Cognitive Health Assessment and Establishment of a Virtual Cohort of Dementia Caregivers

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Key Words
Cognitive performance · Caregivers · Psychosocial factors · Dementia · Virtual cohort

Abstract
Background: Many factors impact caregivers’ cognitive health and, by extension, their ability to provide care. This study examined the relationship between psychosocial factors and cognitive performance among dementia caregivers and established a virtual cohort of caregivers for future research. Methods: Data on 527 caregivers were collected via a Web-based survey that assessed cognitive performance. Caregiver data were compared to corresponding data from 527 age-, race-, gender-, and education-matched controls from a normative database. Caregiver self-reported sleep, stress, health, and social support were also assessed. Results: Caregivers performed significantly worse than controls on 3 of 5 cognitive subtests. Stress, sleep, perceived support, self-rated health, years of caregiving, race, and gender were significant predictors of cognitive performance. Conclusion: In this sample of dementia caregivers, psychosocial factors interacted in complex ways to impact cognitive performance. Further investigation is needed to better understand how these factors affect cognitive performance among caregivers. This could be accomplished by the establishment of a virtual cohort that facilitates the development of digital tools to support the evaluation and management of caregiver needs in a manner that helps them remain effective in their caregiving roles.

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Introduction

In 2014, more than 15 million family members and other unpaid caregivers provided care to people with Alzheimer’s disease and other dementias in the USA [1]. Approximately two thirds of informal caregivers are women, 34% are aged 65 years or older [1], and many experience cognitive decline that can impact their ability to care for themselves and the people who depend on them [2]. The economics of informal caregiving are staggering: a recent report indicated that the value of informal eldercare in the USA is USD 522 billion annually, and USD 218 billion of that figure is accounted for by dementia caregiving [1, 3]. Because the cost of replacing informal care with formal, paid support is so high, research focused on understanding threats to effective caregiving has clear economic and policy implications.

Caregivers face many obstacles as they struggle to balance care provision with their other responsibilities such as caring for their own children, career development, and community engagement. Caregivers of persons with Alzheimer’s disease are also exposed to numerous long-term stressors that stem from care provision itself. These include the demands supporting activities of daily living (e.g., eating, hygiene), aiding with higher-level tasks (e.g., talking, writing), and contending with fluctuations in mood (e.g., depression, anger) and behavior (e.g., agitation, paranoia) [4]. Not only has it been demonstrated that caregivers of Alzheimer’s patients are at increased risk for impaired cognition, depression, anxiety, absenteeism, and increased health care utilization (e.g., physician visits, emergency room usage, and hospitalization), but also, as the severity of Alzheimer’s symptoms increase, the mental and physical health outcomes of their caregivers decrease [5, 6].

The stress associated with caregiving negatively impacts cognitive functioning [4, 7–9]. It has also been shown that cognitive decline occurs as the result of stress and depression, affecting speed, attention, and executive function [10]. Gaugler et al. [11] showed that in-home caregivers of a relative had significant declines in cognitive performance after 3 years of caregiving. Vitaliano et al. [12] reported that people who cared for a spouse with Alzheimer’s disease had a slower cognitive processing speed than non-caregivers. The latter results mirror a report from Caswell et al. [8] that showed caregivers of spouses with Alzheimer’s disease have lower levels of complex attention and speed of information processing than non-caregivers. Recently, Corrêa et al. [13] demonstrated that caregivers exhibit significantly worse performance on information processing speed and working memory than healthy controls.

Not only does this growing body of research suggest a relationship between caregiving and cognitive decline [12], it also points to the idea that the effect of caregiving on cognitive function may depend in part on the degree to which caregivers experience psychosocial risk factors. Poor cognitive function among dementia caregivers may be mediated by depressed mood [12], distress [8], hostility, metabolic factors [4], and the quality and quantity of sleep [4, 8]. A better understanding of how psychosocial factors impact the relationship between caregiving and cognition could help inform the design of programs and guide therapies to address these detrimental effects, thereby helping caregivers to maintain their important role [10].

Given previously observed relationships between chronic stress, psychophysiological functioning, and cognitive factors, we established the Health-eBrain Study (www.Health-eBrainStudy.org). This is the first large, Web-based study of informal dementia caregivers. The purpose of this report is to describe the Health-eBrain cohort and report the relationship between caregivers’ psychosocial factors and their cognitive performance. We recruited dementia caregivers to the study website and administered both a lifestyle survey and a Web-based cognitive assessment. These protocols allowed us to assess caregivers’ demo-
graphic and psychosocial profiles, and to examine relationships between these factors and cognitive performance. Our working hypothesis was that caregivers would exhibit more cognitive impairment than matched controls, and that among caregivers, psychosocial and lifestyle factors such as sleep and stress would have mitigating effects on cognitive performance.

**Methods**

This study was approved by the AnthroTronix Institutional Review Board.

**Participant Recruitment**

This cross-sectional study targeted caregivers of people with dementia. Caregivers were recruited through nonprofit foundation networks including the BrightFocus Foundation and the Geoffrey Beene Foundation Alzheimer’s Initiative, as well as through an invitation to Lumos Labs’ database. Participants were directed to the study website (www.Health-eBrainStudy.org). Eligible respondents were 18 years or older and caring for someone with dementia at the time the respondent (caregiver) completed the online survey. Consent was obtained with an online form, which was followed by administration of the lifestyle survey. Once the lifestyle survey was completed, caregivers were offered a second online consent form for Lumos Labs’ Brain Performance Test (BPT), which consists of 5 subtests. After administration of the BPT, caregivers were given the option to remain in the Health-eBrain Study registry and to be contacted for future studies. Five hundred and twenty-seven caregivers completed the lifestyle survey and the 5 subtests of the BPT, and 378 opted to remain part of the registry for future studies.

**Lifestyle Survey**

Caregivers were given a survey that collected information on demographics, duration of caregiving, and perceptions concerning stress and the availability of social support. Demographic information included age, gender, race, and education (some high school, high school graduate, some college, associate’s degree, bachelor’s degree, or graduate or professional degree). Duration of caregiving was assessed with the question: 'How many years have you been providing care?' Caregivers’ self-reported stress level was assessed with the question: 'On a scale of 1 to 5 with 1 being "no stress," and 5 being "a great deal of stress," how much stress would you say you have in your life?' Self-reported sleep was assessed with the question: 'How much sleep do you typically get each night?' The following response options were offered: 4, 5, 6, 7, 8, 9, or 10 h. To assess social support, caregivers were asked: 'On a scale of 1 to 7 with 1 being "not supported at all" and 7 being "extremely supported," how supported do you feel in your personal life?' Finally, caregivers were asked to rate their overall health by answering the question 'How would you describe your health?' with the following response options: 'poor', 'fair', 'very good', and 'excellent'.

**Cognitive Assessment**

The BPT is a Web-based battery of cognitive assessments that measures function across several domains. The BPT was designed to replicate pencil-paper tests (where applicable) for which a shift to computerized administration does not negatively impact the test mechanics. BPT results are consistent with corresponding pencil-paper testing [14–16]; the test is repeatable, changes predictably with age, is highly intercorrelated, and captures multiple, distinct cognitive abilities [14, 16, 17].

The BPT is optimized for online administration in an unsupervised environment, and was therefore well suited to this research environment. The BPT battery used in this study...
included 5 subtests: Digit Symbol Coding, Forward Memory Span, Reverse Memory Span, Trail Making A, and Trail Making B. The battery took between 20 and 30 min to complete.

**Digit Symbol Coding.** This subtest is similar to the Wechsler Adult Intelligence Scale's Digit Symbol Substitution Task. This test measures information processing speed and relies heavily on visual search and memory [18]. The subtest lasts 90 s, and caregivers used the keyboard to match a series of numbers that correspond to randomly generated symbols as quickly as possible. The primary measure is the number of correct responses minus the number of incorrect responses.

**Forward and Reverse Memory Span.** These subtests are derived from the Corsi block-tapping test [19], which assesses visuospatial short-term working memory. Forward Memory Span assesses the storage and maintenance components of working memory, whereas Reverse Memory Span evaluates information storage and information processing [20]. In computerized versions of these tasks, circles flash on the screen and caregivers reproduce the sequences by clicking on the circles [21, 22]. The subtest concludes when 2 consecutive errors on 1 sequence length are made. The primary measure is 'numbers correct'.

**Trail Making A and B.** Trail Making A involves connecting 25 encircled numbers that are randomly distributed across the computer screen by clicking on them in sequence. Task requirements are similar for Trail Making B, except that the participant alternates between numbers and letters (e.g., 1, A, 2, B, 3, C, etc.). Trail Making provides information on attention, visual search, scanning, processing speed, mental flexibility, and executive functions. Originally part of the 1944 Army Individual Test Battery [23], Trail Making was later incorporated into the Halstead-Reitan Neuropsychological Battery [24]. For both Trail Making A and Trail Making B, the primary measure is 'completion time' (there is no time limit).

Each BPT subtest included clear, interactive tutorials for which respondents were required to demonstrate a minimum level of proficiency prior to being presented with the task. The cycle of tutorial/subtest was repeated until all subtests were completed. All BPT subtest scores are scaled following a percentile rank-based inverse normal transformation, a protocol used in widely accepted measures of cognitive function such as the Wechsler Adult Intelligence Scale [25]. Normative tables are created for each BPT subtest, and the raw score from each subtest (e.g., the number of correct responses, completion time, etc.) is ranked to obtain a percentile for each raw score. The position of that percentile on a normal distribution is used to convert the raw score into a scaled score where the distribution has a mean of 100 and an SD of 15.

**Data Analysis**

To assess differences in cognitive performance between caregivers and controls, each caregiver was matched on age, gender, race, and education with a unique control from Lumos Labs’ reference database of individuals who completed the same BPT battery, for a total of 527 controls and 527 caregivers. For each BPT subtest, two-sample t tests were used to assess differences in mean responses between caregivers and controls. Subtests on which we observed significant differences in performance between caregivers and controls were examined further in regression analyses. Stepwise linear regression was used to explore associations between psychosocial factors and cognitive performance among the dementia caregivers. We ran separate models for each relevant BPT subtest outcome. In each set of models, gender, race, education, years of caring, stress, hours of sleep per night, perceived social support, and perceived health were examined as independent variables. We specified an α criterion of ≤0.09 for entry into the models. The regression analyses could only be completed for the caregiver data, as we did not have psychosocial survey data for the controls obtained from the deidentified database.
Results

Data were collected from 527 dementia caregivers. These caregivers had an average age of 60.1 years, the vast majority (88%) was female, and over half were employed at the time of the study (table 1). Most respondents reported caring for a loved one (family member: 51.2%; spouse/significant other: 38.0%), and nearly 60% of the care recipients had a diagnosis of Alzheimer’s disease.

Table 2 shows that the caregivers’ cognitive performance was significantly lower than that of the matched controls on 3 of the 5 subtests of the BPT: Digit Symbol Coding, Forward Memory Span, and Reverse Memory Span (p < 0.001). These tests were the focus of our regression analyses among the dementia caregivers. The results of these analyses for each BPT subtest are shown in table 3. These results show adjusted effects of individual variables as well as variables with interaction effects. Only variables that were significant at p < 0.05 are presented. It is important to note that despite the statistical significance of the predictor variables shown in the table, the regression models in table 3 explained less than 7% of the variance for all 3 subtests.
For Digit Symbol Coding, reported hours of sleep per night was a positive predictor of performance such that for each additional hour of reported sleep per night there was a 10.5-point increase in performance. Similarly, increasing stress predicted performance on this test such that for each additional level of reported stress there was a 10.7-point increase in performance on Digit Symbol Coding. However, there was a significant interaction between sleep and stress such that above 5.5 h of sleep the effect of stress on performance was negative. The impact of self-rated health on Digit Symbol Coding performance depended on gender such that female caregivers scored 11.6 points higher on Digit Symbol Coding as the level of self-rated health increased.

Perceived support was a positive predictor of performance on Forward Memory Span such that for each increased level of reported support, there was an 8.3-point increase in performance. Similarly, each additional hour of reported sleep was associated with a 3.5-point increase in performance on this test. However, there was a small but significant interaction between sleep and perceived support such that as the number of hours of sleep and the level of perceived support both increased, test performance decreased by 0.8 points. Years of caregiving predicted performance such that for each additional year of caring there was a 1.3-point increase in Forward Memory Span performance; however, due to a small but significant interaction with education, as educational level and years of caregiving both increased, performance decreased by 0.3 points.

Non-White caregivers scored 5.9 points lower than Whites on Reverse Memory Span, and stress was inversely associated with performance on this test such that each increasing level of reported stress was associated with a 5.7-point decline in Reverse Memory Span performance. Perceived support had a negative impact on performance such that for each increasing level of perceived support there was a 4.7-point decline in Reverse Memory Span performance.

**Table 3. Regression results (n = 527)**

<table>
<thead>
<tr>
<th></th>
<th>Digit Symbol Coding</th>
<th>Forward Memory Span</th>
<th>Reverse Memory Span</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>β (SE)</td>
<td>t</td>
<td>β (SE)</td>
</tr>
<tr>
<td>Race (non-White vs. White)</td>
<td>–5.9 (3.0)</td>
<td>–2.0*</td>
<td>–5.8 (2.9)</td>
</tr>
<tr>
<td>Stress</td>
<td>10.7 (5.4)</td>
<td>2.0*</td>
<td>3.5 (1.4)</td>
</tr>
<tr>
<td>Hours of sleep</td>
<td>10.5 (3.2)</td>
<td>3.3**</td>
<td>1.3 (0.6)</td>
</tr>
<tr>
<td>Years of caregiving</td>
<td>–2.1 (0.8)</td>
<td>–2.6*</td>
<td>–0.8 (0.3)</td>
</tr>
<tr>
<td>Perceived support</td>
<td>8.3 (2.6)</td>
<td>3.2**</td>
<td>–4.7 (2.0)</td>
</tr>
<tr>
<td>Stress × sleep</td>
<td></td>
<td></td>
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<tr>
<td>Sleep × support</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gender × health</td>
<td>11.6 (4.1)</td>
<td>2.8**</td>
<td>–0.3 (0.1)</td>
</tr>
</tbody>
</table>

* p < 0.05, ** p < 0.01. a White = 0, non-White = 1. b No stress = 1, great deal of stress = 5 (mean = 3.67, SD = 0.87). c 4, 5, 6, 7, 8, 9, or 10 h (mean = 6.48, SD = 1.23). d 1–41 years (mean = 5.03, SD = 4.27). e Not supported at all = 1, extremely supported = 7 (mean = 3.83, SD = 1.69). f Male = 0, female = 1. g Poor = 1, fair = 2, very good = 3, excellent = 4 (mean = 2.81, SD = 0.66).

When comparing cognitive performance between dementia caregivers and matched controls, our data are consistent with previous results showing that dementia caregivers perform worse than non-caregivers on tests of attention, information processing, processing...
speed, and short-term working memory [8, 9, 12, 13, 26]. Our results also support other studies showing that, in general, better psychosocial functioning such as with reduced stress improves cognitive performance [27]. Notably, our large sample permitted an examination of how psychosocial factors such as stress, sleep, perceived health, and perceived level of support – alone and in combination – have complex effects on multiple cognitive domains among dementia caregivers. Our data also show that among dementia caregivers, gender, race, education, and years as a caregiver are associated with various cognitive performance domains.

The public health implications of our findings are tied to the fact that poor cognitive performance among caregivers has the potential to negatively impact the availability and quality of informal care [2, 12]. As caregivers of spouses with dementia are typically older themselves, they may experience difficulty performing complex activities of daily living for themselves as well as the recipient of their care. For example, caregivers typically oversee medication management, a complex task that is particularly challenging among older adults in whom polypharmacy is common [28]. Medication management requires understanding of interrelationships between multiple concepts such as dosing, whether medicines should be taken with food, and at what time of day they should be taken. These organizational demands are augmented by the fact that older adults often take multiple medications, these medications often resemble one another, pill bottles are easily lost, the writing is small, and prescription renewals are often managed by multiple care providers. When caregivers experience cognitive decline, they may be less capable of managing medications as well as myriad other tasks that are necessary to keep a care recipient at home rather than in a nursing home or other residential care facility [12]. Caregivers’ inability to remain in their role drives up the cost of dementia care and increases its social burden such that even if dementia rates remained stable, health expenditures for dementia could come close to double the amount by 2040 because of the aging of the population [29]. Finding new and effective means of monitoring and managing the health and welfare of the caregiving population has obvious implications for controlling these costs.

Although the differences in cognitive performance between caregivers and controls were statistically significant, caregiver performance in our sample was still within 1 SD of the normed average. Thus, our caregivers’ cognition was ‘normal’ relative to the general population average. However, because we had access to matched controls, we were able to detect differences in cognitive performance between caregivers and controls that may have been due to differences in psychosocial risk factors like stress and sleep. We speculate that these factors contribute in complex ways to caregivers’ cognitive decline, a hypothesis that suggests the need for a better understanding of ways to assess and manage these aspects of caregiver burdens.

Chronic stress associated with caring for someone with dementia may explain some or all of the poorer cognitive performance observed in our caregiver sample [26]. Stress elevates cortisol levels in dementia caregivers [30], and high cortisol is linked to impaired cognitive functioning, particularly reduced attention and memory [31, 32]. Stress also stimulates arousal and suppresses sleep [33, 34]. We found that stress was generally a negative predictor of performance except at very low hours of sleep, at which point higher stress increased performance (Digit Symbol Coding), and at higher levels of education, which mediated the negative effect (Reverse Memory Span).

Although longer sleep generally increased cognitive performance for both Digit Symbol Coding and Forward Memory Span, it interacted in complex ways with stress and perceived support such that for high levels of stress, longer sleep decreased test performance. The impact of perceived social support was positive for Forward Memory Span; however, increased perceived social support had a negative effect on Reverse Memory Span perfor-
mance. The latter finding contrasts with a previous study that linked increased social ties with favorable caregiver health outcomes [35].

It is important to note that although our regression analysis revealed statistically significant predictors of cognitive performance, none accounted for more than 7% of the variance in the data. One explanation for the modest performance of the models could be linked to the fact that the caregivers in this sample had cognitive profiles that were well within the normal range, and the distributions of many of their individual-level characteristics were also clustered. Our regression models, which focused only on caregivers, relied on outcome data that were clustered in a relatively small portion of the overall distribution of cognitive performance. Had our sample included larger numbers of caregivers with markedly poorer performance, or more detailed psychosocial measures, it is possible that the models would have explained more of the variability in the data.

Another explanation for the low model performance could be residual confounding by unmeasured factors such as depression [12, 27, 36–38]. It has been estimated that between 11 and 37% of caregivers suffer from depression [39–41], and previous studies have tied impaired sleep [10] and decreased social ties [35] to depression. Given these links, our findings suggest that future studies should target caregiver depression in parallel with other, related factors. In fact, caregiver interventions are most effective when they are comprehensive or multidimensional [42, 43] and targeted to meet the needs of individual caregivers [44]. Technological innovations that allow for virtual engagement are well suited to addressing this problem by bringing services to caregivers when and where they are needed.

This study has a number of important limitations. We did not have lifestyle data for controls, precluding an examination of these factors across the two study groups. The caregiver population in this study was self-identified and, by definition, consisted of individuals who have access to and were comfortable with computers and who had time to complete the assessments. However, all Web-based research targets individuals with computers, a consideration that will become less relevant over time as secular trends related to computer use in older age become more prevalent. An additional consideration concerns the fact that although most of the caregivers in our sample were informal, others were paid. The distinction between paid and unpaid caregivers is linked to age, with paid caregivers typically being younger than informal caregivers, who are often adult children of the care recipient. Although this heterogeneity may have impacted our results, it reflects the caregiving landscape and is therefore important to recognize and understand. Our results are also limited in that they are cross-sectional and focused on cognitive testing at a single time point. Despite the cross-sectional nature of the data, our cohort of dementia caregivers represents one of the largest samples of its kind, and the fact that many of these individuals consented to future study indicates that we will be able to explore longitudinal relationships among the same individuals going forward. The study also did not include measures of attention, cognitive flexibility, or problem solving, key cognitive domains that contribute to performing complex functional activities, which in future studies could provide insight into the relationship to cognitive performance and activities of daily living. Finally, despite relatively low $R^2$ values for our models, many of the relationships we observed indicated that even among the relatively healthy caregivers in our sample, there are measurable decrements in cognitive function. This suggests that caregivers with early decline may be identified with inexpensive screening technologies that, in turn, may facilitate provision of support that can help caregivers remain effective in their roles.
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Disclosure Statement

The authors declare the following potential conflicts of interest with respect to the research, authorship, and/or publication of this article: Lumos Labs is the developer of the BPT.

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