

A Comparison of Well-Being of Carers of People with Dementia and Their Ability to Manage Before and During the COVID-19 Pandemic: Findings from the IDEAL Study

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Abstract.

Background: Social restriction measures imposed to curb the spread of COVID-19 in the United Kingdom impacted on carers of people with dementia, limiting access to support services and increasing perceived burden of caring. Few studies have compared data collected both during and before the pandemic to examine the effect of these changes.

Objective: To explore whether the COVID-19 pandemic affected the well-being of carers of people with dementia living in the community, and their ability to cope with their caring responsibilities.

Methods: Analysis was conducted on two groups of carers who were enrolled in the IDEAL programme; the 'pre-pandemic group' ($n = 312$), assessed at two time points prior to the pandemic, and the 'pandemic group', assessed prior to and several months into the pandemic ($n = 156$). For the pre-pandemic group, carers were matched 2:1 to carers in the pandemic group on certain characteristics. Differences in change over time between the two groups on self-reported well-being, quality of life, coping, perceived competence, and role captivity, were investigated using mixed effect modelling.

Results: Compared to the pre-pandemic group, those in the pandemic group appeared to cope better and had more stable self-rated competency and role captivity. They did not differ in terms of well-being or quality of life.

Conclusion: Despite reports of negative impacts on carers early in the pandemic, the findings suggest the pandemic had little negative longer-term impact on carers of people with dementia, and in fact they appeared to have a more positive attitude towards coping several months into the pandemic.

Keywords: Alzheimer's disease, competence, coping, quality of life, role captivity, well-being

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INTRODUCTION

The COVID-19 pandemic has had a detrimental effect on people with dementia and their carers [1, 2]. From a health perspective, people with dementia have a high risk of contracting COVID-19 and, if infected, have a high risk of disease-related morbidity and mortality [3–5]. To protect those most vulnerable, governments put in place restrictions such as social distancing, isolation, and limitations on household mixing to help curb the spread of the virus [6, 7]. While important for public health, this disrupted the lives of people with dementia and their carers, with many carers reporting a faster than expected cognitive decline in the person with dementia following the prolonged social restrictions [8–10]. As a result of the pandemic and associated restrictions, carers have been faced with new and largely unfamiliar challenges including a reduction in access to health and social care, increased caring responsibility, lack of respite, changes in daily routines and activities of the person with dementia, and management of worsening behavioral and psychological symptoms of dementia [2, 11–19].

There is emerging evidence of the impact of the COVID-19 pandemic on the health and well-being of carers. At the start of the pandemic, many carers described increased uncertainty, worry about adapting to the ‘new normal’ and loss of control [11]. Others experienced difficulties coping with their daily living activities [14] and approximately 90% of carers reported at least one symptom of stress [20]. One study reported that carers described feelings of being trapped in their situation, having no choice but to take on all the responsibility for care [21]. While these studies clearly demonstrate the initial challenges some carers faced in navigating the pandemic, they do not explore the longer-term effects on health and well-being. Additionally, many studies utilized retrospective accounts, thus relying upon the accuracy of these accounts and carers’ perceptions of change since the pandemic [11, 14, 20, 22, 23]. This recall may be influenced by current events, prior psychological condition, and the timing of data collection in relation to the pandemic. Consequently, there are contradictory findings with some carers reporting no significant changes in their anxiety and well-being throughout the pandemic [22] and others reporting an increase in carers’ stress, burden, anxiety, sadness, and fear [16, 23, 24].

Few studies have investigated the impact on carers both before and during the pandemic, and those

that have done so report equivocal findings. A study that collected data before and during lockdown found that carers reported an increase in carer burden and a decline in their well-being [25]. Meanwhile, other studies found no significant changes in carers’ anxiety, well-being, quality of life, or distress [26, 27]. Such discrepancies may be explained in part by the timing of these studies and carers learning to adapt and becoming better-positioned or more equipped to cope with the unfolding situation. For example, although carers in one study reported an increase in stress, anxiety, and depression following a lockdown period, this had started to decrease after two months [28]. While these studies begin to explore the longer-term effects of the pandemic on carers, they still only cover the first few months of the pandemic and do not control for the normal progression of dementia as a cause of increasing burden rather than the pandemic. An initial study of ours found that, when compared to a single time-point pre-pandemic, several months into the pandemic carers were more lonely, experienced less life satisfaction and felt more trapped in their roles, but were in fact more optimistic and most reported that they were coping well [29]. They reported little change in wellbeing. However, it was not possible to determine if these findings were a result of the pandemic or due to the increasing burden of caring for someone with dementia over time.

The present study therefore aims to further understand the longer-term impact of COVID-19 on carers of people with dementia by comparing changes in well-being and coping across two groups: the ‘pre-pandemic group’—a matched comparison group assessed at two time points prior to the pandemic, and the ‘pandemic group’—assessed prior to and during the pandemic. We hypothesized that carers in the pandemic group were more likely to report a reduction in their well-being, quality of life, and ability to cope compared to their matched counterparts pre-pandemic. We expected that carers would feel more trapped in their caregiving role and feel less competent following the COVID-19 restrictions, than carers whose data were collected during two time-points prior to the pandemic.

MATERIALS AND METHODS

IDEAL

The Improving the experience of Dementia and Enhancing Active Life (IDEAL) programme is a longitudinal cohort study comprising of, at baseline,

1,537 community-dwelling people with a clinical diagnosis of mild-to-moderate dementia (Mini-Mental State Examination (MMSE) score ≥ 15), and 1,277 carers (family members or friends) recruited through memory services and specialist clinics in 29 National Health Service sites throughout England, Scotland, and Wales and via the Join Dementia Research portal (<http://www.joindementiaresearch.nihr.ac.uk/>). Exclusion criteria included if the person with dementia had any other terminal illness or were a potential risk to the interviewer during home visits. Those who expressed interest in the study were sent further information and informed consent was obtained. Where possible, a family member or other informal carer was recruited alongside the person with dementia, both to act as informant and to provide information about experiences of caregiving; however, participation of a carer was not mandatory. People with dementia, and carers where available, were assessed by trained researchers during three home visits at baseline (Time 1; T1; 2014 - 2016). Participants were assessed again 12 (Time 2; T2; 2015–2017) and 24 (Time 3; T3; 2016–2018) months after their initial assessments; two home visits were conducted on each occasion. For T2, 1,183 people with dementia and 988 carers returned and for T3, 851 people with dementia, and 759 carers returned. Full details are in the published protocol [30]. A further study, IDEAL-2, began in 2018 two years after T3 and was designed to follow-up the IDEAL cohort for three more time-points (Time 4 – Time 6; T4-T6) at yearly intervals [31]. However, the COVID-19 pandemic interrupted IDEAL-2 data collection as restrictions meant that assessments could no longer be conducted in participants' homes. For T4 (2018–2020), 253 people with dementia and 242 carers took part, and for T5, scheduled for 2019–2021, 31 people with dementia and 35 carers were assessed before data collection ceased.

INCLUDE

INCLUDE (Identifying and mitigating the individual and dyadic impact of COVID-19 and life under physical distancing on people with dementia and carers) was funded to look at the impact of COVID-19 on IDEAL participants, and focused on the experiences of people with dementia and their carers from the IDEAL cohort during the pandemic [32, 33]. People with dementia and their carers who had previously participated in the IDEAL cohort were eligible for INCLUDE if they had either participated at the most

recent assessment point or indicated willingness to be contacted again at the next assessment point. As there were very few participants from Scotland remaining in the cohort at this stage, and separate ethical and governance procedures applied in Scotland, the decision was taken to focus on participants in England and Wales for this rapid-response study. Potential participants for INCLUDE were contacted by letter, telephone, or email, and trained researchers provided information about the study and answered any questions. A follow up call was arranged to take informed consent. The carers' questionnaire could be administered by a researcher either over the telephone or by videoconference, or could be self-completed online. Self-completed questionnaires were checked by the researchers who would contact the carer if there were any missing or unclear answers to ensure a fully completed questionnaire. The study yielded both quantitative data and responses to open-ended questions. INCLUDE data were collected in the eight months between September 21, 2020 and April 30, 2021.

IDEAL was approved by Wales Research Ethics Committee 5 (reference 13/WA/0405) and IDEAL-2 by Wales Research Ethics Committee 5 (reference 18/WS/0111) and Scotland A Research Ethics Committee (reference 18/SS/0037). INCLUDE was approved as an amendment to IDEAL-2 for England and Wales (18/WS/0111 AM12). IDEAL and IDEAL-2 are registered with the UK Clinical Research Network (UKCRN), numbers 16593 and 37955. Version 5 of the IDEAL datasets was used for analysis.

Participants and design

This study uses quantitative data from IDEAL and INCLUDE and focusses on carers of people with dementia that live in the community. To compare the impact of COVID-19 restrictions on different well-being and coping-related outcomes of carers of people with dementia, two subsamples were identified in the IDEAL and INCLUDE datasets for case-control matching: the pre-pandemic group ($n = 312$) comprising carers who took part in IDEAL T1 and IDEAL T3, but not INCLUDE, and the pandemic group ($n = 156$) comprising carers who were assessed for both IDEAL T3 and INCLUDE. Both the pre-pandemic and pandemic groups incorporated two waves of data; for the pre-pandemic group, wave 1 (w1) refers to IDEAL T1 and wave 2 (w2) refers to IDEAL T3, and for the pandemic group, w1 refers to IDEAL T3 and w2 refers to INCLUDE (see Fig. 1A).

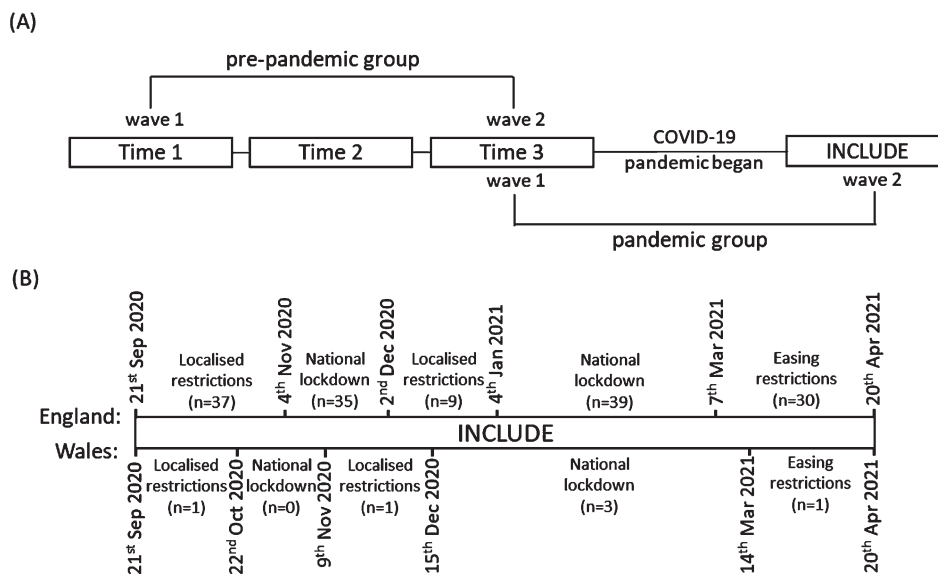


Fig. 1. A) The timeline within the IDEAL programme defining the pre-pandemic and pandemic groups. B) The varying levels of restrictions that took place during the assessment period between the 21 September 2020 and 20 April 2021 in England and Wales, and the number of people participating at each level.

To obtain the pre-pandemic group, carers were matched 2:1 to carers in the pandemic group based on carer status, carer age group, carer sex, and the diagnosis type and time elapsed since diagnosis of the person with dementia (see Supplementary Figure 1 for an overview of the matching procedure). For the pre-pandemic group, those who took part in both T1 and T3 of IDEAL, had data on the outcome measures, and were not a part of the pandemic group, were available for matching ($n=481$). For the pre-pandemic group, average time between waves was 24.5 months (range 21–34 months) and for the pandemic group, average time between waves was 39.7 months (range 27–55 months).

For the 156 people in the pandemic group, assessments took place during varying national lockdowns, local restrictions, and easing restrictions; 77 assessments took place during a period of national lockdown, 48 took place during periods of localized restrictions, and 31 took place during the gradual easing of restrictions. A timeline of the varying levels of restrictions and the numbers interviewed at each time point is shown in Fig. 1B.

Measures

Single items from standardized measures were used for well-being and quality of life. For well-being, the question was ‘in the last two weeks, how

much of the time have you felt cheerful or in good spirits?’ [34], and for quality of life the question was ‘how would you rate your quality of life?’ [35]. A single question for coping was used: ‘do you feel you cope well as a carer?’ [36]. A composite score was used for carer competence and role captivity; in each case, three questions generated a total score out of 12 [37, 38]. The details of these questions and response options are found in Supplementary Table 1. Matching variables included carer status (spouse/partner or family/friend), age group (<65, 65–69, 70–74, 75–79, 80+ years), sex, diagnosis type (Alzheimer’s disease, vascular dementia, mixed Alzheimer’s or vascular dementia, frontotemporal dementia, Parkinson’s disease dementia, dementia with Lewy bodies, or other/unspecified), and length of diagnosis (<1 year, 1–2 years, 3–5 years, 6+ years). The number of hours spent providing care (<1 hour per day, 1–10 hours per day, or 10+ hours per day) was included as a covariate.

Statistical methods

To test whether there were differences in the pre-pandemic and pandemic groups, mixed effect modelling was used, grouping waves within participants, and an interaction between group and wave was modelled. Mixed effect multinomial logistic regression models were used for categorical vari-

Table 1
 Characteristics of the pre-pandemic and pandemic groups on matching variables

		Pre-pandemic		Pandemic	
		N	%	N	%
Carer status	Spouse/partner	280	89.7	140	89.7
	Family/friend	32	10.3	16	10.3
Age group	<65	73	23.4	44	28.2
	65–69	78	25.0	39	25.0
	70–74	73	23.4	28	18.0
	75–79	42	13.5	20	12.8
	≥80	46	14.7	25	16.0
Sex	Men	115	36.9	58	37.2
	Women	197	63.1	98	62.8
Dementia subtype	Alzheimer's disease	178	57.1	89	57.1
	Vascular dementia	25	8.0	19	12.2
	Mixed (Alzheimer's and vascular)	67	21.5	24	15.4
	Frontotemporal dementia	13	4.2	11	7.1
	Parkinson's disease dementia	11	3.5	4	2.6
	Dementia with Lewy bodies	10	3.2	1	0.6
Length of time since diagnosis	Unspecified/Other	8	2.6	8	5.1
	<1 y	129	43.4	0	0.0
	1–2 y	127	42.8	74	49.3
	3–5 y	37	12.5	61	40.7
	≥6 y	4	1.3	15	10.0
	Missing	16		6	

ables (quality of life, well-being, and coping) with the random effects for grouping constrained to be equal for all outcome categories. For continuous outcomes (competence and role captivity) a mixed effect linear regression model was used. Matching variables and carer hours were included in the models as covariates. Including months between waves as either a covariate or an interaction was explored. Analyses were conducted in Stata 16. The *grid* package available in R statistical software was used to generate transition plots for categorical outcomes.

RESULTS

In both the pre-pandemic (N = 312) and pandemic (N = 156) groups, 90% of carers were spouses or partners, almost two-thirds were women, and the average age was approximately 69 years at w1 (Table 1). Over half of carers were caring for someone with Alzheimer's disease. Over 90% of the carers lived with the care recipient in both groups (pre-pandemic group, 92.6%; pandemic group, 90.7%). During the pandemic the number of carers spending over 10 hours per day caring increased compared to the pre-pandemic group (pre-pandemic group w1:34.3%, w2:53.2%; pandemic group w1:37.0%, w2:66.2%). All models were adjusted for number of hours spent providing care. Time between waves was not

important when included as either a covariate or interaction in our models, so was excluded from further analyses.

Quality of life

In the pre-pandemic group, 73% reported high (*good* or *very good*) quality of life at w1 compared to 59% at w2, while 75% of carers in the pandemic group reported high quality of life at w1 compared to 62% by w2 (Table 2). The proportions in each class, and how they change between w1 and w2, are shown in transition plots in Fig. 2A. For both pre-pandemic and pandemic groups, fewer carers report quality of life being *good* or *very good* at w2, with a large number of carers transitioning from *good* to *neither poor nor good*. In the pandemic group, there were more carers moving from *good* to *very good* than in the pre-pandemic group. As shown in Table 3, there is little evidence of a difference between pre-pandemic and pandemic groups at w1. There was evidence that, in the pre-pandemic group, more carers endorsed *neither poor nor good* or *very poor/poor* at w2 than at w1. However, there was no evidence of an interaction between group and wave, suggesting that there is no difference in how carers of people with dementia reported their quality of life between the pre-pandemic and pandemic groups.

Table 2
Outcomes at wave 1 and wave 2 in the pre-pandemic and pandemic groups

	Pre-pandemic				Pandemic			
	Wave 1		Wave 2		Wave 1		Wave 2	
	N	%	N	%	N	%	N	%
Quality of life								
Very poor	3	1.0	4	1.3	0	0.0	3	2.0
Poor	8	2.6	23	7.4	7	4.6	14	9.1
Neither poor nor good	74	23.7	101	32.4	31	20.4	41	26.6
Good	173	55.5	146	46.8	82	54.0	68	44.2
Very good	54	17.3	38	12.2	32	21.1	28	18.2
Missing	–		–		4		2	
Well-being								
At no time	3	1.0	1	0.3	1	0.7	3	2.0
Some of the time	30	9.6	42	13.5	7	4.6	20	13.0
Less than half the time	41	13.1	56	18.0	24	15.9	20	13.0
More than half the time	93	29.8	90	28.9	43	28.5	39	25.3
Most of the time	132	42.3	109	34.9	68	45.0	66	42.9
All the time	13	4.2	14	4.5	8	5.3	6	3.9
Missing	–		–		5		2	
Coping								
Never	2	0.6	4	1.3	1	0.7	0	0.0
Sometimes	66	21.2	103	33.0	44	29.1	36	23.5
Often	156	50.0	155	49.7	79	52.3	77	50.3
Always	88	28.2	50	16.0	27	17.9	40	26.1
Missing	–		–		5		3	
Carer hours								
<1 h	75	24.3	30	9.7	30	19.2	15	9.7
1–10 h	128	34.4	115	37.1	67	43.5	37	24.0
10 + h	106	34.3	165	53.2	57	37.0	102	66.2
Missing	3		2		2		2	
	Mean (sd)	N	Mean (sd)	N	Mean (sd)	N	Mean (sd)	N
Competence	9.3 (1.6)	312	8.9 (1.6)	312	9.1 (1.5)	152	9.3 (1.4)	153
Role captivity	5.3 (2.0)	312	6.3 (2.6)	312	5.6 (2.5)	151	6.1 (2.7)	154

sd, standard deviation.

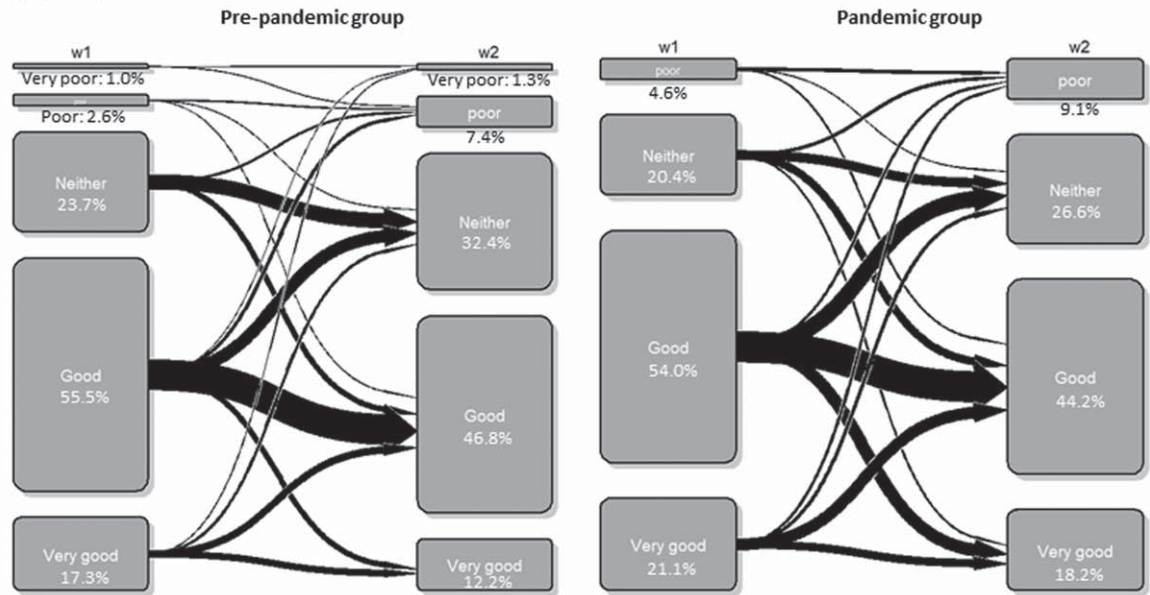
Well-being

At w1, 47% in the pre-pandemic group endorsed high (*most of the time* or *all of the time*) well-being, and this decreased to 39% at w2. In the pandemic group, 50% at w1 and 47% at w2 endorsed high well-being (Table 2). As shown in the transition plots (Fig. 2B), fewer carers at w2 endorsed high well-being for both pre-pandemic and pandemic groups, and there were no obvious differences in transitions between categories for the two groups. This was supported in the regression model (Table 3). There was little evidence of a difference between the two groups at w1, and there was limited evidence of an increase in the number of carers endorsing lower (*at no time/some of the time*) well-being compared to high (*most of the time/all of the time*) well-being for the pre-pandemic group between w1 and w2. There was no evidence of an interaction between group and wave, suggesting no difference in change in perceived well-being in the pre-pandemic compared to the pandemic group.

Coping

Half of the carers stated they *often* coped well in both pre-pandemic and pandemic groups, and this remained stable between w1 and w2. However, as shown in the transition plot in Fig. 2C, more carers in the pandemic group transitioned from *often* to *always*, and from *sometimes* to *often*, compared to the pre-pandemic group. This indicates that carers in the pandemic group may have felt they were coping better compared to the pre-pandemic group. As shown in the regression analysis in Table 3, there was little evidence of a difference at w1 between pre-pandemic and pandemic groups. There was evidence that for the pre-pandemic group, more carers endorsed *never/sometimes* and fewer carers endorsed *always* at w2 compared to w1. There was a strong interaction between group and wave suggesting that more carers in the pandemic group endorsed *always* relative to *often* compared to those in the pre-pandemic group, and there was some evidence that they were less likely to endorse *never/sometimes* in

A) Quality of life



B) Well-being

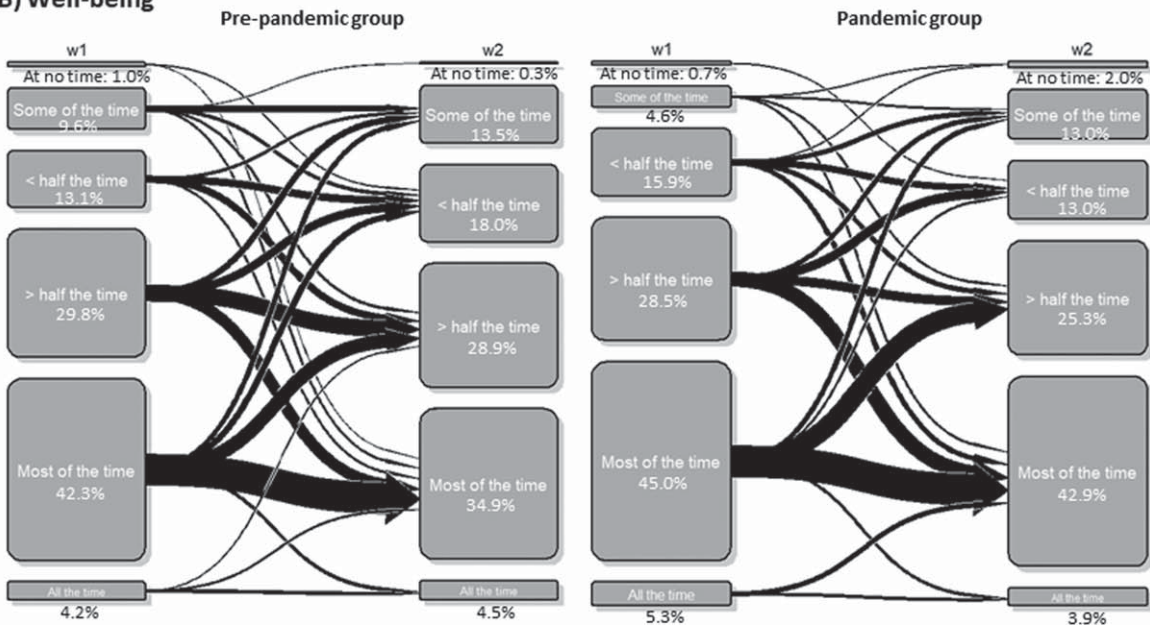


Fig. 2. (Continued).

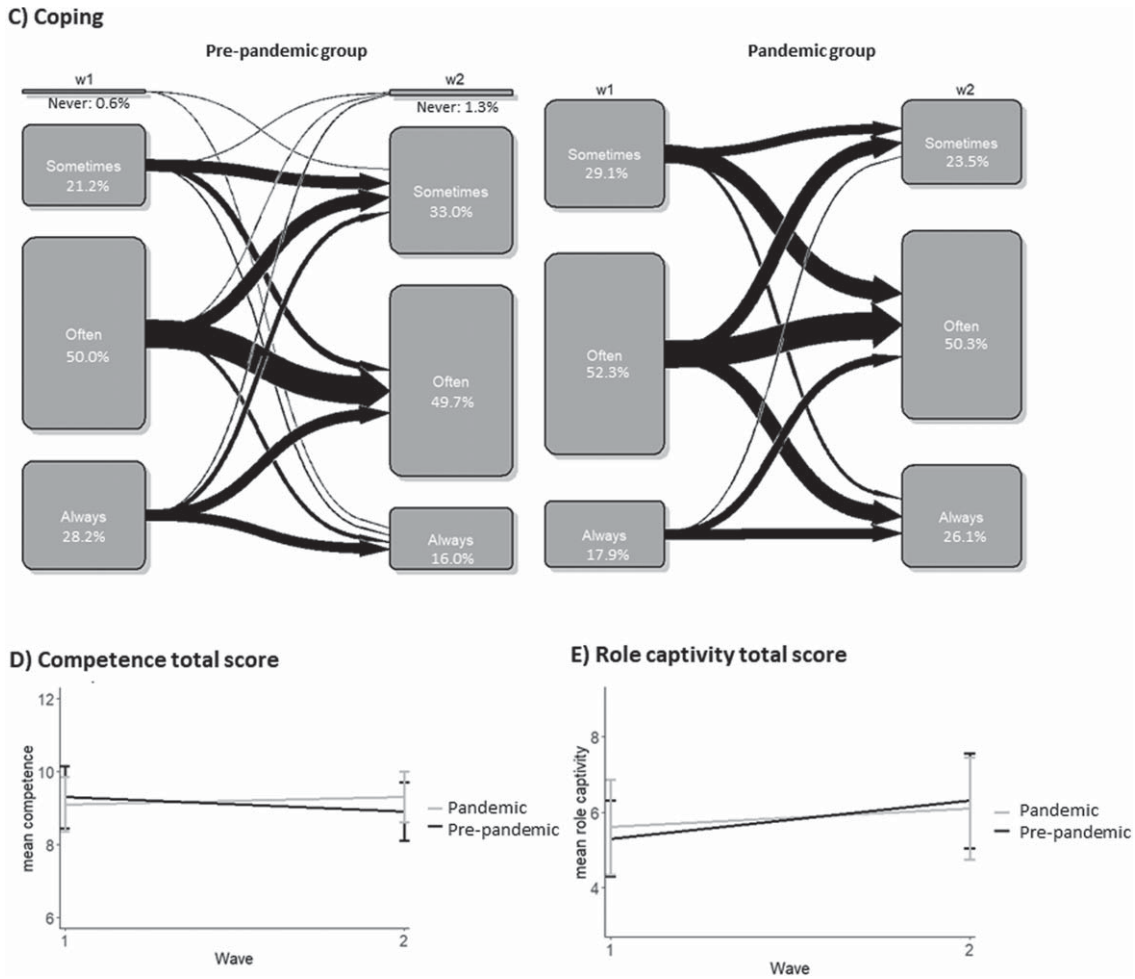


Fig. 2. A-C) Transition plots of quality of life, well-being, and coping. D, E) Plots for competence and role captivity total scores.

the pandemic group compared to the pre-pandemic group. These findings suggest that carers felt they coped better during the pandemic.

Competence

As shown in Table 2 and Fig. 2D, the mean competence score was similar across both waves for pre-pandemic and pandemic groups and so any differences found were small. The scores for pre-pandemic and pandemic groups were similar at baseline with a slight decrease in competence between w1 and w2 in the pre-pandemic group (Table 3). There was an interaction effect indicating less of a decrease in competence in the pandemic group, meaning the competence score in the pandemic group is more

stable across the two waves compared to the pre-pandemic group.

Role captivity

As shown in Table 2 and Fig. 2E, the mean role captivity score increased slightly between w1 and w2 for pre-pandemic and pandemic groups but again differences were relatively small. As shown in Table 3, there was no difference between pandemic and pre-pandemic groups at w1 with an increase in role captivity score between w1 and w2 for the pre-pandemic group. There was an interaction effect, suggesting that the role captivity score, whilst still increasing, increased to a lesser extent for the pandemic group when compared to the pre-pandemic

Table 3
Multilevel modelling to investigate differences in pre-pandemic and pandemic groups

Measures		Pandemic versus pre-pandemic at Wave 1		Wave 2 versus Wave 1 for pre-pandemic group		Interaction between pandemic group and Wave	
		OR	95% CI	OR	95% CI	OR	95% CI
Quality of life	Very poor/poor	1.38	0.37 – 5.22	3.45	1.43 – 8.31	0.90	0.22 – 3.72
	Neither poor nor good	0.60	0.30 – 1.19	1.54	0.99 – 2.40	0.98	0.45 – 2.15
	Good	Ref		Ref		Ref	
Well-being	Very good	1.15	0.56 – 2.36	0.97	0.56 – 1.67	1.32	0.55 – 3.15
	At no time/some of the time	0.49	0.16 – 1.47	1.43	0.77 – 2.66	1.83	0.57 – 5.89
	Less than half the time/more than half the time	0.81	0.39 – 1.68	1.30	0.83 – 2.05	0.67	0.32 – 1.44
	Most of the time/all of the time	Ref		Ref		Ref	
Coping	Never/sometimes	1.20	0.66 – 2.15	1.74	1.14 – 2.65	0.56	0.27 – 1.15
	Often	Ref		Ref		Ref	
	Always	0.71	0.38 – 1.33	0.61	0.38 – 0.97	3.10	1.43 – 6.74
Competence		Est	95% CI	Est	95% CI	Est	95% CI
Role captivity		0.00	–0.34 – 0.34	–0.30	–0.48 – –0.13	0.44	0.14 – 0.74
		0.06	–0.46 – 0.57	0.73	0.51 – 0.95	–0.54	–0.92 – –0.17

OR, odds ratio; CI, confidence intervals; Est, estimate.

group. This suggests that carers felt less trapped in their role as a carer during the pandemic than they did pre-pandemic.

DISCUSSION

To help counter the spread of COVID-19, since March 2020 there have been three national lockdowns in England and Wales, as well as local lockdowns with varying restrictions on social activities in different areas. Here, we followed up on carers from the IDEAL cohort several months into the pandemic, and investigated whether their experience of living through the pandemic and the associated social restrictions affected their well-being, as well as their beliefs about how they managed their caring responsibilities. We used longitudinal data to determine whether the changes in response over time were different during the pandemic compared to before the pandemic, allowing us to distinguish whether any change was due to the COVID-19 pandemic itself or attributable to other possible causes, such as higher levels of carer burden as dementia progressed. Contrary to our hypotheses, we found no effect of the COVID-19 pandemic on either quality of life or well-being of the carers and, interestingly, we found that the carers responded more positively to questions asking about their abilities to cope, their competence, and their feelings of being trapped in their roles compared to how they responded before the pandemic. This sug-

gests that several months after the initial lockdown, the impact of the pandemic did not negatively affect the carers of people with dementia.

Most studies have considered the early stage of the pandemic when there were the greatest restrictions and older people or those with medical conditions were advised to stay at home and shield as much as possible. A number of studies have found that the restrictions and social isolation measures associated with the COVID-19 pandemic negatively impacted on perceived cognition, increased the severity of neuropsychiatric symptoms, and led to perceived functional decline and decreased independence of people with dementia [9, 25, 39]. Several studies found that carers experienced a higher level of stress or worsening well-being, and increased burden and exhaustion as a result of reduced access to formal care services and respite care, or due to anxiety about the person with dementia catching COVID-19 and refusing support to reduce the risk [9, 11, 12, 17, 22, 25, 40, 41]. The primary carer, often a spouse, spent more hours caring and had additional responsibilities, particularly when caring for someone with severe dementia [22]. Alzheimer's Society reported a strong negative emotional impact of the pandemic on the mental health outcomes of carers and strained relationships between the carer and person with dementia [41]. Other studies reported greater anxiety, depression, irritability, and helplessness of the carer [17] and high stress and burden, and low quality of life

[42] but had no pre-pandemic controls. Another study reported that even before the pandemic, spousal carers found responsibilities overwhelming, and as a result of declining help to reduce risk of contracting COVID-19, felt more trapped and had more conflicts with their partner [21].

However, other studies found less of a negative impact of the pandemic. A qualitative study reported some negative consequences of lockdown on carer well-being, but some carers reported no impact because they continued to engage in meaningful or valued activities, or because they already led a restricted lifestyle before the pandemic [16]. Qualitative interviews of 11 carers from the IDEAL cohort during the initial lockdown, with follow-up in July 2020 when restrictions were easing, found that some carers were coping well and this corresponded with good health, strong relationships and community support [43]. Several studies found that carers reported a strengthened relationship and deeper connections with the care recipient due to spending more time together [1, 16, 44]. In a qualitative study focusing on positive aspects of caring during COVID-19, carers also reported finding a greater connection with their values and what was important in their lives, such as the act of caregiving and the way they viewed themselves [44]. Several carers reported gratitude for their experience, which enabled them to develop close relationships with not only the care recipient, but other family members and paid carers, which they wanted to continue. They reported a strengthening of commitment to the person with dementia and recognized the importance of asking for and accepting help with looking after themselves in an ongoing way and of seeking opportunities to connect with others in the same situation [44]. Some studies found no changes in carer-reported neuropsychiatric symptoms of the person with dementia and no difference in carer stress in the first couple of months of lockdown [27], whereas another study reported worsening neuropsychiatric symptoms but no change in carer well-being [26]. Therefore, the findings from the evidence-base present a mixed picture on the impact of the pandemic on carers.

In our study, more carers reported poorer quality of life at wave 2 compared to wave 1 in both pre-pandemic and pandemic groups, suggesting a decline in quality of life over time. However, the decline was similar for those in the pandemic group compared to those in the pre-pandemic group. In addition, well-being was stable across time for both groups. Our findings therefore suggest the COVID-19 pandemic

had little impact on the well-being of the carer, consistent with our previous evidence [29]. Almost all of the studies to date were conducted in the first few months of the pandemic, whereas ours was conducted from 8 months onwards, reflecting a longer-term impact of the pandemic on the carer. One study did find increased stress, anxiety, and depression in the carers at the beginning of lockdown which improved after 2 months [28] and another study found that, coinciding with a drop in social support and particularly in day services shortly after lockdown measures were introduced, an upward trend emerged for carers receiving support over three time points (April-May 2020, followed up at 6 weeks and 12 weeks) which corresponded with a decrease in depression, anxiety, and an increase in quality of life [22]. While timing of studies is the most likely explanation for discrepant findings, most studies also did not have appropriate pre-pandemic control groups or did not compare carer measures to pre-pandemic levels at all. To our knowledge, no other study has compared changes in these measures during the pandemic to the change in a similar time frame under normal circumstances in the way that ours did. Our findings for quality of life and well-being suggest the changes experienced by carers during the pandemic are parallel to those experienced by carers with similar demographic characteristics under normal circumstances. In addition, many of the studies relied on how the carer remembered things before the pandemic, which may have been impacted by current emotional state, whereas we took data collected before the pandemic as our comparison. Another reason that the pandemic did not impact on well-being or quality of life of the carers could be related to theories that older people have an age advantage in relation to positive emotions. It is well documented that they describe more positive emotional experience and react less to negative situations than younger people [45, 46]. Carstensen et al. investigated whether this is still true following prolonged stress in the case of the COVID-19 pandemic [47]. They found that the age advantage of emotion did persist and that participants still reported positive emotions more frequently than negative emotions, with no evidence that the pandemic altered the widely documented age patterns of emotional well-being [47]. This preference for positive emotions in older people may explain at least in part why, despite the increased stress and risk to health, we did not see a decline in well-being or quality of life for carers. However, it is noteworthy that a follow up study that included a pre-pandemic control found that whilst

positive emotions were still higher for older people, the positive effect was weaker during the pandemic compared to immediately before the pandemic [48].

As far as we know this is the first quantitative study to look specifically at carers' feelings of coping, competency, and role captivity during the pandemic, with the exception of our previous study using IDEAL data which lacked a matched comparison group [29]. While there was a small but positive effect on perceived competency and role captivity, our most striking finding was for coping. While pre-pandemic data showed a pattern towards a decrease in coping between wave 1 and wave 2 of data collection, there was an increase in the number of carers endorsing better coping between wave 1 and wave 2 of data collection during the pandemic. In a qualitative study, carers expressed feelings that they were struggling to cope early on in the pandemic [14], but a study conducted later in the pandemic, between June 2020 and March 2021, found that three-quarters of the carers believed they were coping well [23]. Our initial study using IDEAL data also reported that carers were generally coping well [29].

There are several reasons why carers may have begun to feel more positive several months into the pandemic. It is likely that with the changing circumstances and disruptions to daily routines and the loss of external support, carers struggled at the beginning of the pandemic. After overcoming the initial shock of the pandemic, they may have adapted to new routines and coped with the changes. They may have been feeling more optimistic due to the rollout of the vaccination program which began in December 2020, the easing of restrictions, and reintroduction of support services, although this might be dependent on restrictions at the time of assessment. They may have more self-belief and, as several qualitative studies indicate, where their relationships with the care-recipient was already good and reciprocal, their relationship may have strengthened [43]. They may also have responded more positively due to greater negative feelings early on in the pandemic meaning they had a lower starting point than those asked pre-pandemic. Finally, in our study, the majority of carers were spouses so there may have been less of a disruption to normal routines particularly for those who did not previously rely on external support services, which is also evidenced in O'Rourke et al. [43]. Perhaps the small but positive effect we saw on role captivity arose because restrictions meant that these carers did not feel that they were missing out on other things in life because everyone was in the same posi-

tion, and, as evidence suggests, the confinement may have resulted in a strengthened relationship meaning they saw their caring responsibilities as less of a burden [1, 16, 44].

The number of carers spending over 10 hours caring increased by 18.9% between w1 and w2 for the pre-pandemic group, and 29.2% for the pandemic group, suggesting that the carers in the pandemic group were spending more hours caring for the person with dementia. This increase in carer hours is not surprising since support services such as visits from paid carers were reduced during the pandemic. However, while carer hours were controlled for in our analysis, similar results were obtained if carer hours were not controlled for, indicating that the findings we observed were not affected by the number of carer hours. While this might seem surprising, perhaps the burden of increased care may have been offset by the strengthening of the carer and care recipient relationship, or perhaps because carers had to stay at home, they felt as if they were providing more hours of care than they were previously, but were actually providing similar levels of care.

Our study had a number of limitations. While we matched two pre-pandemic carers to each pandemic carer, more carers in the pandemic group were caring for people for whom more time had elapsed since diagnosis, because most of them had been in IDEAL since T1. Time since diagnosis was adjusted for to help mitigate this limitation. Another limitation is the timing of the waves. Because no one was prepared for the COVID-19 pandemic, we had to use pre-pandemic time points that were closest in terms of time between waves for the pre-pandemic and pandemic groups, but the gap between T3 and INCLUDE (pandemic group) was larger than the gap between T1 and T3 (pre-pandemic group). However, we investigated whether including months between waves as a covariate or an interaction impacted on results and found little difference. A further limitation was the relatively small sample size recruited for the pandemic group as part of INCLUDE which can impact on statistical power, particularly when outcomes are categorical. Some categories contained few people and had to be combined for analysis. Our study was conducted between September 2020 and April 2021. There was significant easing of restrictions over summer 2020 (June to September), followed by a return to strict local and national restrictions in England and Wales from October 2020 to March 2021, and the introduction of the vaccination program in December 2020, all of which may have impacted on how

positive carers felt at the time of the assessment. In addition, quality of life and well-being are usually measured with composite measures in IDEAL, but for INCLUDE only single-item measures were administered. This was to reduce the burden for participants as the assessment was conducted over the telephone or online. This may have meant that the measures did not reflect the constructs as effectively. Finally, carers were invited to take part in the study even if the care recipient did not, so the study should encompass those who were caring for someone with more severe dementia. However, those carers who were not coping well may have been less inclined to take part, and so the group who were likely to respond most negatively may have been missed.

Our study suggests that, several months into the pandemic, COVID-19 was not having a negative impact on the well-being of carers, and in fact we found a higher level of beliefs that the carer could manage compared to before the pandemic. This study shows a more positive understanding of how carers managed during the pandemic and suggests that six to twelve months in, carers were resilient and able to cope with the changes brought about by the pandemic. At this point the pandemic was far from over, however, and the impact of living through further stages of the pandemic and the eventual return to normality on both carers and care recipients remains to be determined.

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DATA ACCESS

IDEAL data were deposited with the UK data archive in April 2020 and will be available to access from April 2023. Details of how the data can be accessed after that date can be found here: <http://reshare.ukdataservice.ac.uk/854293/>

INCLUDE data will be deposited with the UK data archive in June 2022 and will be available to access from June 2023.

SUPPLEMENTARY MATERIAL

The supplementary material is available in the electronic version of this article: <https://dx.doi.org/10.3233/JAD-220221>.

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