The experiences of caregivers of patients with delirium, and their role in its management in palliative care settings

Citation for published version:

Digital Object Identifier (DOI):
10.1002/pon.4140

Link:
Link to publication record in Edinburgh Research Explorer

Document Version:
Publisher's PDF, also known as Version of record

Published In:
Psycho-Oncology

General rights
Copyright for the publications made accessible via the Edinburgh Research Explorer is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

Take down policy
The University of Edinburgh has made every reasonable effort to ensure that Edinburgh Research Explorer content complies with UK legislation. If you believe that the public display of this file breaches copyright please contact openaccess@ed.ac.uk providing details, and we will remove access to the work immediately and investigate your claim.
Delirium is a serious and distressing neuropsychiatric syndrome frequently experienced by patients in palliative care settings. It affects up to 62% of patients during a palliative care inpatient admission and up to 88% in the days or hours preceding death [1]. It is the most common and distressing neuropsychiatric complication experienced by patients with advanced cancer and is often under-diagnosed and undertreated [2]. Delirium is characterized by disturbed consciousness, with reduced ability to focus, sustain or shift attention; altered cognition or perceptual disturbance; and an acute onset that occurs over a short period of time and fluctuates throughout the day [3].

Evidence gathered from patients who could recall experiencing delirium during a hospital admission reveals that memories of having had delirium are distressing [4–6]. In particular, patients recalled experiences of reality and unreality, day–night disorientation, clouding of thought, lack of control, strong emotions and misperceptions, hallucinations and delusions. O’Malley et al. [5] identified a number of studies indicating that patients recall being aware of their inability to communicate with family caregivers and healthcare professionals, compounding their feelings of distress and humiliation. Despite this, retrospective patient reports suggest that the presence of family caregivers is beneficial, helps orientate the confused patient and protects against fear, anxiety and isolation [4,7–9].

Complex issues arise in relation to the treatment and management of patients with delirium in palliative care settings. Reversibility of a particular delirium episode can be difficult to ascertain, and decision-making relating to delirium management can be challenging for clinicians and caregivers acting as proxy decision-makers [10]. One study in a hospital palliative care unit estimated that approximately 50% of delirium episodes in patients receiving palliative care cannot be reversed [11]. Patient and family distress because of impaired communication and decision-making capacity in the terminal phase is
common, and concerns often arise in relation to balancing the benefits of investigating the precipitating factors with treatment burden and sedation [12]. In the advanced stages of illness, goals of care should be clarified with the patient, or primary caregiver if the patient is unable to participate. The greater likelihood of irreversible or terminal delirium and impending death distinguishes the experience of delirium in palliative care settings from other contexts.

Caregivers play a vital role in supporting the patient approaching end of life. Two domains of support for caregivers of people approaching end of life have been identified: (i) support for the caregiver themselves and (ii) support for the caregiver to support the patient [13]. National Institute for Health and Care Excellence clinical guidelines advocate the involvement of caregivers in the management of patients with delirium and recommend that information and support in relation to delirium should be offered to them [14].

The current review draws on qualitative, quantitative and mixed-method studies to better understand the role and experience of caregivers of terminally ill patients with delirium and to identify helpful forms of support. Caregivers are broadly defined as family members, relatives or friends who are involved in the practical or emotional care of the patient. By integrating evidence relating to all aspects of the caregiver experience, this review provides a comprehensive overview of the evidence base to inform the development of caregiver interventions. The following questions form the basis for the review:

1. What are the experiences of caregivers of terminally ill patients with delirium?
2. What is the role of caregivers in the identification or management of delirium in terminally ill patients?
3. What type of support improves the experience of caregivers of terminally ill patients with delirium or helps the caregiver to support the patient?

While we are primarily concerned with delirium in palliative care settings, our literature search includes studies relating to experiences of caregivers of patients with delirium in other settings such as Medicine of the Elderly and Intensive Care. Such studies contain findings relevant to caregivers of terminally ill patients with reversible delirium and may include findings regarding caregivers of patients with an advanced illness who have not yet been formally identified as receiving or benefitting from palliative care.

**Method**

**Design**

An integrative review was undertaken. This approach is increasingly recognized as appropriate to inform evidence based practice—an important purpose of this study. An integrative review synthesizes findings from a diverse range of primary experimental and non-experimental research methods, thus providing a breadth of perspectives and a comprehensive understanding of a healthcare issue [15]. The approach reported here was modelled on key aspects of the systematic review methods advocated by the Cochrane Collaboration and the Scottish Intercollegiate Guideline Network (SIGN) [16], PRISMA standards for reporting systematic reviews [17] and the Critical Appraisal Skills Programme (CASP) for the appraisal of qualitative research studies [18].

**Search strategy**

Four electronic databases were searched by the first author (AF)—Psychnfo, Medline, Cinahl and Scopus using the terms ‘delirium’, ‘terminal restlessness’ or ‘terminal agitation’ combined with ‘caregiver’ or ‘carer’ or ‘family’ or ‘families’. The search included literature published between January 1990 and July 2015.

Papers that met the following criteria were included: (i) written in English, (ii) full papers published in a peer reviewed journal, (iii) primary qualitative and quantitative research studies, (iv) caregivers were participants in the study or were actively involved in delivering the intervention and (v) caregiver outcomes could be distinguished from general outcomes. Papers were excluded if they did not focus, at least in part, on caregiver experience or the role of caregivers in relation to patients with delirium.

**Search outcome**

The initial search yielded 1375 records resulting in 987 distinct records once duplicates were removed. Records were first screened by AF by title, and articles with title words such as ‘delirium tremens’, ‘alcohol’, ‘suicide’ and ‘schizophrenia’ were removed. Next both titles and abstracts were read by two reviewers, AF and JL, and papers that were not relevant were removed, leaving 52 papers in total.

Each of the 52 papers were read and independently assessed by AF and JL for relevance. Nineteen papers were excluded for the following reasons: (i) literature reviews (n = 6), (ii) findings were not focused on delirium or informal caregivers (n = 5), (iii) unable to answer research question (n = 1), (iv) unable to distinguish between caregiver and patient data (n = 3), (v) study protocol only (n = 1), (vi) full text not in English (n = 1), (vii) published abstract only (n = 1) and (viii) comment/discussion paper (n = 1). Where there was any concern whether a paper should be excluded, the article was discussed with all members of the team and a decision was reached by consensus. Of the six literature review papers identified [4,5,19–22], only two were specifically focused on caregivers [20,21]. Day and Higgins [20]
conducted a narrative review of family members’ experiences of older loved ones’ delirium based on three papers, a PhD thesis, a book chapter and a government report, while Halloway [21] reviewed 11 papers outlining family approaches to delirium. Both these reviews were narrower in scope than the present study, and contained papers already identified. Thirty-three papers were included in the final review (Figure 1).

Quality appraisal
We used the SIGN levels of evidence to assess the quality of the 20 papers which were predominantly quantitative and the two mixed-method papers [16]. The SIGN scoring system ranges from a 1++ score for high-quality systematic reviews or randomized controlled trials (RCTs) with a low risk of bias to a score of 4 for expert opinion based evidence (refer to Table S1 in the Supporting Information). AF and JL independently graded each of the papers. In the event of a mismatch, consensus was achieved through discussion. Where uncertainty remained CK provided an opinion and a consensus was achieved.

We drew on the CASP guidelines for qualitative research to appraise the 11 papers which were predominantly qualitative [18]. The CASP consists of 10 questions to guide the evaluation of research – two broad screening questions and 8 general questions. AF and JL independently assessed each qualitative paper with the CASP tool to determine whether each paper provided strong, moderate or weak evidence. In the event of a mismatch, consensus was achieved through discussion with the wider research team.

Data extraction and synthesis
All 33 papers were read by AF and JL, and data were extracted under the following headings: author(s) and publication year, country, setting, caregiver-related aim, participants, study design and main caregiver-related finding. A thematic analysis approach to data synthesis was adopted [23,24]. Themes from each paper relating to the research questions were identified and coded independently by AF and JL. These themes were collated in MS EXCEL and classified by AF and JL into higher order themes. The overarching themes were discussed, reviewed and agreed by all members of the research team.

Results
Characteristics of reviewed papers
Nearly half of all the papers reviewed focused on caregivers of patients receiving palliative care (n=16). The remaining papers were concerned with caregivers of older adults (n=9), caregivers of patients accessing oncology services but not identified as palliative (n=2), and caregivers of patients with delirium or at risk of developing delirium in other hospital settings (n=6). Twelve papers (36%) were specifically concerned with caregivers...
of patients with cancer who had experienced delirium or were at risk of experiencing delirium. Six papers focused on terminal delirium or terminal restlessness [25–30].

DSM criteria were most commonly used to identify delirium (n = 11) [25,27,31–39]. The Memorial Delirium Assessment Scale was used in five papers [25,26,40–42], the Confusion Assessment Method (CAM) was used in 10 papers [37,38,42–49] and the Confusion Rating Scale was used in two papers [44,50]. Three delirium detection studies used the Family Confusion Assessment Method (FAM-CAM) [38,45,46]. The Delirium Experience Questionnaire was used to explore experiences of delirium in two studies [40,41]. Terminal restlessness was assessed by an instrument developed by Jones et al. [51] in two papers [28,29].

Quality of reviewed papers

There was only one RCT study; this was accorded a 1− grade [49] (Appendix 1). Five studies were evaluated as well conducted case control studies with a low risk of confounding bias and were given a 2+ grade [33,44,48,50,52]. Eleven studies were characterized by a potentially high risk of bias and given a 2− grade [25,26,31,36,38,40–42,45–47]. Three studies were graded as Level 3 non-analytic studies [43,53,54]. The quantitative part of the two mixed method studies was graded as 2− and 3 in terms of levels of evidence [47,55].

Eleven predominantly qualitative papers were appraised following CASP guidance and classified as providing strong, moderate, adequate or weak evidence. Two qualitative papers were assessed as providing strong evidence [27,37]. Four were assessed as providing moderate to strong evidence [28–30,32], three provided moderate levels of evidence [34,56,57] while the two remaining papers were assessed as providing adequate levels of evidence [35,58].

Synthesized thematic findings

The main caregiver-related findings identified in the 33 reviewed papers are highlighted in Table S2 in the Supporting Information. The overarching themes are displayed in Figure 2.

Experiences of caregivers of patients with delirium

Eighteen papers included in the final review explored the experiences of caregivers of patients with delirium. Four sub-themes were identified: (i) generalized distress and negative emotions, (ii) difficult or deteriorating relationships, (iii) balancing the need to relieve suffering with the desire to communicate and (iv) helplessness versus control

Generalized distress and negative emotions: Several papers report that moderate to severe levels of distress are experienced by the majority of caregivers of patients with delirium [25,26,32,34,40,41]. In two studies of patients with cancer, distress was found to be greater in caregivers compared with patients experiencing delirium [40,41]. Patient correlates of caregiver distress include poor physical performance status, the presence of hyperactive delirium, hallucinations, agitation, cognitive decline and incoherent speech [25,26,32,34,40].

Figure 2. Themes identified from the literature review
Supporting caregivers of patients with delirium

Caregivers of patients receiving palliative care were particularly worried about caring for the patients alone and were anxious about leaving the patient [25].

Anxiety is frequently reported. Buss et al. [31] found that caregivers of patients with advanced cancer who had recently seen the patient in a confused or delirious state were 12 times more likely to have generalized anxiety compared with caregivers of patients who were not thought to have delirium. Anxieties relating to the loss of the loved one before death emerged in a number of studies involving patients with advanced cancer and/or receiving palliative care [31,37,44,56,58]. Day and Higgins describe the caregiver’s experience of suddenly being with an unrecognized familiar person during delirium, an ‘in-stranger.’ This experience of being with their loved one but encountering a stranger is unsettling and distressing [37].

Specific negative emotions such as fear, embarrassment, anger, sadness and guilt are also experienced. Some caregivers experienced the patient’s behaviour as unnatural and frightening [28] in particular if the patient became aggressive during the delirium episode [44,56]. Not knowing the cause of delirium was also frightening for caregivers [32]. Some caregivers were embarrassed by the patient’s actions during delirium, especially in the presence of others [56]. Anger and disappointment were experienced by caregivers when they could no longer have a meaningful interaction with the patient [29], and caregivers felt sadness relating to the changes in the patient or the premature loss of contact before death [37,44,55]. Guilt often occurred when caregivers felt that patients had died with unresolved issues and when caregivers felt that they had not been able to better care for the patient [25,56].

Difficult or deteriorating relationships: Three types of relationship difficulties were identified: (i) between caregivers and staff, (ii) between caregivers and the patient and (iii) between caregivers themselves. In a qualitative study of terminal restlessness, Brajzman [30] reported disagreements and tension between staff and caregivers which stemmed from differing understandings of the patient’s needs. The caregiver-patient relationship also suffers as caregivers experience the loss of a loved and familiar person before their actual death [37,56]. Caregivers of patients in palliative care settings had concerns about not having the opportunity for final goodbyes and were concerned that they had not talked about end of life issues with the patient [32,56]. Caregivers may experience difficulties with other family caregivers. Some felt isolated and unsupported by other family members, and sometimes felt that the patient acted differently with other family members, adding to the burden of caring for patient [56].

Balancing the need to relieve suffering with the desire to communicate: Caregivers of hospice patients experience tension in relation to decision-making around sedation [29]. Caregivers want the patient’s suffering to end, but express regret over the patient’s inability to communicate because of sedation [56]. Mixed feelings in relation to what is best for the patient are experienced. In a multi-centre questionnaire study involving 242 bereaved caregivers of patients who had experienced terminal delirium, 64% of caregivers simultaneously wanted to relieve the patient from suffering, but also wanted the patient to remain conscious [25].

Helplessness versus control: Feelings of helplessness and loss of control are common. Cohen et al. [32] found that caregivers of patients with advanced cancer in a palliative care inpatient unit felt helpless and had concerns about their own well-being as well as about how best to help the patient. In Morita et al.’s [25] study of bereaved caregivers of patients with terminal delirium, 33% of caregivers reported feeling helpless in relation to how to behave around the patient and 28% reported helplessness in the context of not understanding what was happening. In that same study, up to one third of caregivers reported that they felt a burden in relation to proxy decision-making and felt helpless. When caregivers of patients experiencing terminal restlessness have some control over treatment decision-making, they are more accepting of future difficult decisions [30]. For some caregivers, involvement in decision-making reduces helplessness and increases acceptance and control, whereas for others it increases burden [25].

Enabling an active role for families in caring for the patients can be beneficial for both the caregiver and the patient. Toye et al. [55] found that caregivers of older adults in a hospital delirium unit welcomed any opportunity to inform personalized care of the patient and felt that they could add value to the care provided by staff. There is also evidence that an active role for caregivers has psychological benefits for patients [52].

Role of caregivers in the management of delirium

Fifteen of the 33 papers reviewed discuss the role of caregivers in the management of delirium. Three sub-themes emerged: (i) the role of the caregivers in the prevention and detection of delirium, (ii) the caregiver’s role in symptom management and (iii) the caregiver as advocate for the patient.

Prevention and detection: Eleven studies considered the potential role of caregivers in the prevention and/or detection of delirium in the cared-for person [33,38,42,43,45,46,49,50,52,54,58]. A number of studies have focused on the role of caregivers in early screening
and detection. Kerr et al. [33] describe using family caregiver ‘expertise’ to examine precursors of delirium and found that caregivers of patients with delirium in a hospice inpatient unit (n=20) could identify prodromal symptoms of delirium. In a separate retrospective study of 23 caregivers of patients with head and neck cancer, Bond et al. [43] found that the incidence of caregiver reported delirium was substantially higher than the rate of delirium documented by clinicians, but comparable to the rate of subsyndromal delirium, suggesting a role for caregivers in early detection. Intervention studies have reported mixed results. Gagnon et al. [50] found no effect of an intervention involving family caregivers on delirium prevention in terminal cancer, whereas Martinez et al. [49] found that a non-pharmacological intervention delivered by families reduced delirium occurrence in hospitalized older adults compared with a control group.

There is some evidence that caregivers can administer clinical tools which could improve delirium prevention and detection. Steis et al [45] reported that the FAM-CAM is a sensitive tool for delirium screening when used by caregivers and that caregivers report no difficulty in using it. Martins et al. [38] found evidence to support the use of the FAM-CAM for caregiver detection of delirium, while Sands et al. [42] found that asking family members the Single Question in Delirium, “Do you think [name of patient] has been more confused lately?” performs well in terms of delirium detection and demonstrates potential as a simple clinical tool worthy of further investigation. Feasibility studies provide preliminary evidence that active engagement of family caregivers of hospitalized older adults in preventive interventions is feasible and may lead to improvements in well-being [47,54].

**Role in symptom monitoring:** Six papers suggest a role for caregivers in symptom monitoring. Bruera et al. [41] retrospectively examined delirium symptom recall in patients with advanced cancer and their caregivers (n=99 patient/caregiver dyads). High levels of agreement were evidenced between caregiver and patient-reported delirium symptom frequency and related distress. Bruera et al. suggest that caregivers can play an important role in monitoring patient behaviour or response to treatment. Three other studies show that caregivers can potentially help monitor symptoms because of their close proximity to patients [43,54,58]. Furthermore, there is preliminary evidence that the use of symptom assessment tools or smartphones to track delirium symptoms is feasible and acceptable to caregivers of older adults [45,46].

**Caregiver as advocate:** Caregivers potentially have a role to play acting as the advocate for the patient during delirium. In a focus group study [30], palliative care health professionals (n=12) reported that there is an onus on the caregiver to help interpret the patient’s behaviour given the patient’s inability to communicate and that caregivers’ assessments of the patient’s needs provide valuable information for the clinical team, which could be incorporated into patient treatment plans. Namba et al. [27] found that caregivers could play a role in interpreting patient talk that may appear strange to clinicians and could explain how talk about seemingly unconnected events may be linked with past real events. Having a role in caring or advocating for the patient is valued by some caregivers and staff facilitation of this is considered one aspect of emotional care [55].

**Support or interventions to improve the experience of caregivers of patients with delirium or to help the caregiver support the patient**

Sixteen papers identify support for caregivers of patients with delirium or support for the caregiver to help the patient. Four subthemes were identified: (i) caregivers desire for information, (ii) caregivers wish for advice on how to respond to the patient, (iii) interventions to improve caregiver experience and (iv) caregiver interventions to improve patient outcomes.

**Caregivers desire for information about delirium:** Five studies identify informational needs of caregivers [27,29,35,55,56]. Caregivers would like clearer information about possible causes of delirium as well as what to expect in terms of progression and treatment [27,35,55,56]. Caregivers also desire information about how they can play a role in reducing delirium recurrence [55]. Caregivers of patients with advanced cancer [56] report that information early on or prior to the onset of delirium would be helpful. Some caregivers would like information on how the patient is likely to be feeling during a delirium episode [55]. Bereaved caregivers of patients who had experienced terminal delirium found that information about the causes, pathologies, possible treatments and expected course was helpful, as was reassurance regarding the universality of delirium [27].

**Caregivers wish for advice on how to respond to the patient with delirium:** Caregivers want advice on how to respond to patients with delirium [27,55]. Caregivers and healthcare professionals suggested approaches that seemed effective based on their experiences. Bereaved caregivers of patients with advanced cancer reported that talking to the delirious person, sitting quietly with them and explaining things that were happening provided a calming effect [56]. A calming environment, including quiet music, soothing touch and familiar surroundings were also reported by caregivers to be effective. Toye et al. [55] found that professional care that was calm, understanding or cheerful provided emotional
Supporting caregivers of patients with delirium

support, which reassured families and also provided indications to the caregiver of how to behave with the patient.

There were differing views on re-orienting the patient. Bereaved caregivers of patients with terminal delirium felt that staff should ‘respect the patient’s subjective world’ during delirium [27]. Similarly, bereaved caregivers of patients in a palliative setting felt that challenging the patient about the delirium could exacerbate their condition [56]. Otani et al. [36] advise caregivers of terminally ill patients with advanced cancer to converse with patients in a way that puts them at ease and to avoid ‘correcting mistakes’. In contrast, Gagnon et al. [44] advise caregivers to gently reorient the patient if they have ‘inappropriate thoughts’. Despite some consensus, clear evidence for the use of specific supportive behaviours, in particular reorienting, is lacking.

**Caregiver interventions focused on improving caregiver experience:** Five papers examined interventions to support caregivers. Three of these were leaflet or booklet interventions designed to improve family awareness and understanding of delirium [36,39,44] and two were educational interventions to improve caregiver knowledge of delirium [47,53].

Information leaflets or booklets along with routine discussion with a clinician can improve knowledge and confidence of caregivers. In Gagnon et al.’s study [44] caregivers were given a brochure consisting of a brief definition of delirium, and information on its principle symptoms, causes and treatments. Those who received the brochure were more confident about decision-making and were more likely to know what delirium was compared with those in the control group (35% vs. 21%). However, overall benefit of the intervention was modest, and there were no differences in caregiver mood across both groups. Adopting a similar approach, Otani et al. examined the perceived usefulness of a delirium information leaflet for 113 caregivers of patients receiving palliative care alongside the usual practice of verbal discussion [36,39]. A questionnaire sent to caregivers following bereavement found that 81% of respondents reported that the leaflet had been useful, helped them understand the dying process (84%), helped them identify what they could do for the patient (80%), helped them understand the patient’s physical condition (76%) and was useful in preparing for the patient’s death (72%) [36]. Knowledge was significantly higher in those that had received the leaflet compared with a historical control group [39]. However, as in the Gagnon et al. study, increased knowledge did not translate into emotional benefits.

Two papers reported educational interventions to increase knowledge of delirium with a view to prevention and early detection [47,53]. Keyser, Buchanan and Edge [53] ran a once-off community educational intervention for families of older adults (n=22). Questionnaires designed to assess knowledge of delirium showed some evidence for improvement post-intervention; however, the study was weakened by low participation rates, low questionnaire response rates, poor participation in follow-up interviews and a high risk of bias. Rosenbloom and Fick [47] designed the Nurse/Family Caregiver Partnership for Delirium Prevention educational programme to teach staff and families about delirium and to explore attitudes towards partnership. The results of a pre-test and post-test questionnaire showed improved knowledge of delirium and attitudes towards the caregiver–staff partnership, suggesting that educational interventions involving both staff and caregivers are feasible. However, the effect of these interventions on caregiver outcomes such as distress was not assessed.

**Caregiver interventions focused on improving patient outcomes:** Three intervention studies focused specifically on involving caregivers in interventions to improve outcomes for patients [49,50,52]. Black et al. [52] explored the effects of nurse-facilitated family participation in psychological care of the patient on the extent of delirium and psychological recovery following critical illness. Caregivers in the intervention group received a booklet containing information about delirium and a step-by-step guide to providing psychological care to the patient. The intervention did not reduce the incidence of delirium; however, patients who received the intervention demonstrated better psychological recovery and well-being compared with those in the control group at 4, 8 and 12 weeks post-admission, suggesting that caregiver interventions focusing on psychological care for the patient can have a beneficial impact on the patient that is sustained for some time.

Two intervention studies investigated whether involving caregivers would reduce the incidence and severity of delirium [49,50]. In Gagnon et al [50], a bedside nurse provided education to caregivers and was instructed to orient the patient as early as possible in the work shift. Adherence to the intervention was high; however, the intervention was ineffective in reducing delirium incidence or severity in patients compared with usual care. In contrast, Martinez et al. [49] examined the effectiveness of an intervention delivered by caregivers to reduce delirium occurrence in at-risk hospitalized patients. The intervention consisted of an educational component, as well as the provision of a clock and calendar in the patient’s room, avoidance of sensory deprivation, presence of familiar objects, reorientation of the patient by family members and extended visitation times. Delirium occurred in 13.3% of the control group (n=143) compared with only 5.6% of patients in the intervention group (n=144), suggesting a clear benefit.
These studies suggest that caregiver interventions are acceptable to at least some caregivers and professionals; however, evidence remains mixed in relation to their effect on delirium occurrence.

Conclusions

This integrative review demonstrates the high level of distress and negative emotions experienced by caregivers of patients with delirium. This confirms previous evidence highlighting significant caregiver distress during patient delirium [4,5,20] and extends previous findings by identifying the range of negative emotions that can be experienced including sadness, guilt, shame, anger and embarrassment [25,29,32,44,55,56]. High levels of emotional distress may be linked with the breakdown in the relationship with the patient, confusion and lack of information about the causes and course of delirium, as well as helplessness in relation to how to support the patient. Reducing caregiver distress and anxiety should be an important goal of future intervention.

The findings show that delirium disrupts the relationship between the caregiver and the cared-for person [31,37,56,57]. This disruption is temporary in some settings such as intensive care, but potentially permanent in palliative care settings as delirium may be irreversible. Anxiety is heightened because of uncertainty around whether the relationship between the caregiver and the patient can be re-established before death [37,56,57]. The need for communication with the patient increases as death approaches; caregivers want to say their final goodbye, and to understand what the patient may be trying to tell them [25,29]. Clinicians need to be sensitive to the relationship needs of caregivers and help them to relate to the patient during delirium. Clinicians can support the caregiver by providing advice on how to communicate and maintain aspects of the patient—caregiver relationship in spite of delirium that may be irreversible.

Several papers provide suggestions on the type of information that would be useful to caregivers of patients with delirium. This includes information on the causes of delirium, possible treatments, expected course and advice on how to behave around the cared-for person [27,29,35,55,56]. Intervention studies show that leaflets in conjunction with a discussion with the patient’s clinician can increase caregiver knowledge around delirium [36,39,44] and have the potential to improve patient outcomes [49,52]. Consequently, an informational component is recommended as part of any caregiver intervention.

A few papers describe behavioural strategies that caregivers report as being effective. These include playing quiet music, soothing touch and re-orienting the patient [55,56]. However, there is little evidence to support the effectiveness of particular strategies. Perspectives on whether it is best to re-orient the patient vary with some studies taking the view that re-orienting is helpful [44] and others believing that it is best to respect the patient’s subjective world even in delirium [27,36,56]. Further research on the effectiveness of different strategies that caregivers can use to support the patient during delirium is warranted.

Several papers demonstrate that caregivers can play a role in detection and prevention [33,38,42,43,46,52,54,58], symptom monitoring [41,43,54,58] and acting as an advocate for the patient [27,30]. Preliminary evidence suggests that such interventions are feasible and caregivers are generally positive about playing a more active role [46,54,55], although research on optimal levels of caregiver involvement in patient care is necessary. Given limited availability but growing demand for healthcare resources, interventions that optimize caregiver participation in patient care are vital.

There were shortcomings in the quality of many of the reviewed papers which is unsurprising given the challenging nature of research design, participant recruitment and retention in palliative care [59]. Most studies, apart from those focused on delirium detection and prevention, used retrospective rather than prospective designs—caregivers were asked to report on the delirium experience some time after the event. While this is a pragmatic approach in palliative care, it heightens the risk of recall bias, in particular if the patient has subsequently died. Poor response rates to questionnaires were common causing concern that those responding may differ from the general population of caregivers being studied. Only five quantitative studies included control groups [39,44,49,50,52], and there was only one RCT [49]. Outcome measures were often based on questionnaires with face validity, as opposed to psychometric testing, and some measures may have lacked the sensitivity required to detect differences in the outcome measure being assessed. Many of the qualitative studies were small scale, narrow in scope and focused on particular contexts and settings as is typical for qualitative work. However, such studies provide important insights into experiences of caregivers that inform the development of caregiver-focused interventions.

Recommendation for a future caregiver-focused research

The Medical Research Council identifies four key elements in the development and evaluation of complex interventions [60]: (i) development of an intervention, (ii) piloting and feasibility testing, (iii) evaluation and (iv) implementation. We identified 18 papers focused on the experiences of caregivers of patients with delirium providing a strong evidence base from which new interventions can be developed. We identified a smaller number of pilot and feasibility studies and eight intervention studies. This review reveals a need for further piloting and feasibility testing of caregiver interventions, and
components of interventions, as well as robust evaluation studies. Caregiver interventions in palliative care settings should include informational, emotional and behavioural elements. Given strong evidence of high levels of caregiver distress during patient delirium, reducing caregiver distress should be important outcome of any future intervention. Studies to explore the differences of the impact of delirium according to family dynamics (e.g. cohesive or conflicting) are also recommended.

Strengths and limitations of this review

A strength of this integrative review is that it includes quantitative and qualitative research as well as studies based on a range of designs. The qualitative and questionnaire studies provide important insights into caregiver’s experiences and support needs, while the intervention studies provide evidence of the feasibility and the effectiveness of different approaches to improving caregiver and patient outcomes. While the focus of this review is on palliative care settings, we drew on papers outside palliative care as many of the findings from medicine of the elderly and oncology are also relevant in palliative care. However, there are distinct differences in palliative care, most obviously the imminence of death and the possibility of terminal delirium, that are not central to non-palliative settings. Consequently, we have drawn attention to evidence specific to terminal care at certain points throughout this paper. Finally, given the range of countries, in which these studies were conducted, findings need to be interpreted in view of the cultural context and structure of the healthcare system in each country.

Concluding comment

This integrative review highlights the high levels of distress and negative emotion experienced by caregivers of patients with delirium. In palliative care settings, distress is heightened because of uncertainty around whether the relationship between the caregiver and the patient can be re-established before death. Significantly, we have identified the potential contributions of caregivers to managing this distressing syndrome and the potential for reciprocal benefits for patients and caregivers themselves. Caregiver interventions with informational, emotional and behavioural components are warranted to improve support for caregivers and to help support the patient. Reducing caregiver distress should be an important goal of any future intervention.

Acknowledgements

The posts of AF and JS were funded by Marie Curie: https://www.mariecurie.org.uk.

References

Supporting information

Additional supporting information may be found in the online version of this article at the publisher’s web site.

Table S1. Scottish Intercollegiate Guidelines Network Levels of Evidence.
Table S2 (online Supporting Information). Characteristics of papers included in review.