

What factors are associated with informal carers' psychological morbidity during end-of-life home care? A systematic review and thematic synthesis of observational quantitative studies

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Abstract

What factors are associated with informal carers' psychological morbidity during end-of-life home care? A systematic review and thematic synthesis of observational quantitative studies

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Background: Family carers are central in supporting patients nearing end of life. As a consequence, they often suffer detrimental impacts on their own mental health. Understanding what factors may affect carers' mental health is important in developing strategies to maintain their psychological well-being during caregiving.

Aim: To conduct a systematic review and thematic evidence synthesis of factors related to carers' mental health during end-of-life caregiving.

Method: Searches of MEDLINE, CINAHL, PsychINFO, Social Sciences Citation Index, EMBASE, Cochrane Central Register of Controlled Trials and Database of Abstracts of Reviews of Effects 1 January 2009–24 November 2019. We included observational quantitative studies focusing on adult informal/family carers for adult patients at end of life cared for at home considering any factor related to carer mental health (anxiety, depression, distress and quality of life) pre-bereavement. Newcastle–Ottawa Quality Assessment Scale was used. Thematic analysis with box score presentation, and meta-analysis were done where data permitted.

Results: Findings from 63 included studies underpinned seven emergent themes.

1. **Patient condition** (31 studies): worse patient psychological symptoms and quality of life were generally associated with worse carer mental health. Patient depression was associated with higher depression in carers (standardised mean difference = 0.59, 95% confidence interval 0.32 to 0.87, $I^2 = 77\%$). Patients' other symptoms and functional impairment may relate to carer mental health, but findings were unclear.

2. **Impact of caring responsibilities** (14 studies): impact on carers' lives, task difficulty and general burden had clear associations with worse carer mental health.
3. **Relationships** (8 studies): family dynamics and the quality of the carer–patient relationship may be important for carer mental health and are worthy of further investigation.
4. **Finance** (6 studies): insufficient resources may relate to carers' mental health and warrant further study.
5. **Carers' psychological processes** (13 studies): self-efficacy and preparedness were related to better mental health. However, findings regarding coping strategies were mixed.
6. **Support** (18 studies): informal support given by family and friends may relate to better carer mental health, but evidence on formal support is limited. Having unmet needs was related to worse mental health, while satisfaction with care was related to better mental health.
7. **Contextual factors** (16 studies): older age was generally associated with better carer mental health and being female was associated with worse mental health.

Limitations: Studies were mainly cross-sectional (56) rather than longitudinal (7) which raises questions about the likely causal direction of relationships. One-third of studies had samples < 100, so many had limited statistical power to identify existing relationships.

Conclusions and future work: Future work must adopt a comprehensive approach to improving carers' mental health because factors relating to carer mental health cover a broad spectrum. The literature on this topic is diverse and difficult to summarise, and the field would benefit from a clearer direction of enquiry guided by explanatory models. Future research should (1) further investigate quality of relationships and finances; (2) better define factors under investigation; (3) establish, through quantitative causal analyses, why factors might relate to mental health; and (4) utilise longitudinal designs more to aid understanding of likely causal direction of associations.

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List of supplementary materials

Report Supplementary Material 1 Quantitative observational research evidence: in-depth list of factors affecting caregiver mental health overall

Report Supplementary Material 2 Quantitative observational research evidence: in-depth list of factors affecting different types of mental health outcome

Supplementary material can be found on the NIHR Journals Library report page (<https://doi.org/10.3310/HTJY8442>).

Supplementary material has been provided by the authors to support the report and any files provided at submission will have been seen by peer reviewers, but not extensively reviewed. Any supplementary material provided at a later stage in the process may not have been peer reviewed.

Glossary

Carer Family and informal carers/caregivers are used interchangeably throughout this review to refer to adult lay carers. The term 'lay carer' or 'carer' is defined according to the broad definition adopted by the National Institute for Health and Care Excellence [*Guidance on Cancer Services, Improving Supportive and Palliative Care for Adults with Cancer, The Manual*. NICE guideline; 2004:159. URL: www.nice.org.uk/guidance/csg4/resources/improving-supportive-and-palliative-care-foradults-with-cancer-pdf-773375005 (accessed 27 July 2022)]: 'Carers, who may or may not be family members, are lay people in a close supportive role who share in the illness experience of the patient and who undertake vital care work and emotion management', which relates to unpaid carers who might be a partner, family member, friend or neighbour of the person they are caring for.

List of abbreviations

ALS	amyotrophic lateral sclerosis	FCG	family caregiver
APIM	actor-partner interdependence model	GAD-7	Generalised Anxiety Disorder-7
BDI	Beck Depression Inventory	GBM	glioblastoma multiforme
BDI-II	Beck Depression Inventory, version 2	GDS-SF	Geriatric Depression Scale – Short Form
BCOS	Bakas Caregiving Outcomes Scale	GHQ-12	General Health Questionnaire-12
BSI	Brief Symptom Inventory	GI	gastrointestinal
CES	Carer Experience Scale	GSI	Global Severity Index
CHF	chronic heart failure	GU	genitourinary
CI	confidence interval	HADS	Hospital Anxiety and Depression Scale
CMA	comprehensive meta-analysis	HF	heart failure
CQOLC	Caregiver Quality of Life Index-Cancer	HRQOL	health-related quality of life
CQOL-R	Caregiver Quality of Life-Revised	MND	motor neurone disease
COPD	chronic obstructive pulmonary disease	MH	mental health
CRF	case report form	MLHFQ	Minnesota Living with Heart Failure Questionnaire
CSAI	Caregiver State Anxiety Inventory	MBCBS	Montgomery Borgatta Caregiver Burden Scale
CSNAT	Carer Support Needs Assessment Tool	NOS	Newcastle–Ottawa scale
DSM-IV	Diagnostic and Statistical Manual of Mental Disorders, fourth edition	NSCLC	non-small cell lung cancer
EOL	end-of-life	OECD	Organisation for Economic Co-operation and Development
FACQ PC	Family Appraisal of Caregiving Questionnaire for Palliative Care	OCBS	Oberst Caregiving Burden Score
FACT-G	Functional Assessment of Cancer Therapy – General	PMBT	primary malignant brain tumour
FACT-GP	Functional Assessment of Cancer Therapy – General Population	PHQ-9	Patient Health Questionnaire-9 items
		POMS	Profile of Mood States
		PPI	patient and public involvement

LIST OF ABBREVIATIONS

PSOC	perceived stage of cancer	SF-36	36-Item Short Form Health Survey
PSS	personal social services		
QoL	quality of life	SEIQoL-DW	Schedule for the Evaluation of the Individual Quality of Life-Direct Weighting
QA	quality assessment		
RAP	Review Advisory Panel		
SD	standard deviation	seTE	standard error treatment effect
SMD	standardised mean difference	TE	treatment effect
SPC	specialised palliative care	WHOQOL-BREF	World Health Organization quality of life-BREF
SF-12	12-Item Short Form Health Survey	WTHD	wish to hasten death

Plain language summary

Background

Family carers are central in supporting patients nearing end of life. However, their own mental health may often suffer as a result. It is important to understand what makes carers' mental health better or worse, to support them appropriately and help them stay in good health.

Aim

To synthesise what is known about what can affect carers' mental health during end-of-life caregiving.

Method

We identified research literature (1 January 2009–24 November 2019) that looked at factors that may make carers' mental health better or worse when supporting someone nearing end of life. We focused on adult carers of adult patients cared for at home. Researchers worked with the help of a carer Review Advisory Panel to group similar factors into themes. This report presents research that used numerical measurements (for instance, surveys) to investigate factors related to carers' mental health.

Results

Findings from 63 studies were grouped into seven themes: (1) How the patient was: worse patient mental health and quality of life related to worse carer mental health. (2) How much caregiving affected carers' lives: greater impact, burden and feeling tasks were difficult related to worse mental health. (3) Relationships: good relationships between family members and between carer and patient seemed important for carer mental health. (4) Finance: having insufficient resources may affect carers' mental health. (5) Carers' internal processes (carers' thoughts and feelings): feeling confident and prepared for caregiving related to better mental health. (6) Support: carers' mental health seemed related to support given by family and friends and to getting sufficient, satisfactory support from formal services. (7) Background factors: older carers seemed generally to have better mental health, and female carers worse mental health overall.

Conclusions

Factors that may affect carers' mental health are many and varied. We therefore need a broad strategy to help carers stay in good mental health during caregiving.

Background and introduction

Family and friends (hereafter 'carers') provide vital unpaid support for people at end of life (EOL), including physical and psychological support, co-ordinating care and monitoring. A national survey of carers of people with cancer in England found that they provided a median of 70 hours of care per week in the patient's final months of life.¹ Reviews have consistently shown carers to be a main factor in sustaining care at home at EOL,^{2,3} which is likely to reduce acute inpatient care costs and pressures on care home beds, and to be in accord with patient preferences.⁴ Carers' contributions therefore are likely to be of considerable benefit both to patient care and to health and social care services.

Our dependency on carers is likely to increase, given projected future demographic increases in people over 85 and those with life-limiting illness,⁵ dependency in the final years of life⁶ and number of deaths.⁷ Health and social care services are likely struggle to meet increasing future demands. The COVID-19 pandemic saw increases in deaths at home in England and Wales, between waves of the pandemic, while deaths from leading causes in inpatient health care decreased, indicating an increased reliance on carers to provide home care when healthcare systems are under strain.⁸

However, caregiving for patients at EOL has substantial negative impacts on carers' own health. The greatest and most consistent impacts are on carers' psychological health,⁹ where the greatest gains may be made. The prevalence of carer anxiety and depression during palliative care have been reported as 34–72%^{10–15} and 39–69%,^{14–17} respectively. Moreover, during the patient's final 3 months of life, the prevalence of clinically significant carer psychological morbidity was found to be 83% in a national census study of cancer deaths in England.⁹ An estimated 500,000 carers provide EOL care per annum in England.¹⁸ Given the numbers affected, these high levels of psychological morbidity arguably represent a sizeable public health problem with likely long-term effects. Carers' pre-bereavement psychological health is a main predictor of post-bereavement psychological health.^{19,20} If carers become unable to cope, this is likely to have negative impacts on the quality of patient care and increase the likelihood of inpatient hospital admissions.

Research shows that there is large individual variation in level of psychological morbidity from EOL caregiving. Understanding what predicts this variation provides important opportunities for identifying those at risk and pointers for intervention. An earlier, comprehensive review of the quantitative carer literature from 1998–2008 by Stajduhar *et al.*²⁰ identified potential predictors as: patient characteristics (including disease type and severity); carer sociodemographic characteristics (e.g. age, gender, ethnicity, socio economic status); carers' internal appraisals (e.g. of self-efficacy, preparation) and coping strategies; characteristics of the caregiving context and disruptions and restrictions to activities. The review also noted a lack of research into relational variables and available support, and of features of interaction with the healthcare system and providers. While valuable, this earlier review considered potential predictors only as one part of a wider review and only provided a narrative summary of findings.

A more systematic, detailed synthesis of the potential predictors is needed to give clearer pointers for action and illuminate two broad approaches to reduction in carer psychological morbidity. First, there are factors that cannot realistically be changed (e.g. age and gender), but whose effects can be mitigated through early, targeted support for those at higher risk. Second, there are factors that can be changed, for example self-efficacy, that can be subjected to more direct intervention to reduce likelihood of later psychological morbidity. What is non-modifiable or modifiable will partly depend on the stakeholder using the information – for instance, policy-makers may through legislation help modify work and financial factors that may put carers at risk, while practitioners may improve carers' self-efficacy through information tailored to their individual caregiving situation.

Two points can be made from the above. First, there are likely to be a range of potential predictors that require different strategies; therefore, we need a comprehensive rather than piecemeal understanding of what may predict carer psychological morbidity, to enable a co-ordinated and integrated approach to maximise impact. Second, any findings need to be communicated to different stakeholders in ways that are meaningful and relevant to them, so that they can use this information to help enact change within their own remits.

The review of quantitative, observational studies reported here is part of a larger project to synthesise the qualitative and quantitative literature on potential predictors of carer psychological morbidity and to communicate these to stakeholders with capacity to act on this information through formats and media that they find most useful. The project is novel in its comprehensiveness and detail, and in its focus on engaging with stakeholders.

The present review will help establish whether research indicates that there is a measurable, significant relationship between a potential predictor and carer psychological morbidity. However, it cannot directly establish likelihood of causality, nor can it give insight into carer experiences, or the reasons why a factor may cause distress. This will be covered in further papers on our reviews of the intervention and qualitative literature, respectively. The way the findings are presented here is informed by our patient and public involvement (PPI) work with a carer Review Advisory Panel (RAP), whose role was to assess the validity, relevance and accessibility of findings to carers. The collaboration with the carer RAP and a wider end-of-project stakeholder consultation will be reported in detail elsewhere.

This project focuses on factors associated with carer mental health during home care, as this is the setting where most care takes place, where the carer is most involved in a breadth and depth of care tasks, and where most patients want care to take place.

Aims and objectives

The overall aim of the project is to help reduce psychological morbidity among carers during EOL by

- conducting quantitative and qualitative evidence synthesis of factors that increase or decrease carer psychological morbidity during EOL caregiving
- integrating these syntheses into a coherent framework of factors
- translating the findings into accessible, bespoke information for key stakeholders to help them better target efforts to reduce carer psychological morbidity.

The objective of the current review is to conduct a comprehensive evidence synthesis of observational quantitative studies to identify factors associated with carer psychological morbidity during caregiving at home for adults at EOL, where morbidity is defined as anxiety, depression, distress or reduced quality of life (QoL).

Two additional project reports have been published, one on the qualitative synthesis and one on the intervention synthesis and metasynthesis, respectively:

Bayliss K, Shield T, Wearden A, Flynn J, Rowland C, Bee P, *et al.* Understanding what affects psychological morbidity in informal carers when providing care at home for patients at the end of life: a systematic qualitative evidence synthesis [published online ahead of print September 12 2023]. *Health Soc Care Deliv Res* 2023. <https://doi.org/10.3310/PYTR4127>

Grande G, Shield T, Bayliss K, Rowland C, Flynn J, Bee P, *et al.* Understanding the potential factors affecting carers' mental health during end-of-life home care: a meta synthesis of the research literature [published online ahead of print December 21 2022]. *Health Soc Care Deliv Res* 2022. <https://doi.org/10.3310/EKVL3541>

Remaining reports on stakeholder involvement and a synopsis of the project as a whole will be published in *Health and Social Care Delivery Research* and will also be available at <https://arc-gm.nihr.ac.uk/carers-project>.

Methods

We conducted a systematic search and evidence synthesis of the literature. To accommodate the wide-ranging literature, findings were synthesised thematically using box scores, supported by meta-analysis where data permitted. The review was registered with PROSPERO (PROSPERO 2019 CRD42019130279) and was carried out in accordance with the reporting guidelines: Preferred Reporting Items for Systematic Reviews and Meta-analyses (PRISMA) and Meta-analysis of Observational Studies in Epidemiology (MOOSE).

Search and selection strategy

Studies were identified through an electronic search of the literature from 2009 to 2019 in the following databases:

- MEDLINE (Ovid online)
- CINAHL Plus (EBSCO)
- PsycINFO (Ovid online)
- Social Sciences Citation Index (Institute for Scientific Information; Clarivate Analytics platform)
- EMBASE (Ovid)
- Cochrane Central Register of Controlled Trials (CENTRAL)
- Database of Abstracts of Reviews of Effects (DARE; University of York Centre for Reviews and Dissemination).

Following piloting, searches were completed in December 2019, using medical subject headings (MESH) terms relevant to caregivers supplemented with string carer terms, including variations on '*family care giver*' and '*informal carer*'. These were combined with MESH terms for '*palliative care*' supplemented by string terms '*end-of-life*' and '*end of life*'. The search strategy can be viewed in full in [Appendix 1](#).

Study inclusion was based on the following inclusion criteria:

Population: adult informal/family carers caring for adult patients at EOL (EOL was defined as the likelihood that the patient would die within a year). Focus was on home, community and outpatient settings. Only Organisation for Economic Co-operation and Development (OECD) countries were included, to ensure healthcare structures were comparable with the UK.

Intervention: factors associated with psychological morbidity in EOL carers; studies which reported on the relationship between factors and outcomes.

Outcome: mental health outcomes in carers focused on anxiety, depression, distress and QoL (whether self-reported or clinically defined) in home, community and outpatient settings. Psychological well-being was defined as the primary outcome for QoL, with general QoL used as a proxy measure where a psychological well-being QoL score was not available.

Study: observational studies.

Studies were excluded based on the following criteria:

1. factors or outcomes related to bereavement only
2. inpatient settings, given the focus on factors associated with carer mental health during home care
3. in languages other than English or Scandinavian, which would require further translation
4. systematic reviews.

Finally, the review was limited to published peer-reviewed empirical studies.

Ten per cent of both titles/abstracts and full texts were screened independently by two reviewers. Over 90% agreement was established in each case, indicating that no further modifications to the inclusion and exclusion criteria were required. Subsequent studies on title/abstract and full texts were screened by one reviewer.

The above represents some tightening and simplification of the search and selection process due to time pressures, and on advice from the project's external Study Steering Committee (and notification to NIHR HSDR), including limitation to most recent decade, fewer databases, OECD country and English or Scandinavian publications, omission of dissertations and grey literature and single screening once consistency was established. Similar simplification was applied to the data extraction below.

Data extraction and quality assessment process

Data extraction

A data extraction template to extract information on both factors and mental health outcomes was developed jointly by two reviewers and subsequently tested independently by the two reviewers on a 10% sample of included studies. Differences were resolved by discussion and the data extraction template subsequently clarified to mitigate for any further inconsistencies between reviewers. Data extraction was then carried out by one reviewer and a random sample of 10% of remaining studies checked by another. No discrepancies between reviewers were identified in the checking process.

Where a study reported findings for both the overall outcome measure of QoL and the mental health/emotional subdomain of QoL (psychological well-being), only findings related to the mental health/emotional subdomain of QoL were extracted, to reflect the focus on psychological morbidity.

Where a study reported findings for the overall domain of a factor as well as the individual subdomains of the factor (e.g. caregiver burden), findings were reported for the overall scale only to avoid 'over-representing' factors as much as possible (i.e. providing 'multiple counts' of the same factor). However, where only subdomain findings were reported by the study, these were extracted.

Findings relating to the relationship between individual mental health outcomes were not extracted, in keeping with the project aims to identify factors associated with carers' mental health.

Statistical information was only extracted for bivariate relationships to avoid potential collinearity. Where studies reported multivariate analysis only, a narrative summary of the findings was documented.

Quality assessment

An adapted version of the Newcastle–Ottawa Quality Assessment Scale for cohort and case–control studies²¹ was used to perform quality assessment of cohort/longitudinal studies and cross-sectional studies of included studies (see [Appendix 2](#)). This modified version was adapted from the Newcastle–Ottawa Scale (NOS) used in another study²² to appropriately assess the quality of cross-sectional studies.

Quality assessment was carried out independently by two reviewers on 10% of the studies. Over 90% agreement was achieved, so subsequent studies were quality assessed by one reviewer and a random sample of 10% of studies checked by another. No discrepancies between reviewers were identified in the checking process.

Thematic synthesis with PPI

Individual factors were synthesised thematically into sub-themes using box scores.²³ This was conducted in ways that were meaningful to the carer RAP in order for them to assess the relevance of findings. For

example: (1) renaming factors reported in studies in language that made sense to carers; (2) reporting findings from correlation studies so that they referred consistently to improved or worsened mental health to allow easier interpretation; and (3) thematic groupings of factors.

Each sub-theme was then synthesised further by mapping individual sub-themes under one of the overarching thematic groupings identified in the qualitative synthesis (see <https://arc-gm.nihr.ac.uk/carer-project->): patient condition, impact of caring responsibilities, relationships, finances, carer internal processes and support. These were informed by the carer RAP as useful ways of presenting the evidence.

Meta-analysis

The outcome data were converted to standardised mean difference (SMD) using comprehensive meta-analysis (CMA) software. Effect sizes were then pooled using DerSimonian-Laird random-effects model.²⁴ Results of each mental health outcome (i.e. anxiety, depression, distress or QoL) were presented in the forest plots with the SMD calculated using Hedges' *g* and then interpreted according to Cohen's criteria.²⁵ Where data from five or more studies were pooled in a meta-analysis, a random-effects model was performed. For pooled data of fewer than five studies, a fixed-effects model was calculated. Heterogeneity was quantified using the I^2 statistic with values 25%, 50% and 75% indicating low, moderate and high heterogeneity, respectively.²⁶ If more than 10 studies were included in a meta-analysis, funnel plots and Begg's and Egger's test were used to examine potential for publication bias.²⁷ All meta-analyses were conducted in R version 4.0.3 (R Foundation for Statistical Computing) using the 'meta' or 'metafor' packages.^{28,29}

The opportunity for meta-analysis was limited due to the wide range of factors and the range of mental health outcomes considered. There were therefore few instances where studies considered sufficiently similar factors and their relation to the same outcome to permit meta-analysis.

Results

Hits and paper selection

The PRISMA diagram details the study identification and selection process ([Figure 1](#)).

Sixty-three studies met the study inclusion criteria for observational studies. Characteristics of the 63 included studies are specified in [Table 1](#). Studies were excluded where a substantial proportion of the patient population were considered unlikely to be EOL, for example a study which reported metastases

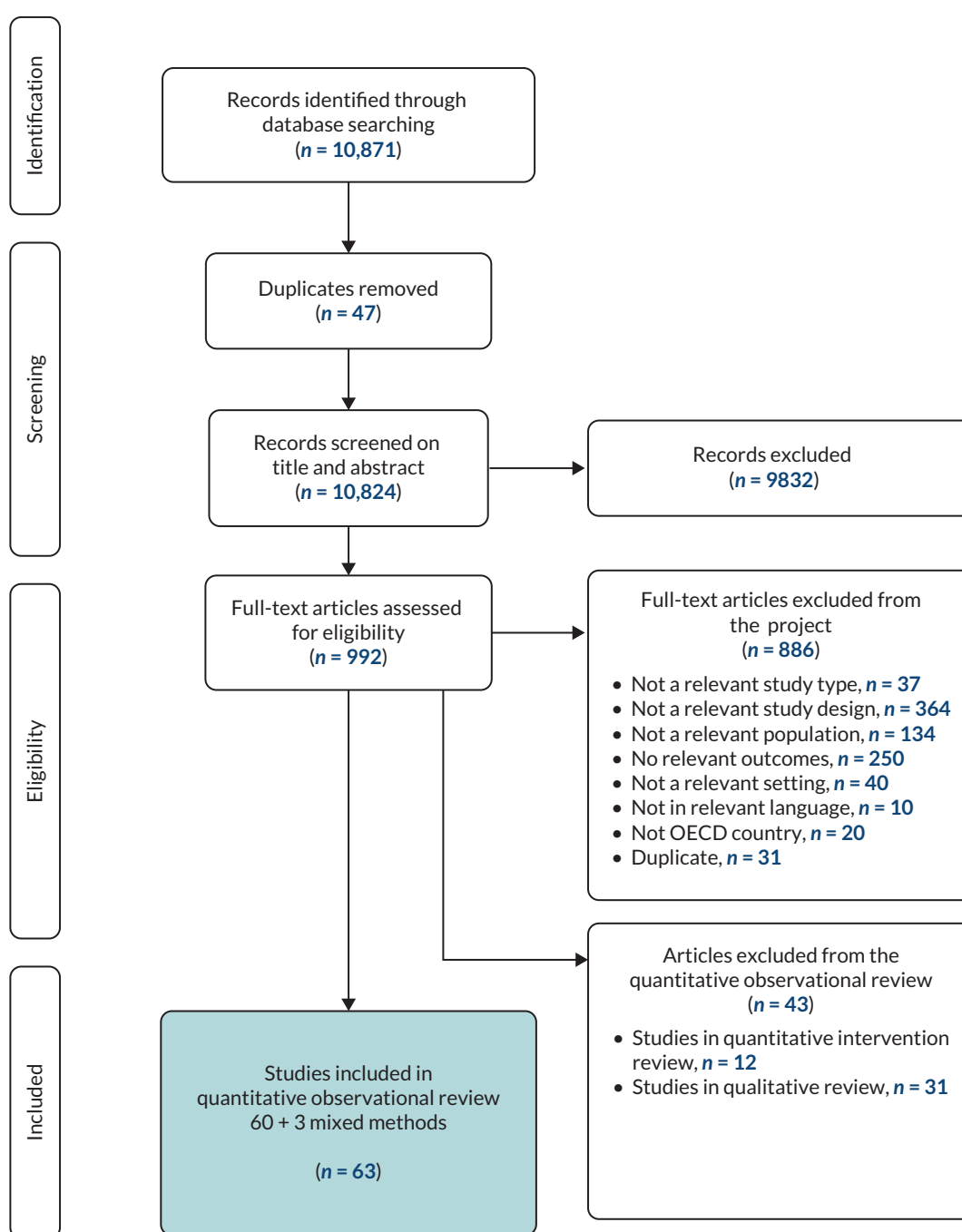


FIGURE 1 PRISMA diagram of study identification and selection.

TABLE 1 Observational review study summary table

Reference and country	Study aims	Study design and data collection; QA score	Participants (number; demographics; carer-patient relationships)	Patient condition	Factors investigated in bivariate analysis	Outcomes (anxiety, depression, distress, QoL)
Aoun <i>et al.</i> (2015) ³⁰ Australia	(1) Compare the profile and differences in well-being outcomes of family caregivers of people with brain cancer with those of people with all other cancers who participated in the CSNAT Intervention. (2) Assess the feasibility of the CSNAT Intervention to identify and address support needs of family caregivers of people with brain cancer in home-based palliative care.	Cross-sectional secondary analysis; 4	N = 500; Mean age, years = 60 73% female; 69% spouse/partner, 21% son/daughter	Mixed cancer: Lung (22.4%) Breast (9.6%) Colorectal (8.8%) Prostate (8%) Pancreas (7.2%) Primary brain cancer (6%)	Patient condition (primary brain cancer vs. other cancers)	QoL: SF12v2-MH
Areia <i>et al.</i> (2019) ¹⁵ Portugal	(1) Determine the prevalence of psychological morbidity with regard to psychological distress, depression, anxiety, somatisation and complicated anticipatory grief in family caregivers of people living with terminal cancer. (2) Describe which factors may be related to psychological distress, depression, anxiety, somatisation and complicated anticipatory grief.	Cross-sectional survey; 6	N = 112 Mean age, years (SD) = 44.5 (15.3) 82.1% female; 42.9% other (not spouse or child), 37.5% adult child	Mixed cancer: GI (24.5%) Respiratory (20.9%) Other solid tumours (20.9%) Central nervous system (11.8%) Breast (9.1%)	Unmet needs in caregiver	Anxiety: BSI anxiety subscale Depression: BSI depression subscale Distress: GSI
Bachner and Carmel (2009) ³¹ Israel	(1) Assess caregivers' perceived level of open communication about illness and death (= mortality communication) with their terminally ill relatives. (2) Examine the contribution of different caregiver characteristics and situational variables to explain open communication.	Cross-sectional survey; 3	N = 236 Mean age, years (SD) = 55.4 (13.7) 77.5% female; 47.9% son/daughter 44.9 % spouse	Cancer	Quality of patient-caregiver relationship (communication)	Depression: BDI-II (modified)

TABLE 1 Observational review study summary table (*continued*)

Reference and country	Study aims	Study design and data collection; QA score	Participants (number; demographics; carer–patient relationships)	Patient condition	Factors investigated in bivariate analysis	Outcomes (anxiety, depression, distress, QoL)
^a Bachner <i>et al.</i> (2009) ³² Israel	Compare response levels as well as the relative strength of association between mortality communication (candid discussion of the terminal illness and impending death between caregivers and their loved ones) and psychological distress among caregivers of terminal cancer patients within two distinct care contexts (i.e. home hospice vs. inpatient hospital settings).	Cross-sectional survey; 3	N = 126 Mean age, years (SD) = 56.61 (14.38) 79.2% female 48.5% spouse, 44.6% son/daughter	Cancer		Depression: BDI-II (modified)
Bachner <i>et al.</i> (2011) ³³ Israel	Compare the relative strength of association between mortality communication (candid discussion of the terminal illness and impending death between caregivers and their loved ones), fear of death, and psychological distress (depressive symptomatology, emotional exhaustion) among secular (non-religious) and religious Israeli Jewish caregivers of terminal cancer patients.	Cross-sectional survey; 2	N = 236; Mean age, years (SD) = 55.37 (13.69); 77.5% female; 47.9%, son/daughter 44.9% spouse	Cancer	Caregiver coping pattern (secular vs. religious)	Depression: BDI-II (modified)
Boele <i>et al.</i> (2013) ³⁴ The Netherlands	(1) Determine whether apart from tumour characteristics, HRQOL and neurological symptoms of the patient as perceived by the caregiver are related to the informal caregiver's HRQOL and feelings of mastery. (2) Investigate if a structured intervention consisting of psychoeducation and cognitive behavioural therapy leads to improvements in the mental component of HRQOL and mastery of caregivers.	Cross-sectional secondary analysis; 4	N = 56; Mean age, years = 50.66; 64.3% female; carer–patient relationship not reported	Cancer: 30.4% Grade 3 glioma 69.6% Grade 4 glioma	Patient condition Patient QoL Patient symptoms	QoL: SF-36 MH

continued

TABLE 1 Observational review study summary table (*continued*)

Reference and country	Study aims	Study design and data collection; QA score	Participants (number; demographics; carer-patient relationships)	Patient condition	Factors investigated in bivariate analysis	Outcomes (anxiety, depression, distress, QoL)
Burrige <i>et al.</i> (2009) ³⁵ Australia	Examine how carers' and patients' PSOC changes over the patients' final year in comparison with the perceptions of carer-patient dyads; whether carers' anxiety and depression scores are correlated with their PSOC, and whether these scores differ by gender.	Cohort Survey; 4	N = 57 Mean age, years (SD) = 57 (12.7); 76% female; 76% spouse	Cancer: Lung (30%) Digestive tract (26%) Other (26%) Breast (18%)	Patient stage of disease (PSOC)	Anxiety: HADS-A Depression: HADS-D
Burton <i>et al.</i> (2012) ³⁶ USA	Expand understanding of caregiver burden and psychosocial-spiritual outcomes among under-studied groups of caregivers – cancer, congestive HF and COPD caregivers – by including differences in outcomes by disease in a diverse population.	Cross-sectional survey; 7	N = 139; Mean age, years (SD) = 57 (14.88) 81.29% female 69.78% white; 56.83% spouse/partner, 43.17% other (child, friend or sibling)	Mixed: cancer COPD CHF % composition not given	Patient disease burden Patient disease severity Caregiver coping patterns Caregiver support Caregiver age Caregiver employment status Caregiver gender Caregiver marital status	Anxiety: POMS-anxiety Depression: CES-D 10
Buscemi <i>et al.</i> (2010) ³⁷ Spain	Analyse the possible relationship between the needs of primary caregivers of patients with terminal cancer and burden, stress and anxiety.	Cross-sectional survey; 2	N = 59 Mean age, years (SD) = 53.35 (15.66); 81.4% female; 57.6% spouse, 35.6% son/daughter	Mixed cancer: Lung (33.9%) Other (33.8%) Colon (13.6%) Breast (11.9%) Liver (6.8%)	Unmet needs in caregiver Caregiver burden (BCOS)	Anxiety: HADS-A Depression: HADS-D
Butow <i>et al.</i> (2014) ³⁸ Australia	(1) Describe the QoL of caregivers of women with ovarian cancer. (2) Describe the most frequent and severe unmet needs in caregivers. (3) Identify caregiver and patient predictors of caregiver QoL.	Cohort Survey; 8	N = 99; Mean age, years (SD) = 59 (13.2); 20% female; 78% husband/partner, 16% child	Ovarian cancer (100%)	Patient stage of disease Caregiver gender Relationship to patient Rural location	QoL: SF12v2 MH Distress: HADS (combined score)
Catt <i>et al.</i> (2012) ³⁹ UK	To evaluate and compare oncologist-led follow-up with a multidisciplinary group follow-up method from the perspective of patients and caregivers after patients' radical treatment for high-grade glioma.	Cohort survey; 6	N = 32; Mean age, years = 51; 56.25% female; 87.5% spouse/partner, 62.5% offspring	Single cancer: High-grade glioma	Caregiver education Caregiver employment status Caregiver gender Relationship to patient Patient treatment Caregiver lifestyle adjustments Caregiver workload	Distress: GHQ-12

TABLE 1 Observational review study summary table (*continued*)

Reference and country	Study aims	Study design and data collection; QA score	Participants (number; demographics; carer–patient relationships)	Patient condition	Factors investigated in bivariate analysis	Outcomes (anxiety, depression, distress, QoL)
Duimering <i>et al.</i> (2019) ⁴⁰ Canada	To assess carers of their patient population, evaluate their expressed caregiving burden and QoL, and determine baseline engagement with support services.	Cross-sectional survey; 6	N = 200; Mean age, years (SD) = 58.7 (14.0); 60.8% female; 60.6% spouse, 28.8% child	Mixed cancer: Lung (25%) Prostate (19.3%) Breast (18.8%) Colorectal (5.7%) Renal (4.5%) Bone (4.5%)	Caregiver gender Relationship to patient Caregiver employment status Caregiver socio economic status Rural location Caregiver lives with patient Patient disease burden Patient treatment Caregiver workload Caregiver support	QoL: CQOLC
^a Ellis <i>et al.</i> (2017) ⁴¹ USA	To examine the relationship between the number of co-existing health problems (patient comorbidities and caregiver chronic conditions) and QoL among patients with advanced cancer and their caregivers, and assess the mediating and moderating role of meaning-based coping on that relationship.	Cross-sectional secondary analysis; 9	N = 484; Mean age, years (SD) = 56.5 (13.4) 56.8% female 79.6% white; 70% spouse, 15.3% son/daughter	Mixed cancer: Breast (32.4%) Lung (29.1%) Colorectal (25.4%) Prostate (13.0 %)		QoL: FACT-G (version 4)
Exline <i>et al.</i> (2012) ⁴² USA	To examine the relevance of forgiveness to carers, and relation to unresolved offences and depression.	Cross-sectional survey; 3	N = 142; Mean age, years (SD) = 60.3 (13.8) 74% female 82% white; 44% child, 42% spouse	Mixed: Cancer (43%) Dementia/Alzheimer's (32%) Heart disease (23%) Lung disease (18%)	Quality of patient–caregiver relationship	Depression: CES-D
Fasse <i>et al.</i> (2015) ⁴³ France	(1) Describe depressive symptoms and depression among spouses who care for palliative cancer patients. (2) Identify the most important factors explaining depression among these individuals.	Cross-sectional survey; 9	N = 60 (all spouses); Mean age, years (SD) = 62.39 (12.99) 36.7% female	Mixed cancer: Breast (36.6%) Lung (16.7%) Cervix (10%) Other (36.6%)	Quality of patient–caregiver relationship Caregiver coping patterns Patient disease severity Caregiver gender	Depression: BDI-short form

continued

TABLE 1 Observational review study summary table (continued)

Reference and country	Study aims	Study design and data collection; QA score	Participants (number; demographics; carer-patient relationships)	Patient condition	Factors investigated in bivariate analysis	Outcomes (anxiety, depression, distress, QoL)
Flechl <i>et al.</i> (2013) ⁴⁴ Austria	To investigate the experiences of 52 caregivers of deceased GBM patients treated in Austria.	Cross-sectional survey; 1	N = 52; Mean age, years = 60 67% female; 88% partner	Glioblastoma (100%)	Caregiver finances Caregiver age Patient age Duration of care	QoL: measure from researchers' own questionnaire
^a Franchini <i>et al.</i> (2019) ⁴⁵ Italy	To investigate impact of possible predictors of carers' QoL.	Cross-sectional survey; 6	N = 570; Mean age, years (SD) = 58.8 (13.9) 77.4% female 46.1% partner, 38.4% offspring	Mixed cancer: GI (33.5%) Thoracic (16.1%) Genitourinary (16.3%) Breast (10.5%)		QoL: CQOLC
^a Francis <i>et al.</i> (2011) ⁴⁶ USA	(1) Investigate how relationships between advanced cancer caregivers and others – specifically family members and the patient's health-care providers – are associated with the emotional distress or well-being experienced by caregivers during the early treatment phase of cancer care. (2) Examine moderating effects by race (African American and white) in the association between these relationships and emotional distress.	Cross-sectional secondary analysis; 7	N = 397; Median age, years = 51 78.3% female 68.3% white; 100% family members (no further breakdown)	Mixed cancer: Stage IV (or stage III lung, pancreatic or liver cancer) % composition not reported		Anxiety: POMS – Tension/anxiety Depression: POMS – Depression-dejection subscale
Götze <i>et al.</i> (2014) ¹² Germany	(1) Examine the level of psychological distress and the QoL of palliative cancer patients and their family caregivers. (2) Investigate the impact of socio-demographic and care-related factors on psychological distress of palliative patients and their family caregivers.	Cross-sectional survey; 8	N = 106; Mean age, years (SD) = 64.1 (11.1) 67.9% female; 75% partner, 16% son/daughter	Mixed cancer: Prostate cancer (17.0%) Lung cancer (14.2%) Pancreas (13.2%) Colon (11.3%)	Patient symptoms Quality of patient-caregiver relationship	Anxiety: HADS – A Depression: HADS – D Distress: HADS (combined score)

TABLE 1 Observational review study summary table (*continued*)

Reference and country	Study aims	Study design and data collection; QA score	Participants (number; demographics; carer–patient relationships)	Patient condition	Factors investigated in bivariate analysis	Outcomes (anxiety, depression, distress, QoL)
Govina <i>et al.</i> (2019) ⁴⁷ Greece	To determine the factors associated with the anxiety and depression of family members caring for patients undergoing palliative radiotherapy.	Cross-sectional survey; 7	N = 100; Mean age, years (SD) = 53.3 (12.6) 76% female; 59% spouse, 27% child	Mixed cancer: Lung (48%) Breast (22%) Urogenital (20%)	Patient gender Caregiver gender Patient condition Patient treatment Patient lives with caregiver Previous experience of informal caregiving Caregiver mode of transport Caregiver marital status Patient educational level Caregiver educational level Additional caring responsibilities Relationship to the patient Caregiver employment status Caregiver age Caregiver burden Patient medical history	Anxiety: HADS-A (Greek) Depression: HADS-D (Greek)
Grant <i>et al.</i> (2013) ⁴⁸ USA	Describe burden, skills preparedness, and QoL for caregivers of patients with NSCLC and describe how the findings informed the development of a caregiver palliative care intervention that aims to reduce caregiver burden, improve caregiving skills and promote self-care.	Cohort survey; 8	N = 163; Mean age, years = 57.23 64% female 71% white; 68% spouse/partner, 16% daughter	NSCLC (100%)	Patient stage of disease	Distress: Psychological distress thermometer QoL: City of Hope QoL Scale-Family version – psychological well-being domain

continued

TABLE 1 Observational review study summary table (*continued*)

Reference and country	Study aims	Study design and data collection; QA score	Participants (number; demographics; carer-patient relationships)	Patient condition	Factors investigated in bivariate analysis	Outcomes (anxiety, depression, distress, QoL)
Hampton and Newcomb (2018) ⁴⁹ USA	To determine the relationship between self-efficacy and perceived stress in adult carers providing EOL care.	Cross-sectional survey; 4	N = 78; Mean age, years (SD) = 61.21 (13.91) 74.4% female 74% white; carer-patient relationship not reported	Mixed: Cancer (37.2%) Heart problems (17.9%) Dementia (11.5%)	Self-efficacy	Anxiety: PSS
Hannon <i>et al.</i> (2013) ⁵⁰ Canada	(1) Assess satisfaction with care in patients with advanced cancer and their caregivers in an ambulatory oncology setting and to examine whether there were important differences between patient and caregiver reports. (2) Identify factors associated with patient and caregiver satisfaction with care.	Cross-sectional secondary analysis; 7	N = 191; Mean age, years (SD) = 56.1 (12.1) 66% female; 84.3% spouse/partner, 11.5% son/daughter	Mixed cancer: GI (37.7%) Genitourinary (17.8%) Breast (17.3%) Lung (16.2%) Gynaecological (11%)	Quality of care	QoL: CQOLC
Henriksson and Arestedt (2013) ⁵¹ Sweden	Explore factors associated with preparedness and to further investigate whether preparedness is associated with caregiver outcomes.	Cross-sectional secondary analysis; 8	N = 125; Mean age, years (SD) = 57.7 (15.8) 60.8% female; 58.4% spouse, 22.4% adult children	Mixed: Cancer (88.8%) Other (11.2%)	Preparedness for caregiving	Anxiety: HADS-A Depression: HADS-D
Hoefman <i>et al.</i> (2015) ⁵² Australia	To study construct validation of the CES and the carer QoL and to investigate the effect of caregiving on caregivers in EOL care.	Cross-sectional Survey; 6	N = 97; Mean age, years (SD) = 62.3 (11.9) 71% female 98% white; 59% partner, 29% child	Not reported	Time for respite Caregiver support Positive aspects of caregiving Control over care situation Quality of patient-caregiver relationship Caregiver burden Additional caring responsibilities Caregiver finances	QoL: CarerQoL-7D dimension-MH question

TABLE 1 Observational review study summary table (*continued*)

Reference and country	Study aims	Study design and data collection; QA score	Participants (number; demographics; carer–patient relationships)	Patient condition	Factors investigated in bivariate analysis	Outcomes (anxiety, depression, distress, QoL)
Huang and McMillan (2019) ⁵³ USA	To apply the Actor–Partner Interdependence Model (APIM) to elucidate importance of mutual effects within dyads with advanced cancer examining contribution of depression on their individual (own) QoL and their carers' QoL.	Cross-sectional secondary analysis; 6	N = 660; Mean age, years (SD) = 65.49 (13.81) 74% female 96% white; 57% spouse, 11% daughter	Mixed cancer % composition not reported	Patient symptoms	Depression: CES-D QoL: SF-12 MH
Hudson <i>et al.</i> (2011) ⁵⁴ Australia	To examine the psychological and social profile of family caregivers on commencement of receiving palliative care services.	Cross-sectional secondary analysis; 6	N = 301; Mean age, years (SD) = 56.52 (13.89) 73.1% female; 47.8% spouse, 37.2% adult children	Mixed cancer: GI tract cancer (20.3%) Lung cancer (13.6%) Head and neck cancer (10.6%) Urogenital cancer (10.6%)	Impact on caregiver's schedule Self-esteem Optimism	Anxiety: HADS-A Depression: HADS-D
Ito and Tadaka (2017) ⁵⁵ Japan	To identify the associated factors of QoL among family carers of patients with terminal cancer at home in Japan.	Cross-sectional survey; 8	N = 74; Mean age, years (SD) = 63.6 (12.2) 79.7% female; 35.1% wife, 28.4% daughter	Mixed cancer: Lung (29.7%) Colon (18.3%) Liver (14.9%) Brain (12.2%) Prostate (12.2%) Stomach (8.1%) Pancreas (8.1%)	Patient age Patient gender Patient condition Patient symptoms Patient treatment Patient disease burden Caregiver support Duration of care Caregiver age Caregiver gender Relationship to patient Family dynamics Caregiver finances Caregiver employment status Caregiver health status Caregiver sleeping hours Self-efficacy Caregiver support Quality of care Accessible information	QoL: CQOLC– Japanese

continued

TABLE 1 Observational review study summary table (continued)

Reference and country	Study aims	Study design and data collection; QA score	Participants (number; demographics; carer-patient relationships)	Patient condition	Factors investigated in bivariate analysis	Outcomes (anxiety, depression, distress, QoL)
Jacobs <i>et al.</i> (2017) ⁵⁶ USA	Understand the prevalence of psychological symptoms (depression and anxiety) in patients and carers and to determine whether their distress is interdependent.	Cross-sectional survey; 10	N = 275; Mean age, years (SD) = 57.37 (13.61) 69.1% female 93% white; 66.2% spouse/partner, 18.4% son/daughter	Mixed cancer: Lung (54.6%) Non-colorectal GI (45.4%)	Patient symptoms	Anxiety: HADS-A Depression: HADS-D
Janda <i>et al.</i> (2017) ⁵⁷ Australia	To address research gap in quantifying association between patients' and their immediate carers' well-being	Cross-sectional survey; 6	N = 84 Age: 38% ≤ 60 years 37% 61–70 years 21% 70+ years 73% female; 81% spouse/partner, 12% son/daughter	Pancreatic cancer (100%)	Patient QoL Relationship to patient Caregiver age Caregiver gender Caregiver education Caregiver support Patient symptoms	Anxiety: HADS-A Depression: HADS-D QoL: FACT-GP
Janssen <i>et al.</i> (2012) ⁵⁸ The Netherlands	To assess caregiver burden and positive aspects of caregiving in family caregivers of patients with advanced COPD, CHF or CRF	Cross-sectional survey; 8	N = 159; Mean age, years (SD) for each condition group COPD = 62.9 (11.5), CHF = 67.3 (11.5), CRF = 59.1 (15.2) 73% female; 83% spouse, 12% child	Mixed non-cancer: COPD (45.9%) CHF (28.3%) CRF (25.8%)	Patient condition (COPD vs. CHF vs. CRF)	Distress: FACQ PC
^a Kapari <i>et al.</i> (2010) ⁵⁹ UK	To identify the risk factors for poor caregiving and bereavement outcomes by assessing both patients and caregivers on a range of measures	Cohort Survey; 9	N = 100; Mean age years = 65.3 75% female 89% white British; 85% spouse/partner	Mixed: Cancer 96% (lung 22% prostate 12% breast, ovarian and colon 5% bowel 5% bladder 2% other cancer 40%) MND (2%) COPD (1%) Liver failure (1%)		Distress: CIS-R

TABLE 1 Observational review study summary table (*continued*)

Reference and country	Study aims	Study design and data collection; QA score	Participants (number; demographics; carer–patient relationships)	Patient condition	Factors investigated in bivariate analysis	Outcomes (anxiety, depression, distress, QoL)
^a Kenny <i>et al.</i> (2010) ⁶⁰ Australia	To investigate associations between health and a range of caregiving context variables	Cross-sectional survey; 5	N = 178; Mean age, years (SD) = 61.7 (13.5) 71% female; 59% spouse, 29% child/grandchild	Mixed: Cancer 89% (main categories: colorectal 15% lung 14% prostate 13%) Non-cancer 11% (main categories: cardiac failure 2.3% chronic airway limitation 1.7% pulmonary fibrosis 1.7%)		QoL: SF-36
Kershaw <i>et al.</i> (2015) ⁶¹ USA	To investigate actor and partner effects of advanced cancer patients' and their family caregivers' mental health, physical health and self-efficacy over time, and to investigate the effects of patients' and caregivers' self-efficacy on their own and the other dyad members' mental health and physical health over time	Cohort secondary analysis; 11	N = 484; Mean age, years (SD) = 56.7 (12.6) 57% female 83% white; 74% spouse, 19% relative	Mixed cancer: Breast cancer (37%) Lung (24%) Colorectal (23%) Prostate (16%)	Caregiver age Caregiver gender Relationship to patient Patient disease burden Patient condition	Distress: FACT-G emotional well-being
Kobayakawa <i>et al.</i> (2017) ⁶² Japan	To determine the prevalence of delirium and suicidal ideation among patients with cancer and determine whether these and other factors influence caregivers' psychological distress.	Cross-sectional survey; 6	N = 532; Mean age, years (SD) = 61.8 (12.1) 74% female; 53% spouse	Mixed cancer: lung (21%) stomach, oesophagus (17%) colon, rectum (12%) liver, bile duct and pancreas (23%) breast (5%) prostate, kidney and bladder (8%)	Caregiver gender Relationship to patient Caregiver educational level Caregiver finances Caregiver health status Caregiver support Patient symptoms Health professionals' understanding of patient needs Control over care situation Patient treatment Acceptance of patient condition	Depression: Single, self-created question
continued						

TABLE 1 Observational review study summary table (continued)

Reference and country	Study aims	Study design and data collection; QA score	Participants (number; demographics; carer-patient relationships)	Patient condition	Factors investigated in bivariate analysis	Outcomes (anxiety, depression, distress, QoL)
Loggers and Prigerson (2014) ⁶³ USA	Authors interested in whether, and how, the EOL experiences of adult patients with rare cancers differed from that of individuals with common cancers.	Cohort interviews; 7	N = 618; Age and gender not reported; Spouse and adult child % composition not reported	Common cancers N = 423: lung (35.2%) colorectal (18.2%) breast (18.0%) pancreatic (11.6%) Rare cancers N = 195: gastroesophageal (19.0%) ovarian and cervical (15.4%) hepatocellular, biliary and gallbladder (13.3%) head and neck (11.3%) sarcoma and GI stromal tumour (10.8%) leukaemia, multiple myeloma and Hodgkin lymphoma (10.3%) central nervous system (7.7%) Other (12.3%)	Patient condition	QoL: SF-36 Distress: SCID
Malik <i>et al.</i> (2013) ⁶⁴ UK	(1) Compare experiences of caring between HF and lung cancer caregivers looking after breathless patients. (2) Determine which factors are associated with caregiver burden and positive rewards to caring in those looking after breathless patients.	Cross-sectional survey; 4	N = 101; Mean age, years (SD) for each condition group HF = 65.8 (12.7) lung cancer = 59.9 (12.8) 78% female; 72% spouse/partner, 20% child	HF (50.5%) lung cancer (49.5%)	Patient condition Caregiver burden	Anxiety: HADS-A Depression: HADS-D QoL: SF-36 – MH
McIlpatrick <i>et al.</i> (2018) ⁶⁵ UK and Ireland	To identify modifiable psychosocial factors associated with caregiver burden and to evaluate the support needs of caregivers when caring for people living with advanced HF at the end of life.	Cross-sectional survey and semi-structured interviews; 5	N = 84; Mean age, years (SD) = 63.9 (14.3) 80% female; 52% spouse/partner, 22% son/daughter	HF (100%)	Patient symptoms Patient QoL Preparedness for caregiving Caregiver age	Anxiety: GAD-7 Depression: PHQ-9 QoL: MLHFQ

TABLE 1 Observational review study summary table (*continued*)

Reference and country	Study aims	Study design and data collection; QA score	Participants (number; demographics; carer–patient relationships)	Patient condition	Factors investigated in bivariate analysis	Outcomes (anxiety, depression, distress, QoL)
Mollerberg <i>et al.</i> (2019) ⁶⁶ Sweden	Determine whether family's sense of coherence was associated with hope, anxiety and symptoms of depression in persons with cancer in the palliative phase and their family members. ['Sense of coherence' consists of comprehensibility (ability to understand the situations clearly), manageability (belief that one has access to sufficient resources to manage challenging situations) and meaningfulness (belief that all challenges are worthy of engagement).]	Cross-sectional survey; 7	N = 165; Mean age, years (SD) = 62.1 (13.6) 64.8% female 67.9% spouse/partner	Mixed cancer: Breast (16.2%) Colon (15.1%) Prostate (10.6%) Kidney (10.6%) Other (47.5%)	Family dynamics	Anxiety: HADS-A Depression: HADS-D
Nielsen <i>et al.</i> (2017) ⁶⁷ Denmark	To investigate pre-loss grief symptoms and the associations with situational, intrapersonal and interpersonal factors in family caregivers of EOL cancer patients.	Cross-sectional survey; 7	N = 2865; Mean age, years = 60.5 69% female; 63.6% spouse/partner 29% son/daughter	Mixed cancer: Lung (25.1%) Colorectal (13.1%) Breast (7.0%) Prostate (7.5%) Haematological (3.7%) Other (43.6%)	Pre-loss grief	Depression: BDI
^a Nipp <i>et al.</i> (2016) ⁶⁸ USA	To describe rates of depression and anxiety symptoms in family carers of patients with incurable cancer and identify factors associated with family carer psychological distress.	Cross-sectional survey; 8	N = 275; Mean age, years (SD) = 57.4 (13.6) 69.1% female 93.1% white; 66.2% spouse, 18.5% son/daughter	Mixed cancer: Lung (54.2%) Non-colorectal GI (45.8%)		Anxiety: HADS-A Depression: HADS-D
Nissen <i>et al.</i> (2016) ⁶⁹ USA	To identify family-type clusters in an American sample of carers of terminally ill cancer patients and to examine the relationship between these clusters and carer QoL, social support and carer burden.	Cross-sectional secondary analysis; 5	N = 598; Mean age, years (SD) = 52.89 (14.05) 72% female 82% white; 58% spouse	Mixed cancer: % composition not reported	Family dynamics	QoL: SF-36-MH
continued						

TABLE 1 Observational review study summary table (continued)

Reference and country	Study aims	Study design and data collection; QA score	Participants (number; demographics; carer-patient relationships)	Patient condition	Factors investigated in bivariate analysis	Outcomes (anxiety, depression, distress, QoL)
O'Hara <i>et al.</i> (2010) ⁷⁰ USA	Not directly stated, but paper considers whether patient intervention affected carer outcomes, and whether patient measures affected carer outcomes (univariate correlations for latter).	Cohort survey; 7	N = 198; Mean age, years (SD) for each group Intervention = 59.9 (13.0) Control = 58.0 (11.9) 77% female; 96% white; 71% spouse/partner	Mixed cancer: GI (42.4%) Lung (36.4%) Genitourinary (13.1%) Breast (8.1%)	Patient QoL Patient symptoms Quality of care	Distress: MBCBS emotional subscale
Owensworth <i>et al.</i> (2010) ⁷¹ Australia	To investigate the association between functional impairments of individuals with cancer and caregiver psychological well-being, and examine the moderating effect of social support.	Cross-sectional survey; 7	N = 29 Mean age, years (SD) = 60.1 (11.7) 71.4% female; 88.8% spouse/partner	Brain tumour (100%) benign (stage 1 or 2 tumour) (52%) malignant (stage 3 or 4) (48%)	Patient disease burden	QoL: WHOQOL-BREF- psychological domain
^a Parker Oliver <i>et al.</i> (2017) ⁷² USA	To explore potential variables affecting depression and anxiety in informal hospice caregivers.	Cross-sectional secondary analysis; 6	N = 395 Mean age, years (SD) = 60.6 (12.6) 81.52% female; 49.1% adult child, 30.4% spouse	Mixed: Cancer (37.22%) Dementia (30.13%) No additional information on the remainder of the sample given		Anxiety: GAD-7 Depression: PHQ-9
Perez-Ordóñez <i>et al.</i> (2016) ⁷³ Spain	To identify the relationship between coping and anxiety in primary family caregivers of palliative cancer patients treated in a pain and palliative care unit.	Cross-sectional interviews; 6	N = 50 Mean age, years (SD) = 55 (13.9) 94% female; 52% daughter, 28% spouse	Mixed cancer: Others (46%) Lung (14%) Prostate (12%) Bladder (12%)	Caregiving coping patterns Patient disease burden Caregiver burden	Anxiety: Anxiety subscale of Goldberg Scale
^a Reblin <i>et al.</i> (2016) ⁷⁴ USA	To describe relationship quality categories among EOL caregivers and to test the effects of relationship quality categories on caregiver burden and distress within a stress process model.	Cross-sectional survey; 7	N = 131 Mean age, years (SD) = 65.3 (10.74) 65% female 97% white; 100% spouse	Cancer % composition not reported		Anxiety: HADS-A Depression: GDS-SF

TABLE 1 Observational review study summary table (*continued*)

Reference and country	Study aims	Study design and data collection; QA score	Participants (number; demographics; carer–patient relationships)	Patient condition	Factors investigated in bivariate analysis	Outcomes (anxiety, depression, distress, QoL)
Rivera <i>et al.</i> (2010) ⁷⁵ USA	To examine predictors of depression symptoms in caregivers of hospice cancer patients.	Cross-sectional secondary analysis; 7	N = 578 Mean age, years (SD) = 64.95 (14.01) 73.7% female 95.8% white; 42.4% wife, 25.2% other	Mixed cancer: Lung/mesothelioma (36%) Pancreas (8.7%) Colorectal (6.6%)	Patient psychological symptoms Patient QoL Caregiver gender Relationship to patient Caregiver ethnicity Caregiver support Patient condition Patient disease burden Caregiver age Caregiver health status	Depression: CES-D 10
Seekatz <i>et al.</i> (2017) ⁷⁶ Germany	To determine screening-based symptom burden and supportive needs of patients and caregivers with regard to the use of specialised palliative care (SPC).	Cohort survey; 4	N = 46 Mean age, years (SD) = 53.3 (14.1) 56.5% female; 57% spouse/partner, 22% child	Mixed cancer: Glioblastoma (68.4%) Brain metastases (31.6%)	Patient treatment	Distress: Hornheider Questionnaire (adapted)
^a Shaffer <i>et al.</i> (2017) ⁷⁷ USA	To examine correlates of mental and physical health among caregivers of patients with newly diagnosed, incurable lung or non-colorectal GI cancer.	Cross-sectional secondary analysis; 9	N = 275 Mean age, years (SD) = 53.37 (13.61) 69.1% female; 66.4% spouse	Mixed cancer: Lung (54.2%) Non-colorectal GI cancer (45.8%)		QoL: SF-36 mental health
Siminoff <i>et al.</i> (2010) ⁷⁸ USA	To investigate depressive symptomatology in stage III or IV lung cancer patients and their identified caregiver.	Cross-sectional secondary analysis; 7	N = 190 Mean age, years (SD) = 55 (13.4) 75% female; 54.7% spouse, 18.9% child	NSCLC (100%)	Patient psychological symptoms Family dynamics	Depression: CES-D -20
Stutzki <i>et al.</i> (2014) ⁷⁹ Germany	To determine the prevalence and stability of wish to hasten death (WTHD) and EOL attitudes in ALS patients, identify predictive factors and explore communication about WTHD.	Cohort survey; 8	N = 35 Mean age, years (SD) = 56.4 (12.7) 61% female; 79% partner, 14.5% son/daughter	ALS (100%)	Patient disease severity	QoL: Numerical Ratings Scale–individual QoL

continued

TABLE 1 Observational review study summary table (continued)

Reference and country	Study aims	Study design and data collection; QA score	Participants (number; demographics; carer-patient relationships)	Patient condition	Factors investigated in bivariate analysis	Outcomes (anxiety, depression, distress, QoL)
Thielemann and Conner (2009) ⁸⁰ USA	To examine the role of social support as a mediating factor between caregiver demands and caregiver depression in spousal caregivers of patients with advanced lung cancer.	Cross-sectional secondary analysis; 8	N = 164 Mean age, years (SD) = 61.9 (10.8) 60.4% female 98.2% white; 100% spouse	Lung cancer (100%)	Caregiver age Caregiver gender Caregiver ethnicity Caregiver educational level Length of patient-caregiver relationship Duration of care Caregiver burden Caregiver support	Depression: CES-D
^a Trevino <i>et al.</i> (2019) ⁸¹ USA	To conduct secondary exploratory analyses of the relationship between individual and dyadic estimations of the patient's life expectancy and patient and caregiver QoL.	Cross-sectional secondary analysis; 8	N = 162 Age, years 65+ (42.6%) less than 65 (57.4%) 66% female 89.5% white; 64.2% spouse/partner	Mixed cancer: Aggressive cancer (50%) – this includes lung, GI cancers (except colon) and GU cancers (except prostate). Less aggressive (50%) – this includes breast, prostate and colon cancers		Depression: DSM-IV (SCID) QoL: SF12v2 -emotional
^a Valeberg <i>et al.</i> (2013) ⁸² Norway	To examine the level of symptom burden in a sample of cancer patients in a curative and palliative phase. In addition to determine (a) whether the patients' symptom burden and patients' demographic variables, and (b) the caregivers' demographic variables impact on the caregivers' QoL and mental health.	Cross-sectional secondary analysis; 6	N = 159 Mean age, years (SD) = 57.0 (12.3) 39% female; 89% spouse, 6% friend	Mixed cancer: Breast (46%) Prostate (18%) Other (18%) Colorectal (13%) Gynaecologic (5%)		Anxiety: HADS-A Depression: HADS-D QoL: SF36-MH

TABLE 1 Observational review study summary table (continued)

Reference and country	Study aims	Study design and data collection; QA score	Participants (number; demographics; carer-patient relationships)	Patient condition	Factors investigated in bivariate analysis	Outcomes (anxiety, depression, distress, QoL)
Wadhwa <i>et al.</i> (2013) ⁸³ Canada	To evaluate the QoL and mental health of caregivers of patients with advanced cancer who are receiving ambulatory oncology care and associations with patient, caregiver and care-related characteristics.	Cross-sectional secondary analysis; 7	N = 191 Median age, years (range) = 57 (22–83) 64.9% female 83.2% white; 83.8% spouse/partner, 5.2% son/daughter	Mixed cancer: GI (37.7%) Genitourinary (17.8%) Breast (17.3%) Lung (16.2%) Gynaecology (11%)	Caregiver gender Caregiver age Caregiver employment status Relationship to patient Caregiver burden Impact on work Patient treatment Patient symptoms Patient QoL Caregiver health status Patient lives with caregiver Caregiver ethnicity Caregiver education Caregiver finances Caregiver workload Caregiver support Patient age Patient gender Patient disease burden	QoL: SF36-MH
^a Washington <i>et al.</i> (2015) ⁸⁴ USA	To generate an in-depth understanding of the extent to which informal hospice caregivers experience symptoms of anxiety and to identify the characteristics of caregivers who experience clinically significant (i.e. moderate or higher) levels of anxiety.	Cross-sectional secondary analysis; 7	N = 433 Mean age, years (SD) = 60.8 (12) 77.1% female 91.5% white; 50.8% adult child, 30.7% spouse/partner	Not reported		Anxiety: GAD-7
Washington <i>et al.</i> (2018) ⁸⁵ USA	To examine the relationships between sleep problems, anxiety and global self-rated health among hospice family caregivers.	Cross-sectional secondary analysis; 7	N = 395 Mean age, years (SD) = 60.6 (12.6) 81.52% female; 49.10% adult child, 30.40% spouse	Mixed: Cancer (37.22%) dementia (30.13%) No additional information on the remainder of the sample given.	Caregiver sleep problems	Anxiety: GAD-7
continued						

TABLE 1 Observational review study summary table (*continued*)

Reference and country	Study aims	Study design and data collection; QA score	Participants (number; demographics; carer-patient relationships)	Patient condition	Factors investigated in bivariate analysis	Outcomes (anxiety, depression, distress, QoL)
^a Washington <i>et al.</i> (2018) ⁸⁶ USA	To evaluate mediational relationships among burden experienced by hospice caregivers because of symptom management demands, caregivers' coping responses and caregivers' psychological distress.	Cross-sectional survey; 5	N = 228 Mean age, years (SD) = 61.35 (12.65) 80.5% female Carer-patient relationship not reported.	Not reported		Anxiety: GAD-7 Depression: PHQ-9
Wasner <i>et al.</i> (2013) ⁸⁷ Germany	The personal experience-based approaches toward QoL, burden of care and psychological well-being of primary malignant brain tumour (PMBT) patients' caregivers are examined.	Cross-sectional survey and interviews; 4	N = 23 Mean age, years (range) = 51 (34-80) 81.5% female; 70.4% spouse, 29.6% parents or adult child	Primary malignant brain tumour (100%)	Patient disease burden Patient QoL Caregiver burden Caregiver gender	Anxiety: HADS-A Depression: HADS-D QoL: SEIQoL-DW
Wilkes <i>et al.</i> (2018) ⁸⁸ USA	To determine the extent to which burden related to patients' symptom subtypes (emotional/psychological and physical) could predict informal hospice caregiver depression, and to illustrate the differences between caregivers who experience suicidal ideation and those who do not.	Cross-sectional survey; 7	N = 229 Mean age, years = 61.4 80.5% female; 43.8% adult child, 33.6% spouse/long term partner	Mixed: heart disease, lung disease, cancer, dementia included. % composition not reported	Patient symptoms	Depression: PHQ-9
Wittenberg-Lyles <i>et al.</i> (2013) ⁸⁹ USA	To investigate the features of oral literacy in recorded care planning sessions between informal caregivers and hospice team members as they related to the caregiving experience.	Cross-sectional secondary analysis; 5	N = 18 Mean age, years (range) = 64.5 (49-86) 78% female 94% white; 77% adult child, 11% spouse	Not reported	Communication with care professionals	Anxiety: CSAI QoL: CQOL-R

TABLE 1 Observational review study summary table (*continued*)

Reference and country	Study aims	Study design and data collection; QA score	Participants (number; demographics; carer–patient relationships)	Patient condition	Factors investigated in bivariate analysis	Outcomes (anxiety, depression, distress, QoL)
Wittenberg-Lyles <i>et al.</i> (2014) ⁹⁰ USA	To compare how caregivers in pairs (informal collective caregivers) experience anxiety and depression compared to solo caregivers and how these outcomes changed over time. Specifically, after controlling for social support and QoL, does being in a caregiver pair affect anxiety or depression?	Cross-sectional survey; 6	N = 304 Age 45.07% ≥ 61 years 47.04% 41–60 years 7.89% 21–40 years 76% female 91.4% white; 67.7% adult child, 24% spouse	Not reported	Caregiver support	Anxiety: GAD-7 Depression: PHQ-10
<p>ALS, amyotrophic lateral sclerosis; APIM, actor-partner interdependence model; BDI, Beck Depression Inventory; BDI-II, Beck Depression Inventory, version 2; CES, Carer Experience Scale; CRF, case report form; CSNAT, Carer Support Needs Assessment Tool; DSM-IV, Diagnostic and Statistical Manual of Mental Disorders, fourth edition; GAD-7, Generalised Anxiety Disorder-7; GBM, glioblastoma multiforme; GI, gastrointestinal; FACT-G, Functional Assessment of Cancer Therapy – General; HADS, Hospital Anxiety and Depression Scale; HRQOL, health-related quality of life; MND, motor neurone disease; NSCLC, non-small cell lung cancer; PHQ-9, Patient Health Questionnaire; PSOC, perceived stage of cancer; PSS, personal social services; QA, quality assessment; SD, standard deviation; SPC, specialised palliative care; WHOQOL-BREF, World Health Organization quality of life-BREF; WTHD, wish to hasten death.</p> <p>^a Denotes studies that only reported multivariate analysis results.</p>						

in less than 50% of a cancer study population; factors or outcomes related to bereavement only; the outcome measured was anticipatory grief; or the outcome was a composite measure encompassing mental health outcomes included in our review, but where it was impossible to extrapolate findings specifically related to our outcomes, for example a study with the outcome measure Profile of Mood States (POMS), which captures the mood states of anger, depression, fatigue, tension and vigour together; and a substantial proportion of the patient population were unlikely to be cared for at home at the time of the study, for example a study looking at the impact on carers of patient stay in an intensive care unit. Finally, due to the large volume of primary research papers returned, dissertations and conference abstracts were excluded on 'study type'; systematic reviews were excluded on 'study design'.

Narrative summary of evidence

The evidence is synthesised under seven themes ([Table 2](#)). The order of themes does not imply importance. Rather, themes are presented in the same order across all syntheses in the project for consistency. The first six themes correspond with and provide quantitative evidence for all the themes identified in the qualitative synthesis (see <https://arc-gm.nihr.ac.uk/carers-project/>). Additionally, the quantitative evidence identified a further, broad theme of contextual factors. This included, for example, age, gender or socioeconomic status, which are factors that carers are perhaps less likely to consider in qualitative reflections on their own carer experience. [Table 2](#) shows a summary of the bivariate evidence synthesised under each of the seven themes, along with the studies underpinning each theme and the corresponding overall quality assessment score per theme.

[Report Supplementary Material 1](#) shows the total number of bivariate investigations (tests for relationships both within individual studies and across studies) which found a statistically significant positive, a significant negative or a non-significant relationship between a factor and a carer mental health outcome (anxiety, depression, distress or QoL). A 'positive' relationship means that the factor is statistically associated with improved mental health, that is lower anxiety, depression, distress or better QoL. Similarly, a 'negative' relationship means a factor is statistically associated with higher anxiety, depression, distress or worse QoL. Results for the outcomes of anxiety, depression, distress or QoL have been grouped in this table to provide a general overview of factors that may have a positive or negative impact on carer mental health. [Report Supplementary Material 2](#) shows bivariate findings reported for

TABLE 2 Summary of overarching themes from bivariate evidence

Sub-themes	Studies underpinning overarching theme
Patient condition	Overall Quality Assessment Score (mean ± SD): 6.65 ± 1.78
Patient condition	Aoun <i>et al.</i> , ³⁰ Boele <i>et al.</i> , ³⁴ BurrIDGE <i>et al.</i> , ^{35,a} Burton <i>et al.</i> , ³⁶ Butow <i>et al.</i> , ^{38,a} Catt <i>et al.</i> , ^{39,a}
Patient disease burden	Duimering <i>et al.</i> , ⁴⁰ Fasse <i>et al.</i> , ⁴³ Götz <i>et al.</i> , ¹² Govina <i>et al.</i> , ⁴⁷ Grant <i>et al.</i> , ^{48,a} Huang
Patient disease severity	and McMillan; ⁵³ Ito and Tadaka; ⁵⁵ Jacobs <i>et al.</i> , ⁵⁶ Janda <i>et al.</i> , ⁵⁷ Janssen <i>et al.</i> , ⁵⁸ Kershaw
Patient QoL	<i>et al.</i> , ^{61,a} Kobayakawa <i>et al.</i> , ⁶² Loggers and Prigerson; ⁶³ Malik <i>et al.</i> , ⁶⁴ McIlpatrick <i>et al.</i> , ⁶⁵
Patient stage of disease	O'Hara <i>et al.</i> , ^{70,a} Ownsworth <i>et al.</i> , ⁷¹ Perez-Ordóñez <i>et al.</i> , ⁷³ Rivera <i>et al.</i> , ⁷⁵ Seekatz
Patient symptoms	<i>et al.</i> , ⁷⁶ Siminoff <i>et al.</i> , ⁷⁸ Stutzki <i>et al.</i> , ^{79,a} Wadhwa <i>et al.</i> , ⁸³ Wasner <i>et al.</i> , ⁸⁷ Wilkes <i>et al.</i> , ⁸⁸
Patient treatment	
Impact of caring responsibilities	Overall Quality Assessment Score (mean ± SD): 5.57 ± 2.10
Caregiver workload	Buscemi <i>et al.</i> , ³⁷ Catt <i>et al.</i> , ^{39,a} Duimering <i>et al.</i> , ⁴⁰ Flechl <i>et al.</i> , ⁴⁴ Govina <i>et al.</i> , ⁴⁷ Hoefman
Caregiver lifestyle adjustments	<i>et al.</i> , ⁵² Hudson <i>et al.</i> , ⁵⁴ Ito and Tadaka; ⁵⁵ Malik <i>et al.</i> , ⁶⁴ Perez-Ordóñez <i>et al.</i> , ⁷³ Thielemann
Caregiver sleeping hours	and Conner; ⁸⁰ Wadhwa <i>et al.</i> , ⁸³ Washington <i>et al.</i> , ⁸⁵ Wasner <i>et al.</i> , ⁸⁷
Caregiver sleep problems	

TABLE 2 Summary of overarching themes from bivariate evidence (*continued*)

Sub-themes	Studies underpinning overarching theme
Relationships	Overall Quality Assessment Score (mean ± SD): 6.00 ± 2.20
Family dynamics Quality of patient–caregiver relationship	Bachner and Carmel; ³¹ Exline <i>et al.</i> ; ⁴² Fasse <i>et al.</i> ; ⁴³ Götze <i>et al.</i> ; ¹² Hoefman <i>et al.</i> ; ⁵² Mollerberg <i>et al.</i> ; ⁶⁶ Nissen <i>et al.</i> ; ⁶⁹ Siminoff <i>et al.</i> ⁷⁸
Finances	Overall Quality Assessment Score (mean ± SD): 5.83 ± 2.48
Caregiver finances Caregiver mode of transport Impact on work	Flechl <i>et al.</i> ; ⁴⁴ Govina <i>et al.</i> ; ⁴⁷ Hoefman <i>et al.</i> ; ⁵² Ito and Tadaka; ⁵⁵ Kobayakawa <i>et al.</i> ; ⁶² Wadhwa <i>et al.</i> ⁸³
Carer internal processes	Overall Quality Assessment Score (mean ± SD): 6.23 ± 1.83
Acceptance of patient condition Coping patterns Control over the care situation Self-efficacy or self-esteem Positive aspects of caregiving Pre-loss grief Preparedness for caregiving Previous experience of informal caregiving Time for respite	Bachner <i>et al.</i> ; ³³ Burton <i>et al.</i> ; ³⁶ Fasse <i>et al.</i> ; ⁴³ Govina <i>et al.</i> ; ⁴⁷ Hampton <i>et al.</i> ; ⁴⁹ Henriksson and Arestedt; ⁵¹ Hoefman <i>et al.</i> ; ⁵² Hudson <i>et al.</i> ; ⁵⁴ Ito and Tadaka; ⁵⁵ Kobayakawa <i>et al.</i> ; ⁶² McIlpatrick <i>et al.</i> ; ⁶⁵ Nielsen <i>et al.</i> ; ⁶⁷ Perez-Ordóñez <i>et al.</i> ⁷³
Support	Overall Quality Assessment Score (mean ± SD): 6.27 ± 1.44
Accessible information Caregiver support Communication with care professionals Health professionals' understanding of patient needs Quality of care Unmet needs in caregiver	Areia <i>et al.</i> ; ¹⁵ Burton <i>et al.</i> ; ³⁶ Buscemi <i>et al.</i> ; ³⁷ Duimering <i>et al.</i> ; ⁴⁰ Hannon <i>et al.</i> ; ⁵⁰ Hoefman <i>et al.</i> ; ⁵² Ito and Tadaka; ⁵⁵ Janda <i>et al.</i> ; ⁵⁷ Kobayakawa <i>et al.</i> ; ⁶² O'Hara <i>et al.</i> ; ^{70a} Rivera <i>et al.</i> ; ⁷⁵ Thielemann and Conner; ⁸⁰ Wadhwa <i>et al.</i> ; ⁸³ Wittenberg-Lyles <i>et al.</i> ; ⁸⁹
Contextual factors	Overall Quality Assessment Score (mean ± SD): 6.63 ± 2.22
Caregiver age, education or gender Caregiver employment, health or marital status Caregiver ethnicity Caregiver socio economic status Composition of household Length of patient–caregiver relationship Patient age, educational level or gender Patient lives with caregiver Relationship to patient Rural location	Burton <i>et al.</i> ; ³⁶ Butow <i>et al.</i> ; ^{38,a} Catt <i>et al.</i> ; ^{39,a} Duimering <i>et al.</i> ; ⁴⁰ Fasse <i>et al.</i> ; ⁴³ Flechl <i>et al.</i> ; ⁴⁴ Govina <i>et al.</i> ; ⁴⁷ Ito and Tadaka; ⁵⁵ Janda <i>et al.</i> ; ⁵⁷ Kershaw <i>et al.</i> ; ^{61,a} Kobayakawa <i>et al.</i> ; ⁶² McIlpatrick <i>et al.</i> ; ⁶⁵ Rivera <i>et al.</i> ; ⁷⁵ Thielemann and Conner; ⁸⁰ Wadhwa <i>et al.</i> ; ⁸³ Wasner <i>et al.</i> ⁸⁷

SD, standard deviation.

^a Cohort or longitudinal study.**Notes**

Maximum score for quality assessment of cohort or longitudinal studies = 12.

Maximum score for quality assessment of cross-sectional studies = 10.

each type of outcome separately, along with references to the research studies that looked at each individual factor and identified a positive impact, negative impact or no change on carer mental health for each different type of mental health outcome (anxiety, depression, distress or QoL).

Studies that only reported multivariate analysis results are briefly summarised separately under each theme. Their reporting is more complex because the significance of each factor in this case is highly dependent on the other factors considered in the same analysis (and their collinearity) and with the variable set varying widely from study to study, making comparisons difficult. However, it is important that these results are also reported. For consistency, we report the results for the final model presented. Further, we only report significant results, as the volume of non-significant relationships in this part of the literature was large and their presentation became unwieldy with little gain in information for the reader.

Narrative summary of themes

Patient condition

The largest body of research relates to patient condition: 31 studies (see [Table 2](#)) reported on 95 bivariate investigations across all four mental health outcomes. Individual factors that contribute to this theme include the patient's diagnosis, patient disease burden (i.e. physical and cognitive functioning, QoL, stage or rate of decline, physical and psychological symptoms) and treatment.

Some studies indicated that a diagnosis of primary brain cancer (one investigation³⁰), rare cancers (one investigation⁶³) or lung cancer (two investigations⁴⁷) is related to worse carer mental health compared to other cancer diagnoses. However, one investigation comparing rare cancers with other cancers found no difference,⁶³ and three investigations considering a range of cancers, including lung and brain, found no difference between cancer diagnoses.^{55,61,75} Further, no differences were reported in three investigations comparing lung cancer with heart failure (HF).⁶⁴ One investigation found patient diagnosis of chronic obstructive pulmonary disease (COPD) to be associated with worse carer mental health when compared with chronic heart failure (CHF) or renal failure.⁵⁸ Findings on diagnosis, however, are likely to be highly dependent on what the comparators are, and whether two large comparison groups are considered or a range of smaller size diagnostic groups. Further, diagnosis in itself may mean little without added knowledge of patient stage or disease burden.

Three investigations found a relationship between greater patient functional impairment and worse carer mental health.^{40,71,73} However, a further nine investigations of functional impairment showed no association.^{36,55,75,83,87} There was no relationship identified between patient cognitive impairment and carer mental health (three investigations⁸⁷).

Three investigations indicated that a more advanced patient stage of disease is related to worse carer mental health,^{38,48,79} while a further four investigations found no relationship with carer mental health.^{35,38,48} These findings include factors related to patient disease trajectory and patient rate of decline, so may tell us little without considering the impact of these factors on patient stage of disease.

Two investigations into patient disease severity found no relationship with carer mental health.³⁶

In six investigations, better patient general QoL was related to significantly better carer mental health.^{57,65,70,83} In a seventh investigation, general QoL was reported to be associated with carer mental health, although the direction of the relationship was not clarified.⁷⁵ One investigation found that better patient psychological QoL was also associated with better carer mental health.³⁴ Three investigations found no significant relationship, however.⁸⁷

Two investigations found patients' overall symptoms to relate to worse carer mental health,^{70,88} but one of these incorporated an element of the carer's stress into the patient symptom measure, thus making

an association with mental health outcomes more likely.⁸⁸ A third investigation found no relationship.⁵⁵ Physical symptoms show a mixed picture: greater drowsiness, fatigue and pain were related to worse carer mental health,⁸³ whereas loss of appetite, breathlessness and nausea showed no relationship (one investigation each per symptom).⁸³

Patients' psychological symptoms appear to show a consistent relationship with carer mental health. Higher patient anxiety and depression were related to worse carer mental health in 8^{12,56,57,65,83} and 10 investigations,^{12,53,56,57,65,78,83} respectively. Only three investigations of patient depression found no relationship.^{53,70,75} Worse patient global distress,⁷⁵ psychological and psychiatric symptoms⁶² also related to worse carer mental health (one investigation each). In contrast, one investigation of patient sense of well-being showed no association.⁸³

Regarding patient treatment, carers had worse mental health if the patient had been admitted to hospital or long-term care within the previous 7 days,⁴⁰ had received no cancer therapy⁸³ and no surgery⁴⁷ (one investigation each), which could imply, respectively, deterioration or that 'nothing could be done'. However, other investigations found no association with receiving no surgery⁴⁷ (one investigation), with receipt of chemotherapy⁴⁷ (two investigations) or medical care provided⁵⁵ (one investigation). Other treatment variables showing no relationships were patients awaiting a new line of treatment,⁸³ frequent visits to emergency outpatient clinics,⁶² type of oncology follow-up³⁹ and patient receipt of specialist palliative care⁷⁶ (one investigation each).

Some corresponding findings were reported in studies only reporting multivariate analyses. Patient QoL⁴¹ and better functioning⁴⁵ were related to better carer QoL, and patients' need for help at night⁶⁰ and problems sleeping⁸² to worse carer mental health. A perceived lower life expectancy was associated with worse carer emotional QoL⁸¹ and worse patient mental health was related to worse carer mental health.^{68,77} There was also worse carer depression where patients had worse social well-being, patients used more emotional support seeking, less acceptance coping and perceived that the primary goal of their cancer treatment was 'to cure my cancer',⁶⁸ whereas patients' use of less emotional support seeking was associated with higher carer anxiety.⁶⁸

Impact of caring responsibilities

A smaller body of research, based on 14 studies (see [Table 2](#)) and 36 bivariate investigations across all four mental health outcomes, concerns the impact of caregiving in terms of life changes and care demands, a construct similar to objective burden. Where studies investigated impact using carer burden measures, we need to exercise some caution, due to the wide variety of these measures, some of which incorporate emotional impact. In our selection and synthesis, we therefore sought to avoid studies using burden measures that essentially measure subjective burden or psychological impact, as these may in effect be synonymous with the outcomes we were investigating.

Studies consistently indicated that the impact of caring responsibilities is associated with worse mental health. Five investigations found that negative changes to carers' lives from caregiving were associated with worse mental health (using Bakas Caregiving Outcomes Scale).^{37,47} Two investigations each found that difficulty of caregiving tasks and time spent on tasks were also related to worse health [using Oberst Caregiving Burden Score (OCBS)-D and OCBS-T, respectively].⁴⁷ One investigation found that the impact on carers' schedules (using the Caregiver Reaction Assessment) had a similar relationship with mental health.⁵⁴ In terms of overall burden, three investigations using the Zarit Burden Inventory,⁶⁴ three using the Burden Scale for Family Caregivers⁸⁷ and one using the Caregiver Strain Index,⁷³ all found increased burden to be associated with worsening mental health.

Studies have also found that making greater lifestyle adjustments,³⁹ greater demands on the carer,⁸⁰ assistance with activities of daily living⁴⁰ and medical tasks,⁴⁰ number of days spent caregiving,⁸³ physical strain from caregiving⁵² and sleep problems⁸⁵ relate to worse mental health (one investigation each), although one investigation found no relationship with carer sleeping hours.⁵⁵

Other demands on carer time⁵² or childcare responsibilities³⁹ may relate to worse mental health, found by one investigation each. However, two further investigations that considered if carers had children of minor age⁴⁷ and one whether they cared for others⁸³ found no relationship. No relationships with mental health were found in one investigation of the number of caregiving hours per week⁸⁰ and three considering duration of care.^{44,55,80}

Studies only reporting multivariate analyses also found that higher carer burden was associated with worse QoL (Caregiver Burden Inventory)⁴⁵ and mental health (Caregiving Burden Interview – Zarit⁵⁹; Caregiver Reaction Assessment⁷⁴), and similarly that more impairment to daily life was associated with worse mental health.⁵⁹

Relationships

There is evidence that the family dynamics and the quality of the carer-patient relationship are related to carer mental health, although this is based on a relatively small number of studies (eight) (see [Table 2](#)) reporting only 16 bivariate investigations across all four mental health outcomes.

Two investigations within the same study found better carer mental health where carers felt that the family had high ability to cope with stressors (measured by Family Sense of Coherence Scale).⁶⁶ Investigations in another study using the Family Environment Scale found carer mental health to be worse both when the patient and when the carer perceived there to be low family cohesion (i.e. low commitment, help and support that family members give to one another);⁷⁸ low family expressiveness (i.e. low encouragement of direct expression feelings);⁷⁸ and high family conflict (i.e. openly expressed anger and conflict).⁷⁸ Correspondingly, one further study also reported worse carer mental health both when the patient and when the carer perceived there to be unresolved family conflicts,⁴² whereas another found better mental health when supportiveness of family relationships was high.⁶⁹

Looking specifically at the patient-carer relationship, one study found that carer dissatisfaction with the relationship was associated with worse carer mental health,¹² whereas a second found no relationship in terms of the carer getting on with the patient.⁵² Good carer communication with the patient about their illness and approaching death was related to better carer mental health.³¹

Finally, one study found worse carer mental health where the carer had an insecure-anxious attachment style,⁴³ whereas no relationship was found if they had an insecure-avoidant attachment style.⁴³

Studies only reporting multivariate analyses have also found that carers with good family relationships had better mental health,⁴⁶ and one study considering mediators concluded that carers with supportive relationships had better mental health through decreased carer burden.⁷⁴

Finances

Although there were relatively few studies considering the role of financial factors (six) and only eight bivariate investigations relating to three of the four mental health outcomes (QoL, anxiety and depression), the majority of studies indicate a relationship between finances and carer mental health.

Having a sufficient family budget was related to better carer mental health (one study),⁵⁵ whereas having financial difficulties due to the patient's disease⁴⁴ or to providing informal care⁵² were related to worse carer mental health (one study each). Changes to work situations in terms of reduction, change or ending of work (one study)⁵⁵ were also associated with worse mental health.

However, level of income in itself (two studies)^{62,83} showed no relationship. Having a private car as a means of transport was, perhaps surprisingly, related to worse mental health in one investigation, but showed no relationship with another mental health measure within the same study.⁴⁷ Level of income or possessions may in themselves be less informative; what matters may be whether they provide sufficient or insufficient resources during caregiving. Findings may also depend on the populations studied. For

example, a study population in which everyone is generally affluent may show different patterns of association with carer mental health compared with study populations with a range of incomes.

Carer internal processes

Thirteen studies (see [Table 2](#)) reporting 36 bivariate investigations relating to QoL, anxiety and depression have considered how carers' internal, psychological processes and coping strategies are related to their mental health, and they have investigated a wide range of variables.

In terms of coping strategies, the picture is quite mixed and mostly showing little association with mental health, which may reflect the challenge of using questionnaires to ask carers about dispositions to cope with hypothetical situations. Difficulty accepting the patient's condition⁶² or 'dysfunctional' coping strategies⁷³ (including lack of acceptance and avoidance) were associated with worse mental health in one study each. Worse mental health was also found in relation to disengagement through substance misuse in one investigation.⁴³ However, other investigations considering denial (one investigation),⁴³ cognitive avoidance (two investigations)³⁶ or mental disengagement (one investigation)⁴³ found no relationship.

Being optimistic was associated with better mental health (one study),⁵⁴ whereas using humour,⁴³ having a 'fighting spirit' coping style³⁶ or using emotion-focused strategies⁷³ (e.g. seeking a positive outlook and acceptance) showed no relationship (one study each). Having a secular outlook was related to better mental health in one study,³³ while religious coping showed no significant association in a second.⁴³

Suppression of competing activities (staying focused on the problem) has been found to relate to worse mental health (one study).⁴³ Conversely, problem-focused coping strategies⁷³ or active coping to solve a problem⁴³ was found to be unrelated to mental health (one study each).

Finally, in terms of coping strategies, seeking emotional social support⁴³ or venting of emotions⁴³ was associated with worse mental health in one study, although it may be important to consider here which is cause and which is effect. Seeking information support was unrelated to mental health in the same study.⁴³

Three investigations found that carer self-efficacy (confidence in one's ability to carry out a task) was related to better carer mental health.^{49,55} Conversely, if carers felt helpless or guilty because they could do nothing for the patient, they had worse mental health (one investigation).⁶² However, carers' sense of control over the care situation was not found to relate to mental health (one investigation).⁵²

Two investigations found that preparedness for caregiving was also associated with better health,^{51,65} although one investigation found no relationship.⁵¹ Further, if carers had provided care to a loved one in the past, they reported worse health (two investigations),⁴⁷ indicating that the experience gained from past caregiving may not be protective.

Pre-loss grief⁶⁷ and, perhaps surprisingly, higher carer self-esteem⁵⁴ were related to worse mental health (one study each), whereas fulfilment from caring and being happy to care (both investigated in the same study)⁵² showed no relationship.

Having enough time for oneself was associated with better mental health in one study,⁵² but activities outside caring measured within the same study showed no association.⁵²

Studies that reported only multivariate analyses have also found higher carer preparedness to relate to better QoL⁴⁵ and also report mixed results for coping. Carer meaning-based coping was associated with better QoL,⁴¹ and carers' use of escape/avoidance coping with worse mental health.⁸⁶ Active coping was in fact associated with worse mental health, and substance abuse with better mental health in a further study.⁵⁹ Carers with stronger religious/spiritual beliefs had better mental health.⁵⁹ Among

studies considering coping strategies as mediators, Washington *et al.*⁸⁶ concluded that the relationship between patients' psychological symptoms (reported above) and carers' mental health was partially explained by carers' increased use of escape/avoidance coping, whereas Ellis *et al.*⁴¹ reported that the number of carers' chronic conditions had an indirect negative effect on their QoL mediated by meaning-based coping.

Support

The third largest body of research has been conducted on support, based on 15 studies (see [Table 2](#)) reporting on 42 bivariate investigations across all four mental health outcomes.

Accessible information for patients and for carers are both related to better carer mental health (one study).⁵⁵

In terms of support for carers themselves, there is some evidence that the presence of informal support is positive. Carers who have social support from family and friends (two studies),^{52,80} who have a sub-caregiver (one study)⁵⁵ and who are satisfied with physical, emotional and informational support (one study)⁷⁵ have better mental health. However, no relationship with mental health was found for carers who were in receipt of informal help (one investigation),⁸³ availability of someone to stay with the patient (one investigation),⁶² who worked in pairs (two investigations)⁹⁰ or where support was perceived (two investigations).³⁶

In terms of formal support for carers, one study found better mental health for carers who received support services⁵⁵ or requested home care for the patient.⁵⁵ However, other studies have found no relationship for formal^{40,83} or institutional help.⁵² One investigation within one study found that carers who had professional psychological help, in fact, had worse health, while two further investigations found no relationship.⁵⁷ We need to consider what may be cause or effect here, as carers with higher distress may be more likely to seek psychological help. Carers interested in accessing future support services,⁴⁰ and those who received no help from home-visit practitioners in managing symptoms,⁶² had worse mental health (one study each). Type and frequency of formal support services showed no association in one study.⁵⁵

Unmet needs in the carer appears to be important. Three investigations relating to carers' unmet psychological, social and physical needs in one study³⁷ and one investigation considering number of carers' unmet needs by health professionals in another study¹⁵ found that they were related to worse carer mental health.

Features of communication with practitioners during care planning sessions made little difference. An investigation in one study found that a faster dialogue pace was related to worse carer mental health,⁸⁹ whereas another investigation found no relationship.⁸⁹ No associations were found for language complexity, length of interaction or the team taking turns to speak.⁸⁹

Carer satisfaction with patient care (two studies)^{50,55} and patient satisfaction with care (one study)⁵⁰ were associated with better carer mental health, while carer perception of problems with patient unmet needs was related to worse mental health (one study).⁷⁰ Perhaps counterintuitively, carers in the same study, who perceived more problems with the patient's emotional and spiritual support, had better mental health.⁷⁰ No associations were found for practitioners' lack of understanding of patient symptom severity⁶² or whether services received were considered necessary by the carer⁵⁵ (one study each).

One study reporting only multivariate analysis found that carers with good healthcare providers had better mental health.⁴⁶

Contextual factors

The second largest body of research relates to contextual factors (16 studies reporting 104 bivariate investigations across all four mental health outcomes). Studies have considered a range of carer and patient contextual factors. Older carers were found to have better mental health in seven investigations,^{47,57,61,65,83} while eight investigations found no relationship.^{36,39,44,47,55,75,80} Female carers in general,^{39,43,47,83,87} or wives,⁷⁵ had worse health in seven investigations, although 14 investigations found no similar gender differences.^{36,38,40,55,57,61,62,75,80}

Otherwise, there are no clear patterns for contextual variables. Being unemployed,⁸³ or employed but on leave,⁴⁰ was related to worse carer mental health (one study each). Conversely, these two studies found retirement to be associated with better mental health,^{40,83} although this variable may be closely associated with age. However, six investigations found no association for employment status.^{36,39,47,55} Poor carer physical health was found to be related to worse mental health in one study,⁶² but four studies found no relation for physical or general health.^{55,61,75,83} In terms of race or ethnicity, being white was associated with better carer mental health in one study,⁷⁵ while two other studies did not find an association between ethnicity and mental health.^{80,83} While carers with higher socioeconomic status were found to have better mental health in one study,⁴⁰ nine investigations found no relationship between education level and health.^{39,47,57,62,80,83}

One study found higher patient age to be associated with better carer mental health,⁵⁵ but two others found no association.^{44,83} The patient being male was related to worse health in two investigations,⁴⁷ but two studies found no relationship with patient gender.^{55,83} If the patient was living with the carer, this has been found to both show a relationship^{40,47} and no relationship with worse mental health^{47,83} (two investigations each). The carer's relationship with the patient has shown quite mixed results. While being a spouse has been found to relate to better mental health⁸³ and not being a spouse/partner to worse health³⁹ (one study each), being a son/daughter⁴⁰ and not being a spouse/daughter⁷⁵ have also been associated with better mental health. Further, 11 investigations found no association for relationship with the patient.^{38,47,55,57,61,62,75}

No association with carer mental health was found for carer marital status (four investigations),^{36,47} composition of the household (one investigation),⁵⁵ length of patient-carer relationship (one investigation),⁸⁰ patient education level (two investigations)⁴⁷ or living in a rural area (three investigations).^{38,40}

Studies reporting only multivariate analyses also found that younger carers had worse mental health,^{68,72,77} and carers of younger patients had worse QoL.⁸² Caring for a patient with a diagnosis other than cancer was also associated with worse mental health.⁷² Again female carers reported worse mental health,^{68,82} although one study found that the relationship between gender and anxiety was age dependent, with females reported to have significantly higher probability of being anxious than males until about the age of 60 years.⁸⁴ One study reported that carers who reported poorer self-rated global health had worse mental health.⁷² Three studies found worse mental health among spousal carers compared with other relationships.^{68,72,77} One study reported employed carers had better mental health than unemployed carers.⁸⁴ One study each reported that African Americans had better mental health compared with white carers;⁴⁶ that English speakers had better mental health compared with those who did not speak English at home;⁶⁰ and that those with a Catholic faith had worse mental health than other denominations/religions.⁶⁸

Meta-analyses

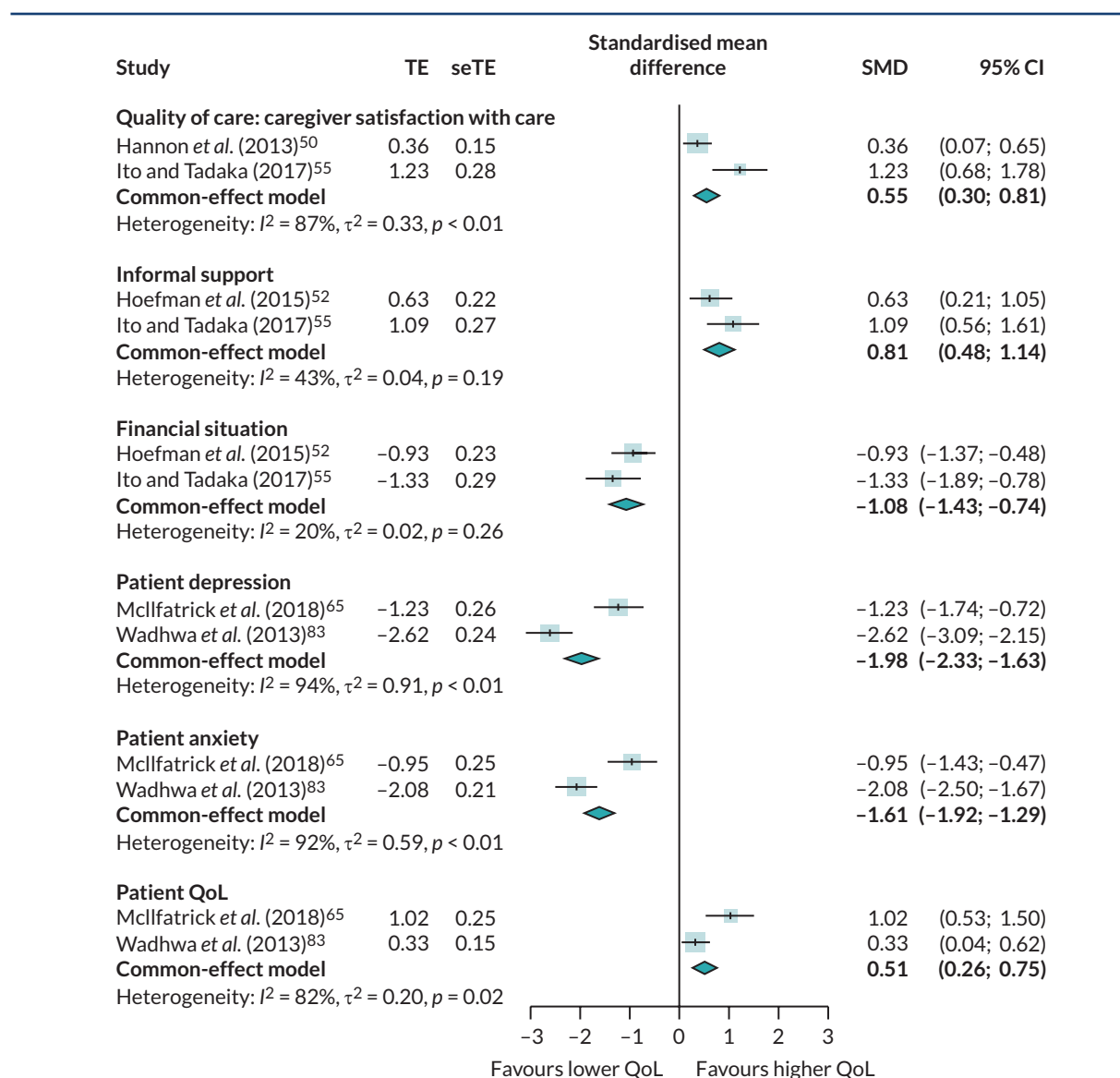
As reported earlier, the opportunity for meta-analysis was limited due to the wide range of factors and the range of mental health outcomes considered. While we felt justified in grouping findings for different mental health outcomes for a narrative thematic summary, stricter criteria had to be applied for meta-analysis. We therefore only performed meta-analysis on studies that considered the same mental health outcome (i.e. anxiety, depression, distress or QoL) to try and avoid introducing a high level of

random variation into the analysis. Similarly, while a wider interpretation of similarity of factors should be permissible for thematic grouping in a narrative summary, we needed to be stricter in ensuring that studies included in a meta-analysis were indeed considering comparable factors. There were therefore few instances where studies considered sufficiently similar factors and their relation to the same outcome to permit meta-analysis.

Quality of life

For the studies in the meta-analysis, higher QoL in carers was highly associated with receiving informal support [SMD = 0.81, 95% confidence interval (CI) 0.48 to 1.14, $I^2 = 43$, $n = 2$ studies]^{52,55} and moderately associated with carer satisfaction (SMD = 0.55, CI 0.30 to 0.81, $I^2 = 87\%$, $n = 2$),^{50,55} along with higher patient QoL (SMD = 0.51, 95% CI 0.26 to 0.75, $I^2 = 82\%$, $n = 2$).^{65,83} Financial strain (SMD = -1.08, 95% CI -1.43 to -0.74, $I^2 = 20\%$, $n = 2$ studies),^{52,55} patient depression (SMD = -1.98, 95% CI -2.33 to -1.63, $I^2 = 94\%$, $n = 2$ studies)^{65,83} and patient anxiety (SMD = -1.61, 95% CI -1.92 to -1.29, $I^2 = 92\%$, $n = 2$ studies)^{65,83} were all found to be highly associated with lower QoL (Table 3).

TABLE 3 Meta-analysis of factors associated with QoL



seTE, standard error treatment effect; TE, treatment effect.

Depression

The studies in the meta-analysis indicated that negative changes to carers' lives from caregiving (using Bakas Caregiving Outcomes Scale) were highly associated with higher depression levels in carers (SMD = 1.36, 95% CI 0.97 to 1.74, $I^2 = 0\%$, $n = 2$ studies).^{37,47} Being a female carer (SMD = 0.40, 95% CI 0.21 to 0.60, $I^2 = 87\%$, $n = 3$ studies),^{43,47,62} patient anxiety (SMD = 0.56, 95% CI 0.34 to 0.79, $I^2 = 69\%$, $n = 2$ studies)^{56,65} and patient depression (SMD = 0.59, 95% CI 0.32 to 0.87, $I^2 = 77\%$, $n = 5$ studies)^{12,53,56,65,78} were all moderately associated with higher depression in carers. Carers caring for patients with lung cancer were found to be moderately associated with higher depression levels than carers of those with other conditions (SMD = 0.42, 95% CI 0.10 to 0.73, $I^2 = 81\%$, $n = 2$ studies),^{47,64} although these results should be interpreted with caution as the comparison group in each study related to a different patient condition (patients with other cancers⁴⁷ and patients with HF⁶⁴) (Table 4).

Anxiety

For the studies in the meta-analysis, negative changes to carers' lives from caregiving (using Bakas Caregiving Outcomes Scale) were highly associated with higher anxiety levels in carers (SMD = 1.10, 95% CI 0.73 to 1.46, $I^2 = 0\%$, $n = 2$ studies).^{37,47} Patient anxiety (SMD = 0.60, 95% CI 0.41 to 0.80, $I^2 = 65\%$, $n = 3$ studies)^{12,56,65} and patient depression (SMD = 0.44, 95% CI 0.22 to 0.66, $I^2 = 90\%$, $n = 2$ studies)^{56,65} were found to be moderately associated with higher carer anxiety. Carers caring for patients with lung cancer were found to be moderately associated with higher anxiety levels than carers of those with other conditions (SMD = 0.38, 95% CI 0.07 to 0.70, $I^2 = 79\%$, $n = 2$ studies),^{47,64} although as previously reported, these results should be interpreted with caution as the comparison group in each study related to a different patient condition (patients with other cancers⁴⁷ and patients with HF⁶⁴) (Table 5).

Distress

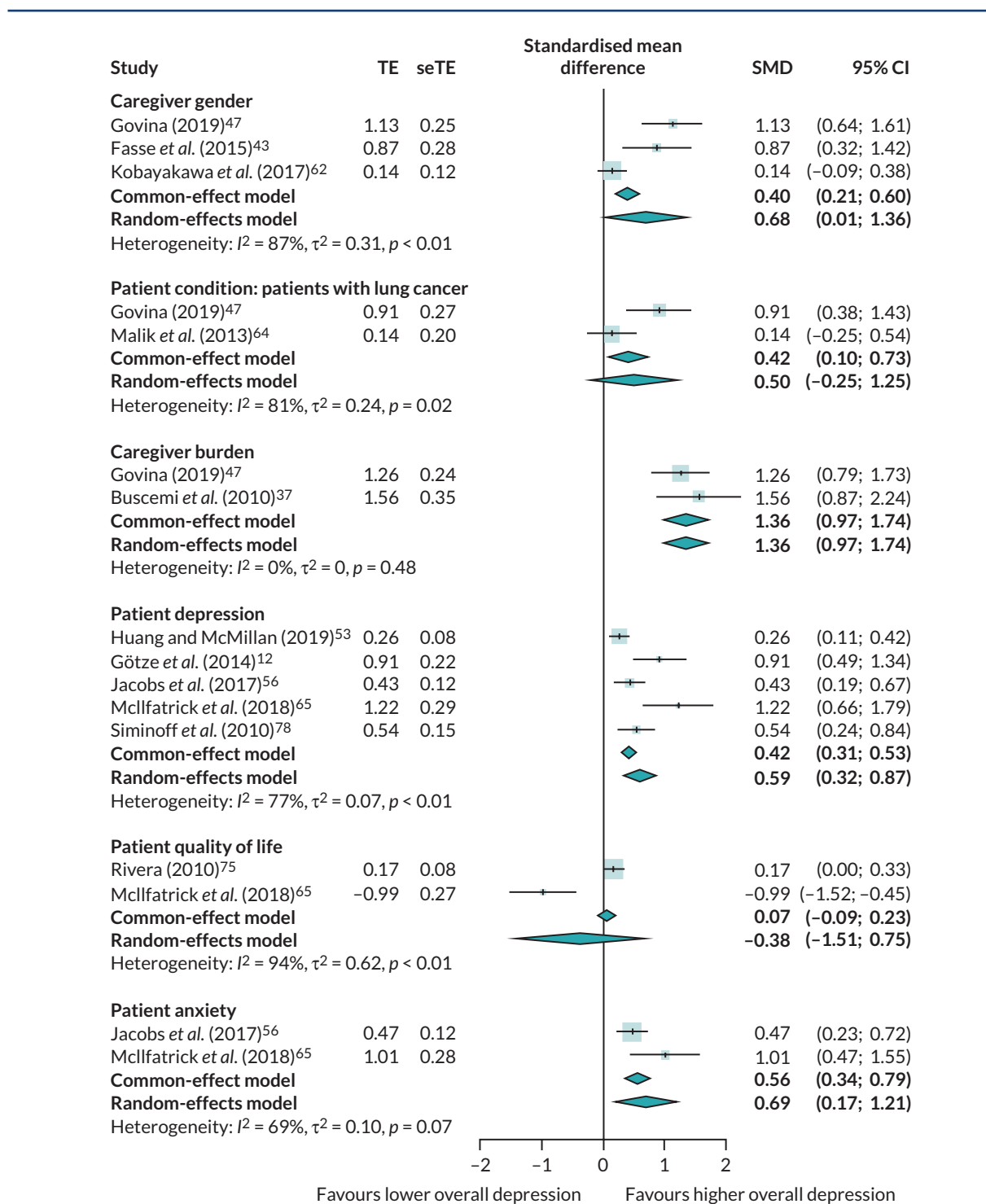
Studies in the meta-analysis indicated that unmet needs in carers was highly associated with higher carer distress (SMD = 0.67, 95% CI 0.32 to 1.00, $I^2 = 64\%$, $n = 2$ studies).^{15,37} Being a female carer was also associated with higher carer distress (SMD = 0.18, 95% CI 0.01 to 0.36, $I^2 = 60\%$, $n = 2$ studies)^{39,61} (Table 6).

Quality assessment

The NOS quality assessment criteria adapted for cross-sectional studies were applied to 56 of the 63 included studies; the remaining 7 studies were assessed using the NOS quality assessment criteria adapted for cohort/longitudinal studies. The predominance of cross-sectional studies over longitudinal studies means we can be less certain overall of the causal direction of any relationships found.

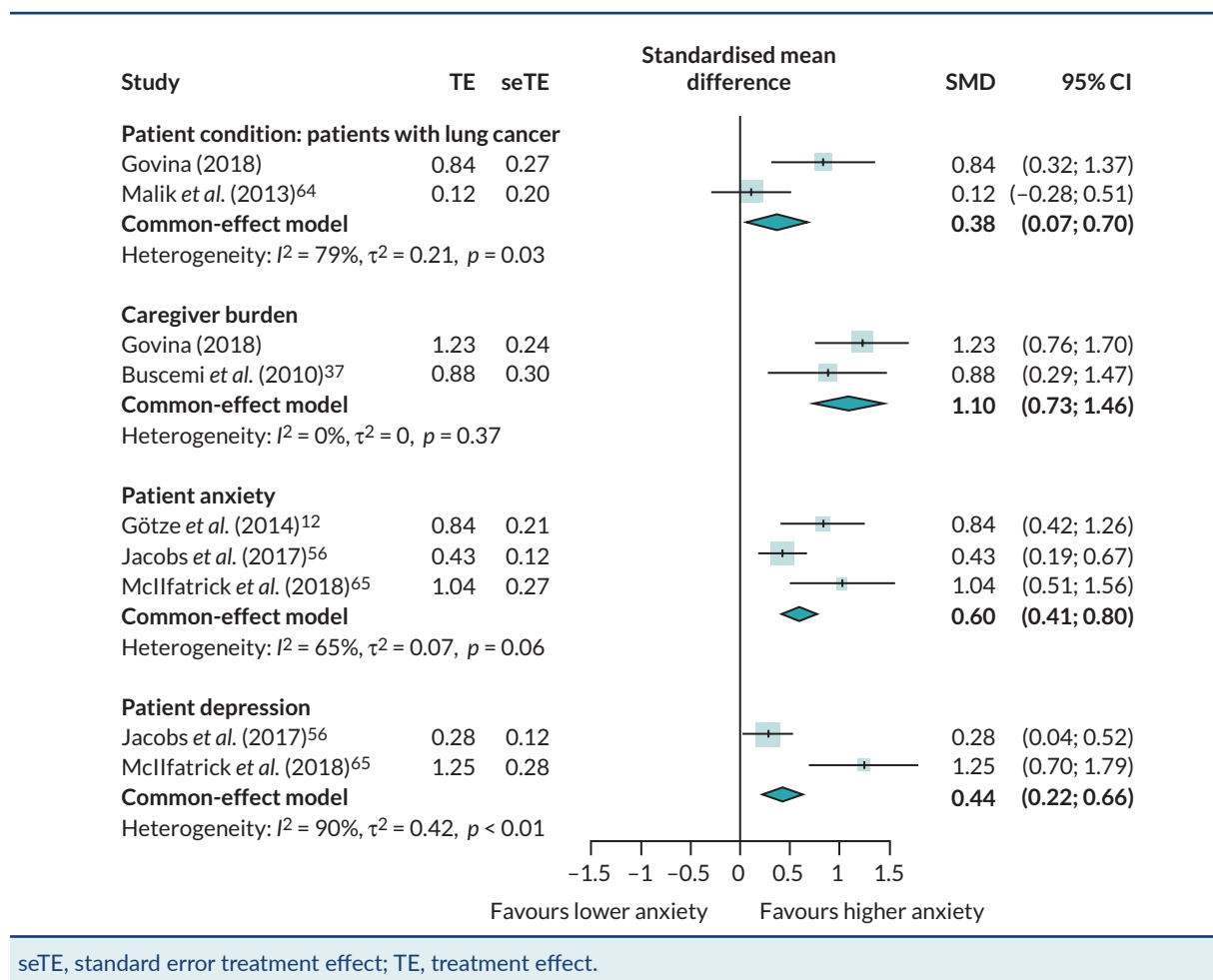
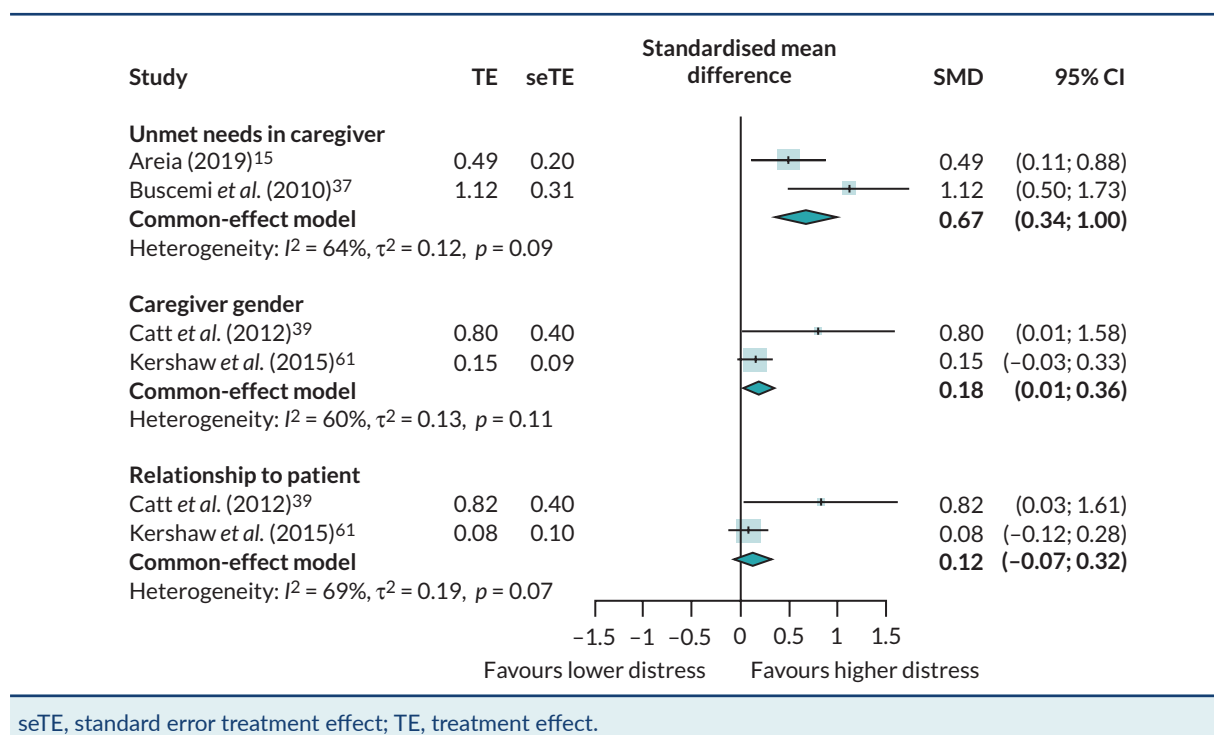
Cross-sectional studies were most likely to meet criteria relating to assessment of validated outcomes (49/56 studies – 87.5%); ascertainment of validated predictors (47/56 studies – 83.9%); adequacy of statistical tests applied (46/56 studies – 82.1%); and selection of sample (42/56 studies – 75.0%). They were least likely to meet criteria relating to reporting an a priori hypothesis (26/56 studies – 46.4%); sampling frame (19/56 studies – 33.9%); and the degree to which non-respondents were adequately reported (11/56 studies – 19.6%).

Cohort/longitudinal studies were most likely to meet criteria relating to representativeness of sample/exposed cohort (7/7 – 100%); selection of exposed and control cohorts (7/7 – 100%); assessment of validated outcomes (6/7 studies – 85.6%); and ascertainment of validated predictors (6/7 studies – 85.7%). They were least likely to meet criteria relating to sampling frame (3/7 – 42.9%); reporting of non-respondents (3/7 – 42.9%); adequacy of follow-up (28.6%); and having an a priori hypothesis (1/7 – 14.3%).

TABLE 4 Meta-analysis of factors associated with depression

seTE, standard error treatment effect; TE, treatment effect.

Table 2 presents the overall quality assessment score for each overarching theme. The quality of studies underpinning each of the seven overarching themes was very similar, although there was some degree of variation. The overarching themes 'patient condition' (mean \pm SD: 6.65 \pm 1.78) and 'contextual factors' (mean \pm SD: 6.63 \pm 2.22) were underpinned by the highest quality studies overall, followed by 'support' (mean \pm SD: 6.27 \pm 1.44), 'carer internal processes' (mean \pm SD: 6.23 \pm 1.84), 'relationships' (mean \pm SD: 6.00 \pm 2.20) and 'finances' (mean \pm SD: 5.83 \pm 2.48) and with 'impact of caring responsibilities' (mean \pm SD: 5.57 \pm 2.10) having the lowest quality studies overall.

TABLE 5 Meta-analysis of factors associated with anxiety**TABLE 6** Meta-analysis of factors associated with distress

Discussion

General discussion of findings

The literature in this review is very diverse, and it is difficult to give a simple summary of findings. However, some patterns emerged. Regarding the patient condition, when patients' psychological symptoms are worse, this seems fairly consistently related to worse carer mental health, while better patient QoL seems related to better carer mental health. A concern here may be if patient scores were provided by carers by proxy, as scoring may then be influenced by the carers' own mental health, making correlations more likely to be an artefact of study design. Findings for other patient condition variables are mixed and less clear.

The impact of caring responsibilities (in terms of impact on carer lives, task difficulty and general burden) showed clear associations with worse mental health. However, these factors have predominantly been measured using carer burden measures, and while we were careful to exclude measures that essentially assessed psychological impact, there is a possibility that some remaining measures still retained a subjective, emotional element that made them more likely to correlate with mental health as an artefact of the measurement tools. However, further studies that did not rely on burden measures also indicated that greater impact on carers' lives is associated with worse mental health.

Family dynamics, the quality of the carer-patient relationship and finance have not been extensively studied, but findings suggest that they show sufficient relationship to carer mental health to warrant further investigation.

Where carers' psychological processes are concerned, self-efficacy and, possibly, preparedness appear related to better mental health. However, having provided care to a loved one in the past may be detrimental rather than positive. Research on coping strategies shows limited or mixed associations with mental health and may need more consistency and direction to become useful.

Having unmet needs appears related to worse mental health, while satisfaction with care may relate to better mental health, but more research is required to better understand where and how informal and formal support may have an impact.

In terms of contextual factors, older age seems generally to be associated with better carer mental health, and being female with worse health, but it is difficult to draw conclusions from findings on other contextual factors.

Compared with the comprehensive review of the quantitative carer literature by Stajduhar *et al.*,²⁰ this review confirms and expands on the previous findings. It identified similar factors in terms of the patient condition, impact on carers' lives, carer internal psychological processes and context, which indicates consistency in the patterns found. Additionally, the current review captured emerging research on relational variables, available support and features of interaction with healthcare providers which the earlier review noted were missing from the literature. We also identified literature on the association between financial difficulties and carer mental health which was absent from the earlier review.

Quality of the evidence

Study designs

A major problem with the observational quantitative literature in informing predictors of carers' mental health is the predominance of cross-sectional studies (56 studies) and the dearth of longitudinal studies (7).

This considerably limited our ability to be certain about the causal direction of the relationships found. Some factors are clearly likely to be precursors to carers' state of mental health, such as elements of the patient condition, age and sex, although we are left to hypothesise as to why they may affect mental health. However, many other variables may plausibly be the effect of, rather than a contributing cause of, carers' mental health. For instance, the carer with poor mental health may feel caregiving tasks are more onerous, have a lower sense of self-efficacy and preparedness and perceive quality of support to be worse, rather than the other way around. Further, many variables may work in both directions; for instance, patient and carer anxiety are likely to influence each other. It is therefore often a matter of judgement whether we believe that factors are precursors and/or contributors to carers' mental health, although mostly it appears plausible that they should be.

A further challenge in assessing causal relationships is that on occasion it can be unclear exactly what a variable measures or what it means, for example, patient diagnosis or type of service (if we know nothing further about the features of the patients or service in a given study context) or 'carer burden' (which is conceptualised somewhat differently within the measures used). It was also not possible to provide any definitive evidence about the strength of the relationships identified, due to the small number of studies identified for each bivariate relationship, which were often based on small sample sizes and may therefore inflate the effect sizes. While some of the factors may be more important in protecting or worsening carers' mental health than others, it is difficult to conclude from the findings in this review which these are.

Gaps within the evidence

It is important to recognise that some of the factors identified within this review were more intensively investigated than others. Although there is evidence that both carer finances and the quality of relationships are related to carer mental health, the evidence base is limited in comparison with patient condition, impact of caring responsibilities, carer internal processes, support or contextual factors.

Similarly, the number of investigations with significant results within each factor also varied. While the highest number of significant investigations were reported for the patient condition (50/95), the lowest number of significant investigations were identified for relationships (14/16) and carer finances (5/8). This is not surprising given the overall body of evidence underpinning each of these factors. What may be of more interest is the proportion of significant interactions identified within the relationship theme, which was the highest across all factors.

The vast bulk of studies within this review investigated depression as an outcome (115 factors across 34 studies), with carer distress investigated the least (32 factors across 12 studies). This may be due in part because distress was limited within our review to measurement scales identified specifically as measuring psychological distress [e.g. General Health Questionnaire-12 (GHQ-12), Psychological Distress Thermometer] and therefore did not capture broader outcomes like anger, frustration and grief (which are included within the qualitative synthesis). Despite this, it is important to recognise that the evidence base in this review is considerably more weighted towards outcomes related to depression, anxiety and QoL.

We can only report on what the observational quantitative research has focused on. This is not the same as saying other factors are unimportant. Further, the review reported primarily on bivariate relationships between factors and mental health outcomes, so there may have been confounding factors which exaggerated or masked the real relationship between the factor/s identified and carer mental health.

Lack of models

Overall, the lack of good models to guide enquiry poses a challenge to synthesis and clear conclusions. Both this review and that of Stajduhar *et al.*²⁰ found little use of models or frameworks within the carer research literature.

There was sometimes an impression that variables were investigated simply because they were available or because validated measurement tools existed for them, rather than because there was a clear theoretical rationale for their inclusion. This is particularly the case for clinical data (e.g. condition, treatment) and contextual data (e.g. age, sex) that are often routinely recorded for other purposes.

Without clear models or frameworks to provide direction and systematic enquiry, research can become piecemeal and sprawling. Accordingly, we found that this research proved difficult to summarise because of the very wide range of factors, carer groups and contexts considered, and little of the evidence lent itself to the meta-analysis.

Limitations to the review

Our review focused on caregiving in a home-care setting. It may therefore not fully capture what gives rise to worse carer mental health in other settings, for example the intensive care unit. However, the home is where most of the care in the patients' final year takes place, where a majority of patients prefer to remain for as long as possible, even to death, and where carers have the greatest responsibility and undertake the widest range of tasks. Making sure carers are supported in this setting is likely to have the greatest impact on patient care, as well as carers themselves.

Our review is mainly representative of the developed world, and it is limited to studies published in English and Scandinavian. Although a range of countries are represented, these are all OECD countries. There is therefore a lack of research from cultures and perspectives beyond the developed world. Further, within the countries represented in the review, there is little research encompassing perspectives of ethnic minorities. We also focused on adult carers during caregiving, and the review may not reflect relevant factors for younger carers and outcomes in bereavement. The literature itself typically focuses on carers of people with cancer and may not fully represent carers of people with other conditions, for example longer term conditions where duration of care and service provision may be different.

The search was limited to 2009–19 and did not cover dissertations or grey literature, nor did we scan bibliographies or contact key authors direct. Important studies may therefore have been missed. However, comparison with overview reviews of the carer literature from 1998 to 2008^{20,91} indicates that the factors identified remain fairly consistent across studies and over time.

Our review did not apply GRADE to provide an overall assessment of the certainty of evidence and strength of the findings as was proposed in the protocol. GRADE is better suited to interventional trials rather than observational research. Furthermore, the diversity in measurement of factors, imprecision and different summary statistics used by studies hampered attempts to transform the data. Therefore, to assess validity and importance of research findings, rather, the project invested time and resources on PPI through carer RAP and stakeholder consultations. This confirmed that the seven identified themes were perceived as important and as meaningful causal factors (see [Relevance of findings: patient and public involvement from carer Review Advisory Panel and stakeholder consultation](#)).

While psychological well-being was defined as the primary outcome for QoL, this was not consistently reported across studies assessing QoL as an outcome measure. Given that we used general QoL as a proxy measure where psychological well-being was not available, it is important to consider that QoL in this review reflects a combination of both overall QoL and psychological well-being measures.

Relevance of findings: patient and public involvement from carer Review Advisory Panel and stakeholder consultation

Carer RAP members felt the quantitative factors investigated had relevance to them, although they suggested amendments in the presentation of some factors. For instance, the coping literature often refers to 'maladaptive' versus more adaptive coping. However, what is termed 'maladaptive' may be the most constructive way for a carer to cope at a given time. Members therefore suggested that coping strategies should be presented simply as having positive or negative impact on carer mental health and with less value-laden labels attached. The RAP also highlighted the incongruence between how important the members considered sufficient finances to be in protecting carer mental health with the dearth of research studies found addressing this factor. Furthermore, the carer RAP highlighted the absence of evidence relating to carer self-identification⁹² as a gap in the observational review evidence. In addition to the themes emerging from the qualitative synthesis and carer RAP work, the observational quantitative synthesis also identified contextual factors that may relate to carer mental health. The RAP agreed that these were an important additional consideration, which would warrant further focus.

Feedback from our wider stakeholder consultation with additional carers, a patient, practitioners, commissioners and policy-makers confirmed that findings within all the resulting themes were considered relevant and informative in understanding the carer experience and what may help in supporting carers. More detailed involvement of the carer RAP and wider stakeholder consultation, along with the lessons learnt, is reported elsewhere (see <https://arc-gm.nihr.ac.uk/carers-project/>).

Implications and future research

Improved models, designs and definitions

In order to move the field of carer mental health research meaningfully forward, the development of clear and comprehensive explanatory models and frameworks are needed, to guide enquiry and develop testable theories to investigate the relationships between caregiving factors as primary stressors, mediators and moderators in relation to carer psychological morbidity. Future research requires more hypothesis-driven longitudinal and larger study designs incorporating quantitative causal analyses to provide a more comprehensive understanding of the range and interaction of factors affecting the mental health of EOL caregivers.

In addition, improvements in how factors within studies are defined/described and the reporting of a priori hypotheses of why factors have been investigated is required. This will prove useful in obtaining a more detailed understanding of the factors investigated and how they relate to carer mental health.

Knowledge gaps

Given the paucity of research studies focused on relationships, finances and different cultural perspectives, further studies exploring the impact of these factors on carer mental health are needed in order to further understand how these factors contribute to carer psychological morbidity within a broader context, for example using political theory.

Comprehensive strategy for carer support

The fact that a range of factors are implicated in carer mental health means that we are likely to need a comprehensive, co-ordinated strategy to improve the mental health of UK EOL carers that encompasses several factors, rather than focusing on one or two. The range of factors captured in this synthesis can help inform such a comprehensive strategy. They may inform decisions about legislation, allocation and distribution of funding and the fiscal incentives to control quantity and quality of services among policy-makers and commissioners, for example to improve work and benefits legislation, boost respite provision or mandate provision for carers within services. Overall findings may guide services in the design of operational procedures to enable more effective carer and patient support through earlier, targeted

carer intervention to prevent later crises, in particular through knowledge of carer protective and risk factors for psychological morbidity. For example, it may help to identify those carers at higher risk and to work with protective factors that build carer resilience and strength to help those at risk within existing resources. Review findings may also help carers identify options for self-help to boost protective factors, and support carer organisations in terms of where to focus their resources and advocacy.

In general, we need to recognise carers as a vital resource and provide better cross-society initiatives to support carers and prevent adverse health outcomes from caregiving.

Dissemination

Project findings have been reported to carer RAP members and stakeholder groups throughout the project through meetings, workshops and focus groups. All components of the project will be written up as NIHR HSDR peer-reviewed publications. The project has been presented at the European Association for Palliative Care Congress 2021. In response to stakeholder recommendations, project findings are disseminated via posters and leaflets, podcasts, webinars and a website. Awareness of the findings will be raised via Twitter and through stakeholder networks of NIHR ARC Greater Manchester and co-applicants. The current report, reports for additional project components and all project materials will be available through the project website: <https://arc-gm.nihr.ac.uk/carers-project->.

Conclusions

It is clear that factors associated with carers' mental health during EOL caregiving are wide-ranging and we cannot focus on one single factor to reduce psychological morbidity among family carers who care for patients at the EOL, whether it be patient condition, impact of caring responsibilities, relationships, finances, carer internal processes, support or contextual factors. We therefore need a comprehensive rather than a narrow approach to improving carers' mental health.

The literature on this topic is very diverse and difficult to summarise, and the field would benefit from a clearer direction of enquiry guided by explanatory models and frameworks. The impact of quality of relationships and finance warrant further investigation.

Factors need to be better defined and it needs to be better established, through quantitative causal analyses, why they should relate to mental health. More longitudinal research is required to help understand the likely causal direction of associations.

Working throughout the research project alongside a carer RAP was important as RAP members were able to act as a lens to validate, present and interpret research findings from the quantitative observational review.

Synthesis of the existing evidence on factors associated with carers' mental health during EOL caregiving provides a comprehensive understanding of factors affecting psychological morbidity of EOL carers. It is anticipated that the findings from this review will inform the development of future initiatives and interventions to improve the mental health of EOL carers and lead to better targeting of carers at risk of poor mental health.

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Author contributions

Gunn Grande (<https://orcid.org/0000-0003-2200-1680>) developed the project protocol; supported the design of the search strategy; supported the search and selection of the literature; supported the analysis and synthesis of the observational review data and drafted the manuscript.

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All authors had access to all study data, contributed to the analysis plan, discussed the interpretation of findings, provided critical revision of the manuscript and take responsibility for data integrity and analysis.

Ethics statement/approval(s)

This is a report of an evidence synthesis of existing literature, and no research data were collected for the project. Consultation with the University of Manchester Research Practice Governance Office established that the project did not require ethics approval.

Information governance

There were no personal data involved in the production of this report.

Data sharing

All data for this report and the project as a whole are accessible via our website: <https://arc-gm.nihr.ac.uk/carer-project->. As this is an evidence synthesis project, data are mainly contained within the referenced literature, tables and appendices of our reports. For more information, please contact the corresponding author.

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Disclaimer

The views expressed are those of the authors and not necessarily those of the NHS, the NIHR or the Department of Health and Social Care.

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Appendix 1 Search strategy

Caregiver

MESH terms related to caregiver:

MEDLINE (Ovid online):

- *Caregivers*

EMBASE (Ovid):

- *Caregiver*
- *Caregiver burden*
- *Caregiver burnout*
- *Caregiver strain Index*
- *Caregiver support*

PsychINFO (Ovid Online):

- *Caregivers*
- *Caregiver burden*

CINAHL Plus (EBSCO)

- *Caregiver burden*
- *Caregiver attitudes*
- *Caregiver support*

Social Sciences Citation Index

(Institute for Scientific Information; Clarivate Analytics)

TOPIC

- *Informal caregivers*
- *Family caregivers*

Cochrane Central Register of Controlled Trials (CENTRAL)

- *Caregivers*

Database of Abstracts of Reviews of Effects

(DARE; University of York Centre for Reviews and Dissemination)

- *Caregivers*

Cochrane Qualitative Reviews

- *Caregivers*

Palliative Care

MESH terms related to palliative care:

MEDLINE (Ovid Online):

- *Palliative care*
- *Hospice and palliative care nursing*
- *Terminal care*
- *Terminally ill*
- *Hospice care*
- *Hospice*

EMBASE (Ovid):

- *Cancer palliative therapy*
- *Palliative nursing*
- *Palliative therapy*
- *Terminal care*
- *Terminally ill Patient*
- *Terminal disease*
- *Hospice*
- *Hospice care*
- *Hospice nursing*

- Use MESH term for carer where database allows. For example, 'caregiver' in MEDLINE. Avoid 'home nursing' as a MESH term as this will incorporate healthcare workers.

- Search for additional string carer terms as both a key word and within ti,ab.
 - *family care giv**; *family caregiv**
 - *informal caregiv**; *informal care giv**
 - *family care* or informal care**

- Combine: (MESH term) OR (additional string carer terms)

Rationale:

- Incorporates use of MESH term.
- Looks to capture additional relevant literature on carers not indexed under the database MESH term. Using MEDLINE as the test database, a number of different terms for carer were searched to determine the most relevant terms for capturing additional literature not included within the MESH term 'caregiver'. Using 'family caregiver' as a key word and 'informal caregiver' in a title and abstract search were shown to include two additional relevant references while 'family carer(s)' and 'informal carer(s)' are terms often used in the literature to represent carers.
- By combining appropriate MESH terms for carer along with additional string search terms, the risk of missing papers not captured by the MESH terms is reduced.

<p>PsychINFO (Ovid Online):</p> <ul style="list-style-type: none"> • Palliative care • Terminally ill patients • Hospice <p>CINAHL Plus (EBSCO)</p> <ul style="list-style-type: none"> • Palliative care • Hospice and palliative nursing • Terminal care • Hospice care <p>Social Sciences Citation Index (Institute for Scientific Information; Clarivate Analytics)</p>	<p>Rationale:</p> <ul style="list-style-type: none"> • Incorporates search terms used by Flemming <i>et al.</i> (2019) and MEDLINE MESH search terms used in Candy <i>et al.</i> (2011) systematic reviews.* • Looks to capture additional relevant literature on palliative care not indexed under palliative care as a MESH term. • Each included MESH term has been tested using MEDLINE as a test database to confirm the retrieval of additional relevant papers which would not have been captured by Palliative Care MESH term only, • 'end-of-life' and 'end of life' have previously been tested using MEDLINE as a test database to confirm the retrieval of additional relevant papers which would not have been captured by any of the MESH terms above.
<p>TOPIC:</p> <ul style="list-style-type: none"> • Palliative care • Palliative care nursing/hospice and palliative care nursing • Terminal care • Terminally ill • Hospice care • Hospice 	<p>* Candy B, Jones L, Drake R, Leurent B, King M. Interventions for supporting informal caregivers of patients in the terminal phase of a disease. <i>Cochrane Database Syst Rev</i> 2011; Issue 6, Art No. CD007617. ISSN 1469-493X. https://doi.org/10.1002/14651858.CD007617.pub2</p> <p>Flemming K, Atkin K, Ward C, Watt I. Adult family carers' perceptions of their educational needs when providing end-of-life care: a systematic review of qualitative research [version 1; peer review: 3 approved with reservations]. <i>AMRC Open Res</i> 2019;1:2. https://doi.org/10.12688/amrcopenres.12855.1</p>
<p>Cochrane Central Register of Controlled Trials (CENTRAL)</p> <ul style="list-style-type: none"> • Palliative care • Hospice and palliative care nursing • Terminally ill • Terminal care • Hospice care • Hospices 	
<p>Database of Abstracts of Reviews of Effects (DARE; University of York Centre for Reviews and Dissemination)</p> <ul style="list-style-type: none"> • Palliative care • Hospice and palliative care nursing • Terminally ill • Terminal care • Hospice care • Hospices 	
<p>Cochrane Qualitative Reviews</p> <ul style="list-style-type: none"> • Palliative care • Hospice and palliative care nursing • Hospices • Hospice care • Terminal care • Terminally ill 	
<p>AND</p> <ul style="list-style-type: none"> • 'Qualitative Research' as MESH or 'Qualitative' in Title, abstract or keyword search 	
<p>Databases</p>	<p>MEDLINE, CINAHL Plus; PsychINFO; Social Sciences Citation Index; EMBASE; Cochrane Central Register of Controlled Trials (CENTRAL); Database of Abstracts of Reviews of Effects (DARE); Cochrane Qualitative Reviews.</p>
<p>Year</p>	<p>2009–19</p>

Appendix 2 Newcastle–Ottawa Scale (NOS) adapted Quality Assessment criteria

Depending on the type of study, studies were assessed using either the cohort/longitudinal study criteria or cross-sectional criteria listed below. The origin of each criterion, which relates to the version of NOS the criteria was adapted from, is also included for each criterion.

Scoring system: a maximum score of 1 was available for each criterion, with the exception of comparability criterion, where a maximum score of 2 was possible. Where a starred condition (*) within the criterion was met, the criterion was awarded 1 point, with the exception of comparability criterion, where 1 point was available for each starred (*) condition. This resulted in a maximum score for quality assessment of cohort or longitudinal studies = 12 and a maximum score for quality assessment of cross-sectional studies = 10.

Cohort or longitudinal study	Cross-sectional study	Origin
Selection		
<u>Representativeness of the sample/exposed cohort:</u> (1) Truly representative of carers of patients at end of life* (all subjects or random sampling). (2) Somewhat representative of carers of patients at end of life* (non-random sampling). (3) Selected group of users (e.g. convenience sampling; not sampling of all carers fitting criteria). (4) No description of the sampling strategy.	<u>Representativeness of the sample:</u> (1) Truly representative of carers of patients at end of life* (all subjects or random sampling). (2) Somewhat representative of carers of patients at end of life* (non-random sampling). (3) Selected group of users (e.g. convenience sampling; not sampling of all carers fitting criteria). (4) No description of the sampling strategy.	cross-sectional/cohort
<u>Selection of exposed and control cohort</u> (1) drawn from the same community* (2) drawn from different sources (3) not clear	<u>Selection of the sample</u> (1) drawn from the same community* (2) drawn from different sources (e.g. mixture of hospital and home/outpatient) – where distribution is likely to be unrelated to the predictor/s* (3) drawn from different sources (e.g. mixture of hospital and home/outpatient) – where distribution is likely to be related to the predictor/s (4) not clear	cohort
<u>Non-respondents</u> (1) Comparability between respondents' and non-respondents' characteristics is established, and the response rate is satisfactory.* (2) The response rate is unsatisfactory, or the comparability between respondents and non-respondents is unsatisfactory. (3) No description of the response rate or the characteristics of the responders and the non-responders.	<u>Non-respondents</u> (1) Comparability between respondents' and non-respondents' characteristics is established, and the response rate is satisfactory.* (2) The response rate is unsatisfactory, or the comparability between respondents and non-respondents is unsatisfactory. (3) No description of the response rate or the characteristics of the responders and the non-responders.	cross-sectional
<u>Sampling frame</u> (1) The analytical (final) frame is $\geq 50\%$ of the initial sampling frame* (2) The analytical (final) frame is $< 50\%$ of the initial sampling frame (3) The initial sampling frame is not reported	<u>Sampling frame</u> (1) The analytical (final) frame is $\geq 50\%$ of the initial sampling frame* (2) The analytical (final) frame is $< 50\%$ of the initial sampling frame (3) The initial sampling frame is not reported	

Cohort or longitudinal study	Cross-sectional study	Origin
<u>Ascertainment of the predictor(s) ('exposure'/'risk factor'):</u> (1) Validated measurement tool (cross-sectional); secure record (e.g. surgical records), structured interview (cohort).* (2) Non-validated measurement tool, but the tool is available or described (cross-sectional); written self-report (cohort). (3) No description (of the measurement tool) (cross-sectional/cohort).	<u>Ascertainment of the predictors ('exposure'/'risk factor'):</u> (1) Validated measurement tool (cross-sectional); secure record (e.g. surgical records), structured interview (cohort).* (2) Non-validated measurement tool, but the tool is available or described (cross-sectional); written self-report (cohort). (3) No description (of the measurement tool) (cross-sectional/cohort).	cross-sectional/cohort
<u>Demonstration that predictors preceded outcome and/or that baseline measurement of outcome variable was taken into account</u> (1) yes* (2) no		cohort
Comparability		
<u>Evidence of attempting to control for other/confounding factors in the analysis of relationship between predictor and outcome</u> (1) The study controls for the most important factor (select one).* (2) The study control for any additional factor.*	<u>Evidence of attempting to control for other/confounding factors in the analysis of relationship between predictor and outcome</u> (1) The study controls for the most important factor (select one).* (2) The study control for any additional factor.*	cross-sectional
Outcome		
<u>Was there an a priori hypothesis/hypotheses or was the study exploratory?</u> (1) A priori hypothesis* (2) Exploratory	<u>Was there an a priori hypothesis or was the study purely exploratory?</u> (1) A priori hypothesis* (2) Exploratory	
<u>Assessment of the outcome(s):</u> (1) Independent assessment* (2) Clinical interview* (3) Record linkage* (4) Validated measurement tool* (5) Self-report (6) No description	<u>Assessment of the outcome(s):</u> (1) Independent assessment* (2) Clinical interview* (3) Record linkage* (4) Validated measurement tool* (5) Self-report (6) No description	cross-sectional/cohort
<u>Adequacy of follow-up of cohorts</u> (1) Complete follow-up – all subjects accounted for* (2) Subjects lost to follow-up unlikely to introduce bias – small number lost – > ____ % (select an adequate %) follow-up, or description provided of those lost)* (3) Retrospective cohort so not applicable* (4) Follow-up rate < ____% (select an adequate %) and no description of those lost (5) No statement		cohort
<u>Statistical test:</u> (1) The statistical test used to analyse the data is clearly described and appropriate, and the measurement of the association is presented, including CIs and the probability level (p-value).* (2) The statistical test is not appropriate, not described or incomplete.	<u>Statistical test:</u> (1) The statistical test used to analyse the data is clearly described and appropriate, and the measurement of the association is presented, including CIs and the probability level (p-value).* (2) The statistical test is not appropriate, not described or incomplete.	cross-sectional