The Relationship among Caregiver Burden, Demographic Variables, and the Clinical Characteristics of Patients with Parkinson’s Disease—A Systematic Review of Studies Using Various Caregiver Burden Instruments

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Abstract

Caring for a person with Parkinson’s disease (PD) extends far beyond the ordinary exchange of assistance among people in a close relationship. Caregivers must learn to cope with the patient’s increasing disability and loss of independence. The aim of this systematic review was to critically assess and summarize the evidence of the influence of the demographic and clinical characteristics of patients with PD on caregiver burden by means of a caregiver burden instrument. In order to identify articles, electronic databases and reference lists were searched using the search word “Parkinson’s disease” in combination with “caregiver” or “carer” and with “burden” or “distress” or “stress” or “strain”. Thirty one articles were deemed eligible for inclusion. The methodological quality of the studies was evaluated. No studies were excluded due to low quality. The results revealed similar associations among caregiver burden, demographic variables and patient characteristics, across different caregiver burden instruments and various clinical scales. Higher PD stage and functional disability are the non-motor characteristics that contribute the most to caregiver burden. However, when comparing the impact of patient motor and non-motor symptoms,
several studies found that mental symptoms had a stronger impact on caregiver burden than motor symptoms. No association was observed between caregiver burden and patient and caregiver demographics with the exception of the sub-scale analysis of caregiver burden in various age groups. Interpreting the results of studies that employ a range of different clinical assessment scales and burden instruments makes it challenging to provide a valid summary of caregiver burden in PD. The most commonly used analysis methods contribute little information about burden variation across caregiver groups or which areas are the most burdensome for caregivers. There is a need for a more uniform use of recommended instruments and for longitudinal studies.

Keywords
Caregiver Burden, Distress, Strain, Stress, Parkinson’s Disease

1. Introduction
From birth we depend on other persons’ caring to survive. The need to be cared for presupposes the responsibility and capacity of another person to provide care. Accordingly, caregiving can be viewed as a response to the care needs of the other. Although caregiving is embedded in all close relationships, many people experience its transformation from the ordinary exchange of assistance to a situation that calls for a more extensive form of caregiving when a close person is not adequately taken care of by her/himself or by the professional healthcare service. Parkinson’s disease (PD) is one such situation. PD is a progressive neurological condition characterized by motor symptoms comprising tremor, rigidity, bradykinesia, postural instability and gait difficulties [1]. In addition, most people with PD will experience some non-motor symptoms including neuropsychiatric problems such as psychosis, depression, anxiety, fatigue, apathy and dementia as well as sleep disturbance and autonomic dysfunction with gastrointestinal, urinary and sexual malfunction [1].

Initially, a person with PD responds well to antiparkinson medication. However, the effect usually wears off after several years and the symptoms become more prominent with increasing motor disability and complications such as involuntary and uncontrollable movements [2] and the fluctuating benefit of the medications [3]. Fluctuations are experienced as a switch between mobility and immobility, called the On-Off phenomenon, which is often followed by additional non-motor fluctuations [4]. In the same way as the person with PD, her/his family and significant others have to cope with these challenges and accommodate themselves to the unpredictable On-Off phenomenon, the increasing disability and need for assistance characteristic of advanced PD. Despite limited evidence in the PD population, increased strain [5] and overwhelming caregiver demands [6] have been reported to trigger caregivers’ long term care (LTC) placement decisions. In the UK, LTC facilities for PD patients have been estimated to represent a 4.5 times higher healthcare cost than that of patients who are able to live at home [7]. In a Norwegian study, Vossius et al. [8] found that even a few months delay in admission to a nursing home could reduce these costs considerably. Accordingly, there is a need for knowledge about which factors may undermine informal caregivers’ ability to provide support and continue in their caring role.

Strain or load experienced by those caring for a person with a chronic disease is often referred to as caregiver burden [9], distress [10], stress [11] or strain [12]. In the literature these terms are frequently used interchangeably [13], although caregivers’ subjective and objective burden related to the emotional, physical and social problems that arise from caring for a person with a chronic and disabling disease are covered by most instruments addressing these constructs [14]. A number of instruments have been designed to assess caregivers’ situation [14] [15]. In a review, Durme et al. [14] identified 55 scales (mainly generic) assessing the negative impact of caregiving, of which 42 evaluated burden, strain or stress as the main dimension. The current interest in the impact of caregiving has also resulted in an increasing number of studies focusing on caregiver burden in PD [16]. Furthermore, three reviews related to caregiving outcomes in PD have recently been published (Lau et al. [10], Martinez-Martin et al. [17] and Greenwell et al. [18]). The review by Lau et al. [10] is a meta-analysis of the correlates of caregiver distress. Martinez-Martin et al. [17] reviewed the state of the art regarding concepts, assessments, related factors, costs and intervention programmes on PD caregivers’ quality of life (QoL) and burden. The systematic review by Greenwell et al. [18] aimed to evaluate the evidence of the predictive factors of psychosocial outcomes, including burden, mental health and QoL in PD carers.
Aim and Review Questions

The aim of this systematic review was to critically assess and summarize the evidence of the influence of the demographic and clinical characteristics of patients with PD on caregiver burden by means of a caregiver burden instrument. Two review questions were addressed:

1) How did the demographic and clinical characteristics of patients with PD influence caregiver burden?
2) What instruments and which analyses were used to assess caregiver burden in relation to PD?

2. Methods

This review was guided by recommendations in the Preferred Reporting Items for Systematic Reviews and Meta-Analysis (PRISMA) checklist [19].

2.1. Search Strategy

A search of the literature was performed in mid-January 2014 in the Pubmed, Medline, PsychInfo, Cinahl, Web of Science and Embase electronic databases. The search term employed was “Parkinson’s disease” in combination with “caregiver” or “carer” and with “burden” or “distress” or “stress” or “strain”. The search was limited to English and Scandinavian languages. There were no limitations in terms of publication year. A manual search was also made of the reference lists of the articles included in the review. A total of 581 articles were identified, of which 318 were duplicates.

2.2. Inclusion and Exclusion Criteria

All articles were assessed for inclusion based on the following criteria: 1) original empirical study, 2) published in peer-reviewed journals, 3) with an identifiable and separately analysed group of informal caregivers of persons diagnosed with PD, 4) explored characteristics that are assumed to influence caregiver burden and 5) burden was measured by means of a standardized caregiver instrument. The sample size of eligible studies was ≥50 caregivers, which has been suggested as a reasonable number of cases to ensure statistical power in correlation and regression analysis [20]. Exclusion criteria were intervention studies, medical trials and testing of instruments. Seventy titles were identified as relevant for inclusion. After reading the abstracts, 23 of the 70 articles were excluded in line with the inclusion and exclusion criteria, while another 16 articles were excluded after reading the full text, leaving a total of 31 studies for inclusion (Table 1). The flow chart illustrating study selection and the reasons for exclusion is presented in Figure 1.

2.3. Data Extraction

For data extraction, two authors (IL, UTL) independently identified the characteristics of the studies such as authors, publication year, country of origin, aim, participants, recruitment methods, study design, main definitions, outcome measures, statistical analysis and key findings. The first author (IL) identified and categorized all de-
Table 1. Characteristics of the included studies.

<table>
<thead>
<tr>
<th>Authors</th>
<th>Aim</th>
<th>Sample (relationship)</th>
<th>Statistical analyses*</th>
<th>Key findings related to caregiver burden</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kudlicka et al. (2014)</td>
<td>To evaluate how executive functions contribute to caregiver burden, as well as QoL and subjective health status of non-demented and non-depressed people with early-stage PD</td>
<td>50 caregivers (45 spouses, 3 parents, 7 children, 2 friends)</td>
<td>Multiple regression analysis</td>
<td>Caregiver burden was best explained by caregiver rated executive function of the PD patient, followed by disease severity</td>
</tr>
<tr>
<td>Carod-Artal et al. (2013)</td>
<td>To analyse the main determinants of burden and HRQoL in Brazilian PD patients</td>
<td>50 caregivers (39 spouses, 7 children, 4 others)</td>
<td>Yes</td>
<td>The patient variables sleep disorders and behavioural psychotic symptoms were independent predictors of caregiver burden</td>
</tr>
<tr>
<td>Oguh et al. (2013)</td>
<td>To determine what measures of PD disability, demographics and patient QoL are associated with caregiver strain</td>
<td>2476 caregivers (91% spouses/partners, 9.4% other relatives, 0.5% other non-paid caregivers)</td>
<td>Multiple logistic regression</td>
<td>Patient quality of life impairment, male sex, disease severity, presence of concomitant medications and decreased verbal fluency were factors that predicted the likelihood of high caregiver strain</td>
</tr>
<tr>
<td>Tanji et al. (2013)</td>
<td>To compare caregiver strain in spouses in one region of Japan and one in the US and examine the correlation between caregiver strain and patient/spousal variables</td>
<td>178 caregivers (Spouses)</td>
<td>Spearman’s rho t-test Multivariate regression analysis</td>
<td>Spouses in the Japanese group reported more physical, time and financial caregiver strain. The US group reported more emotional strain. Falls was the most predictive patient variable for caregiver strain in the Japanese group. In the US group, patient depression was the most predictive variable for caregiver strain</td>
</tr>
<tr>
<td>Agrawal et al. (2012)</td>
<td>To find various predictors of caregiver burden in caregivers of persons with PD in India</td>
<td>91 caregivers (45 spouses, 32 children, 7 siblings, 7 others)</td>
<td>Linear regression</td>
<td>Patient depression scores were the best predictor of increased caregiver burden, followed by patient motor scores and the presence of sleep disturbances</td>
</tr>
<tr>
<td>Kelly et al. (2012)</td>
<td>To consider the relationship between HRQoL in non-demented PwPD and their caregivers and to determine the associations between caregiver and patient HRQoL and caregiver strain</td>
<td>97 caregivers (84% spouses, 13 % children, 3% others)</td>
<td>Spearman’s rho</td>
<td>Correlation was found between caregiver strain and the HRQoL of the person with PD</td>
</tr>
<tr>
<td>Leroi et al. (2012)</td>
<td>To compare quality of life, level of disability and caregiver burden among PwPD with mild cognitive impairment (PD-MCI), PwPD with dementia (PDD), and PwPD with no cognitive impairment (PD-NC)</td>
<td>102 caregivers (55 spouses, 47 adult children)</td>
<td>Analysis of covariance Analysis of variance</td>
<td>Caregiver burden as assessed by the ZBI was significantly greater in the PDD group compared to the two groups without dementia when adjusted for age and motor symptom severity. No significant difference in caregiver distress was observed between the three PD caregiver groups assessed by the NPI-CD</td>
</tr>
<tr>
<td>Author(s)</td>
<td>Year</td>
<td>Country</td>
<td>Objective</td>
<td>Sample Size</td>
</tr>
<tr>
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</tr>
<tr>
<td>Ozdilek and Gunal (2012) [38]</td>
<td>Turkey</td>
<td>To examine the impact of motor and non-motor symptoms of non-demented PwPD on the psychological health, burden and QoL of caregivers</td>
<td>50 caregivers (37 spouses, 11 children, 2 siblings)</td>
<td>Person’s correlation coefficient (multiple linear regression-results not given)</td>
</tr>
<tr>
<td>Shin et al. (2012) [63]</td>
<td>South Korea</td>
<td>To determine factors that predict caregiver burden in spousal and offspring caregiver groups</td>
<td>91 caregivers (50 spouses, 41 offspring)</td>
<td>Spearman’s rho</td>
</tr>
<tr>
<td>Leroi et al. (2012) [36]</td>
<td>United Kingdom</td>
<td>To explore the relationship between carer burden and the presence of apathy and impulse control disorders (ICD) in PD patients</td>
<td>71 caregivers (38 spouses, 33 adult children)</td>
<td>Linear regression</td>
</tr>
<tr>
<td>Peters et al. (2011) [56]</td>
<td>United Kingdom</td>
<td>To explore to what extent patient self-reported health status is associated with carer strain and QoL</td>
<td>704 caregivers (626 spouses/ partners, 41 children, 18 other family members, 18 friends or others)</td>
<td>Regression analysis</td>
</tr>
<tr>
<td>Razali et al. (2011) [39]</td>
<td>Malaysia</td>
<td>To investigate clinical and socio-demographic factors associated with perceived burden among PD caregivers</td>
<td>115 caregivers (51 spouses, 56 children, 8 other relatives)</td>
<td>Correlation Analysis of variance</td>
</tr>
<tr>
<td>Carter et al. (2010) [26]</td>
<td>USA</td>
<td>To compare the difference in negative and positive aspects of strain in young versus older spouse caregivers in early stage PD patients</td>
<td>65 caregivers [Spouses, 37 young (40-55 years), 28 old (≥70 years)]</td>
<td>t-test</td>
</tr>
<tr>
<td>Leiknes et al. (2010) [33]</td>
<td>Norway</td>
<td>To investigate caregiver distress associated with neuropsychiatric problems in patients with newly diagnosed PD and a control group</td>
<td>189 caregivers (134 spouses, 31 children, 18 other relatives, 6 others)</td>
<td>Mann-Whitney U test, Chi-square test, Spearman’s rho</td>
</tr>
</tbody>
</table>
Continued

<table>
<thead>
<tr>
<th>Study</th>
<th>Objective</th>
<th>Methodology</th>
<th>Control Variables</th>
<th>Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lyons et al. (2009) [25] USA</td>
<td>To examine the roles of optimism, pessimism, mutuality, and spouse gender in predicting role strain in PD spouses over a 10-year period</td>
<td>Multilevel modeling</td>
<td>Yes</td>
<td>Female gender predicted both higher role strain at Year 10 and a more rapid increase in role strain over the 10-year period when controlled for stage of PD and caregiver spouse age. Higher disease stage predicted role strain at Year 10, but not more rapid increases over this period</td>
</tr>
<tr>
<td>Sarandol et al. (2009) [34] Turkey</td>
<td>To determine factors that increase caregiver burden and depression and to investigate if adult children and spouses differ in their perception of burden</td>
<td>Mann-Whitney U-Test, Stepwise regression analysis</td>
<td>No</td>
<td>Female caregivers compared to male caregivers and adult child caregivers compared to spouse caregivers experienced significantly higher burden. Patient’s depression and behavioural disturbances were determinants of caregiver burden</td>
</tr>
<tr>
<td>Stella et al. (2009) [51] Brazil</td>
<td>To identify the impact of patient neuropsychiatric symptoms on caregiver burden</td>
<td>Pearson’s product moment correlation analysis, Analysis of variance</td>
<td>No</td>
<td>Caregiver burden was proportional with the degree of neuropsychiatric symptomatology. Caregiver burden was significantly lower in caregivers of non-depressed/non-demented patients compared to caregivers of patients with depression or dementia</td>
</tr>
<tr>
<td>Carter et al. (2008) [28] USA</td>
<td>To examine the relationship between specific clinical motor and non-motor symptoms in early and middle stage PD to caregiver strain and depression</td>
<td>Hierarchical regression analyses</td>
<td>Yes</td>
<td>Patient cognitive impairment and depression have a much greater impact on caregiver strain than the motor symptoms in early and middle stage PD</td>
</tr>
<tr>
<td>Goldsworthy and Knowles (2008) [41] Australia</td>
<td>To examine the relationships between caregiver stressors, and protective factors associated with caregiver burden and quality of life</td>
<td>Path analysis</td>
<td>No</td>
<td>The caregiver stressors; patient behavioural problems and functional dependency contributed to caregiver burden. Patient cognitive impairment and hours of caregiving did not</td>
</tr>
<tr>
<td>Martinez-Martin et al. (2008) [9] Spain</td>
<td>To analyse the association between the characteristics of patients with PD and their caregivers and caregiver burden, perceived health and mood status, and to identify their predictors</td>
<td>Multivariate linear regression based on factor analysis</td>
<td>No</td>
<td>All patient-related factors (including mood, autonomic dysfunction, pain, fatigue, disease duration, disability, motor complication, sleep disorders, age and cognition) were predictors of caregiver burden</td>
</tr>
<tr>
<td>Aarsland et al. (2007) [47] Norway</td>
<td>To explore the profile of neuropsychiatric symptoms in PD patients with dementia</td>
<td>Analysis of variance based on cluster analysis</td>
<td>Yes</td>
<td>Based on a cluster analysis of associated neuropsychiatric symptoms in patients, the highest caregiver distress score was observed in the agitation cluster followed by the psychosis and the mood clusters</td>
</tr>
<tr>
<td>Study</td>
<td>Country</td>
<td>Objective</td>
<td>Sample Description</td>
<td>Method</td>
</tr>
<tr>
<td>-------------------------------------------</td>
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<td>-----------------------------------------------------------------------</td>
</tr>
<tr>
<td>Kim et al. (2007) [32]</td>
<td>South Korea</td>
<td>To elucidate subjective and objective burden on primary PD caregivers and identify related factors.</td>
<td>68 caregivers (31 spouses, 21 children, 14 other family members, 2 unknown)</td>
<td>Pearson’s product moment correlation Analysis of variance Kruskal-Wallis Yes Subjective burden correlated with the caregiver variables age (≤ 40 years reported less), and relation to care recipient (spouses reported highest) and with the patient variables motor disability, disease duration (&lt;1 and &gt;10 years reported less) and monthly medical costs. Objective burden correlated with patient motor disability and disease severity</td>
</tr>
<tr>
<td>Martinez-Martin et al. (2007) [37]</td>
<td>Spain</td>
<td>To assess the burden and HRQoL of PD caregivers and to determine the relationship between these and socio-demographics, emotional, functional factors and HRQoL</td>
<td>79 caregivers (61 spouses, 15 children, 3 others)</td>
<td>Stepwise multiple regression based on factor analysis No Clinical aspects of PD were the main patient variables predicting caregiver burden, followed by patients’ mood and HRQoL</td>
</tr>
<tr>
<td>Schrag et al. (2006) [62]</td>
<td>United Kingdom</td>
<td>To assess caregiver burden and associated factors</td>
<td>116 caregivers (Partners)</td>
<td>Pearson’s product moment correlation Spearman’s rho No Increased PD disability and hallucination, confusion and falls were most strongly correlated with caregiver burden. Correlation was also found between caregivers’ burden and patients’ depression, disease duration and QoL</td>
</tr>
<tr>
<td>Marsh et al. (2004) [50]</td>
<td>USA</td>
<td>To determine the prevalence and impact of psychiatric comorbidities in PwPD with psychosis</td>
<td>50 caregivers (Not reported)</td>
<td>Stepwise linear regression No Psychosis was the major predictor of caregiver burden, followed by patient functional ability</td>
</tr>
<tr>
<td>Caap-Ahlgren and Dehlin (2002) [31]</td>
<td>Sweden</td>
<td>To identify various factors in PD patients and their caregivers that are of importance for caregiver burden</td>
<td>65 caregivers (58 spouses, 3 daughters, 2 sons, 1 brother, 1 niece)</td>
<td>Multivariate regression analyses No Patient’s functional status assessed by H&amp;Y was the most important patient variable for caregiver burden in the multivariate analysis</td>
</tr>
<tr>
<td>Thommessen et al. (2002) [30]</td>
<td>Norway</td>
<td>To compare the psychosocial burden on spouses caring for patients with mild dementia, stroke and PD and to identify associated patient characteristics</td>
<td>58 caregivers (Spouses)</td>
<td>Multivariate modelling based on factor and path analysis Yes The regression model disclosed an effect on PD spouses psychosocial burden from the patients’ depressive symptoms and lower levels of cognitive function, but not from their ADL function</td>
</tr>
</tbody>
</table>
To determine the emotional and social distress of caring for a person with PD and to explore the impact of motor and mental symptoms of PD on caregivers’ situation

“Distress and social upset experienced by the relative as a result of having to care for an elderly person with physical and/or behavioural disability” (p. 867)

58 caregivers (Spouses)

Stepwise multiple linear regression

Yes

Patient mental symptoms were the most powerful predictors of caregiver stress in PD spouses. Patient functional impairment also contributed to the model whereas severity of motor symptoms and disease stage did not

To investigate sources of distress in PD caregivers and the relationships between demographic characteristics, stress and family functioning

No definition

50 caregivers (Not reported)

Multivariate analysis of variance

No

Aspects of caregiving related to caregiver’s personal concerns and anxiety were reported to be the most distressing

To examine the experience of spouses who provide care for patients with PD and to determine whether their experiences differed by stage of disease

No definition

380 caregivers (Spouses)

Analysis of variance

Yes

Caregiver strain is experienced across all stages of PD and increases significantly in line with the progression of the disease

To determine the characteristics of PD patients that are associated with stress for their relatives

No definition

65 caregivers (62 spouses, 2 daughters, 1 daughter in law)

Stepwise regression analyses

No

Patient’s behavioural disturbances contributed most to the variation in level of stress, followed by male sex in patient, patient’s self-care and young age. Patient’s functional disability predicted relative’s stress independently of patient cognitive impairment or dementia

2.4. Data Synthesis

Data from the included studies were narratively synthesized. A narrative approach was used in preference to a meta-analysis due to the heterogeneity of the studies with a huge variety of instruments applied to assess clinical aspects of PD and caregiver burden outcomes [21].

2.5. Assessment of Methodological Quality

Two of the authors (ES/IL) independently assessed the methodological quality of the studies using a modified version of the Norwegian Knowledge Centre for the Health Services (NOKC) check-list for cross-sectional studies [22]. The check-list is a tool for assessing the degree to which the methodology of the studies reduces the risk of systemic bias. It contains seven questions assessing criteria such as 1) population, 2) sampling methods, 3) comparison of respondents and non-respondents, 4) response rate, 5) data collection procedures, 6) reliability and validity and 7) statistical methods. For the purpose of this review, the check-list was supplemented by an

mographic and clinical characteristics examined in the studies regarding their possible influence on caregiver burden. The second author (UTL) made some spot tests to cross-check the validity of the latter extraction. In addition, the various instruments used to assess patient characteristics were collected.
additional criterion; ethical considerations (Criterion 8). The response alternatives for all questions were Yes (Y), No (N) and No information (NI) (Table 2). Differences in judgment between the two reviewers were resolved.

Table 2. Methodological quality assessment of the included studies.

<table>
<thead>
<tr>
<th>First author (Year) [Reference]</th>
<th>Quality assessment question*</th>
<th>Total assessment quality**</th>
</tr>
</thead>
<tbody>
<tr>
<td>Caap Ahlgren (2002) [31]</td>
<td>Y Y Y Y N Y Y Y Y</td>
<td>Acceptable</td>
</tr>
<tr>
<td>Calder (1991) [64]</td>
<td>Y Y N Y NI N Y Y</td>
<td>Acceptable</td>
</tr>
<tr>
<td>Carod-Artal (2013) [61]</td>
<td>Y Y N Y NI Y Y Y</td>
<td>Acceptable</td>
</tr>
<tr>
<td>Carter (1998) [27]</td>
<td>Y Y Y Y Y Y Y Y</td>
<td>Acceptable</td>
</tr>
<tr>
<td>Carter (2008) [28]</td>
<td>Y Y Y Y Y Y Y Y</td>
<td>Acceptable</td>
</tr>
<tr>
<td>Carter (2010) [26]</td>
<td>Y Y Y Y Y Y Y NI</td>
<td>Acceptable</td>
</tr>
<tr>
<td>Goldsworthy (2008) [41]</td>
<td>N N N NI Y Y Y NI</td>
<td>Low</td>
</tr>
<tr>
<td>Kelly (2012) [12]</td>
<td>Y Y N N NI Y Y Y</td>
<td>Acceptable</td>
</tr>
<tr>
<td>Kim (2007) [32]</td>
<td>Y Y N N Y Y Y Y</td>
<td>Acceptable</td>
</tr>
<tr>
<td>Kudlicka (2013) [66]</td>
<td>Y Y N Y Y NI Y Y</td>
<td>Acceptable</td>
</tr>
<tr>
<td>Leiknes (2010) [33]</td>
<td>Y Y N Y Y Y Y Y</td>
<td>Acceptable</td>
</tr>
<tr>
<td>Leroi (2012) [49]</td>
<td>Y Y N NI NI Y Y Y</td>
<td>Acceptable</td>
</tr>
<tr>
<td>Martinez-Martin (2007) [37]</td>
<td>Y Y N NI NI Y Y Y</td>
<td>Acceptable</td>
</tr>
<tr>
<td>Oghu (2013) [65]</td>
<td>Y Y N N N Y Y Y</td>
<td>Acceptable</td>
</tr>
<tr>
<td>Ozdilek (2012) [38]</td>
<td>Y Y N N Y NI N Y</td>
<td>Low</td>
</tr>
<tr>
<td>Peters (2011) [56]</td>
<td>Y Y N N Y NI Y Y</td>
<td>Acceptable</td>
</tr>
<tr>
<td>Razali (2011) [39]</td>
<td>Y Y Y Y Y Y Y NI</td>
<td>Acceptable</td>
</tr>
<tr>
<td>Sarandol (2009) [34]</td>
<td>Y Y N N NI NI N Y</td>
<td>Low</td>
</tr>
<tr>
<td>Schrag (2006) [62]</td>
<td>Y N N NI Y NI Y Y</td>
<td>Low</td>
</tr>
<tr>
<td>Shin (2012) [63]</td>
<td>Y Y N N Y NI Y NI</td>
<td>Low</td>
</tr>
<tr>
<td>Stella (2009) [51]</td>
<td>Y Y N N Y NI Y Y</td>
<td>Acceptable</td>
</tr>
<tr>
<td>Tanji (2013) [54]</td>
<td>Y Y Y Y Y Y Y Y</td>
<td>Acceptable</td>
</tr>
<tr>
<td>Aarsland (1999) [29]</td>
<td>Y Y Y Y Y Y Y Y</td>
<td>Acceptable</td>
</tr>
<tr>
<td>Aarsland (2007) [47]</td>
<td>Y Y N Y Y Y Y Y Y</td>
<td>Acceptable</td>
</tr>
</tbody>
</table>

Note: Y = yes; N = no; NI = no information.

*Quality assessment questions
1. Was the population from which the sample was drawn clearly defined?
2. Were sampling methods adequate?
3. Was it explained whether (and how) the participants who agreed to participate differed from those who refused?
4. Was the response rate adequate?
5. Were procedures for data collection standardized?
6. Were measures shown to be reliable and valid?
7. Were the statistical methods appropriate?
8. Were ethical issues considered?

**Studies meeting > 50% of the criteria rated as acceptable in terms of quality, studies meeting ≤ 50% of the criteria rated as low quality [22].
by re-reading the studies and subsequent discussions. When in doubt, a statistician was consulted for a final decision. Studies that met > 50% of the criteria were rated as acceptable in quality, whereas those that met ≤50% of the criteria were rated as being of low methodological quality [23]. Finally, 26 out of the 31 studies were deemed to be of acceptable quality. No studies were excluded due to low quality. Methodological shortcomings mainly concerned criteria 3, 4 and 6. Twenty one of the studies failed to explain whether and how the participants who agreed to participate differed from those who refused (criterion 3). Fifteen studies did not achieve an acceptable response rate of 65% [24], or failed to provide information about the response rate (criterion 4). Fourteen studies did not comment on the validity or reliability of the outcome measure (criterion 6). Ethical issues were considered in 26 of the 31 selected studies (criterion 8).

### 3. Results

#### 3.1. Study Characteristics

Thirty one studies published between 1991 and 2014 in English-language journals were included in the review. The studies represent European, North- and South American, as well as Asian countries. Key features of the included studies are described in Table 1. All but one longitudinal study [25] have a cross-sectional design. Fifteen were designed as single studies and 16 were sub-studies that emanated from an epidemiological study, a clinical trial or an intervention study. In the sub-studies, data collection was conducted in the original study or new information was collected from the respondents who participated in the original study, or in some cases by recruiting additional participants. Some studies partly share a database with one or more of the other included studies [25]-[28] and [29] [30]. However, they all have separate statistical analyses. In most studies, the caregiver was recruited together with the person she/he cares for. The caregiver sample size ranged from 50 to 2476, with the majority of studies having less than 100 participants, of whom spouses/partners constituted the main caregiver group. Fifteen studies provide a definition or description of caregiver burden, distress, strain or stress, referring to the concepts’ multifaceted nature of external, environmental and personal factors [9], subjective and objective burden [31] [32], emotional distress [33], enduring change of well-being [12], psychosocial load as a result of caring [29] [30], impact on social life and health [34] and difficulties in fulfilling the caregiving role [25] [26]. Five studies [35]-[39] refer to the much cited definition of caregiver burden presented by Zarit et al. [40], which defines it as “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” (p. 261). Only one study [41] has an explicitly theoretical approach, based on a stress-appraisal model. Twenty two of the 31 included studies provide multivariate statistical procedures analysing the relationship among three or more variables [24] (multiple regression, analysis of covariance, multivariate analysis of variance, factor analysis, logistic regression and path analysis). Nine studies only provide bivariate statistical tests, analysing the relation between two variables [24] (Person’s product moment correlation, Spearman’s rho, Mann-Whitney U-test, analysis of variance and t-test). Nineteen studies report sub-scores of the caregiver burden instruments, whereas 12 only report the sum scores of the burden instruments. Eleven studies use some stratification of the sample in the statistical analysis.

#### 3.2. Instruments

Eleven different generic instruments for assessing caregiver burden, distress, strain or stress were identified. The instruments cover a wide range of domains with a variety of items and sub-scales (Table 3). Eleven studies used a caregiver burden instrument based on the ones developed by Zarit et al. [42] [43]. The Caregiver Burden Inventory [44] and The Relative Stress Scale [45] were each used in three studies. Four articles [25]-[28] refer to the caregiver section of the Family Caregiving Inventory [46], although the number of scales and items used vary across studies. One study [47] used the original 10 item version of the Neuropsychiatric Inventory (NPI) [48], whereas four studies [33] [49]-[51] employed the 12 item NPI Caregiver Distress Scale (NPI-CD) [52]. The original Caregiver Strain Index (CSI) [53] was used in the study by Tanji et al. [54], whereas the modified version (MCSI) [55] with an amended response set, was employed by Kelly et al. [12] and Peters et al. [56]. The Care Management Stress Scale [57], the Caregiver Burden Scale [58], The Montgomery, Gonyea and Hooyman’s scale [59] and the Multidimensional Caregiver Strain Index [60] were used in only one study each.

Psychometric testing of internal consistency was carried out by means of Cronbach’s alpha for all but one instrument. Except for a few sub-scales the Cronbach’s alpha level was beyond the acceptable border of 0.70, de-
Table 3. Instruments assessing the impact of caregiver burden in the included studies.

<table>
<thead>
<tr>
<th>Name of the instrument (Abbreviation name)</th>
<th>Items Format</th>
<th>Subscales (items)</th>
<th>Reliability [Cronbach alpha (α) Test-retest Inter-rater reliability]</th>
<th>Validity</th>
<th>Reliability tested in PD population [Reference]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Care Management Stress Scale (CMS) [57]</td>
<td>25 items 5-Point scale</td>
<td>No subscales</td>
<td>Not available</td>
<td>Not reported</td>
<td>α = 0.94 [11]</td>
</tr>
<tr>
<td>Caregiver Burden Inventory (CBI) [44]</td>
<td>24 items 5-Point scale</td>
<td>Time-dependence burden (5) Developmental burden (5) Physical burden (4) Social burden (5) Emotional burden (5)</td>
<td>α = 0.73 - 0.86</td>
<td>Construct</td>
<td>α = 0.94 [41] [62]</td>
</tr>
<tr>
<td>Caregiver Burden Scale (CBS) [58]</td>
<td>22 items 4-Point scale</td>
<td>General strain (8) Isolation (3) Disappointment (5) Emotional involvement (3) Environment (3)</td>
<td>α = 0.70 - 0.87</td>
<td>Construct</td>
<td>α = 0.88 [31]</td>
</tr>
<tr>
<td>Caregiver Strain Index (CSI) [53]</td>
<td>13 items Dichotomous (yes/no)</td>
<td>No subscales</td>
<td>α = 0.86</td>
<td>Construct</td>
<td>Not reported [54]</td>
</tr>
<tr>
<td>Family Caregiving Inventory (FCI) [46]</td>
<td>102 items 5-Point scale</td>
<td>Strain from direct care (38) Strain from lack of resources (6) Strain from worry (10) Strain from role conflict (15) Strain from economic burden (4) Strain from mismatched expectations (5) Strain from increased tension (4) Strain from feelings of being manipulated (4) Global strain (4)</td>
<td>α &gt; 0.70 for 6 of 9 scales for which Cronbach’s alpha was reported</td>
<td>Not reported</td>
<td>α = 0.75- 0.98, except for strain from mismatched expectations, α = 0.57 [27]</td>
</tr>
<tr>
<td>Modified Caregiver Strain Index (CSI) [55]</td>
<td>13 items 3-Point scale</td>
<td>No subscales</td>
<td>α = 0.90</td>
<td>Test-retest reliability 0.88</td>
<td>Not reported</td>
</tr>
<tr>
<td>Montgomery, Gona, and Hooyman’s scale [59]</td>
<td>22 items 5-Point scale</td>
<td>Subjective burden (13) Objective burden (9)</td>
<td>α = 0.85 (objective scale)</td>
<td>Not reported</td>
<td>Not reported [32]</td>
</tr>
<tr>
<td>Multidimensional Caregiver Strain Index (MCSI) [60]</td>
<td>18 items 5-Point scale</td>
<td>Physical strain (3) Social constraints (4) Financial strain (2) Time constraints (2) Interpersonal strain (5) Elder demanding/manipulative (2)</td>
<td>α = 0.75 - 0.85 Financial Strain α = 0.58</td>
<td>Construct Criterion</td>
<td>α = 0.855 for total MCSI [65]</td>
</tr>
<tr>
<td>Neuropsychiatric Inventory Caregiver Distress Scale (NPI-CD) [52]</td>
<td>12 (10) items 6-Point scale</td>
<td>No subscales</td>
<td>Test-retest reliability r = 0.92 Interrater reliability 0.96</td>
<td>Criterion</td>
<td>Not reported [33] [47] [49]-[51]</td>
</tr>
<tr>
<td>Relative Stress Scale (RSS) [45]</td>
<td>15 items 5-Point scale</td>
<td>Personal distress (6) Life Upset (5) Negative feelings (4)</td>
<td>α = 0.72 - 0.88</td>
<td>Construct</td>
<td>α = 0.94 for ‘Psychosocial burden’ based on 8 items of the RSS [30]</td>
</tr>
<tr>
<td>Zarit Caregiver Burden Interview (ZBI) [42] [43]</td>
<td>22 items (originally 29 items) 5-point scale</td>
<td>No subscales</td>
<td>Not reported</td>
<td>Construct validity negative</td>
<td>α = 0.93 [37] [9] [34]-[39] [49] [50] [61] [63]</td>
</tr>
</tbody>
</table>
monstrating the internal consistency of the instruments [24]. In addition, construct and/or criteria validation was reported for six instruments. The validity of the instruments for use in a PD population was further confirmed for seven of the instruments with a Cronbach’s alpha > 0.75 for nearly all scales.

The included studies use various terminologies, including caregiver or carer, burden, distress, strain or stress. In this article we generally use the terms caregiver and burden. When referring to results from one particular article we use that article’s own terminology.

### 3.3. Demographic and Clinical Characteristics Related to Caregiver Burden

A range of different instruments was used to assess patient characteristics possibly associated with caregiver burden. These are listed consecutively when mentioned in the text. A summary of the synthesizing of demographics and the association of PD characteristics with caregiver burden is presented in Table 4.

<table>
<thead>
<tr>
<th>Table 4. Demographic variables and clinical aspects of Parkinson’s disease (PD) related to caregiver burden.</th>
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<tbody>
<tr>
<td><strong>Factors</strong></td>
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<td><strong>Demographic variables</strong></td>
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<td>Gender</td>
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<td>Relationship</td>
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<td>Age</td>
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<td>Socio-demographic</td>
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<td>Motor symptoms of patients</td>
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<td>PD stage</td>
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<td>Parkinsonism</td>
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<td>Falls and motor complications</td>
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<td>Duration of PD</td>
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<td>Non-motor symptoms</td>
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<td>Depression</td>
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<tr>
<td>Anxiety and apathy</td>
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<tr>
<td>Cognition</td>
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<tr>
<td>Psychotic symptoms and behavioural disturbances</td>
</tr>
<tr>
<td>Neuropsychiatric comorbidity</td>
</tr>
<tr>
<td>Other non-motor symptoms</td>
</tr>
<tr>
<td>Patient’s Quality of life</td>
</tr>
</tbody>
</table>
3.3.1. Demographic Variables

1) Gender Effects
Several studies have examined the influence of sex on caregivers’ perception of burden in PD. Most studies in the present review, using different caregiver scales and representing European, North- and South American, as well as Asian cultures, did not find a patient or caregiver gender effect on caregiver burden scores [9] [30]-[32] [38] [39] [61]-[63], although some found that female caregivers tend to report more burden [33] [34] [37]. Female sex in spouse caregivers was also found to be a predictor of role strain over a 10 year period, when statistically controlled for the effect of PD stage and spouses’ age [25]. Likewise, male sex in the patient revealed a significant association with higher caregiver strain scores in multivariate analysis [64] [65].

2) Relationship
A few studies explored the association between caregiver burden and caregivers’ relationship with the care recipient. In a study reporting on a quality improvement registry including 2476 patient-caregiver couples in the US, spouses/partners (constituting 91%) were less likely to report higher levels of caregiver strain on the Multi-dimensional Caregiver Strain Index (MCSI) compared to non-spouses [65]. Likewise, in a small-size Turkish study (n = 57), adult child caregivers reported significantly higher sum scores and personal and role strain sub-scores on the Zarit burden Inventory (ZBI) than spouse caregivers, even without any significant differences in terms of caregiving characteristics [34]. Two Asian studies [39] [63] as well as another Turkish one [38], also using the ZBI, found no statistically significant differences in burden scores between spouses and non-spouses, although factors contributing to caregiver burden differed between spouses and offspring in the study by Shin et al. [63]. In addition, in a South Korean study [32] subjective burden, referring to the affective and perceived changes in life due to caregiving activities, was higher in the spousal group compared to non-spouses, whereas objective burden, defined as how caregivers perceive the disruption caused to their life by caregiving, did not differ between spouses and non-spouses. Moreover, in a Norwegian study spouses reported higher levels of caregiver stress on the “Life upset” subscale of the Relative Stress Scale (RSS) compared to non-spouses, whereas there were no differences between the caregiver groups regarding the RSS “Negative feelings” and “Personal distress” subscales [29].

3) Age
Many studies across various continents have found that caregiver burden is unrelated to patient and/or caregiver age [11] [12] [29]-[31] [36] [38] [49] [54] [61] [63] [66] even after adjustment for disease duration [62]. Nevertheless, age may matter in PD caregiving. In a multi-centre North American study [26] younger spouses (40 - 55 years) compared to older spouses (>70 years) reported more strain from the role strain dimension “Lack of personal resources” but not from the dimensions “Strain from worry” and “Global strain”. In this study a significant amount of the variance of strain from ‘Lack of personal resources’ was explained by age [26]. Unlike in the South Korean study by Kim et al. [32], which compared even younger caregivers (≤40 years) with groups of older (41 - 64 years and ≥ 65 years) caregivers, the youngest reported significantly lower subjective burden than the older ones, with the eldest reporting the most burden. In addition, Calder [64] found that young age in patients contributed to the variation in levels of stress in a group of Scottish PD relatives (mainly spouses). No correlation was reported between age at PD onset and burden scores [35] [36] [39] [51], except in a subgroup of caregivers of PD patients with dementia [51] and in a study by Martinez-Martin et al. [37].

4) Sociodemographic Factors
Marital status, educational level, as well as occupational and financial status were seldom addressed in the identified studies and not found to be associated with caregiver burden [34] [38] [63]. Two studies [39] [54] reported cultural variation in caregiver strain. In a Malaysian study a significant difference in mean burden score was found between caregivers representing four different races [39]. In contrast, the caregiver strain sum scores were equal in a comparative study of Japanese and US spouse caregivers. However, the Japanese group disclosed significantly more burden on single items related to time demands and physical and financial strain on the Caregiver Strain Index, while the US group scored higher on emotional strain items [54].

3.3.2. Motor Symptoms

1) PD Stage
The relationship between caregiver burden and severity of PD as assessed by the Hoehn & Yahr (H & Y) disease stage scale [67] was thoroughly documented in the identified studies [9] [29] [31] [32] [35]-[39] [49] [51] [54] [61] [63]-[66] [68]. Caregiver strain has been reported at all stages of PD [27] and burden scores increase
significantly in line with advancing H & Y stage [27] [35] [61]. PD stage was found to be an important predictor of caregiver burden scores in multivariate analysis [31] [65] [66], even after controlling for sex and cognitive deterioration [64]. In addition, caregivers of patients with a higher disease stage at baseline were more likely to report superior levels of certain role strain variables (“global strain” and “strain from worry”) after 10 years [25].

2) Parkinsonism
Correlation analyses between caregiver burden and the severity of Parkinsonism as measured by the Unified Parkinson’s Disease Rating Scales [69]-motor scale (UPDRS III) and the motor scale of the Scales for Outcomes in Parkinson’s Disease (SCOPA-Motor) [70] confirms the association between disease severity of PD and increased burden scores [9] [29] [35] [36] [38] [49] [61] [63]. Likewise, correlation analysis between sum scores on different caregiver burden scales and functional ability as assessed by various activity of daily living (ADL) scales including the UPDRS II, Schwab & England ADL scale (S & E) [71], ADL subscale of The SCOPA-Motor (SCOPA-ADL) [70], Barthel Index (BI) [72] and Reported Self Care Scale (RSCS) [73] supports the reported relationship between caregiver burden and severity of Parkinsonism [30] [32] [37] [38] [51] [54] [62] [63]. Several studies report the effect of functional disability on the overall caregiver burden based on multivariate analysis [9] [41] [50] [64]. In the study by Aarsland et al. [29], functional impairment (S & E) contributed to the spouses’ “Personal distress” scores as assessed by the RSS but not to the ‘Negative feelings’ and ‘Life upset’ subscales. Other studies found no effect of patient ADL function in multivariate analysis [28] [30] [35] [54] [61] [63]. Severity of motor symptoms was found to be an independent predictor of caregiver burden in the study by Agrawal et al. [35] and a subgroup of offspring caregivers in the study by Shin et al. [63].

3) Falls and Motor Complications
The incidence of falls, a key variable of functional impairment in PD [74], is sparsely examined in a caregiving context. However, in one study [62] patient falls were strongly associated with burden in caregiver partners. Likewise, in the large scale US registry study [65], a higher frequency of falls was associated with greater levels of caregiver strain. In the comparative study of strain between caregiver spouses in Japan and the US, multivariate analysis revealed falls to be a predictor of caregiver strain in the Japanese group, but not in the US group [54]. Beyond this, a few studies report a positive correlation between caregiver burden and the presence of motor complications such as motor fluctuations and/or involuntary movements [9] [35] [36] [38] [62] [63] [65]. In the study by Shin et al. [63] a correlation was found in caregiver spouses but not in offspring.

4) Duration of PD
While several studies have observed a positive correlation between duration of PD and higher burden scores [9] [25] [31] [35] [56] [62] [65], Kim et al. [32] found the emotional (subjective) burden to be highest 1 - 5 years after the patients received their PD diagnosis. Additionally, two studies could not identify a significant correlation between PD duration and caregiver burden [37] [51]. When comparing caregiver strain in a Japanese and a US cohort, PD duration correlated with caregiver strain in the US cohort, but not in the Japanese [54].

3.3.3. Non-Motor Symptoms
1) Depression
Depression is the mental symptom in PD most often examined in a caregiver burden context. A range of self-report and interview instruments were used to explore the relationship between patient depression and caregiver burden, including the Beck Depression Inventory [75], Brief Symptom Inventory [76], Center for Epidemiological Studies Depression Scale [77], Geriatric Depression Scale [78], Hamilton Depression Rating Scale [79], Hospital Anxiety and Depression scale [80] and Montgomery-Aasberg Depression Rating Scale [81]. Several studies using various combinations of depression scales and burden instruments report positive correlations [9] [30] [31] [38] [54] [61] [62] [66] and the predictive power of depression on caregiver burden [28]-[30] [34]- [36] [54].

2) Anxiety and Apathy
Patient anxiety and apathy was sparsely addressed in the reviewed studies. Nevertheless, patient apathy was among the most frequently reported neuropsychiatric symptoms to be associated with caregiver distress on the NPI-D in a cohort of newly diagnosed PD patients [33] and a study of PD patients with dementia (PDD) [47]. Three studies report a positive correlation between patient anxiety and caregiver burden scores [37] [38] [61].

3) Cognition
In PD, even early detectable cognitive impairment assessed by testing the patient’s verbal fluency and delayed
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word recall [82], were associated with higher levels of strain in caregivers [28] [65]. The contribution of delayed recall was significant for five of the six role strain variables assessed in the study by Carter et al. [28] (“global strain”, “strain from worry”, “manipulation”, “tension” and “strain from frustration due to communication problems”). In the study by Kudlicka et al. [66] caregiver rated executive function impairment, a common cognitive deficit observed in the early stages of PD [83], was the strongest predictor of caregiver burden. The relationship between impaired global cognition in the patient and higher burden scores is further supported by bivariate [54] [63] and multivariate analysis [9] [29] [30] [49] using the generic Mini Mental State Examination [84] or the PD specific Scales for Outcomes in Parkinson’s disease-cognition scale [85].

4) Psychotic Symptoms and Behaviour Disturbances

Psychotic symptoms as assessed by the Parkinson Psychosis Rating Scale [86] have a moderate correlation [9] and make an independent contribution [61] to caregiver ZBI scores. Likewise, hallucination and confusion in the patient were among the key symptoms that PD partners associated most strongly with caregiver burden as assessed on the Caregiver-burden Inventory in the study by Schrag et al. [62], whereas no correlation was found in a mixed group of PD caregivers (50% spouses) [35] on the ZBI. In an international multicentre study of PDD caregivers who were clustered on the basis of the patient’s NPI symptoms [47], caregivers of the psychosis group (characterized by high scores on visual hallucinations and delusions) exhibited the second highest caregiver distress (NPI-CD) sum score after the agitation group (including irritability and agitation). In addition, behavioural problems in PD, as identified by the sum scores on the CAPE Behavior Rating Scale [87], Behavioral Problems Scale [88], Behavior Rating Inventory of Executive Function [89] and the Behavior Pathology of Alzheimer’s Disease Frequency Weighted scale [90], have been demonstrated by regression analyses to have a significant effect on various caregiver burden measures [34] [41] [64] [66].

5) Neuropsychiatric Comorbidity

Caregivers of PD patients diagnosed with psychosis [50], impulse control disorders [36], dementia [51] and apathy [36] reported significantly higher burden scores compared to caregivers of patients without neuropsychiatric comorbidity. Likewise, patient depression generated a higher caregiver burden compared to non-depressed patients [51] also when controlled for the effect of patient ADL-scores [37]. ADL-scores in addition to patient age were also controlled for in the study by Leroy et al. [49], who found a significantly greater burden on the ZBI scale of the caregivers of PDD patients compared to non-demented patients. However, no difference between the caregiver groups was observed on the NPI-CD scale in the same study. In the study by Calder et al. [64], the observed differences in stress scores (RSS) between caregivers of patients with and without dementia, were non-significant after controlling for the effect of disability as assessed by H&Y stage. Nevertheless, the influence of patients’ neuropsychiatric status on caregiver burden is well documented in the studies, indicating that in regression analyses neuropsychiatric symptoms have a stronger impact on caregiver burden than motor symptoms or functional impairment [28]-[30] [34] [35] [50] [61].

3.3.4. Other Non-Motor Symptoms

Non-motor problems such as patient fatigue, autonomic dysfunction, pain and sleep disturbance are sparsely addressed in relation to caregiver burden in PD. One study [61], reporting the sum scores of the Non-motor Symptom Assessment Scale for Parkinson’s Disease [91], which covers nine non-motor domains, found a moderate association with the ZBI sum score. In the study by Agrawal et al. [35] sleep disturbances in patients, but not autonomic symptoms, were identified as a significant predictor of caregiver burden after regression analysis. In the study by Martinez-Martín et al. [9], a range of patient-related variables were embedded in factor analysis, resulting in 4 factors of which one included autonomic dysfunction, pain and fatigue, while a second was linked to sleep disorders. Both factors had an independent impact on caregiver burden [9]. Several other studies report a positive correlation [38] or an independent contribution of patient sleep problems on caregiver burden [35] [61]. Unexpectedly, in the only two studies reporting correlations between caregiver strain and unspecific patient comorbidity, the latter did not relate to the level of caregiver strain [65] and was not associated with caregiver strain after regression analysis [54].

3.3.5. Patients’ Quality of Life

Several studies [12] [31] [56] [62] [65] demonstrate an association between higher caregiver burden and a decrease in patient self-reported QoL, based on the sum score of the 39-item Parkinson’s Disease Questionnaire (PDQ-39) [92]. In Peters et al. [56] the PDQ-39 subscales Mobility and Social Support were identified as the
most likely to affect the Caregiver Strain Index scores, when adjusted for duration of disease, years spent as a caregiver, hours spent on caring per week and the sex of patient and caregiver. PDQ-mobility was also the subscale associated with a high caregiver strain score on the Multidimensional Caregiver Strain Index in the US registry study [65], followed by the PDQ-39 emotional subscale. The relation between caregiver burden and patient QoL is further confirmed when using the generic EuroQoL [93], [9] [12] [37] [61] and the 12-item Short Form Health Survey, version 2 (SF-12v2) [94] [56], which was expected due to the observed positive correlation of these scales with the PDQ-39 [56] [62]. In a recently published study Kudlicka et al. [66] claimed that questionnaires used for assessing QoL tend to ask about self-assessed health status rather than the person’s subjective satisfaction with life. Accordingly, the authors supplemented the PDQ-39 by the Question on Life Satisfaction scale (QLS) [95], which includes three domains: general life satisfaction, satisfaction with health and satisfaction with health in relation to movement disorders. Despite a strong correlation between PDQ-39 sum scores and the QLS health and movement disorders subscales, caregiver burden assessed by the Caregiver Burden Inventory correlated with the PDQ-39, but was not related to the QLS domains [66].

4. Discussion

Analysis of the included studies revealed great variation in how the impact of clinical characteristics on caregiver burden is studied in a PD setting. Eleven caregiver burden instruments were identified in the 31 reviewed articles, with the ZBI being the most frequently used. In addition, a large number of different scales were employed to identify specific characteristics that may influence caregiver burden. Consequently, many combinations of instruments were involved in the statistical analyses, making it difficult to review the results in a systematic way. Most studies only report on the association of patient related factors with the sum score of caregiver burden instruments. A few provide analyses of the subgroups of a heterogeneous study sample. All studies with the exception of one had a cross-sectional design. Accordingly, there is a lack of knowledge about the progression of burden in PD caregiving and the factors that influence burden over time.

4.1. How Did the Demographic and Clinical Characteristics of Patients with PD Influence Caregiver Burden?

Despite the great variety of instruments used in the analysis, similar associations were found between caregiver burden, demographic variables and certain patient characteristics across the reviewed studies. There is clear statistical evidence of the relationship between the severity of PD and increased caregiver burden. In particular, higher PD stage and patients’ problems performing ADL contribute to caregivers’ burden. Duration of PD, motor symptoms and complications were not identified as predictors of caregiver burden. Associations between caregiver burden and non-motor symptoms of PD, such as depression and psychosis, cognitive impairment and behavioural problems, are well documented in the reviewed studies. Moreover, several studies report patient depression and behavioural problems as predictors of caregiver burden. In addition, comorbidity of depression, apathy, psychosis, dementia and impulse control disorders in patients correlate with significantly greater burden for caregivers, whereas the individual contribution of patient dementia on caregiver burden was inconclusive. Furthermore, when comparing the impact of patient motor and non-motor symptoms on caregiver burden, several studies found that patients’ mental symptoms have a stronger impact on caregiver burden than motor symptoms or functional impairment. This finding differs from the review by Lau et al. [10], who revealed that patient motor symptoms and dependency in ADL were most strongly correlated with caregiver distress, while the weakest association was the patient’s cognition. The different inclusion criteria applied in the review by Lau et al. [10] and the present review, which resulted in an overlap of only 6 studies, might explain the divergent result. The correlation between burden, assessed by various caregiver burden instruments, and patient HRQoL measured by disease specific as well as generic instruments, is clear. The only included study that employed a QoL instrument that goes beyond a health status assessment did not identify a significant relationship between QoL and caregiver burden. As the authors suggest, this indicates the relevance of studying subjective QoL and self-assessed health status as separate concepts [66].

Despite the fact that many of the authors comment on cultural and ethnic family structures, as well as caring traditions that might influence caregiver burden in their sample [32] [34] [35] [38] [39] [51] [61] [63], there is no obvious evidence of variation in caregiver burden between studies representing European, North- and South American, as well as Asian cultures. The only international comparative study [54] observed an equal amount of
total strain in Japanese and US caregivers despite the differences in demographics and social support, although the two groups differed significantly on several single strain items and the factors that contribute to caregiver strain. Contrary to the expectations due to different caring traditions, the included studies reveal little gender difference in caregiver burden. In contrast, in their meta-analysis of 164 studies of different caregiver samples, Pinquart and Sörensen [96] reported a small but statistically and practically significant gender difference in caregiver burden, with women reporting the highest burden. The influence of patient or caregiver age on reported caregiver burden was found in two studies comparing various caregiver age groups. However, when comparing the younger with the older group the two studies were contradictory in terms of which group was the most burdened. There is also conflicting evidence in the studies about whether caregiver spouses are more burdened than non-spouses, as spouses and non-spouses tend to differ with respect to burden domains, how they are affected and which factors contribute to caregiver burden. In addition, other aspects such as the emotional relationship might explain the inconsistent results. In a literature review carried out by Carbonneau et al. [97], the quality of the relationship was found to impact on caregiver stress. This is supported by a few of the reviewed studies, in which it is stated that caregivers’ assessment of the quality of their relationship with the care recipient [25] [41] and mutuality in the relationship [25] [26] had a protective effect on caregiver burden.

4.2. What Instruments and Which Analysis Methods Were Used?

A number of instruments are available for measuring caregiver burden. Durme et al. [14] identified 55 scales (mainly generic) for assessing the negative impact of caregiving, 42 of which evaluate burden, strain or stress as the main dimension. There could be various reasons for choosing one instrument over another. In this review, the ZBI was the instrument used in most studies (11 studies). Durme et al. [14] rated the ZBI as the preferred caregiver burden instrument and good reliability has been reported in a PD setting [37]. According to Deeken et al. [15] and Den Oudsten et al. [98], the choice of outcome instrument in research should be based on a clear understanding of the phenomenon under study. In the present review, 15 of the 31 studies presented a definition or conceptualization of caregiver burden, distress, strain or stress, while one study had an explicitly theoretical approach. The diversity of burden instruments, lack of a specific definition of burden and the absence of validity testing in a PD setting make it challenging to standardize tools for assessing caregiver burden in PD. Lack of conceptual consistency and standardization of burden measures leads to problems comparing the burden outcomes across instruments. This was illustrated in the study by Leroi et al. [49], who included two caregiver burden instruments. Significant differences were found between caregivers of PDD patients compared to those caring for non-PDD patients when assessed by the ZBI, but not by the NPI-CD. To achieve a more unequivocal picture reflecting the burden of PD caregivers, a smaller number of consistent instruments should be used when planning future studies. Recommendations should be made about the most appropriate caregiver burden instruments for a PD setting, as has been done for other factors associated with PD, including depression [99], fatigue [100], psychoses [101] and HRQoL [102].

With regard to the choice of caregiver burden instruments, a wide range of instruments were chosen in the selected studies to measure similar patient characteristics. For instance, five and seven different scales were used to measure patient functional ability and patient depression respectively. Practical or professional reasons within the research team might explain these choices. However, the variation in assessment tools impedes the opportunity to determine whether equal results indicate the same outcome and whether different results mirror real disparities. A lesser number of scales would make it possible to statistically integrate the results of the associations between caregiver burden and patient characteristics in a meta-analysis, which was not done in this review.

Comparison between studies also requires a clearly defined sampling unit in each study. Six of the 31 selected studies only included spouse or partner caregivers. However, in most studies the analyses are limited to the total group of caregivers. The absence of considerations about and stratification of subgroups in the analysis phase of caregiver studies has previously been criticized for limiting the understanding of caregiver burden [103]. This view is supported by the studies in the present review that provide separate analysis for different caregiver groups or control for variables that may influence the results. For instant, patient sex [25] [64] [65], comorbidity [36] [37] [49]-[51] and the patient-caregiver relationship [34] [65] were differently correlated with caregiver burden in distinct groups of caregivers. In addition, multivariate analysis provided further information about how caregiver age [26], presence of dementia in the patient [64], as well as the patient’s depression and UPDRS mentation and motor scores [63] might make a different contribution to burden across caregiver groups.
The various caregiver scales, number of subscales and the different scope of items used in the reported studies demonstrate the multidimensional nature of caregiver burden. However, several of the included studies treat burden as a one-dimensional variable and only present a sum score showing the overall burden without any information about sub-scores. For many years lack of such information has been criticized for not permitting a distinction to be made between various dimensions of burden [44], for masking variation in the correlates between different patient characteristics and the dimensions of strain [60], and for overlooking the relative contribution of specific domains of burden to the overall burden scores [104]. The few included studies that provide information about subscales partly address this criticism. An example is the comparative study of Japanese and American caregivers in which the CSI sum scores were the same, whereas single item scores differed significantly between the two caregiver groups [54]. Furthermore, the study by Kim et al. [32] demonstrates that several patient characteristics were differently associated with caregivers’ subjective and objective burden scores. In addition, patient functional disability [28] [29], caregiver age [26] [32] and the spouses’ relationship with the patient [29] have different associations within various subscales. Another aspect of the multifaceted character of caregiver burden is the assumption that it is influenced by many different factors. The majority of the selected studies considered this problem as the researchers performed a multivariate analysis. However, in contrast to the review by Greenwell et al. [18] who only included studies that identified predictive factors, the present study contains a correlation analysis to include studies that did not meet the criteria for a multiple regression analysis [105].

4.3. Limitations
This systematic review has several limitations. A meta-analysis was not possible due to the diversity of the instruments used in the various studies. In addition, only quantitative studies using a caregiver burden instrument as an outcome measure of caregiver stress and burden were included. Accordingly, studies employing other methods and instruments that could provide further knowledge about the subject were excluded. Finally, to fully understand the complexity of caring experiences and to plan effective interventions, caregiver-related variables as well as those that may be associated with positive outcomes for the caregiver are required. In addition, the risk of bias needs to be discussed with regard to how the systematic review was carried out. Two of the authors independently assessed the methodological quality. Differences in judgment were resolved by discussion between the authors, who agreed on the final version. However, it is important to acknowledge that this review represents the authors’ understanding of the studies analysed. When conducting a review, researchers are in danger of making choices regarding the selection of studies and data extraction that expose the review to the risk of bias. However, we have strived to reduce the risk of bias by means of a comprehensive search in several electronic databases, formulating explicit inclusion and exclusion criteria for these selection of studies and by cooperation between authors in the data extraction.

5. Conclusions and Implications
Many patient characteristics are thoroughly documented as having an impact on caregiver burden in PD. Patient non-motor symptoms seem to have a stronger impact on caregiver burden than motor symptoms or functional impairment. Nevertheless, there is a need to fill a gap in the knowledge about how non-motor symptoms associated with PD such as patient anxiety, apathy, fatigue, sleep problems and the frequency of falls influence caregiver burden. Future research should also give priority to longitudinal studies in order to deepen the understanding of caregiver burden over time, as well as the expected needs in the caregiving situation. Furthermore, when planning studies on caregiver burden, a more uniform selection of burden instruments and scales to assess clinical characteristics would facilitate comparison among studies, thus deepening the knowledge of caregiver burden in PD.

In most of the included studies the analysis is restricted to the sum scores of the caregiver burden instruments. The majority of studies involved caregivers with different relationships to the care recipient, although relatively few studies contained a subgroup analysis of the caregiver respondents. Accordingly, there is a lack of knowledge about which areas cause the greatest burden to caregivers and whether some caregivers are more vulnerable and if so, for what reason. For clinical purposes, it would be appropriate to conduct subscale and subgroup analyses of caregivers’ experiences in order to provide differentiated and targeted approaches to caregiver needs. This could facilitate healthcare professionals to plan appropriate intervention programmes that might reduce the
burden on caregivers and prevent or postpone institutionalization.

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**Competing Interest**

The author’s declare that there are no conflicts of interest.

**Contributors**

The study was designed by I.L. I.L. coordinated the research. All authors participated in the data analysis. The report was written by I.L. E.S. supervised the study.

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