

Predictors of Informal Caregiver Burden in Parkinson's: A Systematic Review

Abstract

Background: Caregivers of people with Parkinson's disease are at risk of experiencing *caregiver burden*. Understanding contributing factors is required to develop appropriate targeted interventions and support for this group. This systematic review provides an updated appraisal and synthesis of quantitative studies assessing predictors of burden among informal caregivers of people with Parkinson's.

Method: Five electronic databases (APA PsycINFO, CINAHL, MEDLINE, Web of Science, and Cochrane Library) were systematically searched (from inception until July 2024), supplemented by hand-searches. Study quality was assessed using the cross-sectional JBI Critical Appraisal Checklist. Results were synthesised narratively.

Results: Forty-one studies were included. Predictors of increased burden included greater impact of motor symptoms on ADLs, greater severity of neuropsychiatric symptoms, poorer quality of life of the person with Parkinson's, and poorer caregiver mental health. Demographics, presence of motor symptoms, motor complications, and general cognitive function did not predict burden. Evidence was inconclusive for several variables including disease stage and duration, motor symptom severity, functional ability, overall non-motor symptoms, mental health of the person with Parkinson's, and caregivers' involvement and protective factors.

Conclusion: Several areas for potential future intervention are indicated, although methodological weaknesses within the literature constrain the robustness of conclusions. Key areas for future research include exploring understudied variables (caregiver personality and coping style, relationship quality, and positive aspects of caregiving) that may be important predictors of burden, specifying and utilising a more consistent definition of "informal caregiver", and recruiting younger and non-spousal caregivers and more diverse samples regarding disease severity.

Key words: Parkinson disease, caregivers, caregiver burden, systematic review

Introduction

Parkinson's is the second most common neurodegenerative condition after Alzheimer's disease,¹ currently affecting an estimated 9.4 million people worldwide.² It is characterised by motor symptoms including bradykinesia, rigidity, rest tremor, and postural instability,³ which can be used to differentiate Parkinson's from other parkinsonian disorders.⁴ Other features of Parkinson's can include sleep problems, psychological difficulties (e.g., depression, anxiety, apathy, and psychosis), impulsivity, cognitive impairment and dementia, and autonomic dysfunction.⁵

As Parkinson's progresses, individuals typically require increasing support and assistance with daily tasks,⁶ including medication management, activities of daily living (ADL), personal safety, care co-ordination, and social activities.^{7,8} Since most people with Parkinson's (PwP) live in the community,⁹ informal caregivers (people who are not financially compensated for providing care, typically spouses/partners or other family members) are usually main providers of their care.¹⁰ This can become challenging as the disease progresses, especially given a lack of services¹¹ and/or perceived lack of co-ordinated support from healthcare services.¹²

Caring for a PwP can affect various aspects of a caregiver's life.^{12,13} This is often termed *caregiver burden* (CB),⁹ although other terms are also used to describe similar phenomena, such as *stress*,¹⁴ *distress*,¹⁵ or *strain*.¹⁶ The concept of CB has been operationalised in various ways (for reviews see Chou¹⁷ and Liu et al¹⁸), most often as a broad, multidimensional construct describing "the extent to which caregivers perceived their emotional or physical health, social life, and financial status as suffering as a result of caring for their relative".^{19(p261)}

Regarding terminology: although commonly-used, "caregiver" and "caregiver burden" are contentious terms, argued to be culturally-biased,^{20,21} negatively-valenced, disregarding of the caring and reciprocal component of relationships,²²⁻²⁵ and bureaucratising of a common human experience.^{26,27} Similarly, "caregiver burden" risks engendering a blaming or dismissive perspective towards the "cared-for",²⁸⁻³⁰ potentially (unintentionally) prioritising caregivers' needs.³¹ While the term "carer" made some headway in promoting recognition of care work,³² it carries inherent biases,²¹ and using relationship-focused labels is generally preferred by the described populations.²⁴ We use "caregiver" and "caregiver burden" cautiously in this review given their ongoing ubiquity, in the absence of a commonly-accepted alternative.

Caregivers in the Context of Parkinson's

Although caregivers of people with other chronic conditions experience some similarities in burden to those supporting a PwP,⁹ the latter brings unique challenges due to its progressive nature,

myriad of motor symptoms and other difficulties, and frequently fluctuating symptoms.^{33,34} Parkinson's motor symptoms usually develop between 65 and 70 years,³⁵ coinciding with the typical time of transitioning to retirement, and causing disruption and loss of long held future plans for the affected person and family.⁸ Furthermore, as the disease progresses, informal caregivers may experience changed roles within relationships, anxiety about the future, grief, frustration, guilt, reduced involvement in work and social activities, financial pressures (predominantly through loss of income), and struggle with access to services and/or poorly co-ordinated healthcare.^{8,11,12,36,37} These circumstances can negatively impact on caregivers' physical, psychological, and social wellbeing^{13,38-43} and reduce their quality of life (QoL).^{44,45}

Ensuring caregiver wellbeing is paramount in supporting PwP. High CB can lead to burnout,⁴⁶ producing an "invisible patient" in the patient-doctor-caregiver system, and reducing a caregiver's ability to continue providing effective care.⁸ This, in turn, adversely affects health outcomes of PwP.⁴⁷ Furthermore, although many PwP wish to live at home for as long as possible, CB is a risk factor for the PwP entering a care home.^{36,48,49} Informal caregivers also contribute substantially to society; the economic value of their care and support is greater than ever and without them, health and social care systems would collapse.⁵⁰ Recently, the COVID-19 pandemic has exacerbated rates of CB in Parkinson's through loss of professional and family support and social isolation.^{51,52} Accordingly, understanding factors that contribute to burden among informal caregivers of PwP is crucial to developing targeted supportive interventions focused on preventing or reducing their experienced burden and supporting them to maintain wellbeing and effectiveness in their role.

Previous Reviews

Numerous studies have investigated factors associated with CB in Parkinson's. Six quantitative reviews have provided overviews of this literature.^{8,15,53-56} In 2015, Greenwell et al⁵⁴ synthesised and evaluated evidence on predictors of psychosocial outcomes (including burden) in Parkinson's caregivers. Since 2015, a substantial number of relevant articles have been published, so a fresh review is timely. Other reviews in the area have either focused on a specific method (longitudinal studies and only searching in PubMed)⁵⁵ or a specific short time period (2017-2022).⁵³ Consequently, an updated comprehensive synthesis of literature regarding factors related to CB for those supporting a PwP is necessary.

Review Aims

Accordingly, this systematic review provides an updated appraisal and synthesis of evidence around factors associated with CB among informal caregivers of PwP. The focus is on predictors

(rather than correlates) of informal CB, examining the direction of the relationship between variables, and identifying implications of clinical relevance for potential supportive interventions.

Method

Search Strategy

The electronic databases APA PsycINFO, CINAHL Plus with Full Text, MEDLINE, Web of Science Core Collection, and Cochrane Library were systematically searched on 25th July 2023 using a combination of subject headings and search terms relating to “Parkinson’s disease”, “burden”, and “caregiver” (Table 1), which were combined using the Boolean operators “OR” and “AND”. The search strategy was developed based on a prior scoping search and in consultation with a specialist university librarian. Searches were limited to peer-reviewed articles, written in English, and where possible a restriction was added to only retrieve papers with human participants. No limits were placed on year of publication. Additionally, reference lists of relevant review articles returned from the initial search and articles meeting the inclusion eligibility criteria were manually searched. An updated search was conducted on 23rd July 2024, limited to articles published since 1st January 2023.

insert Table 1 here

Inclusion and Exclusion Criteria

Inclusion and exclusion criteria for studies are summarised in Table 2. Findings were only included from multivariable analyses i.e., statistical models that have multiple independent or predictor variables and a single dependent or outcome variable.⁵⁷ Bivariate statistical analyses are argued to be inadequate for investigating complex constructs such as CB,⁵⁸ with multivariable statistics offering a better understanding of the unique contributions of many variables that may influence burden.^{59,60}

insert Table 2 here

Study Selection

The initial search identified 4,277 records after limits were applied. Articles were imported into EndNote reference management software and 1,867 duplicates removed. Titles and abstracts of the remaining 2,410 articles were screened for relevance. After this, 109 articles were retrieved for full-text screening against the eligibility criteria, and 38 selected for inclusion. Two further articles were identified through handsearching reference lists of relevant review articles returned from the

initial search. The online platform Rayyan⁶¹ was used to support the screening process and record decision-making. Figure 1 depicts the systematic search PRISMA diagram.⁶²

Figure 1. PRISMA Flow Diagram of the Systematic Search Procedure

insert Figure 1 here

Data Extraction and Quality Appraisal

A data extraction tool was developed to extract information relevant to the review. Extracted data included author, publication year, country of study, aim, main definitions, participant characteristics, recruitment methods, study design, outcome measures, statistical analysis, and key findings.

Since all included studies utilised a cross-sectional design, methodological quality appraisal was undertaken using the JBI Critical Appraisal Checklist for Analytical Cross-Sectional Studies.⁶³ The tool requires reviewers to indicate whether a study meets eight criteria by selecting *Yes*, *No*, *Unclear*, or *Not applicable* for each of eight questions. The first author independently assessed quality of selected studies using the checklist, with ratings for 10% then cross-checked by another author to ensure reliable quality appraisal. Any differences of opinion were resolved via discussion. The checklist does not yield a score to indicate methodological quality; indeed, it has been argued a pure numerical outcome does not best reflect methodological quality and can mask significant issues in specific areas.⁶⁴⁻⁶⁶ Following JBI guidelines,⁶³ the tool was not used to exclude articles based on low methodological quality. Instead, appraisal results were used to highlight quality of the current evidence base and inform interpretation of findings.

Data Synthesis

Due to heterogeneity of the studies with a wide range of outcome measures used to assess potential predictors and CB, it was not appropriate to conduct a meta-analysis. Instead, a narrative approach⁶⁷ was used to synthesise evidence regarding caregiver and PwP-related predictors of burden in informal caregivers of PwP.

Results

Characteristics of Included Studies

Forty-one full-text articles were identified for inclusion (key features summarised in Table 3). They were published between 1997 and 2023, the majority (35 articles) since 2012. Studies were conducted in Germany ($n=5$), India ($n=5$), the UK ($n=5$), Italy ($n=3$), South Korea ($n=3$), the US ($n=3$), Australia ($n=2$), Brazil ($n=2$), Spain ($n=2$), Sweden ($n=2$), Malaysia ($n=1$), Mexico ($n=1$), Netherlands

($n=1$), Poland ($n=1$), Thailand ($n=1$), and Turkey ($n=1$). Three multinational studies were conducted across the US and Canada,⁶⁸ the US and Japan,⁶⁹ and the UK, France, Germany, Netherlands, Portugal, and Sweden.⁷⁰ All studies used a cross-sectional design. Thirty-two were designed as single studies and nine were sub-studies that analysed baseline data.^{47,68,70-76} Two studies shared a database; however, both were included as they contained separate statistical analyses.^{77,78}

Samples ranged from 20 to 2,476 caregivers, with most studies ($n=35$) having fewer than 200 participants. Not all studies included a definition of “informal caregiver” ($n=22$). Across those that did, the definition varied. Five studies applied the definition devised by Martinez-Martin et al^{79(p925)}: “any person who, without being a professional or belonging to a social support network, usually lives with the patient and, in some way, is directly implicated in the patient’s care or is directly affected by the patient’s health problem”.^{74,80-83} Other studies referred to an unpaid person who assists with daily activities or provides emotional support. Most studies did not specify a minimum time caregivers had to spend with the PwP; for those that did, requirements varied. Among studies that reported the caregiver’s relationship with the PwP ($n=34$), spouses/partners dominated the sample (47-100%), with six studies recruiting spouses/partners only.^{69,72,75,84-86} Other relationships included children, children-in-law, siblings, parents, other relatives, friends, and neighbours. Among studies reporting caregiver gender ($n=32$), all but two^{73,87} of the caregiver samples were predominately female (50-88%). Thirty-four studies reported caregivers’ mean age (43.7-70.7 years).

Seven validated measures of CB were used, including six non-disease specific measures: Zarit Burden Interview ($n=25$),⁸⁸ Caregiver Burden Inventory ($n=6$),⁸⁹ Caregiver’s Burden Scale ($n=2$),⁹⁰ Caregiver Strain Index ($n=2$),⁹¹ Feeling of Burden questionnaire ($n=1$),⁹² and Multidimensional Caregiver Strain Index ($n=1$),⁹³ and one disease-specific measure: Parkinson’s Disease Caregiver Burden questionnaire ($n=5$).⁹⁴

All studies used regression to analyse relationships between CB (the specified dependent or outcome variable) and other variables.

insert Table 3 here

Methodological Quality Assessment

Results of the quality assessment are summarised in Table S1. The main methodological shortcomings across included studies related to statistical analysis. While appropriate methodologies were used (i.e., regression), these analyses were under-specified and/or key data (e.g., type of regression, whether assumptions were fully checked and met, and which variables were included in the regression model) were missing in all but one of the studies,⁷⁶ undermining the robustness of

the approach and convincingness of findings. Fifteen studies used stepwise methods of regression, which is problematic as numerous issues have been identified with these methods (see Field¹¹⁶ and Harrell¹¹⁷ for discussion). Only one study justified use of stepwise methods, citing the exploratory aims of the study⁷⁶; however, given existing theoretical literature is available, it remains questionable whether stepwise regressions were the most appropriate approach to use. Furthermore, it was not uncommon across the reviewed studies for variables to be included in multivariable analysis based on their significance in bivariate analysis. This type of variable selection has been deemed inappropriate because it can result in wrongly rejecting potentially important variables that are only significant after controlling for other variables.¹¹⁸ These issues risk development of inaccurate results.

Most studies used valid and reliable tools to measure potential predictors ($n=33$) and CB ($n=39$), sufficiently to limit any potential measurement bias. However, problems were identified relating to the psychometric adequacy of outcome measures used in several studies. In three studies, researchers modified some of the outcome measures used but these adapted versions were not described as validated before use.^{73,85,95} Additionally, reliability and validity of outcome measures used by Martinez-Martin et al⁸⁰ and Sanyal et al¹⁰⁹ had not been clearly established for the language in which they were administered. Two studies used outcome measures originally developed for use in relation to Alzheimer's disease^{68,110}; it is unclear how these were made appropriate for use in relation to Parkinson's. Furthermore, Tanji et al⁶⁹ used different depression measures in their Japanese and US samples, introducing a source of potential bias. Thirty-three studies did not identify important potential confounds; however, strategies to deal with confounding variables were frequently used, given studies used a form of multiple regression analysis.

Inclusion and exclusion criteria for samples were not clearly defined in 15 studies, limiting conclusions about generalisability. Furthermore, 27 studies did not use both specified diagnostic criteria for Parkinson's and a definition of "informal caregiver" to determine eligibility, increasing the risk of bias. Eighteen studies were appraised as not providing sufficient detail about the study sample and setting. For example, demographic information (i.e., caregiver age, gender, or relationship to the PwP), important for understanding the sample and making decisions about generalisability of the results, was missing in 17 studies and one study did not explicitly state the setting, although this could be assumed from other information provided in the article.

Predictors of Caregiver Burden

Evidence for predictors is reported descriptively due to the diversity of measures used across the studies. Factors predicting CB were grouped into caregiver and PwP-related factors, and

thematically categorised. Due to the large number of included studies, individual study results are presented in Table 4 and key findings in relation to the most investigated predictors are described below.

Caregiver-Related Factors

Demographics. Eight studies found caregiver age did not predict CB.^{47,68,69,75,80,81,104,113} Caregiver gender was consistently found unrelated to CB in five studies investigating this relationship.^{47,75,80,84,113} Level^{47,75} and years¹¹³ of caregivers' education did not predict CB.

Caregiver Involvement. Seven studies investigated the association between caregiver involvement and CB. Two studies found more years providing care predicted CB,^{81,102} but three other studies found no association.^{68,97,115} Frequency of providing care and whether care is provided only during the day, or during day and night, did not predict CB.⁴⁷ Hours per week providing care also did not predict CB,⁹⁷ and neither did hours per day according to Zhong et al¹¹⁵; however, Sanyal et al¹⁰⁹ reported hours per day did predict CB.

Psychological Factors. Nine studies explored the contribution of caregiver mental health to CB. Five studies found CB was positively predicted by caregiver depression.^{68,69,73,113,115} Shin et al¹¹² found between-group differences regarding caregiver-PwP relationship, reporting caregiver depression predicted burden in spouses supporting a partner with Parkinson's, but not for those supporting a parent. Two studies found caregiver anxiety positively predicted CB,^{68,81} but one study found no association.¹¹⁵ Johnson et al⁷¹ found the psychological wellbeing of caregivers positively predicted CB; however, Edwards and Scheetz⁸⁵ found the psychological wellbeing of caregivers did not contribute to burden.

Protective Factors. Reviewed studies explored the relevance of various protective factors including personality, coping styles, social support, and relationship quality to CB. Regarding caregiver personality, two studies reported higher CB was predicted by lower sense of coherence in caregivers.^{73,100} Caregivers' self-esteem and social competence did not predict CB.¹⁰⁰

Caregivers' coping styles were explored in three studies. CB was consistently positively predicted by use of problem-focused coping strategies.^{47,84,101} An avoidant coping style positively predicted CB in one study,⁴⁷ but not another.⁸⁴ Emotion-focused coping strategies did not predict CB.^{47,84}

Different types of social support were explored in seven studies. Two studies found higher perceived support from family, friends, and significant others was associated with reduced CB.^{47,85} Shin et al¹¹² assessed social support regarding degree of assistance caregivers received from private

relationships (e.g., family, friends, neighbours) and the community (e.g., organisations or experts). Community support negatively predicted CB in caregivers of a parent with Parkinson's, but private support did not predict burden for caregivers of either a spouse or a parent with Parkinson's. Carrilho et al⁹⁷ found whether the caregiver received help from others did not predict CB. Tanji et al⁶⁹ found receiving help negatively predicted burden in the main caregiver in a group of US-based caregivers, but not for caregivers living in Japan. In two other studies, CB in the main caregiver was negatively predicted by number of caregivers involved in supporting the PwP.^{95,99}

Factors linked to the caregiver-PwP relationship were investigated in four studies. General relationship satisfaction¹⁰¹ and mutuality⁷⁵ negatively predicted CB. However, no association was found between CB and relationship quality⁸⁴ and marital satisfaction.⁸⁵

Person with Parkinson's-Related Factors

Demographics. Eight studies examined PwP age; most found this did not predict CB,^{73,81,97,104,105,108,113} however one study reported a positive predictive effect of PwP age.⁹⁹ Six studies found PwP gender did not predict CB,^{73,81,103,105,108,113} but two studies reported male gender was associated with higher CB.^{70,74} Level^{47,75} and years¹¹³ of the PwP education did not predict CB.

Disease Factors. The relationship between disease stage and CB was explored by multiple studies, with mixed findings. While seven studies found advancing disease stage positively predicted CB,^{73,74,76,80,86,99,109} nine studies found no association.^{83,84,95,97,105,108,112,113,115}

Twelve studies explored the contribution of disease duration to CB. Eight studies did not identify disease duration as a significant predictor.^{47,78,83,97,103,104,108,115} However, three studies found disease duration positively predicted CB.^{80,99,100} Oh et al¹⁰⁵ found disease duration was a predictor when burden was measured using the Caregiver Burden Inventory but not the Zarit Burden Interview.

Motor Symptoms. The relationship between CB and motor symptoms as assessed by the original Unified Parkinson's Disease Rating Scale (UPDRS) and the Movement Disorder Society-sponsored revision of the scale (MDS-UPDRS) was extensively investigated in the studies. Eight studies utilised the UPDRS II or MDS-UPDRS II to assess motor symptoms. UPDRS II scores did not predict CB,^{95,112,113} whereas higher scores on the MDS-UPDRS II did contribute to greater CB.^{47,78,103,108,114}

Twelve studies utilised the UPDRS III and MDS-UPDRS III to assess motor symptom severity. Among studies using the former, six found motor symptom severity did not predict CB,^{68,75,83,104,111,113} two found it did,^{95,109} and Shin et al¹¹² concluded motor symptom severity positively predicted CB in

offspring but not spousal caregivers. Regarding the MDS-UPDRS III, two studies found scores positively predicted CB^{77,114} and one did not identify a relationship.¹⁰⁸

The UPDRS IV and the MDS-UPDRS IV were used by four and two studies respectively, assessing associations between motor complications and CB. No studies found these predicted CB.
83,95,108,112-114

Functional Status. Numerous studies reported on the effect of functional ability (assessed by various ADL scales) on CB. Findings were variable. Four studies reported greater dependence in ADLs predicted higher CB^{47,83,85,102}; five did not identify an association.^{69,81,95,112,113} Lo Monaco et al¹⁰⁴ found caregiver reports regarding the extent to which they provide help in ADLs, but not score on an ADL scale, predicted burden. CB was not predicted by instrumental ADLs¹⁰⁴ and overall functional status.⁶⁸ Hand et al¹⁰¹ found greater difficulty specifically with eating, bathing, and mealtime tasks predicted CB.

Non-Motor Symptoms. Seven studies used various overall measures of non-motor symptoms to investigate their relationship to CB. Three studies reported CB was positively predicted by non-motor symptoms,^{70,113,114} one concluded non-motor symptoms predicted burden in caregivers of a spouse but not a parent with Parkinson's,¹¹² and three found no association.^{75,83,108}

Eleven studies explored associations between neuropsychiatric symptoms and CB. Six studies found more severe neuropsychiatric symptoms overall predicted higher CB,^{70,71,86,99,102,105} while one study found no association.⁹⁵ Furthermore, Goel et al⁹⁹ demonstrated specific neuropsychiatric symptoms (delusions, hallucinations, agitation, depression, anxiety, elation, disinhibition, irritability, motor disturbance, nighttime behaviours, and appetite/eating) except apathy independently predicted CB. Conversely, Hand et al¹⁰¹ found motor disturbance was the only domain to positively predict CB. Two studies found greater severity of mood/apathy symptoms (but not impulse control disorders) predicted greater CB.^{80,98} Martinez-Martin et al⁸⁰ additionally found psychotic symptoms positively predicted CB.

Depression. Thirteen studies explored relationships between PwP depression and CB, and evidence was mixed. Four studies found PwP depression positively predicted CB.^{83,95,110,115} Tanji et al⁶⁹ concluded feelings of depression was a predictor in a US sample of caregivers. Leroi et al⁸⁷ found more severe PwP depression predicted higher CB only in individuals with Parkinson's with impulse control disorders, and not in those with apathy. Seven studies did not identify PwP depression as a predictor.^{47,68,76,78,81,105,111}

Anxiety. PwP anxiety was investigated in only five studies. One found higher PwP anxiety predicted greater CB,⁸² while four found no association.^{47,68,105,115}

Cognition. The impact of cognitive impairment on CB was extensively explored. Global cognitive function was assessed in 11 studies, all finding this did not predict CB.^{68,69,76,81,97,100,105,111-114} Furthermore, Karlstedt et al⁷⁵ reported caregiver perspectives on cognitive decline in the PwP did not predict burden.

Specific cognitive domains including memory (short-term and working memory)^{100,115} and executive functioning¹¹⁵ were also not found to predict CB. However, greater impairment in visuo-constructional skills was predictive of burden.¹¹⁵ Leroi et al⁸⁷ found attentional ability negatively predicted CB in PwP with apathy, but not with impulse control disorders. Oguh et al⁷⁴ found decreased verbal fluency predicted higher CB, while Golińska et al¹⁰⁰ reported better phonemic (but not semantic) verbal fluency predicted less CB.

Sleep. PwP sleep problems were sparsely explored. Two studies identified sleep disturbances positively predicted CB.^{81,95} Furthermore, Viwattanakulvanid et al⁸² found nocturnal akinesia and PwP feeling tired and sleepy upon awakening predicted higher CB.

Quality of Life. Macchi et al⁶⁸ found poorer QoL from the caregiver's perspective, but not the PwP, contributed to higher CB. Self-reported health-related QoL positively predicted CB in five studies,^{68,74,77,96,99} although this association was not found by Klietz et al¹⁰³ and Klietz et al.⁷⁸ In studies investigating specific aspects of health-related QoL, CB was positively predicted by only mobility,^{101,106} social support,¹⁰⁶ stigma,¹⁰⁷ and emotional wellbeing.¹⁰⁷

insert Table 4 here

Discussion

This review provides an updated appraisal and synthesis of quantitative research investigating caregiver and PwP-related predictors of CB in informal caregivers of PwP. Certain variables consistently predicted increased burden including greater impact of motor symptoms on daily living, greater severity of neuropsychiatric symptoms, poorer quality of life of the PwP, and poorer caregiver mental health. Evidence was inconclusive for many variables including disease stage and duration, motor symptom severity, functional ability, non-motor symptoms, mental health of the person with Parkinson's, and caregiver involvement and protective factors. However, demographics, presence of motor symptoms, motor complications, and general cognitive function were consistently not found to be predictors. Crucial points were also identified to improve robustness and consistency across the literature going forward, which may help define the contribution of inconclusive variables.

Across the identified studies, many failed to define “informal caregiver” or applied varying definitions, seven different CB measures were employed, and outcome measures examining predictors of experienced burden varied widely.

Discussion of Main Findings

Disease stage, disease duration, motor symptom severity, motor complications, and functional ability were among the most-explored variables. Of these, only motor complications were consistently non-predictive. Findings were mixed for the other variables. This contradicts Greenwell et al’s⁵⁴ review, which concluded disease stage and disease duration rarely predicted burden – though this review is now significantly outdated, and new evidence included here may have led to different conclusions. The less consistent findings in the current review may suggest the impact of these factors on burden is mediated or moderated by other variables, and that variables cannot effectively be considered in isolation.¹¹⁹ While impossible to tell from the information available in this review, it does indicate the importance of using theoretical models to inform study design and data analysis, and multivariable analysis, as this allows multiple factors to be taken into account and to consider confounds between variables.

Influences of motor symptoms on CB were also commonly explored. Outcomes of studies utilising the UPDRS II versus MDS-UPDRS II to explore their influences were strikingly different. Studies utilising the former found motor symptoms non-predictive of burden; studies using the latter identified it as predictive. This disparity may be explained by differences in the measures themselves. While there is general parallelism between the original UPDRS II and the MDS-UPDRS II, the original focuses on *presence* of motor symptoms whereas the MDS-UPDRS II focuses on their *impact* on ADLs.¹²⁰ It appears, therefore, only the latter influences burden, which stresses the importance of providing support for caregivers caring for PwP whose motor symptoms significantly impact their daily living. Interestingly, the same difference applies to the UPDRS I and MDS-UPDRS I used for assessing non-motor symptoms (the MDS-UPDRS I focuses on impact rather than presence of non-motor symptoms), although findings varied more across studies using these measures.

Most evidence indicated neuropsychiatric symptoms predicted higher burden. The only study that found no association did not use a disease-specific measure,⁹⁵ potentially explaining their differential finding. This contradicts Greenwell et al’s⁵⁴ review findings of limited, mixed evidence for this predictor. Again, the growth in literature has perhaps provided clarity about relationships between these variables. Regarding specific neuropsychiatric symptoms, evidence consistently implies global cognitive function does not predict burden, but specific cognitive deficits may contribute to burden. Further research is required to draw a more certain conclusion, given the

current limited evidence. Overall, these findings indicate CB could be prevented or alleviated by adequate management of neuropsychiatric symptoms. The contribution of PwP QoL to burden was little-explored by Greenwell et al⁵⁴; however, the current literature suggests poorer PwP QoL predicts higher CB.

Studies exploring contributions of caregiver involvement to burden achieved inconsistent results. Although the various ways this variable was conceptualised and measured could somewhat explain the mixed findings, there were also disparities across studies using the same measures. An alternative explanation for these inconsistencies is that effects of caregiver involvement on burden appear mediated by other variables, such as frequency of breaks and perceived social support.¹¹⁹ Furthermore, caregiver involvement was measured subjectively, based on caregivers' interpretations of which activities and roles constitute caregiving⁵⁴; this reflects wider disagreements over what formally constitutes "caregiving" as opposed to non-bureaucratized support of family.^{26,27} Cultural considerations add to this uncertainty, as many activities considered formalised caregiving in one culture might be regarded as typical family support in another.^{20,21}

Multiple studies found caregivers' psychological difficulties predicted increased burden. However, conceptual confounding is likely since emotional problems are an important aspect of burden¹⁹ and relevant questions are often included in CB measures (e.g., Caregiver Burden Inventory- "I feel emotionally drained due to caring for him/her").⁸⁹ Although in reviewed studies caregiver depression and anxiety were entered as predictors in statistical models, it is difficult to disambiguate the relationship between these variables. Nevertheless, results emphasise the importance of ensuring psychological support is available to caregivers.

Caregiver social support was frequently investigated and predicted burden in most studies. However, studies were again inconsistent regarding conceptualisations and measurement of social support, meaning findings should be interpreted cautiously. As for defining "caregiving", social support is multifaceted and includes varied formal and informal support. The type of social support found helpful differed according to the caregiver's relationship with the PwP, and the cultural background of the caregiver and PwP influenced whether social support was a predictor. Further research is required to identify the most effective ways to develop and maintain support (perhaps by subgroup). One systematic review⁵⁴ suggested perceptions, not actual support, is influential, but this was not confirmed here.

This review found factors contributing to perceived burden could vary between subsamples (e.g., people supporting a spouse versus parent, different cultural backgrounds, certain Parkinson's presentations, and caregiver versus PwP perspective on the variable). Although the evidence base is

limited regarding subsample analyses, these findings have important implications for tailoring approaches to reduce CB for people from different subgroups.

Limitations

There were methodological weaknesses in the studies that underpin this review. Caregiving is a dynamic process, but the cross-sectional included studies provide a more static picture and consequently limited insight into the trajectory of CB and its predictors over time.¹²¹ Cross-sectional designs also inhibit conclusions regarding causal relations.¹²² Furthermore, although multivariable analysis assumes cause-and-effect relationships between variables, some associations explored may be multifaceted. Included studies rarely developed predictive models referencing theoretical literature; instead, most variables investigated appeared to be chosen relatively arbitrarily or at least were not explicitly justified. While the use of validated outcome measures is considered a strength of the reviewed studies, it is also important to acknowledge these measures may be biased towards the research perspective and consequently may not capture all aspects of the patient or caregiver experience.

Regarding power, multivariable analysis requires large samples; however, many studies reported relatively small (and potentially underpowered – only three studies reported power calculations^{75,84,107}) samples, reducing their ability to detect relationships. Definitions of “informal caregiver” varied or were lacking, so there was likely between-study variation regarding type and level of care provided. Predictors of burden may differ depending on the caregiving role, but due to ambiguity in the study set regarding caregiver sample definitions, this cannot be determined for the current review. Samples also predominantly comprised older females caring for a spouse/partner, although this may reflect the older age of disease onset, that Parkinson’s disproportionately affects males, and that spouses/partners are common sources of support.

While it is acknowledged a limitation of the present review is that data on disease severity or stage was not extracted from included studies, generalisation of findings to the wider Parkinson’s population is questionable because most studies recruited from neurology outpatient clinics or similar settings. Samples were therefore likely biased towards caregivers supporting individuals earlier in the course of Parkinson’s, as later motor impairments present challenges for attending appointments or continuing to live in the community rather than institutionalised care.¹²³ Experiences of the identified samples may further differ from those in remote areas with restricted access to healthcare, or from those who choose not to access it. Any impact of Parkinson’s severity on CB would consequently be difficult to evaluate across populations less well-represented given this recruitment bias, yet research has shown those supporting people in the palliative stage of

Parkinson's may experience particular physical and emotional stress.¹² Although 21 countries were represented in this review, the vast majority were countries with predominantly WEIRD (Western, Educated, Industrial, Rich, Democratic) populations and so the evidence base provides a biased understanding of CB among those supporting PwP.

Limitations of this systematic review include that the literature search was restricted to articles published in English, and so the findings are subject to a language and cultural bias. Also, only full-text, peer-reviewed published studies were included and therefore evidence from grey literature might have been missed. Finally, it is acknowledged only one author assessed the methodological quality of included studies, which may have affected the reliability of the quality appraisal.

Future Directions

Future research should investigate caregiver-related variables including personality, coping style, and caregiver-PwP relationship, as these are understudied and are potentially modifiable through intervention. It will also be useful to explore perceived positive aspects of caregiving as these are sparsely investigated but may also be important predictors of CB and contribute to developing interventions to support caregivers. New research should specify their definition of "informal caregiver" and ideally employ a more consistent definition across studies. Future research needs to increase inclusion from younger caregivers and other relationship categorisations (e.g., parent-child, or between siblings), as well as recruit a more diverse sample regarding disease severity, to provide a more comprehensive understanding of experiences of all informal caregivers of PwP. Conducting subsample analyses would help to better understand whether factors predicting CB differ depending on demographic characteristics of caregivers, which could also facilitate development of an evidence base to inform tailored interventions for people from different subgroups. Furthermore, future studies would benefit from using objective and validated measures of caregiver involvement. They might also consider using disease-specific outcome measures to assess aspects of burden relevant to the specific population and develop an understanding of needs in this population for future interventions. Finally, new studies should ideally use longitudinal designs to explore the dynamic nature of CB in Parkinson's and its predictors, to evaluate causation between identified variables and burden – though this may be constrained by financial and practical difficulties.

Conclusion

This systematic review provides an up-to-date synthesis and critical evaluation of the evidence base around predictors of CB among caregivers of PwP. There were reasonably clear and well-supported findings for certain risk factors for CB that can inform future care and support, specifically: caregiver mental health, impact of motor symptoms on ADLs, neuropsychiatric symptoms, and PwP QoL. Key methodological weaknesses across the evidence base are also identified with recommendations for addressing them, offering routes to improved robustness and consistency in future relevant research. Findings from this review can contribute to development of targeted interventions to prevent or alleviate possible negative impacts of caregiving and potentially help identify caregivers particularly at risk of higher CB that could be prioritised in the context of under-resourced services.

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Table 1. Terms for Systematic Literature Search

Subject Headings	Title, Abstract, and Keyword Searches
Parkinson's disease	parkinson*
Caregiver Burden	burden or stress or distress or strain
Caregivers	carer* or caregiv* or "care giv*" or family or families or relatives or spous* or husband* or wife or wives or partner or partners

Table 2. Inclusion and Exclusion Criteria for Studies

Inclusion Criteria	Exclusion Criteria
1) The study's sample consisted only of informal (unpaid) caregivers of people with Parkinson's (at any disease stage), with any relationship to the person with Parkinson's (e.g., spouse/partner, sibling, friend), who had been caring for any duration.	1) The study's sample included formal (paid) caregivers of people with Parkinson's, or caregivers of other populations.
2) Main aim of the study was to identify caregiver or person with Parkinson's factors predicting caregiver burden among informal caregivers of people with Parkinson's through multivariable statistical modelling.	2) Mixed-sample studies (e.g., informal caregivers of people with different conditions grouped together, unless subsample analysis enabled discrimination of data regarding those supporting a person with Parkinson's).
3) The study reported a measure of burden among informal caregivers of people with Parkinson's using a total score on a validated caregiver burden scale.	3) The study only investigated correlations.
4) Caregiver and person with Parkinson's factors explicitly defined; self-reported constructs measured with a validated questionnaire or clearly-described question.	4) The study included predictors or outcomes measuring multiple constructs (e.g., using factor analysis; combining depression and anxiety into one overall predictor; combining burden with other constructs into one outcome measure).
5) Quantitative methodology, or mixed-methods with sufficient quantitative component for extraction of quantitative results.	5) Caregiver burden was an independent variable instead of the outcome.
6) Original empirical study in a peer-reviewed journal, full-text article available, in English (funds unavailable for translation).	6) Intervention study, qualitative study, case study, review, editorial, or scale validation.

Table 3. Study Characteristics and Key Findings

Authors (year); country	Study Characteristics			Caregiver Characteristics			Identified predictors of CB
	Caregiver sample size	Recruitment ^a	Statistical analysis	Spouse/partner of PwP (%)	Female (%)	Mean age (years)	
Agrawal et al (2012) ⁹⁵ ; India	91	Medical setting	Multiple regression	49.5	54.9	44.6	Caregiver: Number of caregivers: $\beta = -.311, p = .000$ PwP: Sleep disturbance: $\beta = .206, p = .025$ UPDRS III: $\beta = .255, p = .004$ BDI: $\beta = .352, p = .000$
Bartolomei et al. (2018) ⁹⁶ ; Italy	55	Medical setting	Multiple regression	85.5	NR	62	PwP: PDQ-39: $R^2 = .38, p < .001$
Caap-Ahlgren and Dehlin (2002) ⁷³ ; Sweden	65	Medical setting	Multiple regression	89.2	36.9	67	Caregiver: GDS: $B = 1.30, p = .014$ SOC: $B = -0.35, p = .004$ PwP: HY: $B = 4.67, p = .001$
Carod-Artal et al. (2013) ⁸¹ ; Brazil	50	Medical setting	Hierarchical multiple regression	78	88	55.7	Caregiver: Caregiving duration: $\beta = .38, p = .002$ HADS-Anxiety: $\beta = .34, p = .006$ PwP: SCOPA-Sleep: $\beta = .38, p = .01$ PPRS: $\beta = .29, p = .008$
Carrilho et al. (2018) ⁹⁷ ; Brazil	21	Medical setting	Multiple regression	47	80	53	No significant predictors
D'Amelio et al. (2009) ⁸⁶ ; Italy	40	Medical setting	Stepwise multiple regression	100	NR	63.6	PwP: HY: $\beta = .62, p < .001$ NPI-12: $\beta = .28, p = .03$

Edwards and Scheetz (2002) ⁸⁵ ; US	41	Medical setting; community	Stepwise multiple regression	100	68	66.8	Caregiver: PSS-Fa: $\beta = -.317, p = .037$ PwP: ADL-MS: $\beta = -.435, p = .005$
Eichel et al. (2022) ⁹⁸ ; Germany	84	Medical setting	Multiple regression	94	64.3	66.2	PwP: SEND-PD (Mood/Apathy): $\beta = .444, p = .001$
Geerlings et al. (2023) ⁴⁷ ; Netherlands	504	Medical setting	Multiple regression	90.7	66.9	67.6	Caregiver: MSPSS: $\beta = -.108, p < .001$ COPE-28 (Problem-focused): $\beta = .352, p < .001$ COPE-28 (Avoidant): $\beta = .973, p < .001$ PwP: Work status (working): $\beta = -.254, p < .05$ MDS-UPDRS II: $\beta = .251, p < .05$ SPDDS: $\beta = -.011, p < .05$
Goel et al. (2022) ⁹⁹ ; India	145	Medical setting	Multiple regression	54.8	NR	46.2	Caregiver: Number of caregivers: $\beta = -.506, p < .001$ PwP: Age: $\beta = .388, p < .001$ Disease duration: $\beta = .434, p < .001$ HY: $\beta = .711, p < .001$ PDQ-39: $\beta = .676, p < .001$ NPI-12: $\beta = .883, p < .001$ NPI-12 Domains - Delusion: $\beta = .533, p < .001$; Hallucinations: $\beta = .690, p < .001$; Agitation: $\beta = .493, p < .001$; Depression: $\beta = .209, p = .046$; Anxiety: $\beta = .332, p = .004$; Elation: $\beta = .255, p = .029$; Disinhibition: $\beta = .337, p = .004$; Irritability: $\beta = .318, p = .006$; Aberrant motor behaviour: $\beta = .288, p = .015$; Sleep disturbances: $\beta = .410, p < .001$; Appetite and eating disorders: $\beta = .394, p < .001$

Golińska et al. (2017) ¹⁰⁰ ; Poland	20	Community	Hierarchical multiple regression	NR	80	61.9	Caregiver: SOC-Manageability: $\beta = -.52, p = .01$ PwP: Disease duration: $\beta = .38, p = .02$ Verbal fluency (phonemic): $\beta = -.68, p = .01$
Hand et al. (2022) ¹⁰¹ ; UK	115	Medical setting	Stepwise multiple regression	85.2	66.1	70.7	Caregiver: Health: $B = 7.165, p < .001$ SCOPA-carer nighttime sleep: $B = 0.532, p = .022$ COPE-28 (Active): $B = 1.755, p = .004$ RAS: $B = -0.726, p < .001$ PwP: NPI-12 (Motor Behaviour): $B = 6.997, p = .036$ UPDRS-eating: $B = 3.447, p = .002$ PDQ-39 (Mobility): $B = 0.073, p = .036$ Bathing: $B = 6.676, p < .001$ Mealtimes: $B = 5.948, p = .002$
Johnson et al. (2023) ⁷¹ ; UK	45	Medical setting	Backward stepwise multiple regression	55.6	NR	NR	Caregiver: GHQ-28: $\beta = .425, p = .003$ PwP: NPI-12: $\beta = .319, p = .023$
Jose et al. (2021) ¹⁰² ; India	104	Medical setting	Multiple regression	51	76.9	NR	Caregiver: Caregiving duration: $\beta = .276, p = .001$ PwP: EASI: $\beta = .158, p = .038$ NPI-12: $\beta = .509, p = .001$
Kalampokini et al. (2022) ⁷⁰ ; UK, France, Germany, Netherlands, Portugal, Sweden	506	Medical setting	Stepwise multiple regression	71.2	NR	NR	PwP: NPI-12: $\beta = .331, p < .001$ Gender (male): $\beta = .172, p < .001$ Residential status (home): $\beta = .178, p < .001$ NMSS: $\beta = .176, p < .001$

Karlstedt et al. (2017) ⁷⁵ ; Sweden	51	Longitudinal study database	Multiple regression	100	56.9	70.7	Caregiver: MS: $\beta=-.559, p<.001$
Klietz et al. (2020) ¹⁰³ ; Germany	118	Community	Multiple regression	93.2	66.1	65.4	Caregiver: SF-36: $\beta=-.253, p=.013$ PwP: MDS-UPDRS II: $\beta=.265, p=.046$
Klietz et al. (2020) ⁷⁸ ; Germany	119	Community; medical setting	Multiple regression	NR	65.5	65.4	PwP: MDS-UPDRS II: $\beta=.415, p=.002$
Klietz et al. (2020) ⁷⁷ ; Germany	78	NR	Multiple regression	NR	53.8	64.8	PwP: PDQ-8: $\beta=.294, p=.064$ MDS-UPDRS III: $\beta=.272, p=.086$
Kudlicka et al. (2014) ⁷⁶ ; UK	50	Medical setting	Backward stepwise multiple regression	69.2	NR	NR	PwP: BRIEF-A (caregiver version): $\beta=.754, p=.000$ HY: $\beta=.377, p=.008$
Leroi et al. (2012) ⁸⁷ ; UK	71	NR	Forced entry multiple regression	54	39.4	62.7	Apathy PwP Group: MMSE-Serial Sevens (PwP): $B=-5.100, SE B=1.335, p=.001$ Impulse Control Disorders PwP Group: LEDD (PwP): $B=0.013, SE B=0.005, p=.012$ HADS-Depression (PwP): $B=1.677, SE B=0.763, p=.042$
Lo Monaco et al. (2021) ¹⁰⁴ ; Italy	51	Medical setting	Backward stepwise multiple regression	89	67	69	PwP: ADLs: $B=-6.48, SE B=2.84, p=.027$

Macchi et al. (2020) ⁶⁸ ; US and Canada	175	Medical setting	Stepwise multiple regression	81.7	73.1	66.1	Caregiver: HADS-Anxiety: $R^2=.077, p=.0002$ HADS-Depression: $R^2=.062, p=.0014$ PwP: FACIT-SP (Faith): $R^2=.024, p=.038$ PDQ-39: $R^2=.161, p<.0001$ QoL-AD Caregiver Reported: $R^2=.088, p<.0001$
Martinez-Martin et al. (2015) ⁸⁰ ; Spain	584	Validation study database	Stepwise multiple regression	61.2	70.5	59.6	PwP: SEND-PD (Mood/Apathy): $\beta=.284, p<.001$ SEND-PD (Psychosis): $\beta=.184, p<.001$ CISI-PD: $\beta=.213, p<.001$ Disease duration: $\beta=.112, p=.003$
Oguh et al. (2013) ⁷⁴ ; US	2476	National register	Stepwise multiple logistic regression	91	NR	NR	PwP: PDQ-39 ≥ 47 : OR=5.1 (95%CI=3.2, 8.2) HY: OR=2.0 (95%CI=1.3, 3.1) Concomitant medications: antidepressants OR=2.1 (95%CI=1.5, 3.1); antipsychotics OR=2.5 (95%CI=1.5, 4.2) Social worker visits: OR=1.6 (95%CI=1.2, 2.1) Gender (male): OR=2.3 (95%CI=1.5, 3.5) Verbal fluency: OR=0.95 (95%CI=0.92, 0.98)
Oh et al. (2015) ¹⁰⁵ ; South Korea	48	Medical setting	Multiple regression	NR	NR	NR	PwP: NPI-12: $B=0.619, SE B=0.151, p<.01$ (ZBI) NPI-12: $B=0.758, SE B=0.144, p<.01$ (CBI) Disease duration: $B=1.449, SE B=0.671, p<.05$ (CBI)
Peters et al. (2011) ¹⁰⁶ ; UK	704	Community	Multiple regression	88.9	71.9	67.1	PwP: PDQ-39 (Mobility): $\beta=.21, p<.001$ PDQ-39 (Social Support): $\beta=.15, p=.001$

Rajiah et al. (2017) ¹⁰⁷ ; Malaysia	130	Community	Stepwise multiple regression	NR	69.8	45.1	PwP: PDQ-39 (Stigma and Emotional Wellbeing): $R^2=.486, p=.001$
Rodríguez-Violante et al. (2015) ¹⁰⁸ ; Mexico	201	Medical setting	Multiple regression	53.2	73.1	51.6	PwP: MDS-UPDRS II: $\beta=.54, p<.007$
Santos-Garcia and de la Fuente-Fernandez (2015) ⁸³ ; Spain	121	Medical setting	Multiple regression	66.9	71.9	60.2	PwP: BDI: $\beta=.321, p=.003$ SE-ADLs: $\beta=-.536, p<.0001$
Sanyal et al. (2015) ¹⁰⁹ ; India	150	Medical setting	Stepwise multiple regression	48	79	50.4	Caregiver: Caregiving duration: $\beta=0.441, p=.037$ PwP: HY: $\beta=3.493, p=0.001$ UPDRS III: $\beta=0.898, p<.0005$
Sarandol et al. (2009) ¹¹⁰ ; Turkey	57	Medical setting	Stepwise multiple regression	61.4	59.6	Spouse=61.3; adult child=43.7	PwP: GDS: $R^2=.157, p<.05$ BEHAVE-AD-FW: $R^2=.145, p<.05$
Schmotz et al. (2017) ¹¹¹ ; Germany	20	Community; medical setting	Multiple regression	NR	70	67.1	No significant predictors
Shin et al. (2012) ¹¹² ; South Korea	91 (spouse=50; offspring=41)	Medical setting	Multiple regression	54.9	Spouse=50; offspring=53.7	Spouse=66.4; offspring=45.8	Spousal Group: CES-D (caregiver): $\beta=.754, p<.001$ UPDRS I (PwP): $\beta=.263, p=.002$ Offspring Group: Community relationships (caregiver): $\beta=-.318, p=.034$ UPDRS III (PwP): $\beta=.455, p=.012$

Shin et al. (2012) ¹¹³ ; South Korea	42	Medical setting	Multiple regression	NR	61.9	60	Caregiver: CES-D: $B=1.1, SE B=0.2, p<.001$ PwP: UPDRS I: $B=2.9, SE B=1.3, p=.03$
Tanji et al. (2013) ⁶⁹ ; Japan and US	178 (Japan=83; US=96)	Medical setting	Multiple regression	100	NR	Japan=68.9; US=63.7	Japanese Group: BDI (caregiver): $\beta=.35, p=.003$ Falls (PwP): $\beta=.36, p=.003$ US Group: Help from others (caregiver): $\beta=-.34, p=.001$ BSI (caregiver): $\beta=.25, p=.03$ BSI (PwP): $\beta=.41, p=.001$
Viwattanakulvanid et al. (2014) ⁸² ; Thailand	85	Medical setting	Stepwise multiple regression	49.4	78.8	50.8	PwP: HADS-Anxiety: $\beta=.494, p=.000$ NADCS-Akinesia: $\beta=.237, p=.008$ MPDSS-item 14: $\beta=-.229, p=.009$
Wallhagen & Brod (1997) ⁷² ; US	45	Medical setting; community	Multiple regression	100	69	69	PwP: Perceived control over symptoms: $\beta=-.29, p=.03$
Wandrekar et al. (2014) ¹¹⁴ ; India	50	Community	Multiple regression	76	76	NR	PwP: MDS-UPDRS I: $\beta=.31, p<.05$ MDS-UPDRS II: $\beta=.23, p<.05$ MDS-UPDRS III: $\beta=.55, p<.05$
Zhang et al. (2022) ⁸⁴ ; Australia	39	Medical setting; community	Multiple regression	100	61.5	67.1	Caregiver: COPE-28 (Problem-focused): $\beta=.39, p=.02$
Zhong et al. (2016) ¹¹⁵ ; Australia	50	Medical setting	Forward stepwise multiple regression	88	68	64.7	Caregiver: HADS-Depression: $R^2=.450, p<.001$ PwP: NUCOG-Visuoconstructional: $R^2=.581, p=.006$ HADS-Depression: $R^2=.713, p=.002$

Note. All studies utilised a cross-sectional design. ADL-MS=Activities for Daily Living Self-Care Scale for Multiple Sclerosis; BDI=Beck Depression Inventory; BEHAVE-AD-FW= Behavioral Pathology in Alzheimer's Disease Frequency-Weighted Severity Scale; BRIEF-A=Behaviour Rating Inventory of Executive Function-Adult Version; BSI=Brief Symptom Inventory; CBI=Caregiver Burden Inventory; CES-D=Center for Epidemiologic Studies Depression scale; CISI-PD=Clinical Impression of Severity Index for Parkinson's Disease; COPE-28=Brief Coping Orientation to Problems Experienced Inventory; EASI=Everyday Abilities Scale for India; FACIT-SP=Functional Assessment of Chronic Illness Therapy-Spiritual Wellbeing; GDS=Geriatric Depression Scale; GHQ-28=28-item General Health Questionnaire; HADS=Hospital Anxiety and Depression Scale; HY=Hoehn and Yahr scale; LEDD=Levodopa equivalent daily dose; MDS-UPDRS=Movement Disorder Society-Sponsored Revision of the Unified Parkinson's Disease Rating Scale; MMSE=Mini Mental State Examination; MPDSS=Modified Parkinson's Disease Sleep Scale; MS=Mutuality Scale; MSPSS=Multidimensional Scale of Perceived Social Support; NADCS=Nocturnal Akinesia Dystonia and Cramp Score; NMSS=Non-Motor Symptom Scale; NPI-12=Neuropsychiatric Inventory - 12-item; NR=not reported; NUCOG=Neuropsychiatry Unit Cognitive Assessment Tool; PDQ=Parkinson's Disease Questionnaire; PPRS=Parkinson's Psychosis Rating Scale; PSS-Fa=Perceived Social Support from Family; QoL-AD=Quality of Life in Alzheimer's Disease; RAS=Relationship Assessment Scale; SE-ADLs=Schwab & England Activities of Daily Living Scale; SCOPA=Scales for Outcomes in Parkinson's Disease; SEND-PD=Scale for Evaluation of Neuropsychiatric Disorders in Parkinson's Disease; SF-36=36-Item Short Form Health Survey; SOC=Sense of Coherence scale; SPDDS=Self-Assessment Parkinson's Disease Disability Score; UPDRS=Unified Parkinson's Disease Rating Scale; ZBI=Zarit Burden Interview.

^aCommunity recruitment included caregiver or patient support groups, newspaper adverts, and newsletters; medical settings included hospitals, outpatient clinics, palliative and primary care, and nursing homes.

Table 4. Variables Investigated in Relation to Caregiver Burden

Significant Predictors of Higher Caregiver Burden		Predictors Investigated but Not Found to be Significant
Caregiver-Related Factors		
Demographics		<ul style="list-style-type: none"> • Age^{47,68,69,75,80,81,104,113} • Gender^{47,75,80,84,113} • Education^{47,75,113} • Socioeconomic status⁹⁷ • Work status⁴⁷ • Marital status⁴⁷ • Living with PwP⁹⁷
Caregiver Involvement	<ul style="list-style-type: none"> • Greater years of caregiving^{81,102} • Greater daily hours of caregiving¹⁰⁹ 	<ul style="list-style-type: none"> • Years of caregiving^{68,97,115} • Frequency of providing care⁴⁷ • Time of day care provided⁴⁷ • Weekly hours of caregiving⁹⁷ • Daily hours of caregiving¹¹⁵
Psychological Factors	<ul style="list-style-type: none"> • Greater depression^{68,69,73,113,115} <ul style="list-style-type: none"> - specifically in spousal (but not offspring) caregivers¹¹² • Greater anxiety^{68,81} • Psychological wellbeing⁷¹ 	<ul style="list-style-type: none"> • Anxiety¹¹⁵ • Psychological wellbeing⁸⁵
Protective Factors	<ul style="list-style-type: none"> • Low sense of coherence^{73,100} • Use of problem-focused coping strategies^{47,84,101} • Use of avoidant coping strategies⁴⁷ • Poorer perceived social support^{47,85} <ul style="list-style-type: none"> - specifically from community in offspring caregivers¹¹² • Less help from others in US (but not Japanese) sample⁶⁹ • Fewer caregivers involved in supporting PwP^{95,99} • Poorer relationship satisfaction¹⁰¹ • Poorer mutuality⁷⁵ 	<ul style="list-style-type: none"> • Self-esteem¹⁰⁰ • Social competence¹⁰⁰ • Avoidant coping style⁸⁴ • Emotion-focused coping style^{47,84} • Social support¹¹² • Help from others⁹⁷ • Relationship quality⁸⁴ • Marital satisfaction⁸⁵ • Spiritual wellbeing⁶⁸ • Activation⁴⁷ • Mindfulness¹⁰³ • Psychological flexibility¹⁰³
Other Factors	<ul style="list-style-type: none"> • Current health problems impacting ability to provide care¹⁰¹ • Poorer sleep¹⁰¹ 	<ul style="list-style-type: none"> • Medical co-morbidities⁶⁹ • Health-related QoL^{112,113}

	Significant Predictors of Higher Caregiver Burden	Predictors Investigated but Not Found to be Significant
	<ul style="list-style-type: none"> Poorer health-related QoL¹⁰³ 	
	Person with Parkinson's-Related Factors	
Demographics	<ul style="list-style-type: none"> Older age⁹⁹ Male gender^{70,74} Occupational status (not working)⁴⁷ Living at home (as opposed to a nursing home)⁷⁰ 	<ul style="list-style-type: none"> Age^{73,81,97,104,105,108,113} Gender^{73,81,103,105,108,113} Education^{47,75,113}
Disease Factors	<ul style="list-style-type: none"> Advancing disease stage^{73,74,76,80,86,99,109} Greater disease duration^{80,99,100} <ul style="list-style-type: none"> specifically using Caregiver Burden Inventory (but not Zarit Burden Inventory)¹⁰⁵ Presence of concomitant medications⁷⁴ Higher dopaminergic load in PwP presenting with impulsivity (but not apathy)⁸⁷ 	<ul style="list-style-type: none"> Disease stage^{83,84,95,97,105,108,112,113,115} Disease duration^{47,78,83,97,103,104,108,115} Symptom burden⁶⁸ Presence of atypical parkinsonism⁶⁸ Presence of dementia⁸⁰ Medication use^{68,69} Dopaminergic load¹⁰⁵
Motor Symptoms	<ul style="list-style-type: none"> Greater motor symptomology^{47,78,103,108,114} Greater motor symptom severity^{77,95,109,114} <ul style="list-style-type: none"> specifically in offspring (but not spousal) caregivers¹¹² Greater dependence in ADLs^{47,83,85,102,104} Greater difficulty with eating, bathing, and meals¹⁰¹ Higher frequency of falls in Japanese (but not US) sample⁶⁹ 	<ul style="list-style-type: none"> Motor symptoms^{95,112,113} Motor symptom severity^{68,75,83,104,108,111,113} Motor complications^{83,95,108,112-114} ADL^{69,81,95,104,112,113} Instrumental ADL¹⁰⁴ Overall functional status⁶⁸ Speech, gait, postural instability, freezing, fluctuation, dyskinesia, dystonia, and presence of autonomic symptoms^{69,95}
Non-Motor Symptoms	<ul style="list-style-type: none"> Greater non-motor symptomology^{70,113,114} <ul style="list-style-type: none"> specifically in spousal (but not offspring) caregivers¹¹² Greater severity of neuropsychiatric symptoms^{70,71,86,99,102,105} <ul style="list-style-type: none"> all subscales except apathy⁹⁹ motor disturbance subscale only¹⁰¹ Greater severity on psychotic symptoms subscale⁸⁰ and mood/apathy but not impulse control disorders subscale^{80,98} Greater depression^{83,95,110,115} <ul style="list-style-type: none"> specifically in US sample⁶⁹ and PwP with impulse control disorders (but not apathy)⁸⁷ Greater anxiety⁸² Greater impairment in visuo-constructional skills¹¹⁵ 	<ul style="list-style-type: none"> Non-motor symptoms^{75,83,108} Psychiatric symptoms⁹⁵ Alexithymia⁷⁸ Depression^{47,68,76,78,81,105,111} Anxiety^{47,68,105,115} Cognitive function^{68,69,76,81,97,100,105,111-114} <ul style="list-style-type: none"> specifically caregiver perspective on cognitive decline⁷⁵ Memory^{100,115} Executive functioning¹¹⁵

	Significant Predictors of Higher Caregiver Burden	Predictors Investigated but Not Found to be Significant
	<ul style="list-style-type: none"> • Greater impairment in attentional ability in PwP with apathy (but not impulse control disorders)⁸⁷ • Decreased verbal fluency⁷⁴ • Decreased phonemic (but not semantic) verbal fluency¹⁰⁰ • Presence of levodopa-induced psychotic symptoms⁸¹ • Greater behavioural difficulties¹¹⁰ • Greater caregiver-rated (but not PwP-rated) executive function-related behavioural problems⁷⁶ • Greater sleep disturbances^{81,95} • Presence of nocturnal akinesia and PwP feeling tired and sleepy upon awakening⁸² 	
Quality of Life	<ul style="list-style-type: none"> • Poorer QoL from caregiver (but not PwP) perspective⁶⁸ • Poorer health-related QoL^{68,74,77,96,99} <ul style="list-style-type: none"> - mobility,^{101,106} social support,¹⁰⁶ stigma,¹⁰⁷ and emotional wellbeing¹⁰⁷ subscales only 	<ul style="list-style-type: none"> • Health-related QoL^{78,103}
Other Factors	<ul style="list-style-type: none"> • Less perceived control over Parkinson's symptoms⁷² • Poorer spiritual wellbeing⁶⁸ • Utilisation of social or mental health visits⁷⁴ 	<ul style="list-style-type: none"> • Perceived control over disease progression⁷² • A&E visits⁶⁸ • Receiving home health services⁶⁸ • Medical co-morbidities^{69,97,105}