Assessment of the level of care-burden in informal caregivers of patients with dementia

S Abeywickrema, R Weerasundera, K Ranasinghe

Background

Worldwide, the incidence and prevalence of dementia is currently increasing. Dementia is associated with significant disability due to cognitive impairment, and sufferers often become dependent on their carers. Close family members often provide care for persons suffering from dementia, particularly in the developing world. This care-burden is associated with many consequences such as carer depression, elder abuse and premature placement of the patients in institutions. The consequences of care-burden have not been studied in depth in Sri Lanka. A better understanding of care-burden would be helpful to develop interventions to support carers as well as minimise the economic burden to the state.

Objectives

The objectives of this study were to describe the level of care-burden in carers of patients with dementia, and to explore possible associations between the care-burden and other variables such as socio-demographic factors and psychiatric morbidity.

Method

A total of 77 carers were included in the study. The locally adapted and translated Zarit Burden Interview (ZBI) was used to assess for care-burden, and participants were also clinically assessed for the presence of depression based on the Diagnostic and Statistical Manual – version IV (DSM IV) criteria for a major depressive episode. Socio-demographic details were gathered via a questionnaire.

Conclusion

Over one quarter of carers were experiencing a moderate to severe level of care-burden, and almost half reported mild to moderate care-burden. The level of burden increased significantly with increasing age, and a proportion of participants were also depressed. Further large scale, population based research is needed in Sri Lanka, to explore these associations further.

Key words: dementia, care-burden

Introduction

Dementia is a syndrome of chronic and progressive nature, in which there is disturbance of multiple higher cortical functions, usually against a background of unimpaired consciousness (1). It is estimated that 24.3 million people worldwide are affected with dementia, and 4.6 million new cases are diagnosed each year, most of whom live in the low and middle-income countries (2). Ageing is the most important non-modifiable risk factor for dementia. Approximately 5-8% of individuals over the age of 65 years, and 20% of those over 80 years, are affected by this disease (3). Dementia, due to its chronic progressive nature and its inherent features of failing cognitive ability, leads to progressive disability, warranting a considerable degree of care. Given the trend towards an ageing population, the increasing incidence of dementia and the significant associated disability, increasing dependence on care givers is inevitable.

Currently, in Sri Lanka, most persons with disability from any cause are cared for by family members, who serve as informal caregivers (4). An informal carer includes any person such as a family member, friend or a neighbour, who is giving regular, on-going assistance to another person without payment for the care given (5). They have no regular hours, often work around the clock and have significant subjective feeling so for burden and are more susceptible to depression and anxiety than the normal population (5). About twice as many women are in the caregiving role compared to men (6).

Care-burden is defined as the extent to which the caregivers feel that their emotional or physical health, social life and financial status have suffered as a result of caring for their relatives (7). A high level of care-burden is associated with depression and anxiety in the carer, reduced level of care for the patient, and premature placement of patients in long term care facilities (5). Sri Lankan studies regarding care-burden are minimal and a better understanding of care-burden in this country will help healthcare workers to identify such problems early and will also indicate ways in which to provide counselling and training for carers, in order to minimize carer depression and anxiety. Overall this will help ensure better care for the patient. The objectives of this study were to describe the level of care-burden in carers of patients with dementia, and to explore possible
assessments between the care-burden and other variables such as socio-demographic factors and psychiatric morbidity.

Method

Carers of patients with dementia attending the psychogeriatric outpatient clinic at the National Hospital of Sri Lanka, (conducted by the National Institute of Mental Health) and outpatient clinics of Colombo South Teaching Hospital Sri Lanka, were recruited for the study over a period of 3 months. Carers of patients with dementia diagnosed for less than 6 months, guardian of patients who live in long term care facilities, and those who are unable to read and comprehend Sinhalese were excluded from the study. All those who met inclusion criteria, who gave written informed consent, were included in the study. Ethical clearance for the study was obtained from the Ethical Review Board of the Sri Jayewardenepura University.

Socio demographic data was collected via a questionnaire designed for that purpose. The care-burden was assessed by the use of the self-administered Zarit Burden Interview (ZBI) (5). This is a widely used self-administered scale for measuring the subjective care-burden in carers, which has been adapted and translated into Sinhalese to be used in Sri Lanka (written communication, the 10/66 Dementia Research Group-Sri Lanka, July 2011). This instrument consists of 22 items, each of which is scored from 0-4. A total score of 0-20 indicates ‘little or no burden’, 21-40 ‘mild to moderate burden’ 41-60 ‘moderate to severe burden’ and 61-88 ‘severe burden’ respectively. A total score of 24 or above indicates a significant risk of depression in the carer.

Depression in the carer was diagnosed via clinical interview as by a psychiatrist, based on DSM IV criteria. Data regarding the presence of behavioural and psychological symptoms of dementia (BPSD) among the care-recipients was gathered from the carers through a semi-structured interview. The last recorded mini mental state examination (MMSE) scores of the respective care-recipients were obtained from clinic notes.

Data was analysed using SPSS. Chisquare and Pearson correlation were carried out to explore associations between socio-demographic variables, BPSD, MMSE, carer depression and the level of care-burden.

Results

A total of 77 carers were included in the study. Most were female (n=55, 71.4%) and the mean age of the participants was 49 years, with 11.7% being over the age of 65 years. Of the carers, 83.1% were currently living with their partners. More than half of the participants earned a monthly income below 20,000 rupees. The majority of carers were unemployed, and of those who had jobs, most were self-employed. With regard to relationship of the carer to the patient, 37.7% were daughters and 21.1% were wives.

Over one quarter of carers were experiencing a moderate-severe level of care-burden, and almost half reported mild-moderate care-burden (Table 1).

<table>
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<tr>
<th>Table 1. Levels of care-burden experienced by carers</th>
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<td>ZBI score and level of care-burden</td>
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<td>0-21: Little or no burden</td>
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<tr>
<td>22-40: Mild to moderate burden</td>
</tr>
<tr>
<td>41-60: Moderate to severe burden</td>
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<tr>
<td>61-88: Severe burden</td>
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Increasing age of the carer was significantly associated with a greater level of care-burden (p=0.02), whereas increasing household income had a significant negative correlation with the level of care-burden (p=0.004) (Table 2). Although a majority of care-recipients had features of BPSD, including wandering (71.4%), aggression (63.6%), features of paranoia (51.9%) and disinhibited behaviour (48.1%), there was no significant association between the presence of BPSD and care-burden. Likewise there was no significant association between the degree of cognitive impairment in the care-recipient, and the level of care-burden experienced. Of the carers, 3.9% (n=3) fulfilled the criteria for a major depressive episode. Surprisingly, there was no significant association between the level of care-burden and presence of depression in the carer (p=0.31).

<table>
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<tr>
<th>Table 2. Associations between socio-demographic and patient factors and level of care-burden</th>
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<td>Carer and care-recipient characteristics</td>
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<td>Socio-demographic characteristics of carer:</td>
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<td>Lack of support from other sources</td>
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Discussion

The findings of this study show that 26% of carers were experiencing a moderate to severe level of care-burden. This is similar to findings reported from India, where care-burden has been reported to be 30% (8). It could be queried whether the prevalence of subjective care-burden would be lower in Eastern cultures, compared to the West – it may be argued that looking after ageing relatives might be appraised as less burdensome given the collectivistic nature of these societies. However studies done to explore this hypothesis so far have not confirmed this hypothesis (9).

In this study there was no significant difference between the care-burden reported by female and male carers; this is contrary to the findings in most international studies, which report a higher degree of burden among females (10,11). Females are traditionally cast in the care-providing role in South Asia; therefore female carers may have been reluctant to state the true degree of burden, as ‘looking after the elderly parent’ is a culturally sensitive issue.

According to the results of this study, most carers (61%) were over the age of 45 years, with 11.7% of the carers being over the age of 65 years – a major concern since the carers themselves are susceptible to a host of medical and psychological morbidities. It has been estimated that about half of informal carers are themselves elderly (6). In this study, the age of the carer was positively correlated with the care-burden, suggesting an increased subjective burden with advancing age. The available literature with regard to the age of the carer and the degree of burden felt is inconclusive. A recent study by the 10/66 Dementia group, of 673 participants carried out in multiple sites, report no relationship between carer strain and carer age (12). However other research has identified carer age as a predictor on dimensions such as ‘guilt’ in carer strain (13).

Our findings indicate that increasing monthly house hold income has a significant negative correlation with the care-burden. This is similar to the findings of other international research – expressed financial distress has been shown to increase the care-burden (14,15). It is likely that the availability of the support of paid carers and other informal support leads to a reduction of burden felt by the principal carer. The inability to arrange such support has been identified as a predictor of depression in the primary carer (16). This holds true even in countries where traditional family care is highly valued such as Japan (17). In this study, 71.4% received external help in providing care for their ageing relative, while 27.3% did not have access to any such support. As expected, analysis revealed significant association between the availability of external support and the degree of burden felt.

There was no significant association between the employment status and the care-burden in this study. Existing literature reveals mixed results, which could be due to the fact that being employed may have both positive and negative effects on the care-burden. Some researchers argue that these positive effects may outweigh the negative, and that being employed may reduce the overall level of burden (18). Other studies, however, report that employment does not have an effect on the overall level of burden (19, 20). Similarly, marital status did not impact significantly on the caregiver burden. This is supported by international findings, which indicates that being supported by a partner does not appear to reduce the burden in carers (12).

The last recorded MMSE score was taken as a measure of cognitive impairment of the care-recipients in this study. There was no significant association between the degree of cognitive impairment and the perceived care-burden. Most studies report that while the degree of cognitive impairment does have an effect on the burden, it is the presence of behavioural and psychological symptoms of dementia that significantly worsens care-burden experienced by the carer (21, 22). However, in contrast to the reported work, in this study the presence of BPSD did not establish a significant association with the degree of burden. The number of carers experiencing no BPSD in their patient may have been too small to give a significant difference in the analysis.

Rates of depression among carers of the demented elderly is reported to be higher than among non-caregivers (23). The prevalence of depression among care givers is estimated to be 41%-55%, which contrasts sharply with the prevalence of 8% in non-caregivers (23). In our study 3.9% (n=3) carers fulfilled the criteria for a major depressive episode at the time of assessment. While all depressed carers reported a high degree of burden, statistical analysis did not reveal a significant association with the presence of depression and the level of care-burden. The limited number of depressed carers in this study may have influenced this finding.

Limitations

While the overall study sample size was adequate to describe the proportion of burdened carers, numbers in each sub-category was low. This is a limitation and may have influenced analysis of associations, as mentioned above. The data for this study was collected using the convenience sampling technique – this may have led to an over representation of care recipients with severe dementia and stressed carers when compared to population based sampling. And since the sample was obtained from two state hospitals situated in urban settings, the findings may not be generalizable to the entire population. The MMSE score was derived from records of the care-recipients, and therefore may not have represented the care recipients’ current level of cognitive impairment. However since this study concerned the carers, and since the care-recipients were often unable
or unwilling to attend the clinic, there were practical difficulties in obtaining the current MMSE. All efforts were made to minimize this error by taking the most recently (‘last recorded’) stated MMSE score from the records. Lastly, the assessment of BPSD was based on the carers’ subjective statement, and may have been influenced by recall bias.

Conclusions

It is evident from this study that carers of patients with dementia in this country experience a significant level of care-burden, and this is significantly greater with increasing age of the carer. A proportion of carers were also depressed. The 10/66 Dementia research group states that although two-thirds of those with dementia live in the developing world, only one-tenth of research on dementia is conducted in these countries (24). In keeping with this, further research is needed in Sri Lanka, in order to identify the true magnitude of the problem, and to develop services and supports for carers of patients with dementia. Particular emphasis should be placed on large population based studies to analyse the care-burden in dementia and its determinants, in this country.

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Declaration of interest

None declared

References


