Exploring costs, cost components, and associated factors among people with dementia approaching the end of life: A systematic review

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Registered PROSPERO (CRD42020166337)

Abstract

Introduction: Understanding costs of care for people dying with dementia is essential to guide service development, but information has not been systematically reviewed. We aimed to understand (1) which cost components have been measured in studies reporting the costs of care in people with dementia approaching the end of life, (2) what the costs are and how they change closer to death, and (3) which factors are associated with these costs.

Methods: We searched the electronic databases CINAHL, Medline, Cochrane, Web of Science, EconLit, and Embase and reference lists of included studies. We included any type of study published between 1999 and 2019, in any language, reporting primary data on costs of health care in individuals with dementia approaching the end of life. Two independent reviewers screened all full-text articles. We used the Evers’ Consensus on Health Economic Criteria checklist to appraise the risk of bias of included studies.

Results: We identified 2843 articles after removing duplicates; 19 studies fulfilled the inclusion criteria. 16 were from the United States. Only two studies measured informal costs including out-of-pocket expenses and informal caregiving. The monthly total direct cost of care rose toward death, from $1787 to $2999 USD in the last 12 months, to $4570 to $11921 USD in the last month of life. Female sex, Black ethnicity, higher educational background, more comorbidities, and greater cognitive impairment were associated with higher costs.

Discussion: Costs of dementia care rise closer to death. Informal costs of care are high but infrequently included in analyses. Research exploring the costs of care for people with dementia by proximity to death, including informal care costs and from outside the United States, is urgently needed.

KEYWORDS
Alzheimer’s disease, cost of care, cost of illness, dementia, end-of-life, health care cost, systematic review
INTRODUCTION

Dementia is currently the fifth leading cause of death globally,\(^1\) and the number of people dying with dementia in the world is projected to increase from \(\approx 2\) million deaths in 2016 to more than 7 million in 2060.\(^2\) When approaching death, people with dementia experience a rapid increase in the incidence of fatigue, pain, and dizziness;\(^3\) potentially burdensome hospital admissions;\(^4\) and emergency department visits.\(^5\)

Assessing the economic impact of illness on society is key to understanding resource needs, identifying main cost components, explaining the variability of costs, and identifying management pathways to inform policy.\(^6\) Two published systematic reviews have examined aspects of dementia care costs, including hospital, outpatient, drugs, informal, and indirect costs.\(^7,8\) In Quentin et al., direct and indirect costs for patients with severe dementia were almost twice the costs for people with mild dementia.\(^7\) In Schaller et al., the main drivers of total dementia costs were nursing home expenditures and informal costs due to home-based care.\(^8\) Neither systematic review examined end-of-life care costs.

Understanding the costs of care at the end of life is particularly important as health-care costs have been shown to increase by proximity to death.\(^9\) A systematic review of studies exploring the cost of caregiving among people receiving palliative care at the end of life found informal caregiving could account for 27% to 80% of total costs and have a significant impact on families and carers.\(^10\) Unlike the cost of dementia care, the evidence on costs of care approaching the end of life is not well understood.

We aimed to understand the costs of care for people with dementia approaching the end of life by answering the following questions:

1. Which cost components have been measured in studies reporting the costs of care in people with dementia approaching the end of life?
2. What are the costs of care for people with dementia approaching the end of life and how do these change closer to death?
3. Which factors are associated with the costs of care among people with dementia approaching the end of life?

METHODS

We undertook a systematic review of primary studies reporting costs of care for people with dementia approaching the end of life. The review protocol was registered with PROSPERO (registration number CRD42020166337).

Eligibility criteria

Studies were included if they reported direct or informal health-care costs for individuals who died with dementia (for retrospective studies) or who are likely to be approaching the end of life based on clinical prognosis, such as "patients with severe dementia" or "patients with dementia who are thought to be in the last year of life" (for prospective studies). Studies that included individuals with different conditions were included only if results for people with dementia were reported separately. Any type of study reporting primary data on costs of care was considered, including both incidence and prevalence-based analyses. We did not include modeling studies that empirically estimated costs of care unless based on primary data. We included only articles published between 1999 and 2019 in recognition that articles published before this time are unlikely to have financial relevance today. Table 1 shows the inclusion and exclusion criteria.

Information sources

We searched the electronic databases CINAHL, Medline, Cochrane, Web of Science, EconLit, and Embase and reference lists of included studies from January 1999 to December 2019. Additional references were retrieved through informal searches on Google and consultations with a health-care economist (DY).

Search strategy

The search strategy included the concepts of "dementia," "end of life," and "health-care costs." Medical subject headings (MeSH) index terms were used together with free text to capture relevant articles. The terms recommended by the Cochrane Dementia and Cognitive Improvement Group for dementia were used. The complete search strategy is available in Table S1 in supporting information.
**TABLE 1**  Inclusion and exclusion criteria

| **Inclusion criteria** | **Exclusion criteria** |
|------------------------|------------------------|
| **Type of study**       |                        |
| - Any type of study with primary data or secondary data analysis | - Discussion papers |
| - Articles that used modeling techniques also could be included if they used primary data | - Letters, editorials, or commentaries |
|                        | - Conference abstract  |
|                        | -Protocols             |
|                        | - Trial registration    |
| **Population**         |                        |
| - Individuals with any type of dementia | - Articles reporting only the cost of an intervention |
| - Decedents or people likely to be in their last year of life based on prognosis | |
| - If an article included individuals with different conditions, it was included only if data for dementia patients was reported separately | |
| **Outcome**            |                        |
| - All types of health-care costs (direct, hospital, outpatient services, community, social services, informal care costs, out-of-pocket costs, etc.) | |
| - Cost-effectiveness or cost-utility analysis papers were included only if they reported the cost for the control group, and the control group received usual treatment | |
| - Articles that reported costs for longer than 12 months before death were included | |
| **Intervention**        |                        |
| - Any type of health-care service provision | |
| **Language**           | - Any language         |
| **Years**              | - Articles published between 1999 and 2019 |
person for the entire end-of-life period considered in the study, and the estimated average costs per month. One reviewer (JL) performed the analysis; tables and results were discussed and checked by three other reviewers (KS, DY, IH).

For question 3, studies were grouped based on whether they reported sociodemographic, illness-related, or health-care service-related factors associated with the cost of care for people with dementia at the end of life. A preliminary descriptive paragraph on each study was produced by one reviewer (JL) and discussed with three other reviewers (KS, IH, DY). A summary of the main points was then agreed.15

3 | RESULTS

3.1 | Characteristics of included studies

After removing duplicates, 2843 articles were retrieved from databases and hand searches. One hundred eighty-eight articles were included for full-text screening; 19 fulfilled the inclusion criteria and were considered for the analysis (Figure 1).

Of the 19 included articles, 16 were from the United States,16–31 two were from Europe (United Kingdom and the Netherlands),32,33 and one from Australia.34 Twelve (75.0%) included studies had a retrospective design.16,17,19,20,22,23,25,26,29,31,33,34 All but one study identified costs from claims records,16–20,22–34 while one study used questionnaires.21 Studies focused on different end-of-life periods, ranging from the last month of life to 60 months before death. Three studies included people with Alzheimer’s disease (AD) only,21,28,34 and the other 16 included people with any type of dementia diagnosis.

3.2 | Quality appraisal of included studies

The mean quality score of all included studies on the CHEC checklist was 82.8% (proportion of "yes" answers). The lowest score was 64.3% and 12 studies scored more than 80% (Table S3 in supporting information). The most common source of poor quality was the lack of an explicit mention of the economical perspective (such as government, insurers, patients, or societal) in 16 studies and the absence of an appropriate discussion regarding ethical and distributional issues in eight studies.

3.3 | Question 1: Which cost components have been measured in studies reporting the costs of care in people with dementia approaching the end of life?

Types of costs measured included hospital, community care, hospice, long-term care facility (nursing or residential home) expenses and informal costs (Table 3). Of the 19 included articles, 16 included
| Author             | Country | Study design     | Age             | Type of dementia diagnosis | Sex (% female) | Ethnicity | Residency | Cost perspective | Number of people with dementia | Description                                                                 | Total costs | Mean for total end-of-life period (months) | Mean by month (USD 2019) | What is included in the total costs |
|--------------------|---------|------------------|-----------------|-----------------------------|----------------|-----------|------------|------------------|-------------------------------|-------------------------------|-------------------------|-------------------------------|----------------------------|--------------------------|
| Lamb VL., 2008     | USA     | Retrospective cohort study | >85: 37.5% (5 years BD) | ADRD                        | 65.0%          | White     | Nursing home | Not reported     | 2449                          | Medicare expenditure difference between people with and without dementia from claim records | 60            | 7587.1†                     |                               | √ √ √ √ √                 |
| Ornstein KA., 2018 | USA     | Prospective cohort study | mean 88.8       | ADRD                        | 67.4%          | White     | Not reported  | Not reported     | 86                            | Total Medicare expenditures from claim records | 36            | 159397.9                   | 4427.7                     | √ √ √ √ √                 |
| Goldfeld KS., 2011 | USA     | Prospective cohort study | mean 85.3 (SD 7.5)* | ADRD                        | 85.5%*         | White     | Nursing home | Not reported     | 177                           | Total expenditures (use from clinical records and costs estimated from Medicare services information) | 18            | 8751.7                     | 486.2                      | √ √ √ √ √                 |
| McCormick WC., 2001| USA     | Case control study  | mean 84.3 (SD 6.8) | ADRD                        | 52.6%          | White     | Not reported  | Not reported     | 396                           | Total Medicare expenditures from claim records | 12            | 21448.7                    | 1787.4                     | √ √ √ √ √                 |
| Gozalo P., 2015    | USA     | Retrospective cohort study | mean 85.6*       | ADRD                        | 64.0%*         | White     | Nursing home | Not reported     | 162459                        | Total Medicare and Medicaid expenditures pp from claim records | 12            | 32494.1                    | 270.78                     | √ √ √ √ √                 |
| Zhu CW., 2017      | USA     | Prospective cohort study | mean 85.3 (SD 7.1) | ADRD                        | 85.1%*         | White     | Not reported  | Not reported     | 49†                           | Total Medicare expenditures pp from claim records | 12            | 21884.6                    | 182.37                     | √ √ √ √ √                 |
| Daras LC., 2017    | USA     | Retrospective cohort study | >85: 36.5%*      | ADRD                        | 55.1%*         | White     | Nursing home | Not reported     | 1426                          | Medicare payments for outpatient, ED visits and hospitalizations from interviews and claim records | 12            | 21879.5                    | 182.33                     | √ √                     |
| van der Plas AG., 2017| NDL    | Retrospective cohort study | mean 86.1 (SD 6.9) | ADRD                        | 67.6%          | Not reported | Not reported  | Not reported     | 3586                          | Total cost from billed insurance costs (Hospitals use, out-of-hospital, home care, care homes and nursing homes) | 12            | 35 985.3                   | 2998.8                     | √ √ √ √ √                 |
| Author           | Country | Study design          | Age         | Type of dementia diagnosis | Sex (% female) | Ethnicity | Residency | Cost perspective | Number of people with dementia | Description                                                                 | Total costs         | Mean for total end-of-life period (USD 2019) | Mean by month (USD 2019) | In-hospital care | ED visits | Outpatient | Community care | Long-term care facility | Hospice | informal care | Out-of-pocket |
|------------------|---------|-----------------------|-------------|-----------------------------|----------------|-----------|-----------|------------------|-------------------------------|-----------------------------------------------------------------------------|---------------------|---------------------------------------------|----------------------------|-----------------|-----------|------------|----------------|---------------------|---------|-------------|------------|
| Spilsbury K., 2017 | AUS     | Retrospective cohort study | >80:46.5%* | AD                          | 45.9%*         | Not reported | Nursing home | 21.2%*          | 605 Gr 6 cohort averaged hospital cost per day x4.7 days in the last 12 months of life from claim records |                      | 12                                           | 62.1                      | -               | √         |           |               |                     |         |              |            |
| Pyenson B., 2019  | USA     | Retrospective cohort study | mean 86.2   | ADRD                        | 65.4%*         | Not reported | Not reported | Not reported    | 131855 Total Medicare expenditures from claim records |                      | 12                                           | 34760.2                   | 2896.7           | √ √ √ √   | √         |               |                     |         |              |            |
| Pyenson B., 2004  | USA     | Prospective cohort study | >85%: 13.4%* | AD                          | 46.5%*         | Not clear | Not reported | Not reported    | 151 Total Medicare expenditures from claim records |                      | mean days 183.8 | 46300.9                   | 7716.8           | √ √ √ √   | √         |               |                     |         |              |            |
| Sampson E., 2012 | UK      | Other                 | 64 to 84 years | ADRD                        | 33.3%          | White      | Nursing home | 11.1%           | 9 Total expenditures (use from clinical records and interviews and costs estimated from NHS services information) |                      | 6                                           | 23597.8                   | 3933.0           | √ √ √ √   | √         |               |                     |         |              |            |
| Nicholas LH., 2014 | USA     | Retrospective cohort study | mean 84.8   | ADRD                        | 59.8%          | White      | Nursing home | 56.2%           | 2509 Total Medicare expenditures from claim records |                      | 6                                           | 34000.7                   | 5666.8           | √ √ √ √   | √         |               |                     |         |              |            |
| Zuckerman RB., 2016 | USA    | Retrospective cohort study | >85: 59.9%* | ADRD                        | 69.2%          | White      | Nursing home | 56.9%           | 244674 Total Medicare expenditures from claim records |                      | 6                                           | 42395.9                   | 7066.0           | √ √ √ √   | √         |               |                     |         |              |            |
| Crouch E., 2019   | USA     | Retrospective cohort study | >85: 61.7%* | ADRD                        | 69.0%          | White      | Not reported | Not reported    | 7895 Total Medicare expenditures from claim records among those with an expenditure |                      | 6                                           | 24483.8                   | 4080.6           | √ √ √ √   | √         |               |                     |         |              |            |
| Miller SC., 2004  | USA     | Retrospective cohort study | mean 85.9 (SD 7.1) | ADRD | Not reported | Not reported | Nursing home | 100%           | 2558 Total Medicare and Medicaid expenditures from claim records |                      | 1                                           | 11920.9                   | 11920.9         | √ √ √ √   | √         |               |                     |         |              |            |
| Gozalo PL, 2008   | USA     | Retrospective cohort study | >85: 55.7%* | ADRD                        | 70.7%*         | White      | Nursing home | 100%           | 2556 Total Medicare and Medicaid expenditures from claim records |                      | 1                                           | 11914.2                   | 11914.2         | √ √ √ √   | √         |               |                     |         |              |            |

(Continues)
### TABLE 3 (Continued)

| Author          | Country | Study design          | Age               | Type of dementia diagnosis | Sex (% female) | Ethnicity | Residency | Cost perspective | Number of people with dementia | Description                                                                 | End-of-life period (months) | Mean total end-of-life period (USD 2019) | Mean by month | What is included in the total costs |
|-----------------|---------|-----------------------|-------------------|-----------------------------|----------------|-----------|-----------|-----------------|-------------------------------|-------------------------------------------|--------------------------|---------------------------------|-----------------|---------------------------------|
| van der Plas AG., 2017 | NDL | Retrospective cohort study | mean 86.1 (SD 6.9) | ADRD | 67.6% | Not reported | Not reported | Not reported | 3586 | Total cost from billed insurance costs (Hospitals use, out-of-hospital, home care, care homes and nursing homes) | 1 | 4570.3 | 4570.3 | ✓ ✓ ✓ ✓ ✓ ✓ ✓ ✓ ✓ ✓ ✓ ✓ ✓ ✓ ? |
| Kelley AS., USA, 2013 | USA | Prospective cohort study | mean 84.3 (SD 7.6)* | AD | 57.0%* | White 78.8% | Nursing home 21.2%* | Not reported | 651 | Health-care out-of-pocket expenditures from interviews (insurance, hospital, physician, medication, nursing home, hired helpers, in-home medical care, and other expenses) | 60 | 80118.5 | 1335.3 | ✓ ✓ ✓ |
| Kelley AS., USA, 2015 | USA | Retrospective cohort study | mean 88.4 (SD 6.4) | ADRD | 68.1% | Non-Black 87.0% | Not reported | Societal perspective | 555 | Total Medicare expenditures from claim records + Health-care out-of-pocket expenditures from interviews (insurance, hospital, physician, medication, nursing home, hired helpers, in-home medical care, and other expenses) | 60 | 335966.3 | 5599.4 | ✓ ✓ ✓ ✓ ✓ ✓ ✓ ✓ ✓ ✓ ✓ ✓ ✓ ✓ |

Abbreviations: AD, Alzheimer’s disease; ADRD, Alzheimer’s disease and related dementias; AUS, Australia; ED, emergency department; NDL, Netherlands; SD, standard deviation; pp, per person; UK, United Kingdom; USA, United States; USD, US dollars.

* Data available for the whole sample and not specific for dementia patients.

† Only including severe dementia because the % of decedents in that cohort > 50%.

‡ Total costs for people with dementia USD$7587.1 more than people without dementia.

? The paper is not clear regarding whether hospice care costs were considered.
in-hospital, out-of-hospital, community, and hospice care expenses paid by insurance companies. Three studies also included long-term care facility expenses. Two of the included studies only reported hospital costs, including in-hospital and emergency department expenses. Two studies measured informal costs, and only one of these included the informal costs as a proportion of the total costs of care.

Only two studies analyzed the proportion of the total cost of care explained by each cost component. In Goldfeld et al., hospice expenditures accounted for 45.7% of total costs, followed by in-patient (33.2%) and primary care provider expenditures (9.7%). In van der Plas et al., the highest cost component in the last year of life was nursing home expenses. Neither of these two studies included informal costs among the cost components analyzed.

### 3.4 Question 2: What are the costs of care for people with dementia approaching the end of life and how do these change closer to death?

#### 3.4.1 Studies including only direct costs

Seventeen studies included only direct costs. Three of them reported costs of care for an end-of-life period longer than the last 12 months, seven reported costs of care for the last 12 months, five for the last 6 months, and three for the last month of life. All 17 studies measured hospital and emergency department costs and all but two included out-patient expenses and community care. Thirteen of 17 studies included hospice expenses, and three included long-term care facility costs.

The derived monthly cost was higher in studies reporting costs in the last six or one month of life compared to those reporting costs for a longer end-of-life period (Figure 2). Derived monthly costs varied from $486 to $4428 USD for the final 18 to 36 months of life, $1787 to $2999 USD for the last 12 months of life, $3933 to $7717 USD for the last 6 months, and $4570 to $11921 USD for the last month of life.

#### 3.4.2 Studies including only informal costs

One study explored out-of-pocket expenditures among Medicare beneficiaries over the last 5 years of life. Monthly derived out-of-pocket expenses among people with AD and other dementias was $1335 USD on average and out-of-pocket payments for long-term care facilities accounted for 56% of the average family spending.

#### 3.4.3 Studies including direct and informal costs

Only one study considered both direct and informal costs in the total cost of care. In this study the estimated total cost of care in the last 5 years of life among individuals with dementia was $5599 USD per month.
3.5 | Question 3: Which factors are associated with the costs of care of people with dementia approaching the end of life?

Among the factors explaining the variance in the cost of care were sociodemographic characteristics, illness-related factors such as comorbidity or type of dementia diagnosis, and health-care service-related factors, such as the use of particular health-care services or places of care.

3.5.1 | Sociodemographic characteristics

Three studies explored the association between costs of dementia care and sociodemographic characteristics. Ornstein et al. examined differences in total Medicare expenditures among people with dementia from diagnosis to death by ethnicity. Non-Hispanic Black people had a 41% higher spending compared to non-Hispanic White people from time of diagnosis to death, but the difference was not significant after adjusting for confounding. In Kelley et al., total cost of care among people with dementia in the last 5 years of life were significantly higher for people older than 90 years (vs. 70–74 years old), Black (vs. White), women, married (vs. not married), and with higher than high school education (vs. less than high school). In Goldfeld et al., younger ages and not living in a special care dementia unit were independently associated with higher costs in the last 90 days of life. 

3.5.2 | Illness-related factors

Three studies explored the association among dementia severity, comorbidities, and cost of care at the end of life. In Kelley et al., people with dementia and a history of stroke, diabetes, heart disease, or psychological conditions had higher total costs in the last 5 years of life. In Zhu et al., total Medicare expenditures in people with severe dementia were approximately 10 times higher among those with more than three comorbidities versus no comorbidities. In Goldfeld et al., people with more severe cognitive impairment, chronic obstructive pulmonary disease, and a prior history of acute illness had higher total costs during the last 90 days of life.

Seven studies explored differences in health-care costs between individuals with and without dementia or other conditions. Two of them only included hospital costs in the analysis. In Daras et al. and Spilsbury and Rosenwax, hospital costs in the last 12 months of life were lower for patients with dementia than patients with non-dementia or cancer diagnosis, respectively. When only direct costs of care were considered, the costs of care for people with dementia were higher than for those without dementia in the last 6 months, 12 months, and nonsignificant in one study in the last 5 years of life. When both informal and direct costs were considered, the cost of care was higher for those with dementia than for those without dementia.

We found three studies exploring the cost of dementia care by proximity to death. In Pyenson et al., Medicare costs for people with AD and general dementia in the last year of life was three times higher than 8 years before death. In McCormick et al., costs tended to escalate in the last 3 months of life. In Goldfeld et al., total mean Medicare expenditures increased by 65% by periods of 3 months in the last year of life.

3.5.3 | Health-care service-related factors

Health-care service factors examined among studies included place of care and the use of hospice services.

One study analyzed differences in total Medicare costs by place of care. Patients with mild dementia living in the community had significantly lower costs in the last 6 months of life than those living in a nursing home. However, for people with severe dementia this relationship was reversed.

Five studies explored differences in costs for people with dementia who used hospice services, and found mixed results.

4 | DISCUSSION

This is the first systematic review to examine costs of care for people with dementia approaching the end of life. The review found that costs of care in people with dementia are high and tend to increase toward death. There was a large variation in costs of care across studies. Long-term care facility and informal care costs were among the highest cost components. Only two studies included informal costs of care and most studies were from the United States. Non-White ethnicity, female sex, married status, higher education level, more severe dementia, and higher number of comorbidities were associated with higher costs.

We found a dearth of information on informal costs. A systematic review exploring the financial impact of caring for people at the end of life shows that informal costs are substantial, and have an important impact on caregiver burden. In Kelley et al., out-of-pocket expenditures were a large proportion of the total cost of care in the last 5 years of life in decedents with dementia in the United States. Studies considering only Medicare expenditures show the costs of care at the end of life for people with dementia appeared to be lower than for other conditions, probably explained by lower rates of hospital admissions. In contrast, when including informal costs, the costs were substantially higher for individuals with dementia. Informal costs may be particularly relevant among people with dementia as their needs for formal and informal care are higher than people dying with other conditions. Therefore, studies that do not account for informal costs are likely to greatly underestimate total costs of care for people dying with dementia.

Our results show that monthly derived costs tend to increase in the last months of life, which is consistent with results from the three individual studies included that directly explored costs by proximity to
Implications for policy and research

4.1 Implications for policy and research

This review exposes the urgent need for studies describing and analyzing the cost of care at the end of life for people with dementia, in particular for costs outside the United States and informal care costs. This is not exempt of challenge. The lack of evidence in informal care costs may reflect important barriers in quantifying and analyzing informal care, such as difficulties measuring and valuing the time of caregiving.38,39

Understanding how cost components change toward the end of life is needed to plan effective use of health-care services.40 We found no studies directly exploring how the different cost components change by proximity to death. van der Plas et al. found long-term care facilities contribute proportionally more to the total cost of care in the last month of life compared to the last 360 days.33 More studies exploring how different cost components change by proximity to death, and the extent to which these costs are modifiable, are needed.

Translating evidence from studies on end-of-life care costs into improved clinical services relies on prospective identification of the period before death. This is particularly challenging for people with dementia due to communication difficulties and the pattern of slow decline.41,42 Nevertheless, ≈47% of older adults who died have a record indicating their general practitioner recognized they were close to death,43 and identification of palliative care needs in patients with dementia has been associated with lower chances of multiple hospital admissions in their last 90 days of life.44 Whether better recognition of the end of life among people with dementia could help reduce costs and which types of costs could be an important question to address.

Factors associated with costs of care for people with dementia at the end of life have been poorly studied. There is a need to explore the variability of total costs explained by factors such as ethnicity or level of deprivation to address potential inequities in health-care access at the end of life.

There is evidence suggesting informal costs are high for people with dementia approaching the end of life. Governments should make efforts to address the economic burden that dementia care places on patients’ families, especially considering the large number of people projected to be affected in the next 40 years. Higher social care expenditure has been associated with fewer hospital readmissions, reduced length of stay, and lower expenditure on secondary
health-care services. Informal care use has been shown to be endogenous and a substitute of formal care received at home by older people. However, this tends to disappear as the level of disability of the older person increases and it is not necessarily a substitute for care received in a long-term care facility or other health-care service. Future research needs to investigate the effects of hospital costs on economic burden on family and the reciprocal relationship between informal care and total care costs.

4.2 | Strengths and limitations

To our knowledge, this is the first systematic review on costs of care for individuals with dementia with a focus on the end of life. While other articles have analyzed the cost of care by dementia stage, the severity of the disease is not necessarily representative of a terminal stage, as only 25% of people who die with dementia are at the severe stage of the illness.

This study has limitations. The heterogeneity in end-of-life periods and cost components considered by studies limited the options for analysis. Only 20% of all titles and abstracts were double-screened, though 90% agreement between researchers was reached. Only one reviewer conducted the quality appraisal of studies, though scores were discussed with a second reviewer and no study was excluded based on the quality appraisal. The dearth of studies from countries other than the United States limits the generalizability of results. The approach we used to derive monthly costs assumes the total cost reported by studies is equally distributed across the end-of-life period considered, which is unlikely to be the case as the cost of care rises toward death.

5 | CONCLUSIONS

Published data on costs of dementia care for people approaching the end of life suggest direct costs increase before death, but more information on costs by proximity to death is needed. Informal costs of care represent a substantial component of total costs but have been rarely measured, and there is little information from countries outside the United States. Understanding informal costs and costs from outside the United States must be a priority given the fast-projected increase in the number of people who will die with dementia over the next 40 years.

FUNDING INFORMATION

J.L is funded by a Royal Marsden Partners Pan London Research Fellowship Award and the Programa Formacion de Capital Humano Avanzado, Doctorado Becas Chile, 2018 (folio 72190265). D.Y is supported by Cicely Saunders International. K.E.S. is funded by a National Institute of Health Research (NIHR) Clinician Scientist Fellowship (CS-2015-15-005). I.J.H. is an NIHR Senior Investigator Emeritus. I.J.H. is supported by the NIHR Applied Research Collaboration South London (NIHR ARC South London) at King’s College Hospital NHS Foundation Trust. I.J.H. leads the Palliative and End of Life Care theme of the NIHR ARC South London, and co-leads the national theme in this. The funders did not have any involvement in the study design, collection, analysis, and interpretation of the data. The views expressed are those of the authors and not necessarily those of the NHS, the NIHR, the Department of Health and Social Care or the funding charities.

CONFLICTS OF INTEREST

The authors declare that there are no conflicts of interest.

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SUPPORTING INFORMATION
Additional supporting information may be found online in the Supporting Information section at the end of the article.

How to cite this article: Leniz J, Yi D, Yorganci E, et al. Exploring costs, cost components, and associated factors among people with dementia approaching the end of life: A systematic review. Alzheimer’s Dement. 2021;7:e12198. https://doi.org/10.1002/trc.12198