Which Is More Challenging? To Be An Alzheimer’s Patient, Or To Be A Caregiver For One?

Abstract

**Background**: The behavioral and psychological symptoms of Alzheimer’s disease impose a significant burden on the patient's family and caregiver.

**Objectives**: The purpose of this study was to assess the health of individuals who care for Alzheimer’s patients by using the World Health Organization Quality of Life Questionnaire (WHOQoL-Bref) and the Caregiver Stress Scale, and also to evaluate the relationship between the health and demographic data of these caregivers.

**Methods**: The study was conducted with 55 participants providing care to Alzheimer’s patients who were admitted to Neurology Polyclinic, and with 55 healthy control groups with age and demographic characteristics similar to that of the caregivers. The form distributed to the participating caregivers included questions relating to socio-demographic characteristics, patient information, the caregiver’s current health, and the Caregiver Stress Scale. The WHOQoL-Bref questionnaire was also administered to each participant to assess their quality of life.

**Results**: Compared to the control group, caregivers had statistically significantly lower scores in all domains (Physical Health, Psychological Health, Social Relationships, Environmental Health) of the WHOQoL-Bref questionnaire ($p=0.000$). The mean stress index of the caregivers was $7.31\pm3.31$. A negative correlation was identified between the stress index score and the domain scores of the WHOQoL-Bref questionnaire.

**Conclusion**: The disease especially affects the caregivers of Alzheimer's patients, leading to an increase in their stress index, and to a decrease in their quality of life. When treating Alzheimer's patients, it is important to also consider their caregivers, to inform them about Alzheimer's disease, and to even provide treatment for them when needed.

**Key Words**: Alzheimer Disease; Caregivers; Dementia; Quality of Life; Stress Psychological
Introduction

Is it more challenging to be an Alzheimer’s patient, or to care for one? In the initial stages of Alzheimer’s disease patients begin to experience memory loss and changes in personality. They might experience difficulties in remembering recent events, in recognizing people they know, or in recalling the name of objects. However, patients at this early stage of the disease generally do not require external care or assistance. In the mid-stages of the disease, patients begin to experience delusions, significant changes in personality, denial of memory impairment, and significant impairment of occupational activity. At this stage, patients become increasingly dependent on external care. In the advanced stages of the disease, when patients become unable to perform daily activities of their own, and begin to develop motor skill impairment along with urine and fecal incontinence, thus, external care becomes obligatory [1].

Advances in the treatment of Alzheimer’s diseases, the medical care provided in Neurology Polyclinics and the increase in the ratio of patients seeking treatment has considerably lengthened the average lifespan of Alzheimer’s patients. The 2010 World Alzheimer Report estimates 35.6 million patients across the world with dementia, and that this number will reach 65.7 million in 2030, and 115.4 million in 2050 [2]. In addition to these developments; industrialization, adaption to urban life and economic concerns are leading most family members to work actively, which reduces the number of individuals at home who might provide care to patients. Those who can remain at home are generally the patient’s spouse, children, and daughters-in-law, who assume the role of caregivers either willingly or unwillingly. In Turkey, care for Alzheimer’s patients is generally given at home, although nursing homes for the elderly and those in need of constant care have also appeared in recent years. In addition to the known symptoms of Alzheimer’s patients, the problems experienced by caregivers have also begun to attract attention in recent years. Factors that affect the caregiver burden includes the caregiver’s age, gender, relation with the patient, education level, economic status, and accompanying physical and mental diseases [3].

In this study, the caregiver burden and health of those taking care of Alzheimer’s patients were evaluated by using the Caregiver Stress Scale and the WHOQoL-Bref questionnaire, and comparisons were made between the scores for these questionnaires and the caregivers’ demographic data.

Methods

This study was designed as a prospective study. The study was conducted with the caregivers of Alzheimer’s patients who were admitted to the Neurology Polyclinic between January 2013 and July 2013. An Ethics Committee approval was obtained for the study (Number: B:30.2.HRU.0.20.05.00.050.01.04-166). The aim was to include mainly those who care for the patients at home rather than those accompanying the patients for their clinic visits. The address and telephone numbers of these caregivers were recorded. All participants were informed regarding the purpose of the study, and written informed consents were obtained from each one of them. Information regarding factors that affect the caregiver burden, such as the caregiver’s age, gender, relation to the patient, education level, economic status, and accompanying physical and mental diseases were recorded. Information on the duration of the disease and any other concurrent diseases were also recorded. Questions from the WHOQoL-Bref Questionnaire and Caregiver’s Stress Index were read to the participants one-by-one. The questions were further clarified to the participants whenever necessary, and their answers were recorded. Assistance was provided to illiterate participants for facilitating the completion of the questionnaire forms. The relevant tests were administered to 26 of the participants with the same method. However, it was not possible to gather all 55 caregivers at the same time and place to complete the study forms. Among the study participants, 12 were not available to meet during working hours; these participants were therefore invited to the Neurology service at a later date that was suitable for them, and the forms were completed during their eventual visit. For 17 of the study participants, the study forms were applied on the same day the patient was diagnosed with Alzheimer’s disease, and immediately after the relative to assumed the role of the patient’s caregiver was determined. The control group was selected among healthy individuals admitted to the Neurology Polyclinic who shared similar age and demographic characteristics as the study group, but lacked any history of previous psychiatric diseases, and had no patient or elderly person to whom they provided care. The forms for the control group were performed in the Patients’ Rights unit of the hospital.

Potential participants who had a history of psychiatric disease prior to becoming a caregiver were excluded from the study. Individuals in the control group who had psychiatric diseases were also excluded from the study. Thus, three individuals from the study group and five individuals from the control group (for a total of eight individuals) were excluded from the study due to a history of prior psychiatric diseases. Based on the study exclusion criteria, a total of 55 participants were included in the study group, while 55 participants with age and demographic characteristics similar to the study group were included in the control group. Written informed consent forms containing information regarding the study purpose were provided to the participants in both the study group and the control group for their reading. Signed informed consent forms were then obtained from each participant.

Data Collection Tools

In this study, the Participant Form (used to record the socio-demographic characteristics of the participants), the WHOQoL-Bref questionnaire, and the Caregiver Stress Index were used as data collection tools.

The WHOQoL-Bref questionnaire is a shorter version of the WHOQoL-100 questionnaire. It was designed by including one question from each one of the 24 sections constituting the WHOQoL-100 questionnaire, and two questions regarding general health and quality of life. The WHOQoL-100 is scored in four “domains,” which are: (i) Physical health, (ii)
Psychological Health, (ii) Social Relationships, and (iv) Environmental Health. WHOQoL-Bref also consists of four domains: Psychological Health, Social Relationships, Physical Health, and Environmental Health domains were 69.31±22.29, 66.88±22.92, 66.65±23.55, and 70.43±20.16, respectively. Among participants in both groups, 7 (12.7%) were illiterate, 34 (61.8%) were elementary school graduates, 11 (20%) were high school graduates, and 3 (5.5%) were university graduates.

Within the study group, 49 (89.1%) of the caregivers were female, while 6 (10.9%) were male. No statistically significant difference was identified between the gender of the caregivers and the WHOQoL-Bref scores (p>0.005). The average age in the study group was 47.98±13.65 years. Furthermore, 90.9% (n=50) of the caregivers were married, while 9.1% (n=5) were single. No statistically significant difference was identified between the marital status of the caregivers and the WHOQoL-Bref scores (p>0.005). With respect to education level, 7 (12.7%) of the caregivers were illiterate, 34 (61.8%) were elementary school graduates, 11 (20%) were high school graduates, and 3 (5.5%) were university graduates. No statistically significant difference was identified between the education level of the caregivers and the WHOQoL-Bref scores (p>0.005). The average score for the Physical Health, Psychological Health, Social Relationships, and Environmental Health domains were 58.81±19.46, 56.47±18.08, 52.72±21.37, and 56.60±15.83, respectively. With respect to relationship with the patient, 14.5% (n=8) of the caregivers were the patient’s spouse, 43.6% (n=24) were the patient’s child, and 41.8% (n=23) were the patient’s daughter-in-law. No statistically significant difference was identified between the caregiver’s relationship with the patient and the WHOQoL-Bref scores (p>0.005). Correlation analyses were performed in order to determine whether any correlation existed between the duration of care and the caregiver quality of life. These analyses revealed no correlation between the duration of care and the domains for the quality of life score (p>0.005).

The mean stress index of the caregivers was 7.31±3.31. A negative correlation was identified between the stress index score and the subscale scores of the WHOQoL-Bref questionnaire. Compared to the control group, caregivers had statistically significantly lower scores in all domains (Physical Health, Psychological Health, Social Relationships, and Environmental Health) of the WHOQoL-Bref questionnaire (p<0.005).

Discussion
Taking care of a patient for extended periods of time can have detrimental effects on the caregiver, both mentally and physically. A person often becomes a caregiver due to unplanned circumstances, and the role is generally assumed...
unwillingly. For this reason, many people find themselves suddenly obliged to assume this role after one of their relatives becomes affected with a serious disease or condition. Independent of language, country or culture, being a caregiver has become an integral part of life for millions of people across the world. In fact, the care provided by family members forms the basis of caregiving activities in many countries. In other words, in many countries, care for the elderly and for those with diseases that require constant care is mainly provided not by health institutions, but by family members and close relatives. In a study conducted by Del Pino et al., it was observed that cultural differences affected the caregiver burden for nearly 29% of caregivers. In the present study, the caregivers were all family members of the Alzheimer’s patients.

The relatives of an Alzheimer’s patient can be affected to varying degrees depending on the disease’s stage of diagnosis. When Alzheimer’s disease is diagnosed at a relatively early stage, there is sufficient time to plan and determine the individuals who will take care of the patient. However, regardless of the stage of diagnosis, it is very likely for all members of the family to be affected by the difficulties associated with the care and requirements of an Alzheimer’s patient. The role of caregiver for Alzheimer’s patients is usually assumed by the patient’s spouse, or other female members of the family. Stone et al. previously determined that 23% of all caregivers are the female spouses of the patients, while 13% are the male spouses of the patients. In a multi-centered study with 1497 cases conducted by Reed et al., it was determined that the majority of caregivers were the patients’ spouses. Reed et al. also determined that the children of Alzheimer’s patients are adversely affected to a greater extent when assuming the role of caregiver than the patients’ spouses, even when they become caregivers for a relatively shorter period of time. Certain studies suggest that patients’ children are affected to a greater extent when they become caregivers, while other studies suggest that patients’ spouses are affected to a greater extent. The study of Hong et al. determined that caregiver spouses are adversely affected to a greater extent than caregiver children and daughters-in-law. In the present study, the large majority of the caregivers (89.1%) were female. In addition, 14.5% of the caregivers had a history of systemic disease. An increase in the caregiver’s age was associated with a decrease in the Physical Health domain subgroup scores of the Quality of Life questionnaire. The scores for the Physical Health domain were statistically significantly higher among the patients’ spouse than among the patients’ children and daughters-in-law (p<0.005). These findings were considered to be associated with the relatively more advanced age of the patients’ spouses compared to the other caregiver groups, which increased the caregiver spouses’ chance of displaying systemic diseases. No statistically significant difference was identified between the existing diseases of the caregivers and the WHOQoL-Bref scores (p>0.005). In addition, no statistically significant difference was identified between the Caregiver Stress Index scores and the WHOQoL-Bref scores (p>0.005). The differences between the findings of the current study and other studies might be due to the differences that exist in the cultural aspects associated with the caregiver role between different countries. In fact, such differences might even be observed between different regions within the same country.

Alzheimer’s diseases is often called a “family disease,” since caregivers within a family are potential secondary patients themselves, who might develop complications in association with their caregiver role. Despite of all difficulties they might encounter, caregivers are socially expected to devote themselves tirelessly to the care of the patient. Having to conduct and manage their occupational and family life while also being required to act as a caregiver can result in tremendous strain for many individuals. While handling all of these responsibilities at the same time, individuals might have difficulties in allocating and spending time for themselves. However, any expression by a caregiver of the problems and strains he/she experiences might be perceived by others as a rejection of the caregiver role. Caregivers may thus find themselves unable to manage the effects and strains engendered by the caregiver role. This, in turn, may lead to feelings of helplessness, guilt, anger, fear, and social isolation. In congruence with their cultural values, they tend to consider taking care of their loved ones as a duty, and might even refuse any outside help. For these reasons, the stress levels of individuals taking care of Alzheimer’s patients were very high in the current study, as well. In the present study, no statistically significant difference was identified between the duration of the disease and the Caregiver Stress Index. This finding might be explained with the adaptation shown by caregivers in the times following the initial diagnosis, and with the reduction or elimination of the patient’s behavioral symptoms as a result of treatment. These two factors might have contributed to reducing the effect of disease duration on the caregivers’ stress scores. A negative correlation was identified between the stress index score and the dimension scores for the WHOQoL-Bref questionnaire. In other words, an increase in the stress levels of the caregivers was associated with a negative effect on the quality of life. This might stem from the caregivers’ lack of knowledge on how to cope with the disease and the patient, and the caregivers’ inability or unwillingness to receive assistance.

As those taking care of Alzheimer’s patients are deeply involved with work, family and the care of the patient at the same time, they generally tend to neglect their own physical and emotional health. The mental and physical health of the caregiver affects not only the quality of life of the caregiver him/herself, but also the quality of life of the patient. Many caregivers around the world – especially those who are not receiving adequate support – begin to experience considerable difficulties and adverse effects over time. In a previous study, 60.6% of caregivers expressed that they were unable to spend time for themselves. 78.8% expressed that they constantly felt tired, 84.9% expressed that they had difficulties communicating with the patient, and 56.9% expressed that they faced financial difficulties. In a multi-centered study conducted by Schulz et al., it was determined that the caregivers of Alzheimer’s patients had very high stress levels, and that they exhibited a higher incidence of mood disorders, depression, insomnia, and lower quality of life scores. In certain studies, the progression of the disease, as well as the additional behavioral problems (neuropsychiatric symptoms) this progression engenders, gradually increases the burden of the caregiver. Neuropsychiatric symptoms are commonly observed during the course of Alzheimer’s disease. Neuropsychiatric symptoms are the symptoms that have the greatest negative impact on the quality of life both of the patient and caregiver.
In the present study, the quality of life scores of caregivers group were lower than those of the control group in all four subscales.

In conclusion, the gradual increase in the number of Alzheimer's patients, as well as the increase in the average human life span as a result of modern medicine, will lead to a parallel increase in the number of caregivers worldwide. This, in turn, will cause the medical needs of these caregivers themselves to become an important public health problem in many countries. Thus, it is necessary for physicians around the world that treat Alzheimer's disease to also investigate the mental and physical effects of the disease on caregivers, and to determine effective means and methods for supporting these caregivers.

References


Table 1. Quality of Life Scale-Brief Of Caregivers and Control Groups

<table>
<thead>
<tr>
<th>Subscales</th>
<th>Caregivers’ group Mean ± Std</th>
<th>Control group Mean ± Std</th>
<th>P value 8</th>
</tr>
</thead>
<tbody>
<tr>
<td>Physical health</td>
<td>53.81 ± 19.46</td>
<td>79.31 ± 19.99</td>
<td>0.000</td>
</tr>
<tr>
<td>Psychological health</td>
<td>56.47 ± 13.03</td>
<td>77.29 ± 22.64</td>
<td>0.000</td>
</tr>
<tr>
<td>Social relationship</td>
<td>52.72 ± 21.37</td>
<td>90.68 ± 16.35</td>
<td>0.000</td>
</tr>
<tr>
<td>Environment</td>
<td>56.60 ± 15.83</td>
<td>84.27 ± 13.42</td>
<td>0.000</td>
</tr>
</tbody>
</table>

*Mann Whitney U, p<0.005*