Fear, Guilt and Shame in the Pre-Senior Offspring of Individuals Suffering from Mid Stage Dementia and Beyond

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Abstract

Objectives: With this study, we looked to resolve three primary questions: 1. Do the offspring of individuals suffering from dementia feel guilty about their parent’s circumstances and the steps that offspring have taken in order to deal with the parent’s caregiving needs? 2. Does the offspring fear that they, too, will inevitably develop the disease? 3. Do they, the offspring, feel shame in regards to this genetic tendency?

Methods: Fifty-three participants took part in an online Likert-scale survey that asked each to gauge the intensity of their fear of developing dementia themselves, their guilt in deciding their parent needed formal full-time caregiving aid, and the shame associated with their genetic tendency toward the disease. The questionnaire scored each answer on a scale of 1-5, with 1 being “not at all” and 5 being the most intense fear, guilt and/or shame.

Results: Per fear, participants averaged a 3.30 score. Sixty-nine percent of those individuals replied “occasionally” or greater. In addition, the DLB subgroup averaged a score of 4.36. Per guilt, participants tallied an average score of 3.60. Seventy-four percent answered “occasionally” or greater. Also, the EOD offspring group recorded an average score of 4.19. Per shame, 15% of the overall group answered “occasionally” or higher, with the overall group averaging a score of 1.72.

Conclusion: Our findings suggest that a majority of people with a parent suffering from mid-stage dementia and beyond are quite fearful they, too, will develop the disease. These individuals also feel significant guilt in regards to their parent’s disease and the caregiving decisions they, the offspring, have made in regards to that parent. And while the shame associated with genetic tendencies is not as intense as the fear or guilt, this study finds that stigma surrounding dementia does still exist.

Keywords: DLB fear; EOD guilt; Dementia shame; Family caregiving; Early-onset dementia

Introduction

Dementia's a general term used to describe a decline in mental faculties that interferes with everyday life; not a specific disease, it instead consists of a wide range of symptoms associated with cognitive decline [1]. Examples of dementia subcategories include Alzheimer's disease, Parkinson's disease and Huntington's disease [2]. The term early onset dementia, otherwise known as OED, refers to dementia that first occurs in persons before the age of 65; the dementia may be caused by Alzheimer's disease or another related condition, and may appear in any stage: early, mid or late [3]. Dementia with Lewy bodies, also known as DLB, is a progressive neurodegenerative dementia that leads to a decline in thinking, reasoning and independent function [4]. The disorder affects some 1.4 million Americans [5]. Its symptoms can include, among other things: pronounced variations in attention and alertness; recurrent, detailed visual and auditory hallucinations; spontaneous features of parkinsonism; REM sleep behavior disorder (RBD); and transient, unexplained loss of consciousness [6].

At present, more than 5.4 million Americans are living with some form of dementia, approximately 200,000 of them suffering from EOD, and one in nine Americans aged 65 or older has the disease; in fact, so widespread is the disorder that it now kills more Americans than prostate cancer and breast cancer combined, with one in three senior individuals dying from dementia [7]. A 2013 study even shows that those figures may be understated, as it finds a larger number of U.S. deaths are attributable to dementia than reported on death certificates [8].

A review of mortality rates for the United States and United Kingdom finds that over the past quarter century, deaths related to dementia have increased by 589%, while deaths from cancer have risen 14%, and deaths from heart disease have actually decreased by 34% (Table 1).

Furthermore, current research suggests dementia caregivers often suffer from anxiety, depression and physical ills [9-11]. Research also indicates that those providing informal care for persons stricken with dementia related disorders many times, offspring are also prone toward mental anguish, depression and even suicidal ideations [12-14]. Furthermore, current research suggests that heredity plays a significant role in the development of Alzheimer's disease and other dementia disorders [15,16]. And finally, dementia, to a large degree, remains stigmatized in most cultures [17-22].

Table 1: Mortality rate in United States and United Kingdom.

<table>
<thead>
<tr>
<th>Disease</th>
<th>1989</th>
<th>2013</th>
<th>Net Gain/Loss</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dementia</td>
<td>41,963</td>
<td>289,261</td>
<td>+589%</td>
</tr>
<tr>
<td>Cancer</td>
<td>658,012</td>
<td>746,891</td>
<td>+14%</td>
</tr>
<tr>
<td>Heart Disease</td>
<td>671,436</td>
<td>443,327</td>
<td>34%</td>
</tr>
</tbody>
</table>

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In this study, we look to resolve three primary questions: 1. Do the offspring of individuals suffering from dementia feel guilty about their parent's circumstances and the steps that offspring have taken in order to deal with the parent's caregiving needs? 2. Does the offspring fear that they, too, will inevitably develop the disease? 3. Do they, the offspring, feel shame in regards to this genetic tendency?

Methods

Diagnosis

Each dementia case was diagnosed by a trained physician (MD), either a neurologist or another cognitive specialist, such as a geriatric psychiatrist. Evaluation tools included the Mini State Mental Examination (MMSE), the General Practitioner Assessment of Cognition (GPCOG), and the Short Informant Questionnaire on Cognitive Decline in the Elderly (IQCODE), as well as positron emission tomography (PET) scans and functional magnetic resonance imaging (fMRI).

Medications

Participants reported the use by their parents of various medications for issues related to their disease, such as the dementiatspecific pharmaceuticals Aricept (donepezil) and Exelon (rivastigmine). One participant reported that her mother was taking the NMDA receptor-blocking drug Namenda (memantine). Other drugs associated with dementia symptoms were also being used by participants' parents, including Ativan (lorazepam) for anxiety, Effexor (venlafaxine) and Cymbalta (duloxetine) for depression, Seroquel (quetiapine) for confusion, and Risperdal (risperidone) for hallucinations.

Those that had abstained from medication (seven) did so for various reasons (allergies, seizure history, fear of side-effects), in spite of pharmaceutical compounds and in favor of either a "holistic approach" such as herb and supplement intake or non-ingested approaches, like cognitive behavioral therapy (CBT), art therapy, and even aromatherapy. In all cases, the holistic approach was employed by individuals with in-home care.

Participants

Individuals with at least one parent suffering from midstage dementia or beyond, and with that parent living under fulltime caregiver supervision, were recruited. No other parameters applied.

In all, 53 participants took part in the study. Of that total, 94.34% were Caucasian, 3.78% Hispanic and 1.88% of African descent. The gender split was 86.79% female and 13.21% male. Participants ranged in age from 26 to 59 years old, with the mean age being 46.8 years. The 50 to 59 year old segment was the largest at 55.7%, followed by the 30 to 39 year old segment (28.3%). The 40-49 age segment made up 11.3% of participants reported the use by their parents of various medications for issues related to their disease, such as the dementiatspecific pharmaceuticals Aricept (donepezil) and Exelon (rivastigmine). One participant reported that her mother was taking the NMDA receptor-blocking drug Namenda (memantine). Other drugs associated with dementia symptoms were also being used by participants' parents, including Ativan (lorazepam) for anxiety, Effexor (venlafaxine) and Cymbalta (duloxetine) for depression, Seroquel (quetiapine) for confusion, and Risperdal (risperidone) for hallucinations.

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Results

Fear

In response to the question regarding fear, participants averaged a 3.30 score. Seventy-nine percent of those individuals replied "occasionally" or greater (Figure 1). Those in the 40-49 age group scored the highest per age segment, at 3.67, while those in the 50-59 age group scored the lowest, at 3.24. The 20-29 age group scored 3.33 and the 30-39 age group scored 3.27. The DLB group tallied the highest score of any group at 4.36, while the EOD group recorded a score of 3.43. On average, men scored lower (3.00) than women (3.35) in the fear category (Figure 2).

Guilt

In regards to guilt, participants tallied an average score of 3.60. Seventy-four percent answered "occasionally" or greater (Figure 3). As per this question, the EOD offspring group recorded an average score of 4.19, while the normal onset group registered a score of 3.22. On average, men scored 3.00, while women scored 3.70 (Figure 4).

Shame

In response to the question about shame, 15% of the overall group answered "occasionally" or higher. Twenty-four percent of the EOD offspring group responded "often" or "always" (Figure 5). And while the overall group averaged a score of 1.72, the LBD offspring group averaged a score of 2.64.

Figure 1: Shows that 69% of the total participant group answered "occasionally" or greater; a score of 3 or higher, to the fear question, suggesting that fear of developing the disease is quite prevalent in a majority of participants.
scored a 1.91 and the EOD offspring group averaged 2.10. In addition, men recorded an average score of 1.43, with 88% answering “never,” while women recorded an average score of 1.80, with 35% answering something other than “never” (Figure 6).

**Discussion**

**Summary**

When surveying the topic of dementia, we find research pays considerable attention to the psychological makeup and needs of professional caregivers, both in-home and in-facility. However, current research rarely considers the emotional and/or psychological state of the dementia sufferer’s offspring. We’ve done so with this research.

And while our study is, in regards to the overall size of the dementia community, a relatively small data sample, we believe it’s both a relevant and necessary first step: with a cross-section of age, gender, and nationality, the present study serves as an excellent jumping off point for further research as it pertains to the psychological travails experienced by such offspring.
Our findings suggest that a majority of people with a parent suffering from midstage dementia and beyond are quite fearful, too, will develop the disease. This fear is most intense amongst the offspring of those with DLB, as well as those in the 40-49 age group. This study also finds that individuals generally feel significant guilt in regards to their parent’s disease and the caregiving decisions they, the offspring, have had to make for that parent. The offspring of a parent with EOD are particularly sensitive to this guilt. And while the shame of owning these genetic tendencies is not as intense as the fear or guilt, this study does find that the stigma surrounding dementia still exists, to a degree, across all participant groups.

The hope is that this study’s results demonstrate why there may be a need by memory care facilities and in-home caregiving services, insurance companies, and national healthcare schemes to make more readily available therapy techniques such as Cognitive Behavioural Therapy, Mindfulness based Cognitive Therapy (MBCT), and Dialectic Behavior Therapy (DBT) to the offspring of individuals suffering from midstage dementia and beyond.

One limitation of the present study is that of self-assessment; as with any self-reported data, there can be exaggerations of data sets due to recall bias and/or self-selection. Indeed, it’s possible that extraneous factors like unfamiliarity with self-assessment protocols, lack of exposure to dementia education, and even a dearth of self-confidence caused by unfulfilled family expectations may influence the accuracy of any self-assessment pertaining to this topic.

Secondly, we must also consider that the offspring most willing to participate in such a study are, quite possibly, those most engaged with the parent and his/her dementia issues, suggesting, perhaps, a relatively heightened emotional and/or psychological response to self-survey questions.

With that said, research suggests that the internal consistency of self-assessment is generally high [23]. In addition, data from the present study does remain relatively consistent across genders and national borders, and throughout age groups and dementia subgroups.

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References

5. Lewy Body Dementia Association (2016) What is LBD?