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A cohort study of the impact of COVID-19 on the quality of life of people newly diagnosed with dementia and their family carers

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Abstract

**INTRODUCTION:** COVID-19 has impacted people with dementia and their family carers, yet little is known about effects on overall quality of life.

**METHODS:** In a UK cohort study, pre- and post-pandemic data were collected from 114 carers and 93 recently diagnosed people with dementia. Latent Growth Curve modelling examined change in quality of life.

**RESULTS:** Carers reported significant decline in quality of life, although no change was demonstrated by people with dementia. In multivariable analyses, higher levels of cognitive impairment, deprivation, study site, and lower number of memory clinic contacts were associated with greater decline in carer quality of life.

**DISCUSSION:** Maintaining life quality for people with dementia during the pandemic appears to have come at the expense of their family carers. This inequity has fallen hardest on those caring for people with more severe dementia, in deprived areas, and with least support from memory services. These effects may be prevented or reversed by post-diagnostic care.
1. Introduction

Quality of Life (QoL) is a complex, multidimensional construct that is widely used to examine people’s experiences of living with dementia. The World Health Organisation defines it as the ‘evaluation by an individual of their position in life, assessed in context of one’s culture, values, goals, expectations, standards and concerns” (pg 1405). Research has highlighted a range of modifiable components that differ for people with dementia and family carers, and can enhance or hinder their QoL. These include demographic (e.g. spouse/non-spouse carer status and living situation), personal (e.g. coping strategies), social (e.g. familial networks) and contextual factors (e.g. ability to contribute to their community). Consequently, anything that may impede the access to, and quality of these resources is likely to result in a worsening of QoL for these populations.

In late 2019, a new coronavirus (COVID-19) emerged causing global disruption as countries introduced public health strategies to attempt to control the spread of infection. Since March 2020, the UK has imposed a series of measures including national and regional lockdowns, prolonged periods of physical/social distancing, reduced access to local services and community facilities, 'shielding' for vulnerable groups, and travel restrictions. The pandemic has had particular negative effects on people with dementia and their carers. People with dementia are at high risk of infection and death, accounting for 25%-31% of all UK COVID-19 related deaths. In the US people with dementia were at significantly increased risk for COVID-19 than people without the condition, and this was particularly the case for black people and those with vascular dementia. This is driven by the association of dementia with age and biological vulnerability, and the social vulnerability of the high prevalence of people with dementia living in care homes where mortality from COVID-19 has been particularly high, exacerbated in the UK by the practice of discharging infected individuals from hospitals to care homes. People with dementia also have difficulties in
remembering and understanding restrictions and precautions\textsuperscript{15,16} and may find government guidance inaccessible and difficult to follow, particularly without the support of a carer.\textsuperscript{17}

COVID-19 governmental public health strategies have restricted socialisation and reduced access to health and social care services and community activities, potentially affecting mental health and wellbeing.\textsuperscript{18,19} Researchers have described the ‘shrinking worlds’ of some people with dementia during the pandemic, finding themselves unable to engage in meaningful activities and social interaction that had previously provided a sense of purpose, identity and social connectedness.\textsuperscript{17} However, they also noted that for some the ‘lock-down bubble’ provided a break from the busyness of the outside world, where they could feel safer. Through the pandemic carers in general (i.e. not just those caring for people with dementia) have reported higher levels of depressive symptoms and anxiety than non-carers\textsuperscript{20} and increasing levels of depression.\textsuperscript{21} Researchers, examining the impact of social support service closures on the mental well-being of a combined sample of older people, people with dementia, and carers during the first three months of the pandemic, found levels of self-reported anxiety dropped through this period, but depression rose. Self-reported quality of life (QoL, measured by Warwick-Edinburgh Mental Wellbeing Scale, which is not dementia-specific) increased for people with dementia and older adults.\textsuperscript{22}

These early studies paint a mixed picture of the impact of the pandemic and subsequent restrictions on the QoL of people with dementia and their carers. To date, most studies have been qualitative or used cross-sectional surveys, relying on subjective retrospective ratings, with few studies comparing findings in the pandemic with pre-pandemic measures.\textsuperscript{23} One small Spanish study compared carer-reported neuropsychiatric symptoms of 20 people with mild Alzheimer’s Disease using data collected one month before lock-down and then re-evaluating them after five weeks of lock-down.\textsuperscript{24} The findings showed increases in apathy, agitation and aberrant motor behaviour for their participants with dementia, although no significant decreases were reported in well-being (as measured by carer proxy reports on
the EQ-5D). This study (DETERMIND-C19) aimed to address this evidence gap by examining QoL in a cohort of people newly diagnosed with dementia who were recruited into the DETERMIND programme in the months before the first COVID-19 lock-down. The cohort was comprehensively assessed prior to the pandemic, providing the opportunity to conduct an in-pandemic assessment allowing for the investigation of how participants were affected. We hypothesised that the impact of the COVID-19 pandemic would be to decrease QoL for people with dementia and their family carers.

2. Methods

2.1 Study design
DETERMIND-C19 used a mixed methods design involving quantitative interviews with people with dementia and carers recruited into the DETERMIND programme, followed by in-depth qualitative interviews with a sub-sample. This paper presents the findings of the quantitative element. DETERMIND involves recruitment of a cohort of 900 people with newly-diagnosed dementia and their carers in three areas of England (North-East, South London and Sussex). DETERMIND is inclusive of all participants with a clinical diagnosis of any dementia (rather than Mild Cognitive Impairment or subjective memory complaints) that have been diagnosed within 6 months of baseline assessment. Baseline interviews started in late July 2019 and were paused in March 2020 because of the pandemic. By then 261 people with dementia and 206 carers had been interviewed. The DETERMIND-C19 study reported here examines the impact of the COVID-19 pandemic on this cohort. Whilst all carers were eligible to participate, only those people with dementia who had the capacity to provide informed consent at DETERMIND baseline were approached because of the difficulty of assessing capacity over the telephone. Ethics approval for the DETERMIND and DETERMIND-C19 studies were obtained by the HRA Brighton and Sussex Research Ethics Committee [REC 19/LO/0528. IRAS 261263].
2.2 Procedure

Baseline DETERMIND interviews were completed face-to-face and follow-up DETERMIND-C19 interviews were conducted between July-October 2020 by telephone (because face to face visits were not permitted due to government restrictions) with people with dementia and/or their carers. Data were collected on topics including: QoL; physical and mental health; service use; social connections; and perception of the pandemic.

Prior to all telephone interviews, participants were posted prompt cards, outlining response scales for each of the questionnaires, and encouraged to use them during data collection. It was envisaged that this would ameliorate some of the challenges associated with working memory that may be encountered by people with dementia and so enhance the validity of the data obtained.

2.3 Measures

QoL - We measured self-rated and carer-rated QoL of people with dementia using the 28-item DEMQOL (range 28-112) and 31-item DEMQOL-Proxy (range 31–124) scales. These are interviewer-administered, dementia-specific questionnaires for assessing health-related QoL and include items focussed on emotions, memory, and activities of daily life in the past week. The DEMQOL-Proxy elicits carer’s perceptions of QoL for the person with dementia, with their answering from the viewpoint of the person with dementia rather than themselves. It was used to collect data from all people with dementia who had a carer participating in the study. To assess carer QoL we used C-DEMQOL (range 30-150) which has 30 items. This is an interviewer-administered, dementia-specific questionnaire that captures how people have felt in their caring role over the past four weeks. It includes items concerned with the carer’s responsibilities and needs, well-being, role, support, and feelings about the future. In all these instruments higher scores represent higher QoL.
Participant characteristics – Research has demonstrated that a range of demographic determinants may influence QoL for people with dementia and/or their carers, alongside dementia characteristics and contextual resources such as access and use of services. Consequently, these variables were collected and incorporated within our analysis. These included: study site; co-residence of the person with dementia and the carer; age at baseline; sex; ethnicity; marital status; education; occupational class based on the National Statistics Socio-economic Classification (NS-SEC); work status; home ownership; Office of National Statistics (ONS) rural-urban classification of the post code area; deciles of Index of Multiple Deprivation (IMD) based on post codes and the number of months between the baseline and C19 interview. Some information was collected only for the person with dementia including receipt of social security benefits (Pension Credit and Attendance Allowance or Disability Living Allowance). Severity of cognitive impairment was measured using the Mini-Mental State Examination. Type of dementia (Alzheimer’s disease, vascular, Lewy body, mixed and other) and the number of months since the diagnosis of dementia was given were also included. Number of hours per day the carer provided care for the person with dementia was used to measure intensity of caregiving. Respondents reported the number of memory clinic contacts in the three months before the baseline assessment (DETERMIND) and the three months before the DETERMIND-C19 interview.

2.4 Analysis
Latent Growth Curve modeling examined the level and change in QoL and its associations with characteristics of the carer and person with dementia. In a Latent Growth Curve model, random effects are used to capture individual differences and fixed effects to estimate the average growth of the entire sample. Analyses were carried out with Mplus 8. As there were only two time points, a simple linear change (slope) in QoL could be estimated with the initial level (intercept). Participant characteristics collected at baseline were included as potential predictors of the level and change in QoL. As there was very little change in the characteristics between the baseline and C19 interview, the baseline value
was used. MMSE was only available at baseline. Number of hours of care showed some individual change, but the sensitivity analysis using it time-varying suggested that the baseline value was a stronger predictor of QoL in C19 interview than the concurrent one. Therefore, we used only the baseline number of hours of care. Memory clinic contacts were used as a time-varying predictor. The sample size was too small to include all characteristics in the same model. Therefore we fitted the models first for each type of characteristic separately (bivariate models) to identify those associated with the level or change in QoL. We then included the bivariate predictors that were associated with either interact or slope of QoL in a multivariable model. All multivariable models were adjusted for age, sex, and the number of months between the baseline and C19 interview. The fit of the model was assessed by chi-square analysis (a p-value >0.05 recommended as a good fit), but as this is sensitive to sample size (chi-square p-value <0.05 when samples sizes reach 200+34), we used three other recommended fit indices:35 the Comparative Fit Index (CFI), Root Mean Square Error of approximation (RMSEA) and Standardized Root Mean Square Residual (SRMR). A value at or below 0.05 for the RMSEA and SRMR and at or above 0.95 for the CFI indicated a good fit for the model. Maximum likelihood estimation with robust standard errors (MLR) was used to take into account any sample non-normality. Missing data were handled using the full information maximum likelihood method (FIML)36 which makes it possible to include cases with missing values for any dependent variable in path models such that information on the means and variances of all data are used.

3. Results

3.1 Participants

Interviews were conducted between July-October 2020; of the 261 people with dementia and 206 carers recruited into DETERMIND, 114 carers and 93 people with dementia were interviewed. 74 (80%) of the people with dementia and 107 (94%) of the carers completed
the questionnaires over the phone with a researcher, the remainder completed a hard copy and posted it back to the research team. The characteristics of carers and people with dementia are presented in Table 1. For carers, average age at baseline was 66 years; 76 (67%) were co-resident with the person with dementia. Average age at baseline of the people with dementia was 80. About 90% of the carers and people with dementia were white British ethnicity. About three quarters of people with dementia scored >19 in MMSE, suggesting mild or minimal cognitive impairment. The diagnosis of dementia (62% Alzheimer’s disease, 11% vascular and 4% Lewy Body) was received about 3.4 months (sd=3.27) before the baseline. Carers spent on average 5.9 hours (sd=6.78) a day caring for the person with dementia at baseline. Table 2 presents distributions in whole sample of people with dementia that included 52 individuals who did not have a carer or where the carer did not participate in the study (and were therefore not included in Table 1). There were no statistically significant differences in participants’ characteristics between the subsample completing the C19 interview and the baseline sample (p>0.05 for chi square for categorical variables and t-test for means). The mean number of memory clinic contacts reported in the North-East was maintained (2.2 (sd=1.52) at baseline, 2.5 (sd=3.98) at follow up), while in South London and Sussex there was marked decline (baseline South London 1.4 (sd=1.18), follow-up 0.04 (sd=0.19); baseline Sussex 1.3 (sd=1.17), follow-up 0.14 (sd=0.54)) (Table 3).

In terms of performance, C-DEMQOL showed good internal consistency (baseline Cronbach’s alpha 0.92, also 0.92 in DETERMIND-C19) as did DEMQOL and DEMQOL-Proxy (baseline Cronbach’s alpha for DEMQOL 0.91 and 0.90 at follow-up; with the respective values for DEMQOL-Proxy 0.91 and 0.91). Table 4 shows distributions of QoL in the two interviews. Carer QoL dropped three-points from an average 100 to 97 on the C-DEMQOL. There was little overall change in the QoL of the people with dementia either self-reported (DEMQLQ) or reported by the carer (DEMQOL-Proxy). Characteristics of participants in the DETERMIND-C19 interview were similar to those assessed at
DETERMIND baseline (Tables 1 and 2). In terms of interpreting these changes in QoL scores, there is no established Minimal Important Difference (MID) for C-DEMQOL, but for the DEMQOL system more generally MID statistics ranged between 2 and 6 points.37

### 3.2 Change in QoL between baseline and DETERMIND-C19 interview

In the unadjusted model for carers, the initial level (intercept) of QoL (C-DEMQOL) was 100.4 (standard error, SE=1.3, p<0.001). The slope estimate was -2.7 (SE=1.2, p=0.027) indicating a decline of nearly three points between the baseline and DETERMIND-C19 interviews. The residual variances for intercept and slope were 283.3 (SE=26.7, p<0.001) and 141.4 (SE=25.8, p<0.001) suggesting statistically significant individual differences in the initial levels and slopes between the carers. The model for the carers showed a good fit: \( \chi^2=48.52, \text{df}=1, \text{CFI}=1.00, \text{RMSEA}<0.001, \text{SRMR}<0.001. \)

In the unadjusted model for the people with dementia, the initial level (intercept) of QoL was 88.9 (SE=1.0, p<0.001) for carer-rated (DEMQOL-Proxy) and 85.8 (SE=0.7, p<0.001) for self-rated (DEMQOL) QoL. The slope estimates in both suggested no change in carer-rated QoL between baseline and follow-up (0.7, SE=1.1, p=0.550 for DEMQOL-Proxy) and slight increase in self-rated QoL (2.2, SE=1.1, p=0.038 for DEMQOL). There were individual differences in initial levels with some increasing and some decreasing over time. Both models for people with dementia showed a good fit: \( \chi^2=12.66, \text{df}=1, \text{CFI}=1.00, \text{RMSEA}<0.001, \text{SRMR}<0.001 \) for DEMQOL, and \( \chi^2=53.99 \text{ df}=1, \text{CFI}=1.00, \text{RMSEA}<0.001, \text{SRMR}<0.001 \) for DEMQOL-Proxy.

### 3.3 Association of characteristics of the carer and person with dementia with change in QoL

There was substantial individual variance around the change in carer QoL. To investigate if participants’ characteristics could explain these changes, linear regressions using the intercept and slope parameters as outcomes were carried out as part of latent growth curve
estimation. First, bivariate models with one predictor at a time were fitted. Then those predictors that were associated with either the intercept or slope were included in a multivariable model. All models fitted the data well (CFI>0.95, RMSEA <0.05, SRMR<0.05).

Higher levels of cognitive impairment (lower MMSE scores) of the person with dementia were associated with greater decline in carer QoL (Table 5 and Figure 2). Similarly, higher area deprivation (lower IMD decile) was associated with greater decline in carer QoL (Table 5 and Figure 3). The results showed decline in QoL in carers of people with dementia living in Sussex and London between the baseline and DETERMIND-C19 interviews, while the North-East carers showed no change (Table 5 and Figure 1). After adjusting for the number of memory clinic contacts, the differences in the slopes of carer quality of life between the three study locations diminished. This suggests that each has an effect on carer QoL. Receiving Attendance Allowance, longer time since dementia diagnosis and lower MMSE score were associated with lower carer QoL. Some factors were significant only in the bivariate analysis (carers who were older, homeowners and had lower education showed higher QoL), and became non-significant in the multivariable model (Table 5).

The bivariate and multivariable analyses revealed few relationships between the person with dementia’s change in QoL and their characteristics (Supplementary Tables 2 and 3). None of the characteristics were associated with the change in DEMQOL-Proxy in the fully adjusted model (Supplementary Table 2). Longer time since dementia diagnosis, receiving Attendance Allowance and carer’s higher educational level were associated with lower DEMQOL-Proxy scores in the multivariable model (Supplementary Table 2). Longer time since dementia diagnosis was associated with increased DEMQOL-Proxy scores while carer co-residence was associated with higher and receiving Disability Allowance with lower QoL in the bivariate analysis only. Non-white ethnicity and receiving Disability Living Allowance were associated with decrease in DEMQOL in the multivariable model (Supplementary Table 3). Homeownership, and shorter time since dementia diagnosis and time between the
baseline and C19 interviews were associated with higher baseline DEMQOL scores in the multivariable model.

4. Discussion

This is the first study exploring the impact of COVID-19 and governmental restrictions on the QoL of people newly diagnosed with dementia and their carers. Our results suggest there has been a maintenance of QoL of people with dementia during the pandemic, but that this may have come at the expense of the QoL of their family carers, which declined over this time. This decrease in carer QoL was largest in those supporting people with more severe dementia, and those in more deprived areas. While causation cannot be inferred from the study design, these results suggest the possibility that these negative effects might be mitigated by continued post-diagnostic care since those carers where memory services continued to support people with dementia had better outcomes than those where such support services were largely withdrawn.

Our finding that the QoL of people with dementia did not decrease was unexpected. Given that the DEMQOL system is psychometrically robust and was developed specifically for the measurement of QoL in dementia, it suggests that, with the help of family carers, people with dementia were able to maintain their QoL despite the sudden disruption imposed by COVID-19. The resilience of people with dementia is often underestimated. However, these data were collected between July-October 2020, during the UK summer and a partial lifting of the restrictions. It is possible that participants had come to terms with the initial shock of the pandemic and had begun to develop mechanisms to counter any potentially adverse impacts. Our findings fit with reports that QoL may have been maintained or even have increased for some with dementia. It is possible that, as in other research, having been recently diagnosed with dementia, our participants benefitted from the lock-down bubble and
shrinking of their worlds as a means to come to terms with their dementia diagnosis in a safe environment. We will explore this in the qualitative component of DETERMIND-C19. Experiences are likely to have been very different for those with dementia in care homes.

Our findings for carers are of concern and in line with reports of an increase in depressive symptoms and anxiety in carers generally and in depression among carers of people with dementia specifically as a consequence of COVID-19 and the public health restrictions. During the pandemic, the QoL of the DETERMIND carers decreased. This was measured using an instrument specifically developed for use with family carers of people with dementia that quantifies how caring as a whole, with its positive and negative aspects, has affected the carer. Those carers who were supporting people with higher levels of cognitive impairment experienced a greater decline in QoL in the pandemic. Difficulties accessing formal health and social care services, together with limits on community activities and restricted informal social networks, will have placed a high level of strain on these carers. They may also have opted against (re)employing paid home care or less formal paid and unpaid arrangements for fear that introducing others into their household would increase their risk of infection; thereby resulting in a need for them to provide additional care at a cost to their own wellbeing.

Those carers living in areas of high deprivation were most affected. This demonstrates that there are social determinants of outcomes for carers in addition to the influence of characteristics of those they care for. The variation between the three recruitment sites is striking and offers the hope that these harms to carers might be prevented or reversed by appropriate service provision. Part of the variation between sites was due to differences in deprivation, but more was associated with service delivery. We found that support from memory services in the North East had continued during the pandemic while those in South London and Sussex decreased. While the design of our study does not allow for inference of causality, our modelling suggests that continued contact from memory services during the
pandemic was protective of the QoL of carers. This association seems independent of the effects of deprivation and severity of dementia. Our data provide support for the potential value of post-diagnostic care in supporting carers of people with dementia and preventing these harms. Pre-pandemic, diagnostic and post-diagnostic services were a “postcode lottery" and this is only likely to have widened in the last year. Post-diagnostic care is key to enabling people to live well with dementia and to preventing harms. The majority of post diagnostic care is provided by primary and community care teams with memory clinics and secondary care services providing expert specialist advice when needed. However, what should be provided and by whom is underspecified in commissioning terms in the UK and there is marked variability with very little provided to many. There is a need to develop better post-diagnostic care in dementia and the data presented here suggest how valuable such services may be, even in the most challenging of circumstances.

There are important limitations to this study. The first is inherent in the DETERMIND cohort that was designed to investigate care inequalities and so includes variation in characteristics that might be associated with variations in care such as ethnicity, social class, and region. It is therefore not a representative sample of the UK population of people with dementia. However, this variation in deprivation may have helped us to identify the associations found by increasing the statistical power for subgroup analyses. Second, we were unable to recruit all eligible participants into DETERMIND-C19. This non-response, including people with dementia who lacked capacity at baseline, may have caused bias in our results, but it is positive that DETERMIND-C19 sub-sample was similar to the baseline DETERMIND population in sociodemographic characteristics. It may have been more likely to include those who engaged with a telephone questionnaire and so we may have missed data from participants who found the pandemic particularly challenging. We will attempt to pick this group up in further waves of DETERMIND and re-engage with them to explore retrospectively their experiences during the pandemic. Third, we focussed on QoL which is a broad measure of overall wellbeing and did not investigate more subtle changes in
psychological function, or clinical depression and anxiety. However, in a multifaceted challenge such as the pandemic and in a population as heterogeneous as those with dementia, it is particularly important to look at overall effects. In the absence of MID statistics for C-DEMQOL, the clinical rather than the statistical significance of our findings are subject to question, but the direction of change is clear from our study. Fourth, the design of the project and the nature of the pandemic does not allow us to directly attribute the changes in carer quality of life to the pandemic rather than the normal process of adjustment to caring for someone with dementia following diagnosis. Fifth, the administration of the follow-up questionnaires coincided with a time of eased restrictions in the UK, further limiting the ability to make causal attributions of the current results to effects of the COVID-19 pandemic, we may have obtained different results had we interviewed during the height of the lockdown. Sixth, in our mixed group of diagnoses, the MMSE has limitations as measure of cognitive decline for non-AD diagnoses, as it might under-estimate cognitive function, which could affect the relationship in the data with cognition. Finally, our data do not cover care homes where people with dementia and carers have suffered greatly during the pandemic. In terms of strengths, our study is the first to examine change in QoL using empirical data collected prior to the pandemic and repeat measurement during it, as opposed to relying on cross-sectional or subjective retrospective ratings. The longitudinal design of DETERMIND also means that we can continue to monitor participant QoL over the years to provide insights into the long-term impact of the pandemic.

5. Conclusions

Our data show that the major QoL impacts of the pandemic have been on family carers of people with dementia rather than the people with dementia. The evidence of resilience of people with dementia in QoL terms, at least in the early months of the pandemic, is encouraging and it is important that future research seeks to examine positive outcomes, as well as the negative, and elicit the individual and societal facilitators that supported positive
QoL. With the limitations inherent in the design of this study, these data provide support for the positive value of post-diagnostic care, much of which has been closed by the pandemic or is only working virtually or with a much-reduced service.\textsuperscript{42} Particular attention is needed to address these inequities for those caring for people with more severe dementia and those living in more deprived areas. Our findings should encourage primary and secondary care services providing memory assessment services and post-diagnostic care to re-open them and maintain them as a priority. This demonstration of value should also be of use to service planners in developing and commissioning good quality post-diagnostic dementia care, supporting and training primary care teams to provide generalist support whilst working with specialists in more complex cases.
**Contributors**

SB was the chief investigator for the study and designed and managed the study with input from the group. SR carried out the statistical analyses. All authors had access to data and participated in data interpretation. BH, JR, AK, AR, ET, EMa, EN, AG, BS and ES collected the data and tested the data collection systems. SR, BH and BHu have verified the underlying data. BH, SR, BHu, RW and SB drafted the first and SB subsequent versions of this paper with input and revisions by all authors, who reviewed and approved the final submitted paper.

**Data sharing**

Deidentified participant data will be available with investigator support from 9 months after publication of the last DETERMIND-C19 paper via sube.banerjee@plymouth.ac.uk for researchers whose proposed use of the data has been approved by the DETERMIND Programme Management Board. The study protocol will be available as a supporting document.

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Table 1: Distribution of carer sociodemographic variables at the DETERMIND baseline and DETERMIND-C19 follow-up

|                              | Baseline          | C19 interview     |
|------------------------------|-------------------|-------------------|
|                              | n     | %/mean (sd)     | n     | %/mean (sd)     |
| Location                     | 206   | 114             |
| Sussex                       | 47    | 45              |
| North-East                   | 26    | 30              |
| London                       | 27    | 25              |
| Age at baseline, carer       | 206   | 66.5 (13.86)    | 114   | 66.1 (13.81)    |
| Female, carer                | 206   | 69              | 114   | 67              |
| White British ethnicity, carer | 206   | 91              | 114   | 90              |
| Marital status, carer        | 205   | 114             |
| Married                      | 82    | 82              |
| Widowed                      | 3     | 4               |
| Separated/divorced           | 7     | 4               |
| Single                       | 8     | 11              |
| Education, carer             | 198   | 108             |
| No qualification             | 12    | 7               |
| Lower secondary school (O-level/GCSE) | 26 | 24 |
| Upper secondary school (A/AS level)/vocational degree (NVQ 1-4 levels) | 33 | 34 |
| Higher education degree      | 29    | 34              |
| Occupational class, carer    | 183   | 101             |
| Professional                 | 43    | 44              |
| Intermediate                 | 33    | 36              |
| Routine | 25 | 21 |
|---------|----|----|
| Work status, carer | | |
| Working | 205 | 28 | 113 | 24 |
| Volunteering | 205 | 12 | 113 | 16 |
| Unemployed | 205 | 5 | 113 | 6 |
| Retired | 205 | 61 | 113 | 65 |
| Fulltime carer | 205 | 16 | 113 | 18 |
| Homemaker | 205 | 14 | 113 | 13 |
| Home owner, carer | 206 | 79 | 114 | 80 |
| Rural, carer (vs. Urban) | 203 | 15 | 112 | 13 |
| IMD, carer (higher=less deprived) | 202 | 6.6 (2.79) | 112 | 6.4 (2.85) |
| Coresident with person with dementia | 206 | 67 | 114 | 67 |
| Age at baseline, person with dementia | 204 | 80.3 (8.25) | 114 | 79.8 (8.85) |
| Female, person with dementia | 206 | 55 | 114 | 58 |
| White British ethnicity, person with dementia | 206 | 92 | 113 | 92 |
| Marital status, person with dementia | 206 | 114 |
| Married | 60 | 64 |
| Widowed | 30 | 26 |
| Separated/divorced | 8 | 8 |
| Single | 2 | 2 |
| Education, person with dementia | 186 | 104 |
| No qualification | 32 | 36 |
| Lower secondary school (O-level/GCSE) | 29 | 25 |
| Category                                                      | Lower | Upper |
|---------------------------------------------------------------|-------|-------|
| Upper secondary school (A/AS level)/ Vocational degree (NVQ 1-4 levels) | 23    | 21    |
| Higher education degree                                       | 16    | 18    |
| Occupational class, person with dementia                     | 199   | 108   |
| Professional                                                 | 37    | 35    |
| Intermediate                                                 | 27    | 27    |
| Routine                                                      | 36    | 38    |
| Social benefit, person with dementia                         |       |       |
| Pension credit                                               | 188   | 24    |
| Attendance allowance                                         | 204   | 33    |
| Disability living allowance                                  | 204   | 6     |
| Home owner, person with dementia                             | 206   | 72    |
| Rural, person with dementia (vs. Urban)                      | 206   | 11    |
| IMD, person with dementia (higher=less deprived)             | 206   | 6.3 (2.90) |
| MMSE score baseline, person with dementia                    | 204   | 22.0 (5.22) |
| MMSE score cut-offs baseline                                 | 204   |       |
| None or minimal                                              | 30    | 35    |
| Mild 20-25                                                   | 42    | 39    |
| Moderate 10-19                                                | 25    | 22    |
| Severe 0-9                                                   | 2     | 4     |
| Dementia type                                                | 202   | 110   |
| Alzheimer's disease                                          | 61    | 59    |
| Category         | N  | Mean (SD) | N  | Mean (SD) |
|------------------|----|-----------|----|-----------|
| Vascular         | 11 | 15        |
| Lewy body        | 4  | 5         |
| Mixed            | 17 | 14        |
| Other            | 6  | 6         |

N of months since the diagnosis of dementia at baseline:
- N: 176, Mean: 3.7 (2.67), N: 95, Mean: 3.6 (1.81)

N of months between baseline and C19 interview:
- N: 113, Mean: 8.2 (1.79)

N of hours/day caring for person with dementia at baseline:
- N: 189, Mean: 5.9 (6.78), N: 104, Mean: 6.4 (7.02)

IMD= Index of Multiple Deprivation, MMSE=Mini-Mental State Examination.
Table 2: Distributions of person with dementia sociodemographic variables at the DETERMIND baseline and DETERMIND-C19 follow-up

|                           | Baseline all | C19 interview |
|---------------------------|--------------|---------------|
|                           | n            | %/mean(sd)    | n            | %/mean (sd) |
| Location                  |              |               |              |             |
| Sussex                    | 47           | 261           | 46           |
| North-East                | 25           | 140           | 28           |
| London                    | 28           |               | 26           |
| Carer/person with dementia| 258          | 137           |              |
| person with dementia only | 20           |               | 17           |
| Non-coreident carer and person with dementia | 26 | 528 | 28 |
| Coresident carer and person with dementia | 53 | 137 | 55 |
| Age at baseline, person with dementia | 252 | 80.2 (8.14) | 137 | 79.5 (8.56) |
| Female, person with dementia | 254 | 56 | 137 | 58 |
| White British ethnicity, person with dementia | 255 | 90 | 137 | 91 |
| Marital status, person with dementia | 254 | 137 |     |     |
| Married                   | 52           |               | 58           |
| Widowed                   | 33           |               | 28           |
| Separated/divorced        | 9            |               | 10           |
| Single                    | 5            |               | 3            |
| Category                                                                 | Count | Mean (SD) | Count | Mean (SD) |
|-------------------------------------------------------------------------|-------|-----------|-------|-----------|
| **Education, person with dementia**                                     | 237   | 22.5 (5.09)| 126   | 22.9 (5.33) |
| No qualification                                                        | 31    | 33        |
| Lower secondary school (O-level/GCSE)                                   | 27    | 24        |
| **Upper secondary school (A/AS level)/ Vocational degree (NVQ 1-4 levels)** | 24    | 23        |
| Higher education degree                                                | 18    | 20        |
| **Occupational class, person with dementia**                            | 244   | 40        | 129   | 38        |
| Professional                                                            | 40    | 38        |
| Intermediate                                                            | 26    | 28        |
| Routine                                                                 | 34    | 34        |
| **Social benefit, person with dementia**                                | 229   | 24        | 123   | 21        |
| Pension credit                                                          | 249   | 28        | 133   | 31        |
| Attendance allowance                                                    | 249   | 6         | 133   | 7         |
| Disability living allowance                                             | 249   |           | 133   | 7         |
| **Home owner, person with dementia**                                    | 254   | 73        | 137   | 74        |
| **Rural, person with dementia (vs. Urban)**                             | 255   | 12        | 137   | 13        |
| **IMD, person with dementia**                                           | 255   | 6.3 (2.83)| 137   | 6.2 (2.88) |
| (higher=less deprived)                                                  |       |           |       |           |
| **MMSE score baseline, person with dementia**                           | 259   | 22.5 (5.09)| 139   | 22.9 (5.33) |
| **MMSE score cut-offs baseline**                                        | 259   |           | 139   |           |
| None or minimal 26-30                                                    | 34    | 40        |
| Mild 20-25                                                               | 41    | 38        |
|                          | Moderate 10-19 | Severe 0-9 |
|--------------------------|---------------|-----------|
|                          |               |           |
| Dementia type            | 247           | 133       |
|                          |               |           |
| Alzheimer’s disease      | 62            | 61        |
| Vascular                 | 11            | 15        |
| Lewy body                | 4             | 4         |
| Mixed                    | 15            | 14        |
| Other                    | 8             | 7         |
| N of months since the diagnosis of dementia at baseline | 212 | 3.8 (2.71) | 74 | 3.8 (1.79) |
| N of months between baseline and C19 interview | - | - | 140 | 8.2 (1.86) |

IMD= Index of Multiple Deprivation, MMSE=Mini-Mental State Examination
Table 3: carer-reported number of memory clinic contacts three months before the baseline (DETERMIND) and during the pandemic (DETERMIND-C19) by person with dementia study site

|                      | All participants | Those present in both waves |
|----------------------|------------------|-----------------------------|
|                      | baseline (n)     | C19 (n)                     | baseline (n) | C19 (n)  | Differencea |
|                      | mean(sd)         | mean(sd)                    | mean(sd)     | mean(sd) |
| North-East           | 51 2.2 (1.52)    | 35 2.5 (3.98)               | 33 2.1 (1.55) | 33 2.5 (4.08) | ns          |
| Sussex               | 93 1.3 (1.17)    | 49 0.1 (0.54)               | 46 1.3 (1.20) | 46 0.2 (0.56) | ***         |
| London               | 53 1.4 (1.18)    | 27 0.0 (0.19)               | 26 1.0 (1.00) | 26 0.0 (0.20) | ***         |
| Total                | 19 1.6 (1.32)    | 11 0.9 (2.50)               | 10 1.5 (1.34) | 105 0.9 (2.56) | *           |
|                      | 7 1              | 5                           |              |           |             |
| Differenceb          | ***              | ***                         | **           | ***       |

a Paired t-test for those present in both waves. b ANOVA between the study locations.

ns = non-significant, * p < 0.05, ** p < 0.01 , *** p < 0.001.

Table 4: Distribution of quality of life total scores at baseline (DETERMIND) and DETERMIND-C19 interviews

|                      | All participants | Those present in both waves |
|----------------------|------------------|-----------------------------|
|                      | baseline (n)     | C19 (n)                     | baseline (n) | C19 (n)  |
|                      | mean(sd)         | mean(sd)                    | mean(sd)     | mean(sd) |
| C-DEMQLQ             | 172 100.5        | 101 97.4                    | 85 99.5      | 85 97.1  |
| carer                | (16.75)          | (17.21)                     | (16.41)      | (16.60)  |
| DEMQOL | 245 | 85.8 | 91  | 86.7 | 88  | 84.7 | 88  | 86.8 |
|--------|-----|------|-----|------|-----|------|-----|------|
| Person with dementia | (10.01) | (10.12) | (10.54) | (10.12) |
| DEMQOL-Proxy | 205 | 88.9 | 110 | 88.4 | 106 | 86.6 | 106 | 88.0 |
| Proxy person with dementia | (14.64) | (15.47) | (14.77) | (15.43) |
Table 5: Associations of level (intercept) and change (slope) of carer quality of life (C-DEMQOL) with baseline characteristics of the carer and person with dementia (n = 206).

| Person with dementia and carer characteristics | Bivariate models for quality of life | Multivariable model for quality of life | Multivariable model for quality of life + adjusted for memory clinic contacts |
|-----------------------------------------------|------------------------------------|----------------------------------------|--------------------------------------------------------------------------|
|                                               | Intercept (SE)                      | Intercept (SE)                         | Intercept (SE)                                                          |
| Location (ref. North-East)                    | Intercept estimate (SE)             | Intercept estimate (SE)                | Intercept estimate (SE)                                                 |
| Sussex                                        | 1.8 (3.17)                         | 0.3 (3.39)                             | 0.6 (3.45)                                                             |
| London                                        | 4.4 (3.69)                         | 1.9 (3.62)                             | 2.1 (3.72)                                                             |
| Age at baseline, person with dementia         | 0.0 (0.14)                         | 0.2 (0.18)                             | 0.0 (0.18)                                                             |
| Female, person with dementia                  | -0.5 (2.51)                        | 0.7 (3.15)                             | 1.0 (3.17)                                                             |
| White British ethnicity, person with dementia | 3.9 (6.12)                         | -                                      | -                                                                     |
| Married, person with dementia                 | 0.5 (2.63)                         | -                                      | -                                                                     |
| Education, person with dementia (ref = no qualification) |  |  |  |  |  |  |
|----------------------------------------------------------|---|---|---|---|---|---|
| Lower secondary school (O-level/GCSE)                    | 3.2 (3.35) | -4.8 (3.31) | - | - | - | - |
| Upper secondary school (A/AS level)/Vocational degree (NVQ 1-4 levels) | 1.2 (3.49) | -4.6 (3.64) | - | - | - | - |
| Higher education degree                                  | -1.7 (4.19) | -0.3 (4.40) | - | - | - | - |
| Social benefit, person with dementia                     |  |  |  |  |  |  |
| Pension credit                                           | 2.0 (2.88) | -2.3 (3.42) | - | - | - | - |
| Attendance allowance                                     | -7.7 (2.63)** | -0.2 (2.73) | -6.2 (2.77)* | -3.8 (2.23) | -6.4 (2.78)* | -3.7 (2.43) |
| Disability living allowance                              | 0.5 (4.69) | -0.2 (3.52) | - | - | - | - |
| Occupational class, person with dementia (ref = routine) |  |  |  |  |  |  |
| Intermediate                                              | 2.4 (3.25) | -0.2 (3.02) | - | - | - | - |
| Professional                                              | 2.5 (2.97) | -1.5 (3.00) | - | - | - | - |
| Home owner, person with dementia                          | 3.4 (2.77) | -1.5 (2.82) | - | - | - | - |
| Age at baseline, carer                                    | 0.2 (0.09)* | -0.1 (0.10) | 0.0 (0.11) | -0.1 (0.12) | 0.0 (0.11) | -0.1 (0.14) |
| Category                                    | Coefficients |
|---------------------------------------------|--------------|
| Female, carer                               | -2.4 (2.68)  |
| White British ethnicity, carer              | 4.9 (5.55)   |
| Married, carer                              | 6.2 (3.56)   |
| Education, carer (ref = no qualification)  |              |
| Lower secondary school (O-level/GCSE)       | -5.8 (4.57)  |
| Upper secondary school (A/AS level)/        | -9.3 (4.37)* |
| Vocational degree (NVQ 1-4 levels)          |              |
| Higher education degree                     | -9.6 (4.54)* |
| Occupational class, carer (ref = routine)   |              |
| Intermediate                               | 0.5 (3.75)   |
| Professional                               | 0.4 (3.67)   |
| Home owner, carer                           | 6.6 (2.81)*  |
| Work status, carer                          |              |
| Working                                    | -0.4 (5.20)  |
| Volunteering                               | -2.4 (3.45)  |
| Unemployed                                 | -6.8 (6.04)  |
|                                      | Retired | Fulltime carer | Homemaker | Rural, carer (ref = Urban) | IMD, carer (higher=less deprived) | Carer coresident with person with dementia | MMSE score baseline, person with dementia | Dementia type (ref=Alzheimer’s disease) | N of hours/day caring for person with dementia at baseline |
|--------------------------------------|---------|----------------|-----------|----------------------------|-----------------------------------|------------------------------------------|------------------------------------------|----------------------------------------|----------------------------------------|
|                                      | 5.2 (4.83) | -3.3 (5.04) | -         | -                          | -                                 | -                                        | -                                        | -                                      | -                                      |
| Fulltime carer                       | -5.4 (3.43) | 0.1 (3.66)    | -         | -                          | -                                 | -                                        | -                                        | -                                      | -                                      |
| Homemaker                            | 0.1 (4.24)  | -1.9 (5.35)   | -         | -                          | -                                 | -                                        | -                                        | -                                      | -                                      |
| Rural, carer (ref = Urban)           | 3.0 (2.72)  | -4.0 (3.28)   | -         | -                          | -                                 | -                                        | -                                        | -                                      | -                                      |
| IMD, carer (higher=less deprived)    | -0.4 (0.49) | 0.7 (0.30)*    | -0.6 (0.46) | 1.4 (0.44)**               | -0.6 (0.47)                      | 1.5 (0.41)***                           | -                                        | -                                      | -                                      |
| Carer coresident with person with dementia | 1.1 (2.65)  | -4.9 (2.69)   | -         | -                          | -                                 | -                                        | -                                        | -                                      | -                                      |
| MMSE score baseline, person with dementia | 0.8 (0.22)*** | -0.4 (0.23) | 0.5 (0.23)* | -0.5 (0.21)*              | 0.5 (0.23)*                      | -0.5 (0.20)*                            | -                                        | -                                      | -                                      |
| Dementia type (ref=Alzheimer’s disease) | -2.9 (4.05) | -2.1 (2.81)   | -         | -                          | -                                 | -                                        | -                                        | -                                      | -                                      |
| Vascular                             | 1.6 (4.15)  | 0.5 (4.25)    | -         | -                          | -                                 | -                                        | -                                        | -                                      | -                                      |
| Lewy body                            | -1.5 (3.44) | 1.1 (3.58)    | -         | -                          | -                                 | -                                        | -                                        | -                                      | -                                      |
| Mixed                                | 1.8 (5.39)  | 3.3 (4.30)    | -         | -                          | -                                 | -                                        | -                                        | -                                      | -                                      |
| Other                                | -0.6 (0.19)** | -0.2 (0.15) | -0.4 (0.19)* | -0.2 (0.15)              | -0.4 (0.19)*                      | -0.2 (0.15)                            | -                                        | -                                      | -                                      |
| Time-varying predictors | Baseline | C19 | Baseline | C19 |
|-------------------------|----------|-----|----------|-----|
| N of months since the diagnosis of dementia at baseline | -0.8 (0.45) | 0.2 (0.94) | -0.9 (0.40)* | 1.8 (0.84)* | -0.9 (0.40)* | 1.9 (0.76)* |
| N of months between baseline and C19 interview | -0.5 (0.73) | 0.01 (0.63) | -0.7 (0.76) | -0.1 (0.63) | -0.7 (0.76) | -0.3 (0.61) |

IMD= Index of Multiple Deprivation, MMSE=Mini-Mental State Examination. * p < 0.05, ** p < 0.01, *** p < 0.001. Unstandardized estimates and standard errors (SE) shown.
Legends for figures

Figure 1: Estimated quality of life among carers (C-DEMQOL total score) by site at baseline and C19 interview (n = 206) from the latent growth curve multivariable model (Table 5). Adjusted for age and sex of the person with dementia and carer; marital status, educational level and index of multiple deprivation (carer); whether receives attendance allowance and MMSE score; number of hours caring for person with dementia/day; and number of months between baseline and C19 interview.

Figure 2: Estimated quality of life among carers (C-DEMQOL total score) at baseline and C19 interview by cognitive impairment (MMSE score, higher score=less impaired) at the baseline (n = 206). To illustrate the shape of the interaction the examples of slopes for the MMSE scores 10, 16, 22 (approximate sample mean) and 28 were calculated from the latent growth curve multivariable model (Table 5). Adjusted for location, age and sex of the person with dementia and carer; marital status, educational level and index of multiple deprivation (carer); whether receives attendance allowance (person with dementia); number of hours caring for person with dementia/day; and number of months between baseline and C19 interview.

Figure 3: Estimated quality of life among carers (C-DEMQOL total score) at baseline and C19 interview by area deprivation (IMD score, higher score=less deprived) at the baseline (n = 206). To illustrate the shape of the interaction the examples of slopes for
the IMD scores 2.5, 4.5, 6.5 (approximate sample mean) and 8.5 were calculated from the latent growth curve multivariable model (Table 5). Adjusted for location, age and sex of the person with dementia and carer; marital status and educational level (carer); whether receives attendance allowance and MMSE score; number of hours caring for person with dementia /day; and number of months between baseline and C19 interview