The Effects of Adult Day Programs on the Quality of Life of Individuals with Alzheimer’s disease: A Pilot Study

Authors

Emma Bartfay and Wally J. Bartfay

Faculty of Health Sciences, University of Ontario Institute of Technology (UOIT), Ontario, Canada

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Abstract

Little research has been conducted on examining the relationship between caring interventions such as adult day programs (ADPs) and the quality of life (QOL) of individuals with Alzheimer’s disease. Our study objective was to investigate the merits of attending ADPs on the QOL of these individuals. We hypothesized that individuals with Alzheimer’s disease who attended ADP had higher QOL than those who did not. We also hypothesized that individuals with Alzheimer’s disease who attended ADP enjoyed comparable QOL as their non-diseased counterparts. To explore these issues, we piloted a cross-sectional study in Durham, Ontario, Canada. We recruited 130 participants at five ADPs and at six caregiver support groups in the region. Among them, there were 73 disease-free ADP clients,
28 ADP clients with Alzheimer’s disease and 29 individuals with Alzheimer’s disease who did not attend ADPs. Study procedures involved primary data collection using assisted self-report questionnaires and a 13-item quality-of-life scale. This current paper provided a detailed description of the study process. We also provided results that showed overall QOL scores for individuals with Alzheimer’s disease who attended ADP were comparable to those without the disease (2.7 vs. 2.9, p=0.1), whereas the scores were much lower for individuals with Alzheimer’s disease who did not attend ADP than those who attended ADP (1.9 vs. 2.7, p=0.0001). Individuals who attended ADP consistently provided higher ratings than those who did not attend ADP. Based on our findings, we concluded that attending ADPs may be partially responsible for the observed differences in QOL.
Keywords: Alzheimer’s disease, dementia care, adult day program, quality of life

Introduction

We are living in a rapidly aging world. The Centre for Disease Control and Preventions (2003) asserted that we could expect to see almost 1 billion persons aged 65 and older in 2030 worldwide. In Canada alone, Statistics Canada (2010a) estimated that the number of older persons could increase to 15 million in the next 50 years. That is to say, persons aged 65 or old would represent 28% of the projected Canadian population by 2061. Previous research estimated that 6% to 10% of individuals aged 65 or older, 5% to 15% of those aged 70 and older, and as much as 40% of those aged 85 and older suffer from dementia.
Together, these figures implied that we will face an unprecedented number of age-related medical adversities, including Alzheimer’s disease and other forms of dementia, in a very near future. Furthermore, the latest projections from the US Alzheimer’s Association (2010) suggested that one diagnosis of Alzheimer’s disease was made every 70 seconds presently in America. This time would decrease to every 33 seconds within the next four decades. Canada faced with a very similar situation that the number of individuals living with Alzheimer’s disease and other forms of dementia could reach 1.1 million (CIHI, 2010).

Dementia is not a single disease, but a collection of syndromes that are characterized by a plethora of cognitive, emotional and behavioural impairments (Chapman et al., 2006). To-date, only a
few risk factors are identified, including family history, education, head injury, genetic predisposition and age (The Canadian Study of Health and Aging [CSHA] 1994a; Jorm 1997; Hall et al., 1998; Myhrer, 1998; Lindsay et al., 2002). However, as there is still no cure for dementia and only 9% of the cases are reversible (Chapman et al., 2006), we need to examine how we can care for these individuals and how to improve their quality of life (QOL).

_Dementia Care around the World_

Dementia care is a major global issue that concerns us both medically and socially (Ross-Kerr et al., 2003, Shanley, 2006). It also imposes a tremendous socioeconomic cost on our society (Zhu et al., 2006; Yaari and Corey-Bloom, 2007). The US Alzheimer’s Association (2010) stated that the country’s
medicare payments for individuals with Alzheimer’s disease and other forms of dementia were estimated to be $172 billion in 2010. With our aging society, dementia care is an issue that we cannot afford to ignore.

The majority of the studies that examined dementia care were conducted in the US (Gaugler et al., 2003; Ready et al., 2004; Carroll et al., 2005), Japan (Miyamoto et al., 2002; Suwa 2002; Hosaka and Sugiyama, 2003; Fukushima et al., 2005), Korea (Kim et al., 2002), Australia and New Zealand (Shanley, 2006), UK (Upton and Reed, 2005; Hoe et al., 2005; Hoskins et al., 2005; Reilly et al., 2006) and the rest of Europe, including Italy (Balla et al., 2007), the Netherlands (Droes et al., 2004a; Droes et al., 2004b), Germany (Zank and Schacke 2002; Zank and Frank 2002), Denmark (Vogel, 2006), Sweden (Ericson et al., 2001,
Andren and Elmstahl 2005; Mavall and Thorslund, 2007), France (Thomas et al., 2002; Gramain et al., 2004). In Canada, a landmark study of dementia was conducted by the Canadian Study on Health and Aging (CSHA) working group in the early 1990’s. In their reports (CSHA Working Group 1994b, 2000), the working group estimated that 1 in 13 individuals aged 65 and over had dementia, and the risk doubled for every five years of life after 65. However, only a handful of Canadian studies were conducted to determine the relationship between ADPs and QOL among older individuals and their caregivers. In a study that involved 14 ADPs in Alberta, Warren et al. (2003) examined the impact of ADPs on family caregivers of the elderly. They found that caregiver burden, QOL of caregivers and the perceived health status remained constant over time after attending ADPs. Nonetheless, the lack of improvement in outcome measures, and
particularly the lack of a control group in the study, made it impossible to evaluate the findings.

Baumgarten et al. (2002) conducted a randomized trial on the satisfaction of ADPs for frail elderly in the Province of Quebec, Canada. Although the study found that clients and their family caregivers were generally satisfied with the service, there was no evidence that suggested reductions neither in the clients’ anxiety, depression and functional status nor in the family caregivers’ burden. The authors attributed the non-positive findings to the low participation of ADPs. Other Canadian research studies, Burdz et al. (1988), Chappell et al. (2001) and Bartfay and Bartfay (2013) also focused on caregivers. It is noteworthy to mention that of the aforementioned Canadian studies, only Burdz et al. (1988) was conducted specifically on individuals with dementia,
and all other Canadian studies mainly concerned with the impacts on caregivers.

Adult Day Programs (ADPs) as a Form of Caring Intervention

Caring for the elderly, particularly those with dementia, is physically and emotionally arduous (Lee and Cameron, 2004; Upton and Reed, 2005; Andren and Elmstahl, 2005). In Canada, our ever escalating health care costs have prompted policy makers to put more emphasis on community care in recent health care restructuring as a way to curtail spending (Warren et al. 2003). Indeed, many older individuals prefer to live at home for as long as they can and to delay institutionalization (Zank and Schacke, 2002; Ross-Kerr et al., 2003; Warren et al., 2003). Although this stay-at-home arrangement is favoured by many,
family members often assume the primary care-giving role and suffer from many physical and emotional consequences (Gramain and Malavolti 2004; Gitlin et al., 2006; Yaari and Corey-Bloom, 2007). ADPs have been projected to reduce family strain on caring for the elderly (Gaugler et al., 2003; Gramain and Malavolti, 2004). In fact, interventions are now emerging as an important aspect of dementia care (Chapman et al., 2006). Daytime care was originated in Moscow in the 1930’s and further developed in the UK after the Second World War (Ross-Kerr et al., 2003; Gaugler et al., 2003). In Canada, daytime services came into existence in Montreal in the 1940’s, and it became popular in the US in the 1960’s (Gaugler et al., 2003). Generally speaking, day programs provide out-of-home services during daytime where clients continue to live at home. Many programs are designed to promote cognitive functioning, facilitate
independence and maintain QOL and health of their clients (Edelman et al., 2004; Debelko, 2005; Chapman et al., 2006). There is evidence that suggests rapid decline in cognitive function may decrease survival in individuals with Alzheimer’s disease (Wilson et al., 2006).

Previous research showed that mental exercises, such as reading, playing games, puzzles and musical instruments were associated with decreased risk of dementia (Wilson et al., 2002a; Wilson et al., 2002b; Verghese et al., 2003). Nonetheless, studies designed to examine the outcomes and effectiveness of ADPs remained scarce and lacked rigour (Lee and Cameron 2004; Edelman et al., 2004; Jeon et al., 2005). These studies also have mixed results due to differences in case mix, program emphases, inadequate sample size, low usage of services and the lack of control groups
Gaugler and Zarit, 2001; Ross-Kerr et al., 2003; Gaugler et al., 2003; Zarit et al., 2003). Most of the evaluations also focused on caregivers, rather than on the clients (Bartfay and Bartfay, 2013; Miyamoto et al., 2002; Zank and Schacke 2002; Gaugler et al., 2003; McCann et al., 2005; Marvall and Thorslund, 2007). If ADPs are designed to benefit both clients and caregivers, we must also examine the needs of the clients. 

**Quality of Life (QOL)**

Health-related QOL is an important aspect in evaluating an individual’s overall health status (Schneider, 2001; Hoe et al., 2005). For many individuals with Alzheimer’s disease, improving or maintaining QOL may be the best care that we can offer. It has long been argued that “QOL should be the central goal of our professional activity” (Whitehorse and Rabins, 1992). Hoe et al.
(2005) argued that measuring QOL was more important than the number of symptoms one experienced. They also stated that QOL did not decrease as cognitive functions declined.

Although the issues of QOL in individuals with dementia have received some attention since the 1990’s (Ready et al., 2004), empirical assessment has largely been ignored (Schneider 2001; Logsdon et al., 2002; Edelman et al., 2004). Fortunately, researchers are slowly recognizing the importance of QOL. For example, a Japanese qualitative study found a remarkable change in attitude in individuals with dementia after attending a daycare program (Fukushima et al., 2005).
Objectives and Hypotheses of Study

Empirical studies relating to ADPs have mainly focused on the impact of family caregivers. Little attention is paid on the relationship between QOL of individuals with Alzheimer’s disease and attending ADPs. According to Chapman et al. (2006), the majority of these individuals did not receive appropriate treatments. If we are truly committed to this vulnerable population, every effort must be made to reach out to them. As most dementia, including Alzheimer’s disease, is not reversible, timely intervention is crucial to optimize their QOL. According to many researchers (Gaugler et al., 2003; Edelman et al., 2004), the benefits of ADPs for individuals with dementia needed to be explored and the broader concerns of QOL deserved further
attention. With our aging population, individuals suffer from Alzheimer’s disease will only continue to grow.

Accordingly, the objective of this pilot study was to explore the effect of ADPs on the QOL of individuals with Alzheimer disease. We hypothesized that individuals with Alzheimer disease who attended ADPs had higher QOL than those who did not attend ADPs. We further hypothesized that individuals with Alzheimer’s disease who attended ADPs enjoyed comparable QOL as other ADP clients without Alzheimer’s disease.

Methods

A pilot study was conducted using a cross-sectional design in Durham, Ontario, Canada. The regional municipality of Durham
is located in the southern part of Ontario, with an approximate area of 2,500 square kilometres and a population of just over 560,000 persons (Statistics Canada, 2010b). Participants were recruited at five adult day programs and at six caregiver support groups in the region, involving primary data collection using assisted self-report questionnaires and a 13-item quality-of-life scale.

Sample 1 – Adult day program (ADP) clients with Alzheimer’s disease and sample 2 – Adult day program (ADP) clients without Alzheimer’s disease

ADPs within the area of Durham region were contacted and meetings were arranged with the managers and staff members at
Each site. All day programs had similar program emphases and case mix. The programs provided social, recreational and educational activities to their clients. The clients consisted of frail elderly, individuals with needs relating to physical disability, aging and mental problems as well as individuals with cognitive impairment. All ADP clients were eligible to participate if they attended the programs at the time of data collection. The only ineligibility was when the client was unable to speak, write or comprehend English, where they would be excluded from the study. No one was excluded in our pilot study.
Subject Recruitment and Data Collection Procedure for Sample 1 and Sample 2

To recruit day program clients to participate in our study, we visited the day program meetings and explained the study to the clients with the presence of day program staff members. Letters of invitation were then circulated. For clients who were interested in the study, members of the research team subsequently went over the consent form with these clients individually. For clients who were not able to complete the consent form, we approached the family member for assent when the clients were picked up at the end of the day program session. Upon completion of the consent form, the clients were given a copy of the questionnaire to complete with one-on-one assistance provided by the research team. When any participants felt
uncomfortable in answering questions in public, a separate room was offered to conduct the interview. The questionnaire consists of client’s basic demographic information and a 13-item QOL scale developed by Logsdon (Logsdon et al., 1999; Logsdon et al., 2002). Participants rated each item as “poor” (score=1), “fair” (score=2), “good” (score=3) or “excellent” (score=4). In this study, we decided to adopt only one scale to measure QOL. According to Logsdon, this scale could be used on individuals with or without dementia. Thus, the same scale was used for all participants for consistency. Furthermore, the main reason for this choice was the simplicity of this scale, such that we did not impose too much burden on our subjects. Unlike many other scales (e.g., the dementia QOL scale (DQoL) is a 29-item scale (Ready et al., 2004)), Logsdon’s QOL scale consists of only 13 items and has demonstrated good validity, internal and test-
retest reliability (Logsdon et al., 1999). Hoe et al. (2005) adopted the Logsdon scale and found that it was valid and reliable for use in individuals with dementia with Mini-Mental State Examination (MMSE) scores of 3-11.

Sample 3 – Alzheimer’s Disease Caregiver Support Group Clients

Due to the privacy issue at the Alzheimer’s Society of Durham Region, we were not able to contact the clients directly. As an alternative, we decided to approach caregivers as a proxy to obtain patient information. All caregiver support groups in the Durham region were organized by the Alzheimer’s Society of Durham Region. These support groups were designed to encourage caregivers to talk with others who understood or had
gone through similar experiences about the issues and decisions they were facing regarding dementia care. They also provided attendees with information, emotional support and opportunities for socialization to cope with the changes in their lives. Attendees of the support groups were typically adult children and spouses of individuals with Alzheimer’s disease. Our sample included all family caregivers who attended the support groups at the time of data collection and the person they were caring for (who had been diagnosed with Alzheimer’s disease).

**Subject Recruitment and Data Collection Procedure for Sample 3**

With the permission of the Alzheimer Society of Durham Region, we attended the support group meetings and invited the
attendees to participate in our study. The same procedure described above was followed. At the meeting, we explained the study to the attendees with the presence of a support group facilitator. A letter of invitation and a consent form were then given to the support group attendees. For individuals who agreed to participate, they were given the same questionnaire to complete with regard to the information of the person they were caring for.

**Results**

We collected information on a total of 130 individuals. In reference to AD diagnosis and disease severity, all participants were asked two simple questions: (i) ‘have you ever been told by your doctor that you have Alzheimer’s disease’ and (ii) ‘if so,
what is the severity of the disease?' Based on the answer of either ‘yes’ or ‘no’ to the first question, participants were classified into one of three groups: (1) disease-free ADP clients (n=73), (2) individuals with Alzheimer’s disease who attended ADP (n=28), and (3) individuals with Alzheimer’s disease who did not attend ADP (n=29). For the latter question, participants were asked to select his or her disease severity from mild, moderate or severe. Table 1 summarizes the basic demographic characteristics of our participants. We first compared only participants who attended ADPs. The mean age (74.5 vs. 74.7) was similar regardless of the disease status. There were slightly more females who did not have a diagnosis of Alzheimer’s disease. Furthermore, ADP clients with a diagnosis of Alzheimer’s disease were more likely to have post-secondary education (50.0% vs. 25.0%), to be married (81.3% vs. 44.0%) and living with a spouse/partner
(68.8% vs. 36.0%), as well as to have in-home support (57.1% vs. 39.0%). Next, we compared only participants who had a diagnosis of Alzheimer’s disease. We found that individuals who attended ADPs were slightly younger (74.7 vs. 78.0). They were also more likely to have a post-secondary education (50.0% vs. 34.6%), to be married (81.3% vs. 74.1%), and be living with a spouse/partner (68.8% vs. 29.6%), as well as to have in-home support (57.1% vs. 24.0%). In terms of disease severity, there were equal numbers of moderate stage but ADP clients were likely to have a mild stage (46.2% vs. 19.2%), and less likely to have a severe stage of Alzheimer’s disease (7.7% vs. 34.6%).

Please see Table 1 in the PDF version
In accordance with our hypotheses stated earlier, we performed separate 2-sample t-tests to compare the three groups. First, we involved only individuals who attended ADPs. Our results showed that the overall QOL scores for individuals who attended ADP were similar to those of non-disease ADP clients (2.7 vs. 2.9, p=0.1). We also observed that all specific QOL scores were similar among these individuals, with the exception of the memory category (see Figure 1). In the memory category, approximately 60% of ADP clients without Alzheimer’s disease rated themselves as either good or excellent, whereas only 17% of ADP clients with Alzheimer’s disease rated themselves as either good or excellent.

Second, we involved only individuals with a diagnosis of Alzheimer’s disease. The overall QOL scores were significantly
lower for individuals who did not attend ADP than those who attended ADP (1.9 vs. 2.7, p=0.0001). In addition, we noticed that individuals with Alzheimer’s disease who attended ADPs were more likely to rate each item as either good or excellent, as compared to those who did not attend ADPs. This observation held true for all QOL categories.

Please see figure 1 in the PDF version

Discussion

Due to its subjective nature, assessment of QOL is a highly complex matter. To make matters worse, individuals with severe cognitive impairments may be unable to evaluate their own health status and/or issues relating to their QOL. To compensate
this predicament, many researchers either used proxy measures (e.g., ask a family member or professional caregiver to rate the patient’s QOL), or abandoned the idea of measuring QOL altogether and concentrated on caregivers only. Our research team, however, believed it was of paramount importance to understand the needs and concerns of individuals with Alzheimer’s disease as well. In this study, our first task was to compare the QOL of all ADP clients with or without Alzheimer’s disease. Our results showed that all ADP clients had similarly favourable QOL ratings, regardless of whether they have the disease or not. The lack of a difference in QOL ratings among individuals with or without Alzheimer’s disease may provide support for future studies to further examine the benefits of ADPs.
Our second task was to compare the QOL of individuals with Alzheimer’s disease according to whether they attended ADPs or not. There were two issues relating to this task. First, information was obtained differently for each group. For the ADP attendees who also had Alzheimer’s disease, they filled out the questionnaires themselves with a one-on-one assistance from the research team. For individuals who did not attend ADP, information was obtained by proxy (caregivers). There has been much debate over the use of patient ratings versus proxy ratings. In particular, proxy ratings may sometimes be mixed-in or affected by the proxy’s own feelings. Second, there were slightly more individuals with severe disease stage in our non-ADP groups. Although disease stage may explain some of the differences in QOL, we believed that this difference only represented a portion of the QOL. As pointed out by Hoe et al.
(2005), QOL did not decrease as cognitive function decreased. Thus, our study result provided evidence that suggests that ADPs may be partially responsible for the observed differences in QOL ratings among individuals with Alzheimer’s disease who attended or did not attend ADPs, and the merits of ADP should be further examined. Given the prevailing shift of the Canadian health care system from institutional- to community-based care, future studies are desperately needed to examine all community-based interventions, including ADPs, and their effects on the QOL of individuals living with Alzheimer’s disease and other forms of dementia. As there is still no cure for Alzheimer’s disease, improving and maintaining QOL may be the best course of action that we can provide to this group of vulnerable individuals. In addition, longitudinal studies, that follow participants over time, may provide further evidence that community-based
interventions can provide sustainable improvement. As dementia is a condition that also greatly affects family caregivers, studies that examine the impact of these interventions on caregivers such as caregiver strain and burden are warranted. Indeed, an understanding of how community-based services and resources for client’s with Alzheimer’s disease and their families are critical in order to provide safe and cost-effective services to this growing population.

References


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