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[Anil Gumber](#) and [Smruti Bulsari](#) *

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Article

Financial Distress and Wellbeing of People with Parkinson's and Their Informal Caregivers: Evidence from the United Kingdom

Anil Gumber ¹ and Smruti Bulsari ^{2,*}

¹ Research Centre for Healthcare & Communities, Coventry University, Coventry CV1 5FB, UK

² Institute of Public Health and Wellbeing, University of Essex, Colchester CO4 3SQ, UK

* Correspondence: smruti.bulsari@essex.ac.uk

Abstract: Background: Parkinson's Disease (PD) is a long-term condition, which is more common among people aged 65 years or more. People with Parkinson (PwP) tend to lose motor functions and find themselves in more fatigue and pain, sometimes leading to anxiety, distress and in extreme cases, even depression. So far, there is no cure for PD and they need support from family members, healthcare professionals and carers, as the disease progresses. A large proportion of carers for PwP are their spouses. Progression of PD can result in loss of employment, reduced work hours or premature retirement, for both PwP and their carers. This poses a huge financial burden as income reduces on one hand, and expenses increase on the other. There is a lack of evidence on the societal costs of Parkinson's disease, and this study tries to fill that gap. **Objectives:** To examine the impact of financial distress on quality of life (QoL) and wellbeing of PwP and their carers in the UK. **Methods:** A mix of online and postal, cross-sectional survey was administered, which received responses from 853 respondents, out of which 776 were PwP and 546 were carers. The survey sought data on management of the socioeconomic, health and living conditions of PwP and their carers. Net effect of financial distress (cost of healthcare use, non-health expenditure, societal costs and other direct / indirect cost components) on QoL (and VAS scores) as well as wellbeing scores (life satisfaction and happiness scores) of PwP and their primary carers are estimated using simple linear regression models. **Results:** Financial burden of treatment increase whereas quality of life and wellbeing of PwP decrease with progression of PD. Similar association but of a milder magnitude is observed on quality of life of their primary carers. A strong negative association is found between quality of life of PwP, with financial burden, while holding for demographic and socioeconomic factors, and for disease progression and treatment. The QoL of carers increase up to a financial burden threshold, before it starts declining. This could be because of the increased expenditure in initial phases is associated with improved activities of daily living and lesser reliance on carers. Financial distress is more pronounced among PwP having annual income below £20,000. Despite a small representation of persons from ethnic minority (5 per cent representation), it is observed that the impact of financial distress could be more pronounced on PwPs from other ethnicities. Also, it would be more on women than on men, irrespective of ethnicity. **Conclusion and Policy Implications:** Cost of care cuts both ways, reduced or lost employment as well as increased expenses on healthcare. This results into a considerable financial distress and deterioration in quality of life and wellbeing of both PwP and carers. Most carers are spouses of PwPs, thus their physical and psychological health and wellbeing is found to deteriorate with the progression of PD due to increased burden of care. Therefore, informal / family carers should get support in the form of respite care. This can help revitalise their energy and psychological wellbeing. Financial distress indicators need to be explored further.

Keywords: financial distress; Parkinson's; people with Parkinson's; informal care; quality of life; wellbeing

1. Background

About 153,000 people are living with Parkinson's disease in the UK, and one in 37 persons get diagnosed with Parkinson's in their lifetime. This is expected to increase to 172,000 by 2030 (Parkinson's UK, 2023). Parkinson's is a neurodegenerative disorder that affects movement and

coordination. Some common symptoms of Parkinson's include tremors, stiffness, and impaired balance. All these adversely affects the quality of life (QoL) of the PwP as well as of their caregivers.

Rahman et al., (2008); and Schrag et al., (2000) found that the QoL of a PwP is adversely affected because of sudden unpredictable on/off states, depression, disability, postural instability, difficulty in walking, difficulty in dressing, falls, urinary incontinence, pain, cognitive impairment and confusion. Rahman et al., (2008) further found that hesitation to begin walking, shuffling gait, freezing, festination, propulsion, and difficulty in turning also had influence on their QoL.

Diagnosis of Parkinson's not only affects the quality of life of the person diagnosed with Parkinson's but also that of their carers. In most cases, where the carers are family members (McCrone et al., 2007), the direct and indirect costs of managing the condition are borne by both – the PwP and their carer – more specifically much higher in advanced stages. Gumber et al., (2019) found that there is deterioration in Quality of Life (QoL) of PwP and their carers and family members with the progression of disease. This can be because of family members volunteering their time, altering their employment status, utilising their own resources, developing stress and health problems alongside the deteriorating condition of the PwP.

Informal carers of PwP in advanced stages, especially when the PwP's cognition is adversely affected, exhibit a significant decreased in QoL (Henry et al., 2020; Rosqvist et al., 2022). They also found that the deterioration in QoL of carers of PwP with a high non-motor symptoms (NMS) burden is higher, especially if the carer was a woman. Of all the informal carers, spousal caregivers are most affected. Modugno et al., (2020) observed that the obligations of carers towards PwP care increased with disease progression and is not just limited to assisting with activities of daily living (ADL). This adversely affects the QoL of informal carers. Santos-García et al., (2022) found that burden, strain, mood, and QoL of the principal carer of a PwP worsened after a 2-year follow-up study. The changes in mood of the PwP and their carers with disease progression add to the carers' both economic and social burden.

These are the primary contributors to a worsened QoL of PwP and their carers on financial condition. The economic burden associated with the condition, and its progression, exacerbates the deterioration in the QoL of PwP and their carers. Chaudhuri et al., (2024) found that the economic burden of Parkinson's increased with its progression, and the incremental economic costs for advanced stage condition was higher by an average of £1069 as compared to mild-to-moderate stages. Economic burden of Parkinson's is estimated for PwP and different stakeholders at different points of time. Weir et al., (2018) found that the mean annual health care cost was higher by £2471 (2013 constant prices) for PwP as compared to those who did not have Parkinson's per person in the first year post-diagnosis. This increased to £4004 per PwP by 10 years following diagnosis. Costs for PwP with markers of advanced Parkinson's (i.e., "presence of levodopa-equivalent daily dose > 1100 mg, dyskinesias, falls, dementia, psychosis, hospital admission primarily due to PD, or nursing home placement") were higher by £1069 per PwP as compared to those without it; all these estimates are at 2013 constant prices.

L. J. Findley et al., (2011) estimated that of all the healthcare costs incurred for an PwP, only 7% contributed to direct medical care. The remaining 93% of costs were comprised of direct non-medical professional care (50%) and indirect informal care (43%). L. J. Findley, (2007) estimated that the annual costs of direct and indirect costs of Parkinson's ranged between £449 million and £3.3 billion in the UK. They also found that more than direct costs, which comprises largely of inpatient care, nursing home costs, and prescription drugs, the indirect costs, comprising of productivity losses and carer burden are far higher. McCrone et al., (2007) found that the annual service costs (baseline and follow-up combined) were £13,804 per person, of which 80% component was informal care provided by family / friends. They also found that 42% of the follow up costs were because of disability and depression. Moreover, these costs were higher for men as compared to women. L. Findley et al., (2003) had estimated per PwP costs for different stages of Parkinson's. They further estimated that 38% of direct costs of care is accounted towards NHS costs, 34% towards social services. However, a move from home to residential care for PwP resulted in an approximately 500% increase in the cost of care.

This explains the exponential growth in the economic costs of treatment and care for PwP with the progression of Parkinson's. Effective treatment and care for Parkinson's aims to manage symptoms and improve quality of life. However, the financial distress caused by treatment and care costs can impact the quality of Parkinson's treatment and subsequently, the QoL of the PwP. Addressing financial distress is essential for ensuring access to effective treatment and care of the person with Parkinson's as well as safeguarding the wellbeing of the informal carers. PwPs and their families face increased financial burden due to rising in out-of-pocket to meet healthcare needs including procuring specialized equipment or making modifications in the house / living arrangements to support the PwP with activities of daily living. Rehabilitation, exercise to loosen joint structures and motion exercises to prevent deformities may add to the expenses for using private physiotherapy and occupational therapy together with procuring specialised equipment to improve body movements. This is over and above prescription / medication expenses.

Costs to carers and lost income due to disability adds to the financial strain. Household expenses augmented with loss of household income in providing care, adds to the distress.

Thus, physical and psychological wellbeing which is already affected by the diagnosis of Parkinson's itself, and its progression, has a compounding effect because of financial distress caused by out-of-pocket expenses and loss of income. PwP and their carers often are required to reduce working hours or take premature retirement due to ill health and caring responsibilities, respectively.

The main objective of this paper is to examine the impact of financial distress on PwP households on their quality of life, and psychological wellbeing. We have also explored how financial distress affects the quality of life of their informal caregivers¹. Additionally, the study also assesses the influence of socioeconomic background and demographic profile of PwP on their quality of life, and psychological wellbeing using multivariate regression models. This epidemiological study is undertaken in the United Kingdom.

The paper is organised into six sections: Section 2 explain the health economics evaluation framework in the context of economic costs to PwP, their carer and all other stakeholders. Section 3 explains the methodology adopted to estimate financial distress and the patterns in socioeconomic and demographic profile of PwP and their caregivers. Section 4 discusses the patterns in QoL of PwP and their carers by the magnitude of financial distress. Section 5 presents the regression results, which gives insights into the association of financial distress with that of QoL and physical as well as psychological wellbeing of PwP. The key findings of this study are highlighted in section 6; and the policy prescriptions to improve the QoL and wellbeing of PwP and their caregivers are discussed in the same section.

2. Health Economics Evaluation Framework

Costs to persons living with Parkinson's (PwP), NHS, Local Government and employer / society are estimated using the health economics evaluation framework. This study focuses largely on costs to PwPs and their caregivers. Costs are further segregated into direct costs, indirect costs and implicit costs. Figure 1a,b represent the economic costs to PwP, their caregivers and other stakeholders.

In this study, we focus only on the costs to PwP and their caregivers for estimating the financial distress. Since the objective of our study is to examine the impact of financial distress on PwP and their informal carers, we have not taken into consideration, the costs to NHS, local government or the society / employer.

¹ Informal carers are interchangeably referred to as carers or caregivers in this paper.

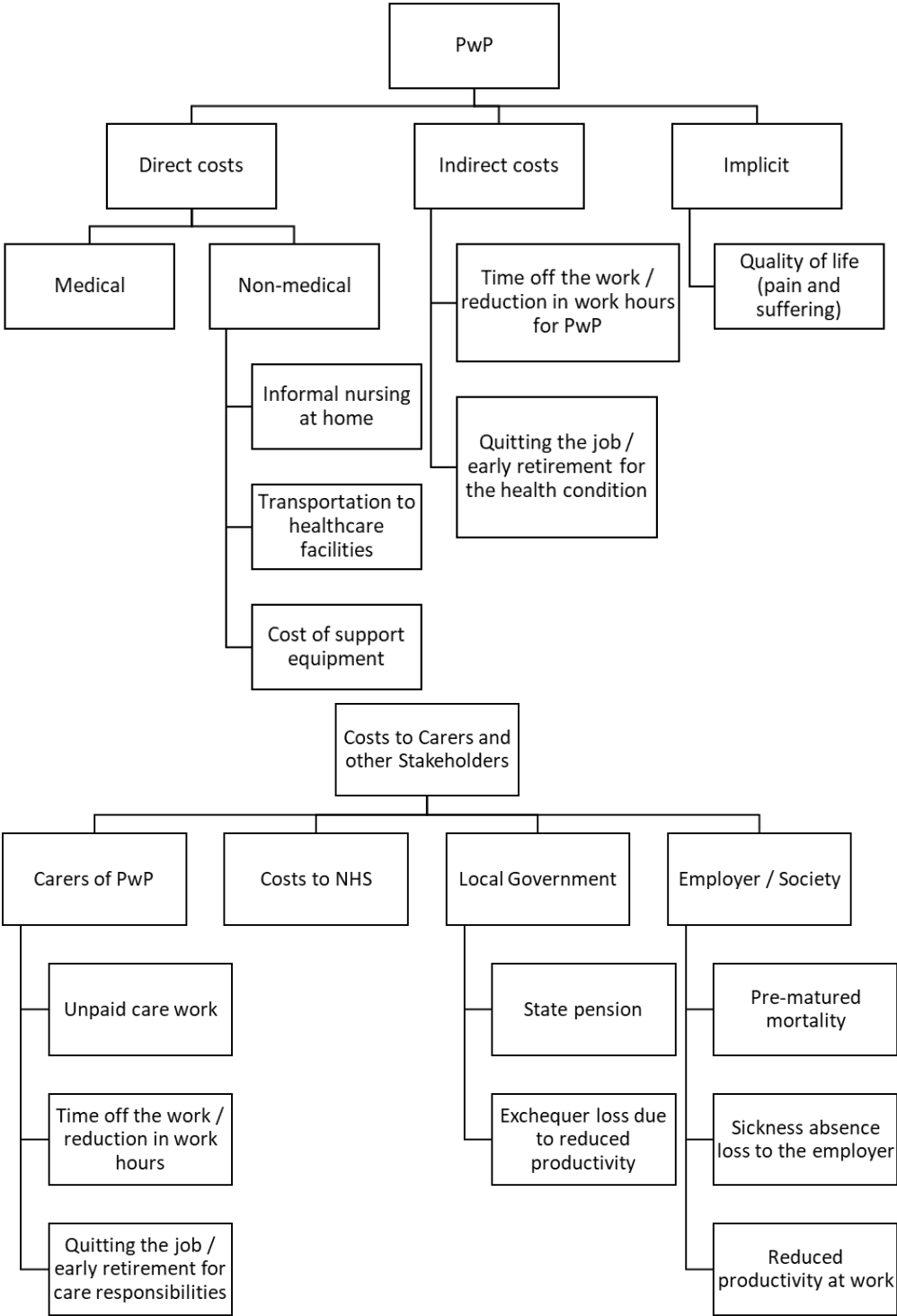


Figure 1. (a): Economic Costs for Persons Living with Parkinson’s; **(b):** Economic Costs for Carers and other Stakeholders.

3. Methods and Data

This study employs a cross-sectional design and administers a questionnaire for collecting data on health and living conditions, socioeconomic status, key items on economic and financial costs and quality of life (QoL) of PwP and their caregivers. QoL is measured using the EuroQoL EQ-5D-5L questionnaire and then Quality Adjusted Life Years (QALY) is estimated using England population weighting (N. Devlin et al., 2016; N. J. Devlin & Brooks, 2017; EuroQol Research Foundation, 2019).

The sampling method and sample size are discussed in details in Gumber et al., (2017)²; this paper is an extension of that study. Their study had adopted a mixed-methods approach, of administering a questionnaire as well as in-depth interviews. However, in this paper, we make use of the data collected using questionnaire only.

A questionnaire was administered on 776 PwPs and 546 carers, across all four countries of the United Kingdom during 2015-16. However, this had resulted in an over-representation of respondents of White ethnicity and therefore, a booster sample of 41 PwPs and their 37 carers belonging to Black and Minority Ethnicity (BME) was drawn from Manchester and Sheffield areas.

This questionnaire was administered to collect data on socioeconomic and demographic profile of PwP and their carers. The data on different components of costs to PwP, their carers and other stakeholders were also collected using this questionnaire. Data were collected, for PwP and their carers, through both, online and postal questionnaires, as per the choice of the respondent. Table 1 gives an overview of the socioeconomic and demographic profile of these respondents:

Table 1. Brief Socioeconomic Profile of PwP and their Carers included in the Study.

Respondent Characteristics	PwP		Carers	
	N	Percentage	N	Percentage
Gender				
Male	472	60.8	174	31.9
Female	298	38.4	364	66.7
Others / Preferred not to say	1	0.1	8	1.5
No response	5	0.6		
Marital Status				
Single / widowed / divorced separated/ other	156	20.1	72	13.2
Married / cohabiting	610	78.6	465	85.2
No response	10	1.3	9	1.6
Education				
All except higher education	420	54.1	296	54.2
Higher education	345	44.5	344	43.4
No response	11	1.4	13	2.4
Ethnic Group				
White	714	92.0	504	92.3
Non-white (including others)	41	5.3	37	4.7
No response	21	2.7	5	0.9
House Ownership				
Owned	569	73.3		
Not owned	102	13.2		
No response	105	13.5		
Employment				
Working	95	12.39	169	31.24
Non-working (including others)	568	74.05	305	56.38
No response	3	0.39	15	2.77
Income Group				
< £10000	114	21.8	116	29
£10000-£19999	180	34.4	124	31
£20000-£29999	100	19.1	59	14.7
£30000-£39999	47	9	37	9.3
£40000-£49999	31	5.9	15	3.8
£50000 or more	25	4.8	23	5.7

² Please refer this report for details on recruitment strategy (overall and for the booster sample) as well as other details of methodology.

Don't know	26	5.0	26	6.5
Duration since Diagnosis of Parkinson's				
Up to 2 years	84	10.82		
2.1-5 years	155	19.97		
5.1-10 years	253	32.60		
10.1-15 years	130	16.75		
15.1-20 years	65	8.38		
More than 20 years	37	4.77		
Not recorded	52	6.70		

Note: Darkened cells mean data are not available / collected as they are not applicable to carers.

Table 1 shows that two-thirds PwP were men and one-third were women. The level of education for a majority is beyond GCSE (65.5%) and they are currently married (78.6%). Out of a total 546 carers, two-thirds of them were women, mainly spouses of the PwP. A large number of PwP (74%) were not working, of which 23.7% took early retirement due to ill health. Also, 56% of their carers had retired from the workforce, with 11.4% due to caring responsibilities or ill health.

A close to 76% PwP lived either with their spouse or with their children or in an extended family; 14% lived on their own. More than 75% PwP and a close to 75% of their carers had income less than £30000. Almost 32% persons were living with Parkinson's had been diagnosed with somewhere between 5-10 years, and a close to 20% between 2-5 years. Apart from the characteristics mentioned in Table 1, the mean age of caregivers was 62.6 years (as compared to 67.1 years for PwP).

Financial distress to PwP and their carers comprised of 10 components, out of which 6 were different types of expenses, and four of them were because of income loss. PwP and their carers had to either reduce the number of work hours or take a premature retirement because of health and caring responsibilities, respectively.

Health economics evaluation framework, described in the previous section, was used to estimate the annual costs to PwP households (PwP + their caregivers, who are largely, their spouses). The components and the cost of each component can be seen in Figure 1. Summing up, the annual costs (financial distress) to PwP households work out to £16582.

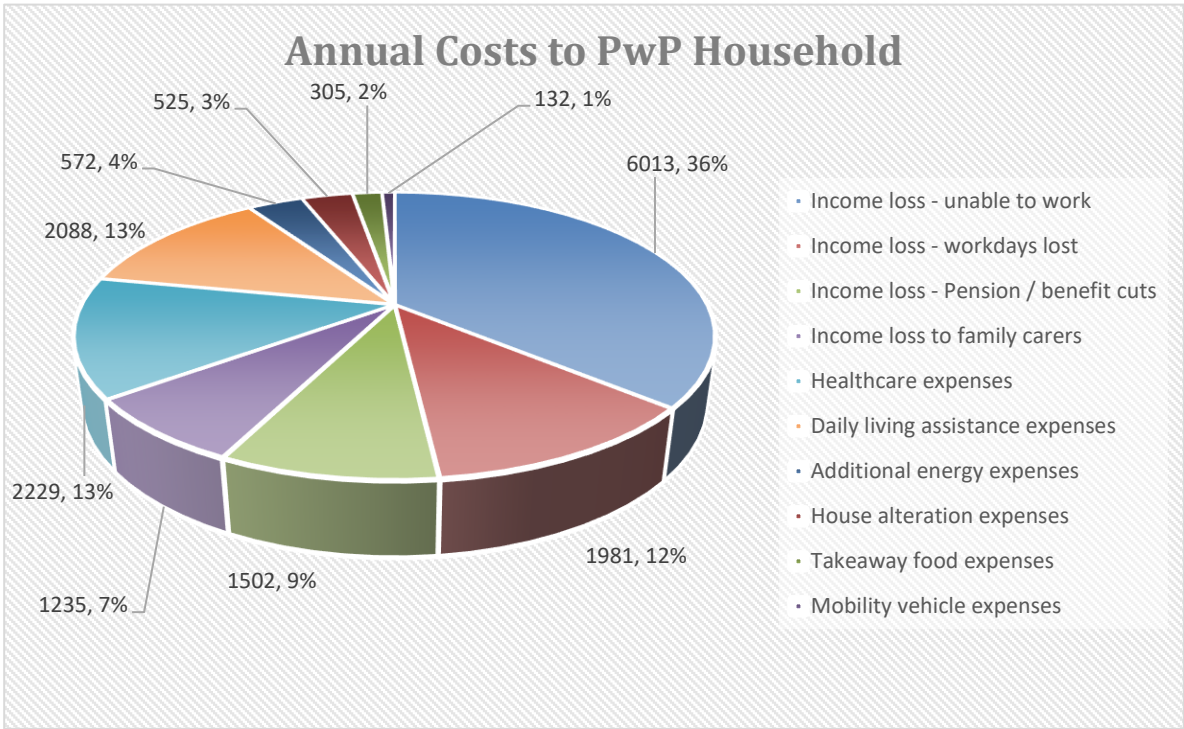


Figure 1. Financial Distress (Average Annual Costs) to a PwP Household.

Psychological wellbeing of PwP and their carers was measured on the following metrics:

- EuroQoL EQ-5D-5L,
- Visual analogue scale (VAS) for overall health state (ranging between 0 ‘worst’ and 100 ‘best’ health imaginable), and
- Wellbeing scores (life satisfaction, worthwhileness, happiness and anxiousness), all measured on 0 to 10 scale and adapted from the Office for National Statistics, (2012).

Individual-level QALY were estimated by combining the responses for the five dimensions of Euro-QoL using the weight suggested by (N. Devlin et al., 2016). The average QALY score for PwP was 0.576 and that of their carers was 0.674; QALY score of 1 means one year of life in perfect health. The mean Visual Analogue Scale (VAS) score was 57.9 for PwP and that of their carers was 70.1 (where, 0 indicates the best health state and 100, the worst health state). The QALY and VAS scores are based on the responses of 650 PwP and 301 carers.

Wellbeing scores are based on the responses of 638 PwP and 294 carers. The mean wellbeing score for PwP was 5.3 for satisfaction with life, 5.5 for how happy they felt yesterday. The corresponding wellbeing scores for their carers were 5.7 for satisfaction with life and 5.8 for happiness.

QoL and wellbeing scores for PwP and their carers are discussed in detail, in the following section.

4. Association of Financial Distress with QoL and Wellbeing of Informal Carers

Financial distress affects not only the quality of life of PwP but also their carers. As discussed, most carers are the spouses and therefore, financial distress is experienced not by the PwP as an individual, but by a household with a PwP. It is, therefore, important to know the impact of financial distress on informal carers of PwP. Figure 2 shows how out-of-pocket expenses affect the QoL of PwP and their carers, and figures 3 and 4 shows how total financial burden is associated with the QoL of PwP and their carers.

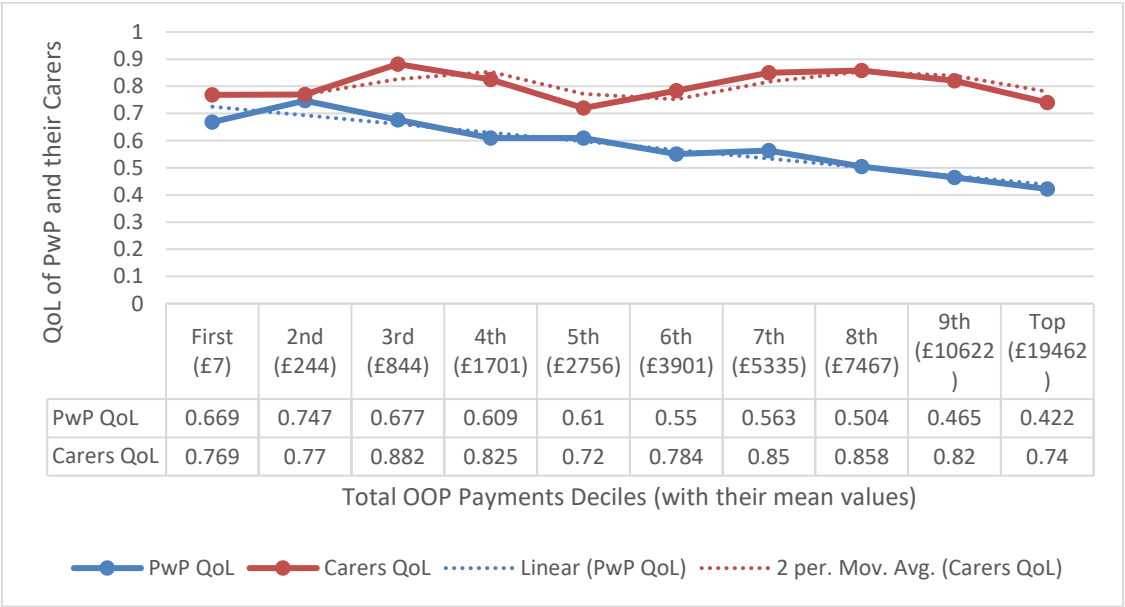


Figure 2. QoL of PwP and their Carers by their out-of-pocket (OOP) Payments Deciles.

It can be seen in Figures 2 and 3, PwP’s quality of life shows a linearly decreasing trend over the deciles of out-of-pocket payments and total financial burden deciles (blue dotted line). However, the carers’ quality of life follows a 2-point moving average trend (red dotted line).

Figure 2 shows that the QoL for PwP declines with increase in out-of-pocket expenses. This could be because the PwP would have to reduce spending on other things to make up for the increased expenditure on Parkinson’s treatment and care. However, the 2-point moving average trend for the QoL of carers show a sinusoidal trend. This implies that the quality of life of carers increase with

increase in out-of-pocket payments up to 4th decile, then decreases until the 6th, and then again increases till the 8th decile before it again starts declining. This is because in the initial phase, the expenditure on physiotherapy and other alterations in the house help the with the ADL (Trend et al., 2002), reducing their dependence on their carers, which results in an improved quality of life for carers. However, with the progression, the expenses increase and so does the dependence of PwPs on carers, which results in a decline in quality of life of carers. The second sine wave of the trend needs further exploration to explain the improved quality of life between the 6th and the 8th decile.

Figure 3 shows a downward sloping trend in quality of life of PwP with total financial burden, which implies a decline in quality of life with increase in overall financial burden. A part of the financial burden is the loss of income due to reduced working hours or early retirement, which augments the financial constraints. However, there is an overall improvement in quality of life of carers until 6th decile and then their QoL falls.

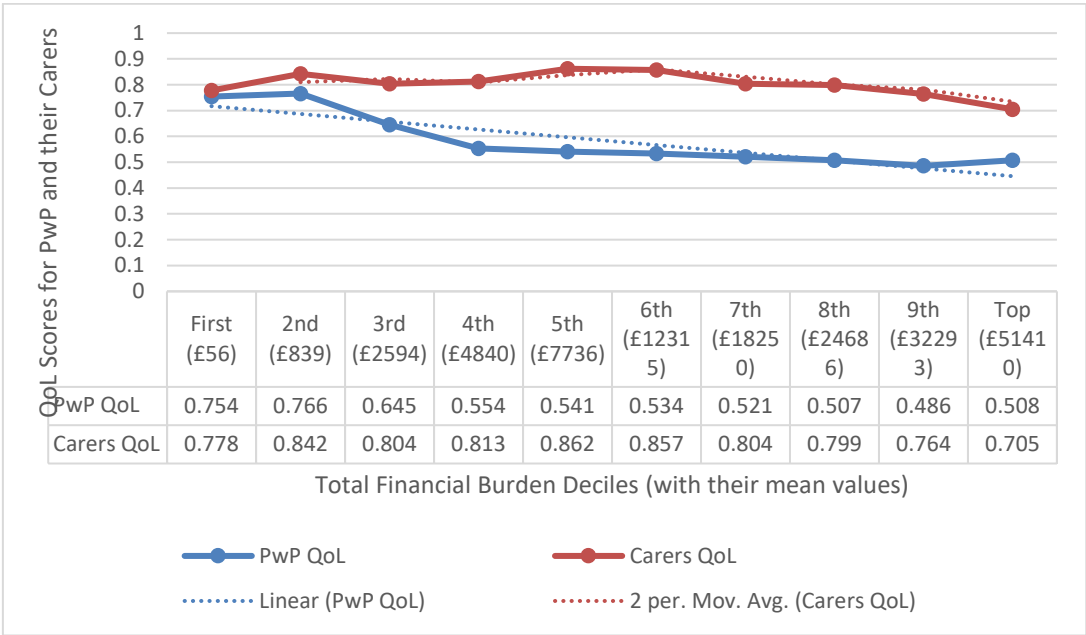


Figure 3. QoL of PwP and their Carers by Total Financial Burden Deciles.

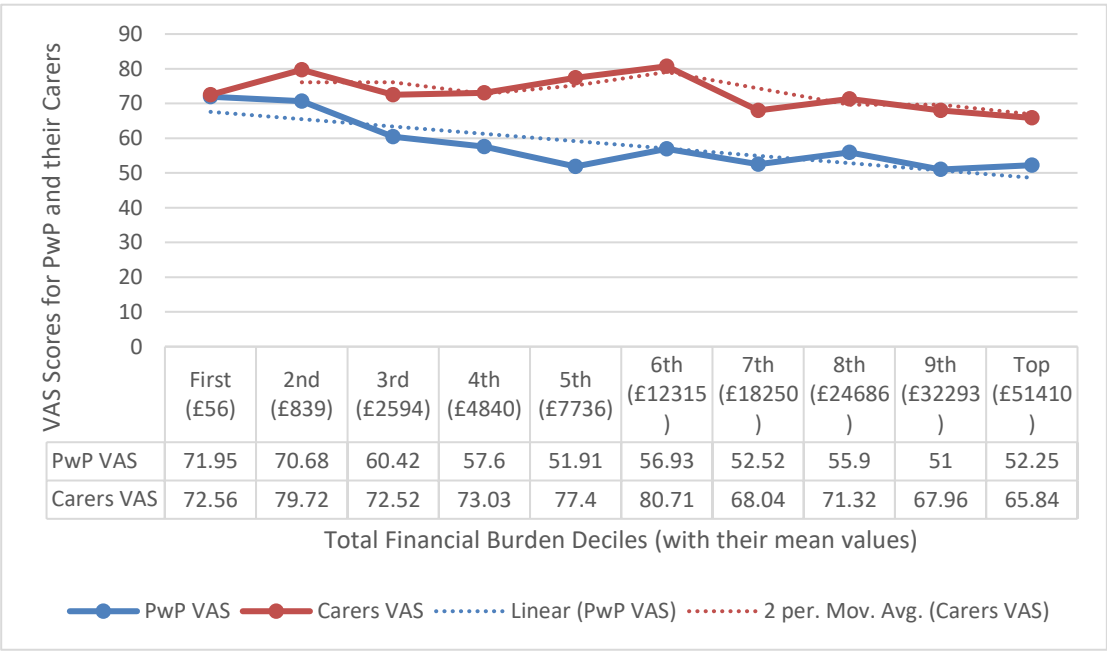


Figure 4. Overall QoL (VAS Scores) of PwP and their Carers by Total Financial Burden Deciles.

Figure 4 shows the overall QoL based on VAS scores of PwP and their carers over the total financial burden deciles. Here too, the VAS score of the PwP declines with the increase in total financial burden, and that of carers largely increase till the 6th decile and then falls.

Thus, financial burden directly impacts the QoL and VAS of PwP, whereas that of their carers improve initially before declining. Also, the QoL and VAS scores of PwP are much poorer as compared to that of their carers.

5. Impact (Determinants) of Financial Distress on QoL and Wellbeing of PwPs

The principal objective of this study is to examine how financial distress and socioeconomic background and demographic profile influences the wellbeing scores of PwPs. Therefore, four separate regressions were undertaken to examine the effect of financial distress and socioeconomic background and demographic profile on (1) QALY, (2) VAS Scores, (3) life satisfaction scores and (4) happiness (wellbeing) scores of PwP only (and not their carers).

The predictors for all the four models are the same to ensure comparability of the predictors on various psychological wellbeing indicators. The outcome variables are: (1) quality of life as estimated using the EQ-5d-5L questionnaire, (2) VAS Scores, (3) Life satisfaction score, and (4) Happiness score. Life satisfaction and happiness scores are adapted from the Office for National Statistics, (2012).

There are 12 predictors to each of these four regression equations. Out of these 12 predictors, demographic characteristics of the PwP are explained by age and gender (male and female). Social characteristics of PwP are represented by marital status, education and ethnicity (Whites or Non-Whites). Economic status of a PwP is represented by house ownership (owned or otherwise), working or not / retired) and their household income categories. Disease specific predictors are number of years since the diagnosis of Parkinson's, number of medications for Parkinson's treatment and number of visits to healthcare professionals / services. Financial distress predictors are out-of-pocket payments, and income loss. However, the actual calculations of out-of-pocket expenditures and income losses are added to arrive at the total estimate of financial distress and then categorised into deciles.

The regression equation is given in Equation (1):

$$y = \beta_0 + \beta_1 age + \beta_2 (age)^2 + \beta_3 gender_f + \beta_4 (marital status_s) + \beta_5 education_o + \beta_6 ethnicity_w + \beta_7 (house ownership_{no}) + \beta_8 working_{no} + \beta_9 (income category_{nr}) + \beta_{10} Parkinson\ duration_{<2} + \beta_{11} healthcare\ visits + \beta_{12} medications + \beta_{13} (financial\ distress\ decile_1) \quad (1)$$

where,

y = QoL scores for the first regression,
 VAS scores the second regression,
 Life satisfaction scores for the third, and
 Happiness scores for the fourth regression equations.

Predictors:

age = actual age of the PwP

$gender_f$ = dichotomous categorical: male and female (reference category: female),

$marital\ status_s$ = dichotomous categorical: married and not married / single (reference category: not married / single),

$education_o$ = dichotomous categorical: higher education or otherwise (reference category: other than higher education),

$ethnicity_w$ = dichotomous categorical: whites and BMEs (reference category: whites),

$house\ ownership_{no}$ = dichotomous categorical: owned and not owned (reference category: house not owned by the PwP),

$working_{no}$ = dichotomous categorical: working and not-working (including retired) (reference category: not working),

$income\ category_{nr}$ = six categories of income ranging from less than £10,000 to £50,000 and more, and a seventh category of PwP who did not disclose their income (reference category: income not reported),

*Parkinson duration*_{<2}= number of years since diagnosis of Parkinson's, categorised into six groups ranging from less than 2 years to more than 20 twenty years (reference category: less than 2 years),
healthcare visits= number of visits to healthcare professional / services in a year,
medications= number of medications for Parkinson's and other long-term conditions,
*financial distress decile*₁= deciles of the total of out-of-pocket expenses + income loss, with lowest mean expenses in the first decile, which increase with the deciles and the highest mean expenses in the 10th decile (reference category: 1st decile)

Beta coefficients

β_0 = intercept coefficient

β_1 to β_{13} are the slope coefficients of the respective predictors.

It may be noted that the quadratic term for age in the equation is used because it is observed that the QoL, VAS and wellbeing would decrease sharply after stage 3 (Parkinson's Community Help, 2021). Moreira et al., (2017) also observed that the quality of life and wellbeing decrease sharply after moderate Parkinson's (that is eight years or more since the diagnosis), whereas mild Parkinson's did not have any effect on the QoL or wellbeing. They further observed that the QoL was affected by stigma, decline in cognition, increased impairment in mobility and ADL. Using a quadratic term implies that we hypothesise improved quality of life in early years of diagnosis and that is because physiotherapy intervention and rehabilitation exercises in early stages would improve the mobility and delays the impairment in ADL (Cholewa et al., 2014; Tomlinson et al., 2012).

The regression results are in Table 2:

Table 2. Results of Regression Analysis: Effect of Financial Distress, Socioeconomic and Demographic profile on QoL, VAS, and wellbeing scores.

Sr. No.	Socioeconomic factors	QoL		VAS		Life Satisfaction		Happiness	
		Beta Coef.	Sig.	Beta Coef.	Sig.	Beta Coef.	Sig.	Beta Coef.	Sig.
1	Gender (Female): Male	0.011	0.77	-0.024	0.53	-0.014	0.71	0.008	0.84
2	Age	0.680	0.05	0.830	0.02	1.021	0.00	0.520	0.17
	Age Squared	-0.745	0.03	-0.955	0.01	-1.068	0.00	-0.528	0.16
3	Marital Status (Single & WDS): Married	-0.038	0.33	0.010	0.80	0.029	0.47	0.046	0.27
4	Education (Not Higher): Higher Education	0.098	0.01	0.090	0.02	0.114	0.00	0.067	0.10
5	Ethnicity (White): Non-White	-0.044	0.24	-0.084	0.03	-0.025	0.52	-0.025	0.53
6	House Ownership (Not Owned): Owned house	0.118	0.00	0.123	0.00	0.112	0.00	0.140	0.00
7	Occupation (Non-Working): Working	0.072	0.09	0.073	0.09	0.115	0.01	0.078	0.09
8	Income (Don't Know): <£10,000	-0.024	0.57	-0.001	0.99	-0.026	0.54	-0.038	0.40
	£10,000 - £19,999	0.075	0.11	0.069	0.14	0.027	0.56	0.017	0.74
	£20,000 - £29,999	0.072	0.11	0.033	0.46	0.020	0.66	-0.009	0.84
	£30,000 - £39,999	0.072	0.10	0.076	0.08	-0.021	0.64	-0.039	0.40
	£40,000 - 49,999	0.049	0.23	0.033	0.42	-0.010	0.81	0.035	0.42
	£50,000 or more	0.093	0.04	0.035	0.42	0.043	0.33	0.025	0.60
9	Parkinson's duration (Not Recorded): 2-5 years	-0.036	0.48	-0.027	0.60	0.065	0.22	0.043	0.44
	5-10 years	-0.091	0.10	-0.087	0.12	-0.020	0.72	-0.054	0.36
	10-15 years	-0.221	0.00	-0.148	0.00	-0.090	0.09	-0.117	0.04
	15-20 years	-0.163	0.00	-0.098	0.03	-0.109	0.02	-0.111	0.02
	> 20 years	-0.111	0.01	-0.130	0.00	-0.119	0.01	-0.123	0.01
10	Health services visits	-0.189	0.00	-0.203	0.00	-0.165	0.00	-0.137	0.00
11	Medication	-0.056	0.16	-0.029	0.47	-0.036	0.38	0.020	0.64
12	Financial Burden (1 st Decile): 2 nd Decile (£839)	0.004	0.93	-0.018	0.72	-0.002	0.97	-0.009	0.86

3 rd Decile (£2594)	-0.065	0.19	-0.086	0.08	-0.113	0.02	-0.103	0.05
4 th Decile (£4840)	-0.184	0.00	-0.144	0.00	-0.134	0.01	-0.150	0.00
5 th Decile (£7736)	-0.156	0.00	-0.186	0.00	-0.209	0.00	-0.227	0.00
6 th Decile (£12315)	-0.178	0.00	-0.128	0.01	-0.178	0.00	-0.154	0.00
7 th Decile (£18250)	-0.154	0.00	-0.153	0.00	-0.196	0.00	-0.198	0.00
8 th Decile (£24686)	-0.204	0.00	-0.147	0.00	-0.195	0.00	-0.155	0.00
9 th Decile (£32293)	-0.210	0.00	-0.192	0.00	-0.161	0.00	-0.172	0.00
10 th Decile (£51410)	-0.198	0.00	-0.204	0.00	-0.192	0.00	-0.181	0.00

Note: Text mentioned in parentheses, next to the socioeconomic factor, refer to the reference category.

Regression results from Table 2 show that:

Demographic indicators: None of the QoL score, VAS and wellbeing scores³ are found significant across gender or marital status. Gender is classified as male and female, and marital status as married / cohabiting and singles (including divorced, widowed, separated). QoL scores, VAS and life satisfaction increases to an extent with age, and then falls. The magnitude of life satisfaction score declines continuously with increase in the duration of Parkinson’s since the diagnosis. This means that Parkinson’s diagnosis influences life satisfaction the most as compared to other QoL and wellbeing scores.

Social indicators: QoL score, VAS and life satisfaction scores are significant for education. PwP with higher education have these scores higher by 0.1 (0.09 to 0.11) standard deviations (sd) as compared to those below this level of education. This include no formal education, attended school but did not complete, completed school with or without qualification, educational qualification beyond school and other levels of education. Only VAS scores are significant across ethnicity (whites and BME). Compared to whites, VAS score of BME is lower by 0.084 sd. BME (Black & Minority Ethnic) group include Asian / Asian British, Black/African/Caribbean/Black British, Mixed / Multiple ethnic groups and other non-whites.

Economic indicators: All four scores – QoL, VAS and wellbeing – are higher for PwP with own house by 0.11 to 0.14 sd as compared to those without, and for working PwP by 0.07 to 0.11 sd as compared to non-working. Own house includes own with and without mortgage and also shared owned. All other are housing association tenants (sheltered or non-sheltered), local authority tenants (sheltered or non-sheltered), private renters, living with parents / partners / children / friends, residing in a residential / nursing home, no fixed abode and other non-owners of the house. Working PwP include all those who are employed full-time, part-time or self-employed. Non-working PwP include all those who are unemployed, not working or retired early because of illness / incapacity or for caring responsibilities or retired because of age, pensioners, and homemakers. Only QoL and VAS scores are significant for income-group: compared to those who have not disclosed their annual income, the QoL score of those with an income of £50000 or more, is higher by 0.093 sd, and VAS score is higher by 0.076 sd for the income-group £30000 - £39000. However, wellbeing scores are not influenced by income.

Parkinson’s specific: All the four scores worsen for persons diagnosed with Parkinson’s for 10 years or more, as compared to those diagnosed for less than 2 years. There is no difference in any of the QoL, VAS or wellbeing scores for persons diagnosed with Parkinson’s for less than 10 years as compared to those diagnosed with it for less than 2 years. QoL scores worsen by 0.22 sd for people diagnosed with Parkinson’s between for 10-15 years, and then relatively improve for those diagnosed with dementia for 15 to 20 years and more. VAS and happiness scores worsen for 10-15 years, relatively improve for 15-20 years and again worsen but not as much, for people diagnosed with dementia for 20 years or more. This corroborates with our finding of inverted-parabolic relationship of QoL and wellbeing with age. However, life satisfaction worsens with every one sd for additional years of living with Parkinson’s since diagnosis. QoL, VAS and wellbeing scores worsen between 0.14 to 0.20 sd with each sd in visit to avail health services. Number of medications is a part of

³ Life satisfaction score and happiness score are jointly referred to as the psychological wellbeing scores.

healthcare expenses, and it used in the regression model in the forthcoming section. Number of medications means the number of prescribed medications; 38% PwP were on one of two medications, while 1.5% were not prescribed any medication. This is despite 6.19% PwP had other long-term conditions (LTCs) but not on medications. On the other extreme, 4% were prescribed more than 10 medications. This is because 39.56% PwP were being treated for other LTCs like diabetes, blood pressure, angina, and other cardiovascular diseases. There is no effect of number of medications on either of these scores.

Financial Distress: Average financial burden to PwP works out to £16,582 per year per household. However, for the regression purpose, we categorised the financial burden into deciles; the mid-points (mean values) of financial burden for each decile is shown against the decile in Table 1.

Beta coefficients also show that age has the strongest impact on all the QoL and wellbeing scores. QoL and VAS are also affected by more number of years elapsed since diagnosis and higher deciles of financial burden. Life satisfaction and happiness are influenced the most by the financial distress.

The effect sizes of all the four QoL as well as wellbeing indicators with financial burden is shown in Figure 5. It can be seen that except for the 2nd and 3rd deciles for both QALY and VAS, the coefficients are statistically significant for all other deciles of financial burden. It can also be observed that the QoL for PwP in 8th and 9th deciles are the worst affected in comparison with that of those in the first decile (the reference group). VAS is worst affected for 9th and 10th deciles. The coefficients for life satisfaction and happiness scores are statistically significant for all deciles of financial burden, except the 2nd. Figure 2 further shows that life satisfaction score is affected the most for PwP in 5th, 7th and 8th deciles of financial burden. Happiness scores are worst affected for the PwP in the 5th decile of financial burden. Overall, it can be seen that all the scores – QoL, VAS and wellbeing – largely worsen with increase in financial burden, 4th decile onwards.

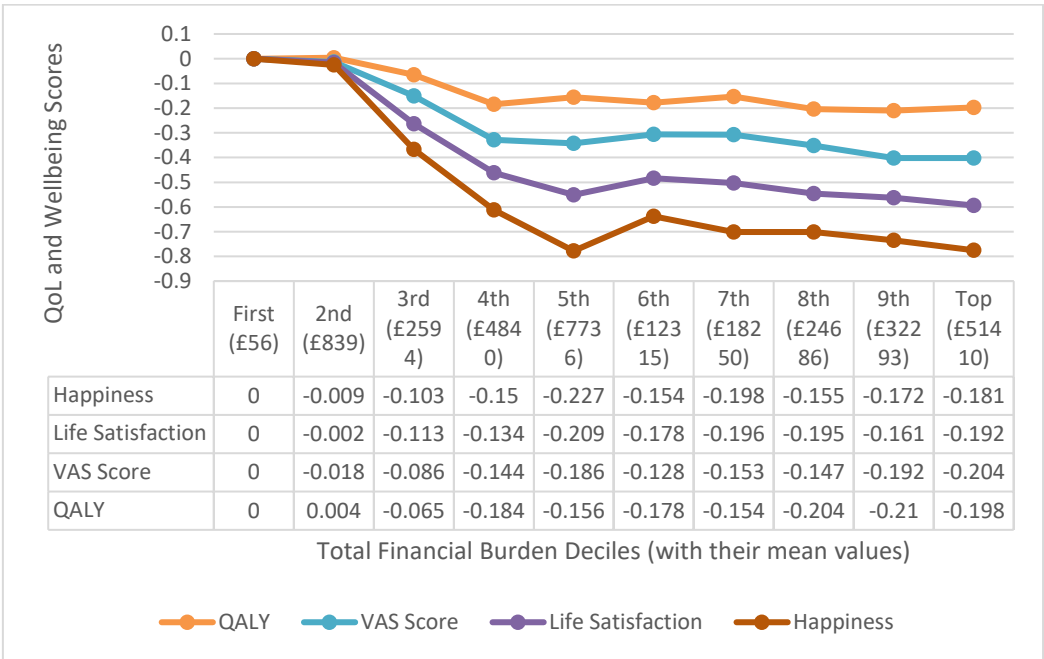


Figure 5. QoL and Wellbeing Scores by Financial Burden Deciles.

6. Limitations of the Study

This study in a community setting has adopted a household-based approach and therefore, PwP living in institutional settings such as rehabilitation centres, hospitals and care homes could not be included. These are largely PwP with severe conditions. Therefore, there is an under-representation of PwP with severe conditions in this study. As a result there is an under-estimation of such associated institutional and personal cost. The study has also not computed any cost of premature mortality due to Parkinson’s and thus the economic and financial burden is entirely based on the survived PwP and the management of their disease condition.

Carers might have an increased need for healthcare services, as compared to them not having the caring responsibility. Carers have reported increased morbidity and deterioration of QoL because of caring responsibilities. This could be because of additional work burden and emotional distress. This can have a negative multiplier effect on costs of healthcare. In this study, we have not taken into consideration, these consequential costs.

7. Key Findings and Policy Recommendations

Annual average cost of Parkinson's care is £16,582, and that is a huge strain on their limited income. Most PwP households surveyed had an annual income under £30,000. Financial distress caused by this income-expenditure gap can significantly affect the treatment and management of Parkinson's in the UK, resulting in further deterioration of quality of life and wellbeing of PwP. The findings of this study show that there is an increased impact on the QoL of PwP with the progression of Parkinson's. Also, both QoL and wellbeing scores have negative association with the financial burden, implying a deteriorated quality of life and wellbeing of PwP with increase in financial burden.

There is no known cure for Parkinson's. Therefore, addressing financial barriers is essential for ensuring equitable access to quality care for PwP. Various support programs like employment and support allowance, and carer's allowance are already available to assist PwP with financial constraints. Charities could be encouraged to offer assistance programs for PwP, by introducing government schemes to provide financial aid. Social workers and healthcare professionals could be encouraged to help PwP navigate available resources.

Financial distress can lead to anxiety, depression, and overall mental health decline in people with Parkinson's. Comprehensive care for Parkinson's should include mental health support alongside medical treatment. Pontone et al., (2019) has found that raising awareness of mental healthcare needs, and promoting integrated speciality mental healthcare would help improve the QoL of PwP. Comprehensive support programs, advocacy efforts, and mental health services are key components in addressing financial distress in Parkinson's treatment.

Advocacy efforts are crucial also for raising awareness about the financial challenges faced by PwP, and the already available benefits for PwP with financial needs. Collaboration between PwP and their carers, advocacy groups, healthcare providers, and policymakers is essential to access the demand and supply of financial support, estimate the gap and take necessary steps to reduce the gap. Policy provisions to improve access to healthcare should aim at reducing out-of-pocket medication costs and costs of healthcare visits.

Considering that Parkinson's is not curable, continued research and innovation in Parkinson's treatment may focus on more affordable, accessible and effective therapies. Dodel et al., (2021) estimated that treatment intervention for Parkinson's had a cost savings between €1800 and €7600 with 0.10 QALY for the United Kingdom. They estimated the incremental cost effectiveness ratio (ICER) of adequate treatment availability to be €15,000 to €32,600 across country⁴ settings. Target adherence to treatment would save €917,000–€2,980,000 (UK) for every 1,000 patients treated adequately. Ensuring effective and adequate treatment, augmented with lifestyle changes can help further slow the progression and eventually, reduce the financial distress. Studies (Janssen Daalen et al., 2022; Marras et al., 2019; Veronese et al., 2024) have examined and identified some lifestyle factors that helps in progression of prodromal to manifest Parkinson's, and progression of Parkinson's.

Provision for person-centered care (PCC) at home is advocated for PwP. This helps people at different stages of Parkinson's, from different ethnicities and cultures to access care suited to their needs. PCC could apparently appear to be expensive, but it could increase the quality of life and wellbeing of a PwP, resulting in slow progression of Parkinson's, which could lead to reduced costs over the lifetime of a PwP.

Bhimani, (2014) further observes the need for reposing trust in palliative care. Caring for PwP in advance stages could affect QoL and wellbeing of carers. Our study shows that a majority of the carers are spouses, who not only get physically distressed but also emotionally disturbed seeing their

⁴ Their study was undertaken for the United Kingdom and Germany.

loved ones in advanced stages of Parkinson's. Since Parkinson's progresses with age, the spouses also age alongside their PwP partners. Affordable and reliable palliative care can help improve the QoL and wellbeing of both PwP and their carers.

While this study is focused on examining the effect of financial distress on QoL and wellbeing of PwP, we also broadly looked at the QoL of carers with increased out-of-pocket payments and overall financial burden. Our study shows that QoL of carers seem to decline after an unbearable threshold level of financial burden experienced. Additionally, carers were also found to have an average wellbeing score lower as compared to that of general population because they too experience financial stress and burden (Gumber et al., 2017). We recommend providing financial resources and support also for carers of PwP. Respite care services and caregiver assistance programmes are required to be implemented, to alleviate financial strain. Giles et al., (2015) have observed that there is a huge demand-supply gap for respite care, especially for PwP in moderate and advance stages. The need for respite care would increase by the number of years and stage of Parkinson's. However, Rosqvist et al., (2022) found that there is a demand for respite care but people prefer the informal care route for respite because of their apprehensions about the quality of care in respite care facilities. Bhimani, (2014) observes that respite care is often not sought for, because of the additional costs. As discussed, PwP already bear the burden of increased expenditure because of their condition. The demand-supply gap for respite care could be bridged partially by making it affordable and accessible. Undertaking measures to reposing trust about respite care could help PwP and their carers avail these services, thereby, improving their QoL and wellbeing.

In nutshell, Parkinson's is a progressive neurodegenerative condition, which deteriorates with age, resulting in reduced quality of life and wellbeing of PwP and their caregivers. This also results in increased financial burden to PwP household. Financial distress arising out of out-of-pocket expenditure can add to deterioration of QoL and wellbeing of PwP and their carers. Appropriate policy measures like financial support programmes, respite care for carers, creating awareness about lifestyle factors for prevention and slowing down the progression of Parkinson's as well as person-centric care could be some responses to address financial distress and improve the QoL and wellbeing of PwP and their caregivers.

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Conflicts of Interest:

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