Does the use of specialist palliative care services modify the effect of socioeconomic status on place of death? A systematic review

Hong Chen¹, Donald J Nicolson², Una Macleod¹, Victoria Allgar³, Christopher Dalgliesh¹ and Miriam Johnson¹

Abstract

Background: Cancer patients in lower socioeconomic groups are significantly less likely to die at home and experience more barriers to access to palliative care. It is unclear whether receiving palliative care may mediate the effect of socioeconomic status on place of death.

Aim: This review examines whether and how use of specialist palliative care may modify the effect of socioeconomic status on place of death.

Design: A systematic review was conducted. Eligible papers were selected and the quality appraised by two independent reviewers. Data were synthesised using a narrative approach.

Data sources: MEDLINE, Embase, CINAHL, PsycINFO and Web of Knowledge were searched (1997–2013). Bibliographies were scanned and experts contacted. Papers were included if they reported the effect of both socioeconomic status and use of specialist palliative care on place of death for adult cancer patients.

Results: Nine studies were included. All study subjects had received specialist palliative care. With regard to place of death, socioeconomic status was found to have (1) no effect in seven studies and (2) an effect in one study. Furthermore, one study found that the effect of socioeconomic status on place of death was only significant when patients received standard specialist palliative care. When patients received more intense care adapted to their needs, the effect of socioeconomic status on place of death was no longer seen.

Conclusion: There is some evidence to suggest that use of specialist palliative care may modify the effect of socioeconomic status on place of death.

Keywords

Socioeconomic factors, palliative care, place of death, review, neoplasms

What is already known about the topic?

- Cancer patients in lower socioeconomic groups are significantly less likely to die at home and experience more barriers to access to palliative care.
- Intensity of home care including palliative care is associated with a home death.
- It is not clear whether and how use of specialist palliative care may modify the effect of socioeconomic status (SES) on place of death in cancer patients.
What this paper adds?

- There is some evidence to suggest that use of specialist palliative care may modify the effect of SES on place of death, thus contributing to the reduction of inequalities in achieving preferred home death.
- There is a need for further observational research that fully utilises the existing service patterns and compares the effect of SES on place of death between users and non-users of specialist palliative care.
- The common methodological challenges arise from the complexities in providing palliative care to diverse populations. These should be addressed in future research.

Implications for practice, theory or policy

- Palliative and supportive care services potentially can help cancer patients to die at their preferred place especially if they adapt level of care to patients’ unique needs.

Introduction

Despite considerable differences in personal, family and cultural meanings associated with dying at home, the overwhelming majority (approximately two-thirds) of cancer patients identify their own home as the preferred place of death; and this trend has been found across all socioeconomic groups. However, the majority of deaths from all diseases/conditions in most Western countries occur in a hospital. In the United Kingdom, although the proportion of people with cancer dying at home has been increasing since 2004, hospital remains the most common place of death, and a great discrepancy still exists between the preferred and actual place of death for cancer patients.

Furthermore, the ability to die at home is unequally distributed among cancer patients. Research evidence repeatedly demonstrates that cancer patients with higher socioeconomic status (SES) (e.g. better education, better occupation, higher income, living in more affluent or less deprived areas) are more likely than patients with lower SES to die at home rather than in an institution, and this pattern exists in countries with and without universal health care programmes.

According to Grande et al., differences in place of death based on SES may be associated with differences in access to palliative care services by different socioeconomic groups. They suggested that cancer patients in higher socioeconomic groups were more likely to die at home possibly because they were more able to access services which improved their chances of dying at home. In contrast, lower socioeconomic groups experience many barriers to access although their health care needs are greater than those of the general health care population. Lewis et al. identified barriers to access in four dimensions: availability, affordability, acceptability and geographical accessibility. For instance, barriers to access may arise from limited availability of services in deprived/poor areas (availability); inequitable distribution of services – services far away from and thus hard to reach by those with greatest need (geographical accessibility); limits of informal care arrangements, stigma and mistrust, and communication and health literacy issues in lower socioeconomic groups (acceptability); and financial burdens for the poor accessing services (affordability). As such, lower socioeconomic groups are more likely to rely on acute care services during illness progression and significantly less likely to die at home, both in countries with or without universal health care programmes.

However, it is not clear whether or not SES has the same effect on place of death when cancer patients receive specialist palliative care (see Box 1), having overcome the barriers to access; or how the effect of SES on place of death differs between those who receive specialist palliative care and those who do not. This systematic review of the worldwide literature therefore aims to examine whether or not and how use of specialist palliative care may modify the effect of adult cancer patients’ SES on place of death.

Methods

Two reviews (Q1 and Q2) were undertaken simultaneously, guided by the same protocol due to the overlap between their research questions. Q1 (this review) examines how use of specialist palliative care may modify the effect of adult cancer patients’ SES on place of death. Q2 examines how use of specialist palliative care may modify the effect of patients’ age, gender and ethnicity on place of death. Search was conducted separately for Q1 and Q2. However, search results were pooled together for a combined study selection because of overlapping data (i.e. some papers reported socio-demographic data which included SES, age, gender and ethnicity). The combined study selection helped to reduce the chance of missing eligible papers for both reviews. Once eligible papers were identified for each review, data extraction, quality appraisal and data synthesis were conducted separately. Q1 has been completed while Q2 is ongoing. This
Box 1.

**Palliative care** is an approach that improves the quality of life of patients and their families facing the problems associated with life-threatening illness, through the prevention and relief of suffering by means of early identification and impeccable assessment and treatment of pain and other problems, physical, psychosocial and spiritual.

*Source: World Health Organisation 2003*

**Specialist Palliative Care** is the active, total care of patients with progressive, advanced disease and their families. Care is provided by a multi-professional team who have undergone recognised specialist palliative care training. The aim of the care is to provide physical, psychological, social and spiritual support …

*Source: Tebbit, National Council for Palliative Care, 1999*

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**Figure 1.** PRISMA flow chart for Q1 and Q2.

Paper reports on Q1 in adherence to the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) Statement (http://www.prisma-statement.org/statement.htm), where applicable.

**Information sources and searches**

MEDLINE, Embase, CINAHL, PsycINFO and Web of Knowledge were searched (January 1997–December 2013). The search terms (MeSH headings and keywords) are summarised in Table 1 and the search strategy for MEDLINE and Embase is presented as an example in Appendix 1. Experts in the field were contacted to identify additional references; and the grey literature was searched using Caresearch (http://www.caresearch.com.au/Caresearch/Default.aspx). The reference lists of relevant reviews8–10,12 and the included studies13–21 were scanned to ensure comprehensiveness of the search. The review was limited to English language papers.

**Study selection (combined for Q1 and Q2)**

Two reviewers (D.J.N. and H.C.) screened all titles and abstracts independently. Full-text papers were retrieved for those eligible or indeterminable from titles and abstracts. Two reviewers (D.J.N. and H.C.) independently assessed the full text of all potentially relevant papers. Disagreement at each stage was resolved by consensus and with recourse to a third reviewer (M.J.) and a fourth (U.M.) if necessary.

**Eligibility criteria**

Papers were included if they reported original, empirical data showing the effect of both SES (e.g. income, education, occupation) and use of specialist palliative care (including hospice care, home care, inpatient or outpatient care) on place of death for adult cancer patients (≥18 years). Papers were excluded if the diagnosis was unknown or if the paper reported only non-malignant disease; if the study focussed exclusively on children and on preferences or attitudes about place of death rather than actual place of death.

**Data extraction**

A data extraction form was designed and piloted using three papers and adjustments made following consensus among the research team. Data were independently extracted by two reviewers (D.J.N. and H.C.) for 10% of the papers using the post-pilot form (see Appendix 2). As there was good agreement, the remaining data extraction was completed by D.J.N. Information was extracted from each included study on design and methods, patient characteristics, measure of SES, type of specialist palliative care, place of death outcome, statistical results demonstrating the relationships between SES, specialist palliative care, place of death and so on. Unreported data or clarifications were requested from first authors.
Quality assessment was conducted independently by two reviewers (D.J.N. and H.C.), using the Item Bank for Assessing Risk of Bias and Confounding for Observational Studies of Interventions or Exposures. Disagreement was resolved by team consensus. Main methodological components assessed include inclusion/exclusion criteria, strategy for recruiting participants, selection of the comparison group, variations in intervention/exposures, validity and reliability of measures, length of follow-up, attrition and control for confounding.

Narrative synthesis

All the included studies are non-randomised, observational studies, and there was considerable heterogeneity in methods, participants and outcomes. Under such circumstances, meta-analysis is not sensible and possible, and therefore, a narrative approach to synthesis is necessary and appropriate. Guidance on the Conduct of Narrative Synthesis in Systematic Reviews was followed, particularly with regard to tabulating results; exploring relationships between characteristics of individual studies and their reported findings, and between the findings of different studies; and overall assessment of the robustness of the evidence.

Results

Altogether, 7196 records were identified through database searching for both reviews (Q1 and Q2). Additional 37 records were identified through reference lists, grey literature search and the experts. After adjusting for duplicates, 6934 records remained. Of these, 273 full-text papers were retrieved and assessed in detail for inclusion for Q1 and Q2 (Figure 1). For this review (Q1), nine papers were included which reported nine different empirical studies (see Table 2).
Table 2. Characteristics of included studies.

<table>
<thead>
<tr>
<th>Ref.</th>
<th>Design and methods</th>
<th>Population</th>
<th>SES</th>
<th>Palliative care</th>
<th>Place of death</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>13.</td>
<td>Prospective cohort study: patients and carers followed up from patient’s admission to death</td>
<td>Spain</td>
<td>Level of education: 9.5% No schooling, 50.5% Primary, 15.3% Secondary, 5.3% High/higher, 19.5% Unknown</td>
<td>100% Home</td>
<td>64% Home</td>
<td>Univariate analysis: after receiving SPC, no differences found in place of death regarding patient’s education level</td>
</tr>
<tr>
<td></td>
<td>Patients from single service</td>
<td>380 Patients</td>
<td>Mean age: 66.76 years</td>
<td>60.5% Male</td>
<td>100% Cancer</td>
<td>19% Hospital, 16% Hospice, 1% Nursing home</td>
</tr>
<tr>
<td></td>
<td>Logistic regression</td>
<td>Spain</td>
<td>64% Home</td>
<td>19% Hospital, 16% Hospice, 1% Nursing home</td>
<td></td>
<td></td>
</tr>
<tr>
<td>14.</td>
<td>Retrospective cohort study: review of administrative and clinical database</td>
<td>United States</td>
<td>Family income: 1.8% &lt; US$20,000, 11.6% &gt; US$20,000–US$30,000, 29.3% &gt; US$30,000–US$40,000, 24.4% &gt; US$40,000–US$50,000, 32.9% &gt; US$50,000</td>
<td>Home: 54.9% routine care, 45.1% continuous care</td>
<td>77.4% Home</td>
<td>Multivariate analysis: among those who did not receive continuous care, the odds of non-home death increased as median annual household income decreased. Among those receiving continuous care, no significant difference was found in rates of non-home death across income levels.</td>
</tr>
<tr>
<td></td>
<td>Patients from multiple services in multiple regions</td>
<td>61,063 Patients</td>
<td>Mean age: 77.7% ≥ 65 years, 48% Male, 70.2% White, 64.2% Cancer</td>
<td>Home: 54.9% routine care, 45.1% continuous care</td>
<td>22.6% Elsewhere</td>
<td>Multivariate analysis: among those who did not receive continuous care, the odds of non-home death increased as median annual household income decreased. Among those receiving continuous care, no significant difference was found in rates of non-home death across income levels.</td>
</tr>
<tr>
<td></td>
<td>Logistic regression</td>
<td>United States</td>
<td>77.7% ≥ 65 years</td>
<td>48% Male</td>
<td>70.2% White</td>
<td>64.2% Cancer</td>
</tr>
<tr>
<td>15.</td>
<td>Prospective cohort study: patients followed up from admission to death or end of study period</td>
<td>United States</td>
<td>Family income (n = 142): 31.7% &lt; US$20,000, 39.4% ≥ US$20,000 and &lt; US$40,000, 28.9% &gt; US$40,000</td>
<td>100% Inpatient</td>
<td>23.7% Home</td>
<td>10.5% Nursing home</td>
</tr>
<tr>
<td></td>
<td>Patients from multiple services in single region</td>
<td>180 Patients</td>
<td>Mean age: 67 years, 49.4% Male, 79.4% White</td>
<td>100% Cancer</td>
<td>40.8% Hospice, 25% Hospital</td>
<td>10.5% Nursing home</td>
</tr>
<tr>
<td></td>
<td>Cox proportional-hazards regression</td>
<td>United States</td>
<td>67 years</td>
<td>49.4% Male</td>
<td>79.4% White</td>
<td>100% Cancer</td>
</tr>
<tr>
<td>16.</td>
<td>Cross-sectional survey of home hospice care agencies</td>
<td>Japan</td>
<td>Financial sufficiency: 40.7% yes, 59.3% No</td>
<td>100% Home</td>
<td>64.8% Home</td>
<td>Univariate analysis: after receiving SPC, no differences found in place of death regarding patient’s financial situation</td>
</tr>
<tr>
<td></td>
<td>Patients from multiple services in multiple regions</td>
<td>528 Patients</td>
<td>Mean age: 75 years, 59.5% Male</td>
<td>100% Cancer</td>
<td>35.2% Hospital</td>
<td>Univariate analysis: after receiving SPC, no differences found in place of death regarding patient’s financial situation</td>
</tr>
<tr>
<td></td>
<td>Logistic regression</td>
<td>Japan</td>
<td>75 years</td>
<td>59.5% Male</td>
<td>100% Cancer</td>
<td>35.2% Hospital</td>
</tr>
<tr>
<td>17.</td>
<td>Cross-sectional survey of home hospice care agencies</td>
<td>Japan</td>
<td>Financial sufficiency: Proportion not reported</td>
<td>100% Home</td>
<td>64.8% Home</td>
<td>Univariate analysis: after receiving SPC, no differences found in place of death regarding patient’s financial situation</td>
</tr>
<tr>
<td></td>
<td>Patients from multiple services in multiple regions</td>
<td>428 Patients</td>
<td>Mean age: 75 years, 57.8% Male</td>
<td>100% Cancer</td>
<td>35.2% Hospital</td>
<td>Univariate analysis: after receiving SPC, no differences found in place of death regarding patient’s financial situation</td>
</tr>
<tr>
<td></td>
<td>Logistic regression for multivariate analysis</td>
<td>Japan</td>
<td>Mean age: 75 years</td>
<td>57.8% Male</td>
<td>100% Cancer</td>
<td>35.2% Hospital</td>
</tr>
</tbody>
</table>

(Continued)
or dedicated units located in hospitals. In another study, patients received home palliative care in addition to outpatient and/or inpatient palliative care. In the remaining seven studies, patients received home palliative care.

Categorisation of place of death differs in the studies. All the studies have home as an outcome of place of death, which is compared with ‘hospital, hospice and aged/residential care home’ in three studies, with ‘other than home’ in one study, with ‘hospital + hospice’ in one study and with ‘hospital’ in four studies.

Common methodological limitations included sample unrepresentative, insufficient controlling for confounding factors and imprecise measurement of SES and use of specialist palliative care, all of which exist in the included studies in varying degrees (see ‘Discussion’).

**Table 2. (Continued)**

<table>
<thead>
<tr>
<th>Ref.</th>
<th>Design and methods</th>
<th>Population</th>
<th>SES</th>
<th>Palliative care</th>
<th>Place of death</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>18.</td>
<td>Cross-sectional survey of home hospice care agencies</td>
<td>Japan 568 Patients Mean age 73 years 59.7% Male</td>
<td>Financial sufficiency: 47.4% yes 52.6% No</td>
<td>100% Home</td>
<td>54.9% Home 45.1% Hospital</td>
<td>Univariate analysis: after receiving SPC, no differences found in place of death regarding patient’s financial situation</td>
</tr>
<tr>
<td>19.</td>
<td>Prospective cohort study: patients, carers and family physicians followed up from patient’s admission to death</td>
<td>Canada 73 Patients Mean age 68 years 48% Male</td>
<td>Financial resources sufficient (n = 67): 77.6% Yes 22.4% No</td>
<td>100% Home</td>
<td>47% home 53% Hospital + Hospice</td>
<td>Univariate analysis: after receiving SPC, no differences found in place of death regarding patient’s financial situation</td>
</tr>
<tr>
<td>20.</td>
<td>Prospective cohort study: primary carers followed up from patient’s admission to death</td>
<td>Canada 110 Patients Mean age 70 years 46.4% Male 100% Cancer</td>
<td>Carstairs Deprivation score: 24.8% ≤ 0.5 25.7% 0.51–0.64 24.8% 0.65–1.07 24.8% ≥ 1.08</td>
<td>100% Home</td>
<td>66.4% Home 33.6% Hospital</td>
<td>Univariate analysis: after receiving SPC, no differences found in place of death regarding patient’s deprivation level</td>
</tr>
<tr>
<td>21.</td>
<td>Retrospective cohort study: review of hospice chart</td>
<td>New Zealand 1268 Patients Mean age 65 years 48% Male 82% European 82% Cancer</td>
<td>Community Services Card: 42% Yes 28% No</td>
<td>100% Home + other</td>
<td>28% Home 46% hospice 8% hospital 16% care home</td>
<td>Multivariate analysis: relative to those people without a CSC, people with a CSC were more likely to die at acute hospital than at home (OR = 2.09, 95% CI: 1.096–3.996)</td>
</tr>
</tbody>
</table>

SES: socioeconomic status; SPC: specialist palliative care; CSC: Community Services Card; OR: odds ratio; CI: confidence interval.

**Does use of specialist palliative care modify the effect of SES on place of death?**

In seven studies (sample size: 73–568 patients), univariate or bivariate analysis found no statistically significant differences in place of death regarding SES among patients who received home care or inpatient care (see Table 2). Two studies did find statistically significant differences in place of death regarding patient’s financial situation. Taylor et al. found SES to be a predictive factor of place of death for patients receiving specialist palliative care. In this study, retrospective administrative data were obtained from a hospice in New Zealand. Although 1268 patients were included, only 890 patients had information recorded about whether or not they had a Community Services Card.
Services Card (CSC). Having a CSC is a crude indicator of lower SES. Multivariate analysis of these patients (controlling for age, ethnicity, marital status, gender and medical condition) showed that relative to those people without a CSC, people with a CSC were more likely to die at acute hospital than at home (odds ratio (OR) = 2.09, 95% confidence interval (CI): 1.09–3.996). There was no significant difference between those with/without a CSC regarding likelihood of dying in an aged/residential care facility relative to home (OR = 1.50, 95% CI: 0.88–2.565). Likewise, there was no evidence for CSC status to differentiate between likelihood of dying in the hospice inpatient unit relative to home (OR = 1.00, 95% CI: 0.71–1.42).

Barclay et al.’s14 was the largest study, which included 61,063 patients (64.2% with cancer). It was a retrospective administrative dataset obtained from a hospice provider that operated 26 hospice programmes in eight states in the United States. Altogether, 77.4% of 61,063 patients died at home. Two levels of home care (routine versus continuous) were provided to these patients based on their needs; 54.9% of the patients received routine home care only (periodic home visits) and 45.1% received continuous home care, that is, a short-term intense period of care that included the presence of hospice staff providing care for a minimum of 8 h in a 24-h period, with at least half the care provided by a nurse. In the multivariate model, the income × level of care interaction was significant in predicting place of death, after controlling for confounding factors including sex, age, ethnicity, marital status, disease type, payment source, enrolment in a health maintenance organisation, relationship of the primary caregiver to the enrolee, days in hospice care and hospice programme location by region. In the routine-care group, the odds of non-home death increased as median annual household income decreased. Those in the lowest income group compared with those in the highest income group had almost twice the odds of dying elsewhere rather than home (OR = 1.76, 95% CI: 1.48–2.09). In comparison, in the continuous-care group, no significant difference was found in rates of non-home death across income levels; and for all income levels, a smaller proportion of those receiving any continuous care (vs no continuous care) died elsewhere rather than home.

Discussion

Gomes and Higginson8 found strong evidence for the association of SES (i.e. education, social class, income) with place of death for patients with cancer. They found that the evidence that supported the influence of SES was stronger than the evidence that showed no effect, suggesting that higher SES increased the odds of home death. They also found strong and consistent evidence to suggest that intensity of home care (more home care input and more frequent visits) was associated with a home death. However, their review did not examine the link between the two associations, that is, whether or not SES and intensity of home care may interact together to impact on place of death in cancer patients. This review is particularly interested in whether or not and how use of specialist palliative care may modify the effect of SES on place of death. The majority of the included studies (seven) found that SES had no effect on place of death among users of home-based specialist palliative care13,16–20 or inpatient specialist palliative care.15 Only one study found that SES had an effect on place of death among users of specialist palliative care (home, outpatient and/or inpatient).21 However, the biggest study14 (n = 61,063) found that the effect of SES on place of death was only significant when patients received standard home-based specialist palliative care. When patients received more intense home care that was adapted to their needs, SES stopped having an effect on place of death. Moreover, intensity of home care adapted to patients’ needs particularly at the end of life increased the chance of home death across all socioeconomic groups. Particularly, for patients in lower socioeconomic groups, more intense home care meant that they had a higher chance of home death despite the fact that they had been identified as having increased needs. These findings thus suggest that use of specialist palliative care may modify the effect of SES on place of death and thus help to reduce inequalities in achieving preferred home death.

Nonetheless, definite conclusions cannot be reached because not only is there a paucity of empirical studies identified but also considerable heterogeneity exists in the study characteristics. Moreover, all the studies had some methodological weakness, which may have contributed to underestimates or overestimates of the actual effect of SES and use of specialist palliative care on place of death.

Selection bias was common. In four studies,13,19–21 patients were recruited from a single service; even in studies where patients were recruited from multiple services in one region15 or in multiple regions,14,16–18 findings may not be generalisable to other populations or geographical areas, since palliative care services and health and social care environments differ in different areas, regions in the same country and between countries.

Another major methodological problem is controlling for confounding variables because there is a complicated network of factors that affect both use of specialist palliative care services and place of death.3,26 These factors fall into three groups: those related to the illness, the individual and the care and social environment.8 Both studies,14,21 in which SES was entered into the multivariate regression model as a significant predictor of place of death, lacked specific patient, clinician, health system or environmental variables that might represent important sources of residual confounding.27
One of the key variables of this review – SES – is a complex concept that often refers to the position of individuals, families, households or other aggregates on one or more dimensions of stratification. These dimensions include income, education, prestige, wealth or other aspects of standing that members of society deem salient. Each of these dimensions is capable of exerting separate effects although they are interrelated. Bollen et al. found that although SES was widely referred to in health and health care research, there is a lack of consensus with respect to its conceptual meaning and measurement. The same problem persists in this review. SES was measured differently across studies – by different singular or aggregated dimensions (i.e. education, household income, financial resources and deprivation). As such, outcomes of SES are not comparable across studies. Also, data availability clearly influenced the ways in which SES was measured, leading to imprecise and ambiguous measurement (e.g. Community Services Card).

With regard to use of specialist palliative care, practice variations prevail in and across the studies. First, the composition of multi-disciplinary team differed across the studies that reported such information: in the Spanish study, it consisted of doctor, advanced nurse, assistant nurse and administrative clerk; in an American study, it consisted of doctor, nurse, home health aide, chaplain and social worker; in a Canadian study, it consisted of nurse, social worker, occupational therapist, physical therapist, respiratory therapist and volunteers; in the Japanese studies, it consisted of home care nurse and patient’s family doctor and/or hospital doctor; in the New Zealand study, it consisted of doctor, nurse, chaplain, social worker, counsellor and music, massage, occupational and physiotherapists. Second, the frequency and intensity of periodic home care varied. In some studies, level of care was adapted to the needs of patients and families; for example, in an American study, a short-term intense period of care was provided to patients at the end of life, which included a minimum of 8-h care in a 24-h period, with at least half the care provided by a nurse; in a Canadian study, 4–5 h/day or more of services can be provided to patients with advanced disease and 24-h nursing and personal care may be provided for a short period at end of life. Third, palliative care services were provided free of charge in Spain, Canada and New Zealand, whereas in United States and Japan, they were mainly covered by medical insurance. Such varying practices were not sufficiently reflected in the measures for the use of specialist palliative care.

In all, this review found some evidence to suggest that use of specialist palliative care may modify the effect of SES on place of death and is the first to synthesise such evidence. The general implication of the findings for practice is that palliative and supportive care services potentially can help cancer patients in all socioeconomic groups to die at their preferred place, especially if they adapt level of care to patients’ unique needs. The review also identified a need for more rigorous empirical evidence in this area and highlighted the methodological issues to be dealt with in future research. Due to the challenges in the use of randomised controlled trial designs, observational data sources and study designs will continue to be strong contributors to building the evidence base for palliative care research and practice. As such, future observational research needs to consider how to (1) reduce selection bias; (2) collect data about a myriad of confounding factors, particularly those related to the individual and the care and social environment; (3) better define and measure SES; and (4) more precisely measure use of specialist palliative care to reflect practice variations and complexities in service provision. To further ascertain that use of specialist palliative care can modify the effect of SES on place of death, a better design would be to utilise existing patterns of practice and prospectively or retrospectively compare the effect of SES on place of death between users and non-users of specialist palliative care.

Conclusion

There is some evidence to suggest that use of specialist palliative care may modify the effect of SES on place of death, thus contributing to reducing inequalities in achieving the preferred home death. This also means that palliative and supportive care services potentially can help cancer patients die at their preferred place by adapting level of care to their unique needs. However, more rigorous empirical studies are needed to further confirm this finding. Future observational studies need to fully utilise the existing service patterns and compare the effect of SES on place of death between users and non-users and also pay more attention to selection bias, controlling for confounding factors and measurement of SES and use of specialist palliative care.

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Declaration of conflicting interests

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References

Appendix 1

Search strategy—MEDLINE and Embase

(MEDLINE and Embase Jan 1997–Dec 2013)

1. exp Neoplasm/
3. cancer*.mp. [mp=ti, sh, hw, tn, ot, dm, mf, dv, kw, nm, kf, ps, rs, ui]
4. or/1-3
5. exp Social Class/
7. Socio-econom*.mp.
8. socioeconom*.mp.
9. Inequalit*.mp. [mp=ti, sh, hw, tn, ot, dm, mf, dv, kw, nm, kf, ps, rs, ui]
10. exp Poverty/
11. poverty.mp.
12. or/5-11
13. exp Hospices/
15. exp Terminal care/
16. Terminal care.mp.
17. exp Palliative care/
18. Palliat$.mp.
19. exp End-of-life/
20. place of death.adj
21. location.mp. [mp=ti, sh, hw, tn, ot, dm, mf, dv, kw, nm, kf, ps, rs, ui]
22. Hospitals/
23. Hospitals.mp.
24. Health Services Access*.mp. [mp=ti, sh, hw, tn, ot, dm, mf, dv, kw, nm, kf, ps, rs, ui]
25. exp Inpatient/
26. Inpatient.mp.
27. exp Outpatient/
28. Outpatient.mp.
29. or/13-28
30. 4 and 12 and 29
31. limit 30 to English language
32. limit 31 to yr=’1997–Current’
33. limit 32 to humans
34. remove duplicates from 33

Appendix 2

Data extraction form

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</tr>
<tr>
<td>Article title:</td>
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<tr>
<td>Article type:</td>
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<td>Reviewer:</td>
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1. NATURE OF STUDY.

1.1 Research aim(s)/question(s):
- Population:
- Exposure:
- Outcome:

1.2 Study design:
- Cross-sectional survey
- Prospective cohort
- Retrospective cohort
- Other:

1.3 Setting of study:
- Country:
- Urban/rural/mixed:
- Other:

1.4 Study period (in months):
- Duration:
- Follow-up (if prospective):

1.5 Sample selection Method:
- Random
- Convenience
- Consecutive
- Other:

1.6 Data source/method of measurement:
- Record review
- Questionnaire
- Other:

1.7 Method of data analysis/statistical tests:

2. PARTICIPANT DETAILS.

2.1 Study Inclusion/exclusion criteria:
- Inclusion:
- Exclusion:
2.2 Participant characteristics:

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3. MAIN FINDINGS.

3.1 Numerical results re association between study variables:

3.2 Summary of narrative results re association between study variables:

3.3 Authors conclusions: