (78.6%) |
|               | No     | 8 (38.1%) | 1 (5.6%)  | 3 (21.4%)  |
| Number of criteria (n = 2) met | 0 | 1 (4.8%) | 0 | 0 |
|               | 1 | 14 (66.7%) | 4 (22.2%) | 3 (21.4%) |
|               | 2 | 6 (28.6%) | 14 (77.8%) | 11 (78.6%) |

### Table 4. Patient treatment according to palliative sedation guidelines (3 criteria) per hospice type.

| Type          | Hospice | VDH | SAH | HU |
|---------------|---------|-----|-----|----|
| Indication for palliative sedation described | Yes | 30 (88.2%) | 31 (96.9%) | 43 (86.0%) |
|               | No     | 4 (11.8%) | 0 | 7 (14.0%) |
|               | Data missing | — | 1 (3.1%) | — |
| Medication according to guidelines | Yes | 28 (82.4%) | 28 (87.5%) | 46 (92.0%) |
|               | No     | 5 (14.7%) | 3 (9.4%)  | 3 (6.0%)  |
|               | Data missing | 1 (2.9%) | 1 (3.1%) | 1 (2.0%) |
| Evaluation of treatment | Yes | 12 (35.3%) | 22 (66.8%) | 25 (50.0%) |
|               | No     | 22 (64.7%) | 10 (31.3%) | 25 (50.0%) |
| Number of criteria (n = 3) met | 0 | 2 (5.9%) | 0 | 1 (2.0%) |
|               | 1 | 5 (14.7%) | 1 (3.1%)  | 5 (10%)   |
|               | 2 | 15 (44.1%) | 11 (34.4%) | 23 (46%)  |
|               | 3 | 12 (35.3%) | 20 (62.5%) | 21 (42%)  |
documented (90%) and received medication treatment following the guidelines (88%). Treatment was routinely evaluated for 51% of patients. In total, 35% of patients in VDHs met all criteria, 63% in SAHs and 42% in HU units. Table 4 describes these results per hospice type. The four cases of missing data included one case where an indication for palliative sedation was not described and three cases when clinical records regarding medication use did not detail the type of medication used.

**Reasons for Medication Treatment Divergent to Guidelines.** 11 patients received medication treatment that did not conform to the guidelines. Four patients were given levomepromazine (step 2 in the guidelines) first line for palliative sedation without the use of benzodiazepines (step 1). Two patients were given escalating doses of midazolam at greater than 20 mg/hr without adding levomepromazine (step 2) or specifying a reason why additional sedatives were not considered, both occurring at the same hospice. For five patients, starting doses of midazolam used exceeded specified doses, with four patients having syringe drivers initiated at >2.5 mg/hr (including three patients from the one hospice), or pro ne rata midazolam being initiated at >10 mg for one patient.

**Treatment According to Guidelines for each Clinical Scenario**

Treatment was provided conforming to the guidelines for 46% of patients who received treatment for pain, 59% for delirium and 48% for palliative sedation therapy. For each hospice type, patient care was conducted in accordance with the guidelines for 33% of clinical practices in VDHs, 65% in SAHs, and 50% in HUs (P < .001). Table 5 describes the percentage of patients who received care for each clinical scenario according to the guidelines, according to hospice type.

**Discussion**

**Main Findings of the Study**

This study describes how routine palliative care processes were provided to a national sample of Dutch hospice patients for the first time. For approximately 50% of patients, they did not receive care in accordance with national guidelines for that specific clinical practice. Patients in VDH units were less likely to receive care conforming to the guidelines than SAH and HU settings. For patients, this equates to discrepancies in care provision, where the care they receive is influenced by the site of care. These variations are likely due to systemic influences; as can be identified from the discrepancies between types of hospice, and from the reasons for medical care divergent from the guidelines, which identify particular sites where the non-conforming clinical practices are repeated. Such factors may be related to individual hospice policies (or lack of) on specific clinical practices, variations in levels of staff experience and training, and differing approaches to clinical decision-making and documentation.

The study identified that the majority of patients (pain 91%, delirium 77%, palliative sedation 88%) received medication treatment in accordance to the guidelines. The clinical practices that demonstrated most inconsistency in our study were non-medication processes, such as signalling and treatment evaluation, processes that are central to quality care provision. Hasselaar et al examined palliative sedation practice in The Netherlands between 2003 and 2005 using self-reported questionnaires, in which 43% of physicians employed medication treatment that did not follow national guidelines. This was in contrast to our study, where medication treatment was in most cases, according to the guidelines. It is hoped that this may represent changing clinical practices in The Netherlands as a result of national guidelines first published in 2002. Whilst the focus of health care professionals is often on practices such as appropriate use of medication, these other processes are equally important, identifying the needs of the patient, guiding treatment, and promoting collaborative care. A retrospective clinical documentation review of six inpatient settings (acute hospitals and palliative care units) from Canada described similar variability in palliative sedation practice. Indications (85-88%) for palliative sedation and informed consent (73%) were commonly present, yet other practices described in guidelines such as documentation of patient goals of care directed toward terminal management (16%) infrequently occurred. This study noted substantial variation between care settings, highlighting how care structures shape processes of care.

**Strengths and Limitations of the Study**

The unique contribution of this study is to provide an in-depth ‘snapshot’ of inpatient palliative care practices across the Netherlands throughout different types of hospices. The study has collected extensive data on diverse range of patients throughout The Netherlands, detailing the specifics of their personal and clinical characteristics, care needs, and how and what care was provided, as longitudinal data throughout the course of admission. Other studies have used survey methods to examine specific clinical practices from the practitioner perspective, mostly focusing on medication management and referral. Whilst these methods are able to access widespread practitioner populations to explore variations in
practice, there may be differences between how care is provided by health professionals in theory and in practice. Additionally, these clinical practices are complex processes that are dependent on many individuals. The main strength of this study is that it has enabled an understanding of how patient care occurs in reality across a national sample of hospices, as longitudinal care processes involving many individuals.

The methods employed in this study were those deemed most appropriate for the aims of the study, which needed to be able to be replicated in each study site. Whilst these methods enabled such a broad range of variables to be collected, they also confer limitations. Foremost was the reliance upon the quality of clinical documentation to collect data, which in some circumstances were insufficient to fill all variables. There were instances where clinical documentation was sub-optimally recorded, with information missing regarding indications for treatment, non-medication interventions, and exact doses and timing of medications. The interpretations we used to determine ‘care in accordance with the guidelines’ were inclusive, only identifying those care practices that definitely deviated from the guidance, and thus is likely an underestimation of variations in care practices. The outcomes related to these variations in practice differ greatly, as some care processes may have negligible impact on the patient, whilst others, such as very high doses of sedatives, can significantly alter patient care.

The clinical practices described are complex processes, involving many individuals, tasks and influences. Clinical documentation only describes aspects of these processes, missing many important elements, such as communication, that are key care processes supported by guidelines, yet lacking any definitive outcome. In this study we were only able to examine outcomes that could be measured from the clinical documentation. Many processes were not routinely listed in the clinical notes (i.e., provision of information) in a manner that could be routinely measured through retrospective clinical dossier review. Medication treatment was routinely well documented, and thus forms a central focus of guideline adherence.

What this Study Adds
The challenge posed by this study is how these variations in care provision can be standardised to promote routine best practice care for each patient. The variations in care observed in this study are substantial and likely to be structural influenced, thus requiring a systematic approach to address these challenges. The logical next step would be consideration of quality measures that could be standardised throughout hospices in The Netherlands. Through measuring and reflecting on these clinical practices, health care professionals and hospice organisations can identify elements of patient care that can be developed and improved. Further research would ideally focus on identifying care process measures that could be implemented through hospices in The Netherlands, to drive quality improvement and promote evidence-based, consistent care.

Central to all these care processes and their measurement is documentation. Documentation is a fundamental and critical care process that communicates the patient’s needs, responses to care, and care team planning, which has been recognised as integral to quality palliative care. Some care processes, such as medication treatment, may be more readily documented, which reflect the important quality and safety issues related to prescribing. Hospices act as complex systems where care is provided by a diverse group of regionally involved GP’s, nurses, volunteers and allied health professionals, and thus documentation of all these processes is central to promoting continuity of care. This is of particular relevance for complex interventions such as palliative sedation, where patient’s needs are often evolving, treatment can have narrow therapeutic windows, and many care providers (both formal and informal) are involved. Care processes should ideally be iterative, transparent and promote continuity between the care team. Developing key quality measures for documentation in palliative care practice may enable hospice services to audit performance, guide hospice teams in which aspects of care should be communicated, and promote its importance as a key driver of care continuity and quality improvement.

Conclusion
Hospice care in the Netherlands is delivered through many small scale institutions with structural variations in staffing, organisation and reporting. This study describes that there are substantial variations in clinical practices focusing on the treatment of pain, delirium and palliative sedation. When interpreted in relation to the national guidelines, patient care adhered to the guidelines for 65% of cases in SAHs, 50% in HUs and 33% in VDHs. Whilst medication treatment in most cases conformed to the guidelines, there were some institutions where clinical practices that deviated from official guidance were routinely provided. Signalling, evaluation of treatment, and documentation of these practices exhibited greater variability in practice. To improve patient care, care practices need to be further standardised in line with best practice so that each patient is able to access the same quality care.

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References
1. Grol R, Thomas S, Roberts R. Development and implementation of guidelines for family practice: Lessons from the Netherlands. J Fam Pract. 1995;40(5):435-439.
2. Lugtenberg M, Burgers J, Westert G. Effects of evidence-based clinical practice guidelines on quality of care: A systematic review. BMJ Qual Saf. 2009;18(5):385-392.
3. Abarshi E, Rietjens J, Robijn L, Caraceni A, Payne S, Deliens L, et al. International variations in clinical practice guidelines for palliative sedation: A systematic review. BMJ Support Palliat Care. 2017;7(3):223-229.
4. Palliactief IK. Netherlands quality framework for palliative care. IKNL; 2017.
5. Margo CE. Quality care and practice variation: The roles of practice guidelines and public profiles. Surv Ophthalmol. 2004;49(3):359-371.
6. Wise PH, Eisenberg L. What do regional variations in the rates of hospitalization of children really mean? Mass Medical Soc; 1989.
7. Spencer BA, Miller DC, Litwin MS, Ritchey JD, Stewart AK, Dunn RL, et al. Variations in quality of care for men with early-stage prostate cancer. J Clin Oncol. 2008;26(22):3735-3742.
8. Bristow RE, Chang J, Ziogas A, Anton-Culver H. Adherence to treatment guidelines for ovarian cancer as a measure of quality care. Obstet Gynecol. 2013;121(6):1226-1234.
9. Du Pen SL, Du Pen AR, Polissar N, Hansberry J, Kraybill BM, Stillman M, et al. Implementing guidelines for cancer pain management: Results of a randomized controlled clinical trial. J Clin Oncol. 1999;17(1):361-361.
10. Higginson I. Quality of care and evaluating services. Int Rev Psychiatr. 1994;6(1):5-14.
11. Ibn El Haj H, Lamrini M, Rais N. Quality of care between donabedian model and ISO9001V2008. International Journal for Quality Research. 2013;7(1).
12. Nelson JE, Puntillo KA, Pronovost PJ, Walker AS, McAdam JL, Ilaoa D, et al. In their own words: Patients and families define high-quality palliative care in the intensive care unit. Crit Care Med. 2010;38(3):808-818.
13. Ferris FD, Gómez-Batiste X, Furst CJ, Connor S. Implementing quality palliative care. J Pain Symptom Manag. 2007;33(5):533-541.
14. De Graaf E, van Klinken M, Zweers D, Teunissen S. From concept to practice, is multidimensional care the leading principle in hospice care? An exploratory mixed method study. BMJ Support Palliat Care. 2020;10(1):e5.
15. Donabedian A. An introduction to quality assurance in health care. Oxford, UK: Oxford University Press; 2002.
16. Van Wijlick E, Verkerk M, De Graef A, Legemaate J. Palliative sedation in the Netherlands: Starting-points and contents of a national guideline. Eur J Health Law. 2007;14(1):61-73.
17. Nederlands IK. General Principals of Palliative Care; National Guideline, Vol 3. IKNL; 2016.
18. De Graaf E. Exploring hospice care in the Netherlands: Current practices and patients’ symptom burden and well-being. Utrecht University; 2018.
19. Koorn RM, van Klinken M, de Graaf E, Bressers REGW, Jobse AP, van der Baan F, et al. Who are hospice patients and what care is provided in hospices? A pilot study. American Journal of Hospice and Palliative Medicine®. 2020;37(6):448-454.
20. Kankercentrum VVI. Palliatieve Sedatie, Vol 236. The Netherlands2009.
21. Kankercentrum VVI. Delirium National Guideline, Vol 326. The Netherlands2010.
22. Kankercentrum VVI. Pain, Landelijk richtlijn, Vol 2. The Netherlands2010:60.
23. Rietjens JA, Heijltjes MT, Van Delden JJ, Onwuteaka-Philipsen BD, Van der Heide A. The rising frequency of continuous deep sedation in the Netherlands, a repeated cross-sectional survey in 2005, 2010, and 2015. J Am Med Dir Assoc. 2019;20(11):1367-1372.
24. Kengetallen SF. Toenmane palliative sedatie in thuissituatie begraagt 5. Pharm Weekbl. 2018;153(39).
25. Van Deijck RH, Hasselaar JG, Verhagen SC, Vissers KC, Koopmans RT. Patient-related determinants of the administration of continuous palliative sedation in hospices and palliative care units: A prospective, multicenter, observational study. J Pain Symptom Manag. 2016;51(5):882-889.
26. Von Elm E, Altman DG, Egger M, Pocock SJ, Gotzsche PC, Vandenbroucke JP. The strengthening the reporting of observational studies in epidemiology (STROBE) statement: Guidelines for reporting observational studies. Bull World Health Organ. 2007;85:867-872.
27. Hasselaar JG, Reuzel RP, Verhagen SC, De Graef A, Vissers KC, Cruy BJ. Improving prescription in palliative sedation: Compliance with Dutch guidelines. Arch Intern Med. 2007;167(11):1166-1171.
28. Abdul-Razzak A, Lemieux L, Snyman M, Perez G, Sinnarajah A. Description of continuous palliative sedation practices in a large health region and comparison with clinical practice guidelines. J Palliat Med. 2019;22(9):1052-1064.
29. Matlock DD, Peterson PN, Sirovich BE, Wennberg DE, Gallagher PM, Lucas FL. Regional variations in palliative care: Do cardiologists follow guidelines? J Palliat Med. 2010;13(11):1315-1319.
30. Cabañero Martínez MJ, Ramos Pichardo JD, Velasco Álvarez ML, Garcia Sanjuán S, Lillo Crespo M, Cabrero García J. Availability and perceived usefulness of guidelines and
protocols for subcutaneous hydration in palliative care settings. *J Clin Nurs*. 2019;28(21-22):4012-4020.

31. Grant M, De Graaf E, Teunissen S. A systematic review of classifications systems to determine complexity of patient care needs in palliative care. *Palliat Med*. 2021;35(4):636-650.

32. Bausewein C, Daveson BA, Currow DC, Downing J, Deliens L, Radbruch L, et al. EAPC white Paper on outcome measurement in palliative care: Improving practice, attaining outcomes and delivering quality services–recommendations from the European association for palliative care (EAPC) task force on outcome measurement. *Palliative medicine*. 2016;30(1):6-22.

33. Ferrell B, Connor SR, Cordes A, Dahlin CM, Fine PG, Hutton N, et al. The national agenda for quality palliative care: The national consensus project and the national quality forum. *J Pain Symptom Manag*. 2007;33(6):737-744.

34. De Roo ML, Leemans K, Claessen SJ, Cohen J, Pasman HRW, Deliens L, et al. Quality indicators for palliative care: Update of a systematic review. *J Pain Symptom Manag*. 2013;46(4):556-572.

35. Kelley TF, Brandon DH, Docherty SL. Electronic nursing documentation as a strategy to improve quality of patient care. *J Nurs Scholarsh*. 2011;43(2):154-162.

36. Tully MP, Cantrill JA. Insights into creation and use of prescribing documentation in the hospital medical record. *J Eval Clin Pract*. 2005;11(5):430-437.

37. Pype P, Mertens F, Helewaut F, Krystallidou D. Healthcare teams as complex adaptive systems: Understanding team behaviour through team members’ perception of interpersonal interaction. *BMC Health Serv Res*. 2018;18(1):1-13.