

Caregiver burden and its prevalence, measurement scales, predictive factors and impact: a review with an Asian perspective

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Abstract

Caregiver burden is a well-recognised global phenomenon. The primary aim of this review is to summarise the prevalence of caregiver burden and its measurement scales, predictive factors and impact in Singapore. PubMed® and Scopus® databases were searched using the key terms ‘caregiver’, ‘burden’, ‘stress’, ‘strain’ and ‘Singapore’. A total of 206 papers were retrieved and 20 were included. This review showed that a significant proportion of caregivers in Singapore experience caregiver burden. Caregiving experiences and outcomes are affected by cultural, behavioural and socioeconomic factors. Being a Malay caregiver appears to be a protective factor for caregiver burden, whereas having negative coping strategies and care recipients with functional, cognitive and behavioural impairments are positive risk factors. As for outcomes, caregiver burden is associated with poorer self-rated health and reduced quality of life. Multiple instruments have been used to measure caregiver burden, and the Zarit Burden Interview is the most widely used.

Keywords: Caregiver burden, caregiver stress, quality of life, Singapore

INTRODUCTION

Caregivers play an important role as part of the healthcare team that co-manages patients with physicians, and informal caregiving accounts for much of home care.^[1] For adults in America, the economic value of informal caregiving has greatly exceeded that of paid services.^[2] The domains of caregiving are multidimensional, diverse and far-reaching, as caregivers often provide not only physical (e.g. basic and instrumental activities of daily living) and medical (e.g. nursing, medication administration and titration) care, but also psychological (e.g. emotional support), financial (e.g. paying for medical bills) and social (e.g. arranging or accompanying for appointments) assistance to patients under their care. With these demands, caregivers are often subject to enormous stressors and are potentially at risk for caregiver burden and declining physical and mental health.^[3]

Zarit and colleagues defined caregiver burden as “*the extent to which caregivers perceive that caregiving has had an adverse effect on their emotional, social, financial, physical*

and spiritual functioning.”^[4] Caregiving burden has negative impact on the well-being of caregivers, including increased mortality,^[5] poorer self-care,^[6] social isolation,^[7] depression,^[8] anxiety^[9] and financial stress.^[10] A recent review suggested that caregiver burden can be the most compelling problem affecting caregivers of chronically ill elderly patients.^[11] The burden of caregiving also has negative impact on the well-being of care recipients, such as abandonment and institutionalisation of the care recipient.^[12-14] Previous research has characterised the various risk factors associated with caregiver burden, including female gender,^[15] low education,^[16] co-residence with care recipient, depression,^[8] social isolation,^[7] financial

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stress,^[10] higher caregiving hours, lack of choice in being a caregiver,^[17] smaller repertoire of coping strategies^[18] and perceived patient distress.^[11]

Many instruments for the measurement of caregiver well-being are available, and they cover various domains recommended for inclusion by the Family Caregiver Alliance.^[19-21] A recent systematic review of tools for measurement of the impact of informal caregiving categorised them according to their degree of integration of dimensions and level of specificity to the caregiving process [Figure 1],^[20] and caregiver burden was found to be the most specific and most integrated. Zarit Burden Interview (ZBI), the most widely employed tool to measure caregiver burden, has good test-retest reliability, internal consistency and construct validity.^[20-22] ZBI is a self-reported caregiver measure that was originally developed by Zarit *et al.* as a 29-item questionnaire to assess the subjective burden experienced by informal caregivers in various aspects (physical, emotional, social and financial burden, and relationship with care recipient); it was subsequently revised to a shorter 22-item version.^[21,23]

Singapore's population is rapidly ageing, with an increasing proportion of elderly residents (12.4% in 2016 and expected to double in number by 2030) and a decreasing support ratio (5.4 residents aged 20–64 years per resident aged ≥65 years in 2016 vs. 13.5 in 1970).^[24,25] With such changing demographics, informal home-based caregiving is expected to be increasingly demanding, and this may further exacerbate the issue of caregiver burdens. Thus, it is imperative to understand the prevalence, predictive factors and impact of caregiver burdens in Singapore.

Singapore is a unique developed city state in Southeast Asia with a multi-ethnic and multicultural population consisting

of 74.3% Chinese, 13.4% Malay, 9.1% Indian and 3.2% other ethnicities as of 2016.^[24] Current evidence suggests that caregiving experiences and outcomes are affected by culture, with different levels of burden reported among Asian caregivers and their Western counterparts.^[26-28] In many Asian societies, including Singapore, traditional Confucian principles such as filial piety may have an impact on the caregiving experience.^[29,30] The validity of instruments measuring caregiver burden developed in the West has also been questioned when they are applied to a different population and cultural setting.^[31] Therefore, it is of great interest to review studies on caregiver burden conducted in a multi-ethnic, multicultural population in Singapore, and to compare these findings with those from international literature. This will enable recognition of the magnitude of problem within the local context and highlight the presence of critical knowledge gaps that may have health policy implications.

The primary aim of this review paper is to summarise research papers on caregiver burden in Singapore, in terms of its prevalence, measurement scales, predictive factors, and its impact on both caregiver and care recipient outcomes.

METHODS

Search strategies and inclusion criteria

We carried out a systematic literature search using PubMed® and Scopus® up to 30 January 2018. In our search strategy, we included these key terms: caregiver, burden, stress, strain and Singapore. Specifically, the search terms for PubMed were: (((caregiver[MeSH Terms] OR caregiver*[Title/Abstract]) AND (stress*[Title/Abstract] OR burden*[Title/Abstract] OR strain[MeSH Terms] OR strain*[Title/

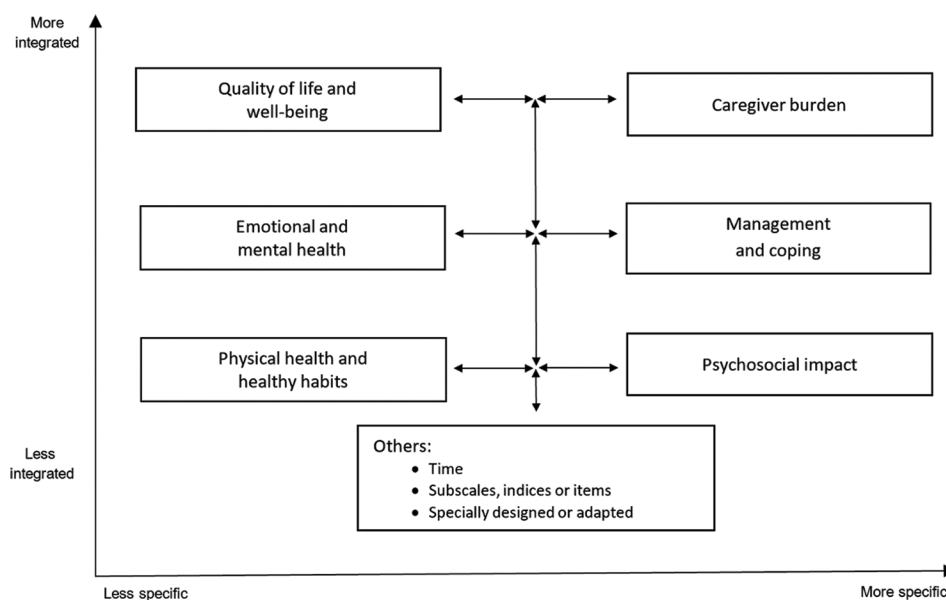


Figure 1: Diagram shows the measurement of impact of informal caregiving according to their degree of integration of dimensions and level of specificity to the caregiving process.

Abstract])))) AND English[Language]) AND Singapore. The inclusion criteria were full text original studies published in English on the topic of caregiver burden in Singapore. Meta-analyses, case series, case reports, commentaries and reviews were excluded. Studies that were not in English, not conducted on human subjects, or entirely qualitative were also excluded.

Selection of studies and data collection

After duplicate articles were removed using EndNote™ (Clarivate Analytics, Philadelphia, PA, USA), two independent researchers (i.e. first and second authors, YXL and SY) screened the titles and abstracts of the retrieved studies and engaged in discussion whenever discrepancies arose. The full text studies were then independently read and assessed for eligibility by the two researchers. In the case of disagreement during the evaluation process, any discord was resolved by discussion with a third researcher (i.e. lead author, LLL). Once an article was deemed to be eligible, data from the eligible article was extracted independently by YXL and SY using a standardised information extraction format that included the authors, year of study, populations studied (including population size and characteristics), instrument for caregiver measures, prevalence, predictive factors and impact of caregiver burden. Risk of bias assessment (e.g. selection, attrition and detection bias) was performed by LLL by means of one of the SIGN checklists that is appropriate for the study design (<https://www.sign.ac.uk/checklists-and-notes.html>).

RESULTS

In total, 206 articles were retrieved from PubMed® and Scopus®. After removing 92 duplicates and 94 articles that did not fulfil the inclusion criteria, the final number of articles included for full text review was 20 [Figure 2]. The percentage agreement between the first and second authors

was 90%. Table 1 summarises the key findings of the included articles.^[32-51]

Studies on caregiver burden in Singapore focused mostly on caregivers of the elderly^[36,38,42,43,48,50] and persons with dementia.^[40,41,46,51] Other studied populations included caregivers of cancer patients,^[35,36] psychiatric patients,^[37,44] people with chronic illness,^[45] discharged hospital patients with long-term care referrals,^[34] hip fracture patients^[47] and patients on peritoneal dialysis.^[39]

Of the 20 studies, nine presented data on the prevalence of caregiver burden. The prevalence reported ranged from 23.0% to 59.2% in studies conducted in different study populations. For example, among caregivers of community-dwelling elderly, one study found that 24.5% of caregivers fulfilled the criterion for caregiver burden,^[38] while another study found 39.0% with mild-moderate burden and 7.3% with moderate-severe burden among caregivers of elderly with dependence in their activities of daily living.^[50] In patients with cancer, two studies found significant burden in 23% and 50% of caregivers of patients with cancer attending an outpatient oncology clinic and patients with advanced cancer, respectively.^[35,36] A study based on national household survey data found that 30.9% of relatives of persons with chronic physical or mental illness were perceived to have high burden.^[45]

In Singapore, the most widely used instrument to measure caregiver burden is ZBI, and it was used in seven out of the 20 studies.^[32,33,35,36,39,46,50] Of these seven studies, three were conducted in caregivers of persons with dementia. The ZBI was also used for caregivers of cancer patients, elderly with care needs and patients on peritoneal dialysis. Most studies used the 22-item version of the ZBI.

Multiple studies reported the association between poorer functional status of care recipients^[36,48] or higher care demands^[35,38,51] and increased caregiver burden. In patients with cognitive impairment, there was increased caregiver burden in those with behavioural and psychological symptoms of dementia (BPSD) or behavioural issues.^[38,41,46,51] The presence of dementia and dementia severity were also identified as risk factors for caregiver burden.^[38,46]

Sociodemographic variables of caregivers showed mixed results in terms of their effects on caregiver burden. One study showed that being middle-aged was associated with worse outcomes compared to being younger than 35 years.^[37] Another study reported that caregivers who were younger than 75 years performed better compared to older caregivers.^[42] However, less burden was seen in older caregivers in another study.^[40] In some studies, caregiver age was not found to be a significant predictor of caregiver burden.^[38,46,48,51] Four studies did not find a significant association between caregiver burden and the gender of the caregiver.^[21,38,46,48] Married caregivers and caregivers without children were correlated

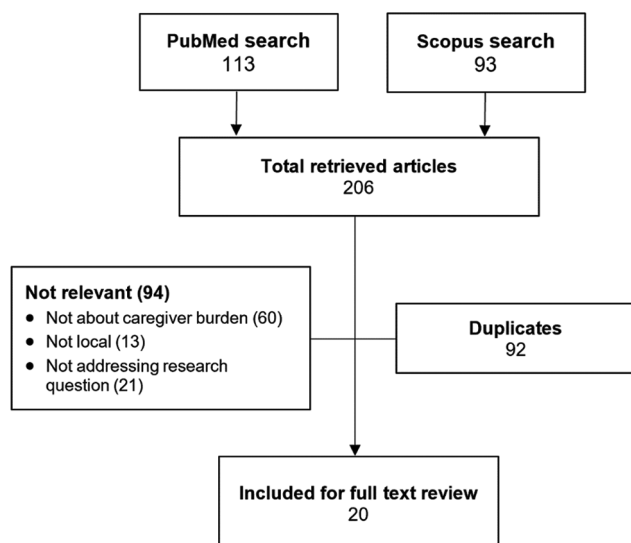


Figure 2: Flow chart shows the retrieval and inclusion of articles.

Table 1. Summary of key findings of the included studies.

Study, year	Study aim	Population	Caregiver measure	Study size	Findings	Remarks
Lim <i>et al.</i> , ^[32] 2018	Evaluate and compare the level of caregiver burden in young-onset dementia (YOD) and late-onset dementia (LOD)	Caregivers of patients with YOD and those with LOD	ZBI (12-item)	183	Prevalence: 52.6% of caregivers of patients with YOD and 32.5% of caregivers of patients with LOD reported a high caregiver burden Predictive factors: 1. Family history of dementia 2. Behavioural symptoms, including disinhibited behavior, delusions and apathy Impact: Nil	Study sample is relatively small
Win <i>et al.</i> , ^[33] 2017	Explore the demographic characteristics and burden pattern among caregivers of dementia in the oldest-old	Caregivers of dementia	ZBI	458	Prevalence: Nil Predictive factors: Caregivers of dementia in the oldest-old associated with higher ZBI scores Impact: Nil	Study sample of oldest-old care recipients is relatively small
Eom <i>et al.</i> , ^[34] 2017	Understand effects of different patient-caregiver relationships and LTC service use on well-being of informal caregivers	Informal caregivers of patients discharged from tertiary hospital with referrals to LTC services	AMA Caregiver Self-assessment Questionnaire (stress); EuroQoL-VAS (self-rated general health)	781	Prevalence: Nil Predictive factors: 1. Spousal and adult children caregivers showed worse self-rated general health and stress compared to non-immediate family caregivers, but difference did not reach statistical significance 2. Family caregivers who utilised home-based services presented poorer self-rated general health than those who did not use any services Impact: Nil	No data on caregiver well-being before patient use of LTC services
Chua <i>et al.</i> , ^[35] 2016	Understand the impact of caregiver burden on QoL, mental health and work capacity	Informal caregivers of advanced cancer patients	ZBI (12-item)	16	Prevalence: 50% of caregivers experienced high burden Predictive factors: Balancing multiple responsibilities, significant physical and non-physical caregiving demands, patients with debilitating symptoms, and poor social support Impact: Higher burden is associated with lower QoL, higher depression scores and greater work impairment	<ul style="list-style-type: none"> • Small sample size (exploratory study) • Non-random sampling; only English-speaking and available caregivers of inpatients were recruited • Breakdown of frequency of predictors not provided
Rajasekaran <i>et al.</i> , ^[36] 2016	Predict increased caregiver burden among elderly cancer patients using CGA-based risk factors	Family caregivers of elderly patients with cancer attending an outpatient oncology clinic	ZBI	244	Prevalence: 23% of caregivers had mild to severe burden. Predictive factors: Significant variables after multivariate analysis • ECOG 3-4, OR 4.47 (2.27-8.8) • Hb < 12 g/dL, OR 2.38 (1.14-4.99) Impact: Nil	<ul style="list-style-type: none"> • All outpatients (potential selection bias for patients who are less frail) • Large number of covariates analysed
Chang <i>et al.</i> , ^[37] 2016	Identify the sociodemographic variables affecting the caregiving experience to relatives with mental illness, and the association between caregiving and psychological distress	Informal caregivers of outpatients attending a tertiary hospital psychiatric clinic	GRA	344	Prevalence: Nil Predictive factors: Significant variables after multivariate analysis • Age: those aged 35-49 yr had greater impact on health and schedule vs. those aged < 35 yr • Education: those with lower education had greater impact on finances vs. those with university degree • Employment: the unemployed had greater impact on finances	<ul style="list-style-type: none"> • Certain patient characteristics that could impact caregiving and distress (e.g. age, neuropsychiatric symptoms) were not explored • Analysis by patient diagnosis was not conducted due to inadequate sample size in some groups

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Study, year	Study aim	Population	Caregiver measure	Study size	Findings	Remarks
Vaingankar <i>et al.</i> , ^[38] 2016	Describe the factors related to care burden of older adults with care needs, and their association with dementia	Informal caregivers of older adults with care needs	ZBI	693	<ul style="list-style-type: none"> Income: those earning <\$2,000/mth perceived greater lack of family support Ethnicity: the Chinese had lower self-esteem vs. Malays and Indians Impact: Higher domain scores of CRA were significantly associated with psychological distress even after controlling for sociodemographic factors 	<ul style="list-style-type: none"> Population-based sampling using data from a national survey Broad definition of informal caregivers, who may not have been the primary caregivers. Did not evaluate role of paid helpers who took care of a sizeable proportion of participants
Griva <i>et al.</i> , ^[39] 2016	Compare the psychosocial outcomes in patients undergoing assisted vs. self-care peritoneal dialysis	Caregivers of patients undergoing peritoneal dialysis	ZBI	111	<p>Prevalence: Nil</p> <p>Predictive factors: Levels of perceived burden were equal and in the mild to moderate range.</p> <p>Impact: Nil</p>	
Tay <i>et al.</i> , ^[40] 2016	Examine the associations of burden, coping and self-efficacy with QoL among family caregivers of PWD	Family caregivers of persons with mild-moderate dementia attending a tertiary hospital outpatient clinic	FBIS	84	<p>Prevalence: Nil</p> <p>Predictive factors: Older caregivers had reduced burden scores but not better QoL</p> <p>Impact: Burden has significant negative correlation with QoL, particularly in the physical health and environment domains</p>	<ul style="list-style-type: none"> Convenience sampling performed. Majority of caregivers was Chinese and may not be representative of other ethnicities. Potential variables such as dementia stage, presence of BPSD, use of community services were not considered.
Khoo <i>et al.</i> , ^[41] 2013	Determine the impact of neuropsychiatric symptoms (NPS) on caregiver distress in PWD	Family caregivers of PWD attending a tertiary hospital memory clinic	NPI-Q	667	<p>Prevalence: Nil</p> <p>Predictive factors: NPS of various types significantly predicted caregiver distress, after controlling for type and severity of dementia, and patient's age, race, gender, education and marital status. The top 3 NPS that predicted the most distress were: agitation, depression and irritability.</p> <p>Apathy caused the least distress.</p> <p>Impact: Nil</p>	<ul style="list-style-type: none"> All were outpatients. Potential variables, such as caregiver characteristics, were not controlled for.

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Study, year	Study aim	Population	Caregiver measure	Study size	Findings	Remarks
Chan <i>et al.</i> , ⁽⁴²⁾ 2013	Estimate the health impact of caregiving to older adults requiring assistance	Informal caregivers of older adults requiring assistance in at least 1 ADL	SRH	1,077	Prevalence: Nil Predictive factors: 1. Odds for poorer SRH lower among: • Caregivers aged <75 yr • Malay 2. Odds for poorer SRH higher among: • Low educational level • Residence in 1-2 room public housing • Increased number of chronic diseases Impact: Caregivers, compared to non-caregivers, had higher odds of: • Poorer SRH, OR 2.45 (1.84-3.26) • Clinically significant depressive symptoms, OR 2.36 (1.44-3.86) • Outpatient visits, OR 1.27 (1.01-1.59)	<ul style="list-style-type: none"> • Large number of covariates included in analysis • Large sample size from a national survey representing the general population and not only those attending healthcare or social services
Østbye <i>et al.</i> , ⁽⁴³⁾ 2013	Assess the impact of support from FDW in moderating caregiving outcomes in informal caregivers of older persons	Informal caregivers of older persons requiring assistance in at least 1 ADL	Modified CRA	1,190	Prevalence: Nil Predictive factors: FDW support is associated with better caregiving outcomes, by moderating the association of: • Physical impairment with caregiver's disturbed schedule and poor health • Memory impairment with caregiver's disturbed schedule, poor health and lack of finances • Behavioural impairment with caregiver's lack of family support Impact: Nil	<ul style="list-style-type: none"> • Large sample size from a national survey of community-dwelling elderly • Considered many sociodemographic variables of caregiver and recipient as covariates
Tan <i>et al.</i> , ⁽⁴⁴⁾ 2012	Examine the level of burden experienced by caregivers of persons with schizophrenia in the community	Caregivers of persons with schizophrenia attending outpatient mental health clinic	BAS	150	Prevalence: 31.3% of caregivers felt distressed, 24.7% experienced increased social burden and 26.7% had disturbed routines. Predictive factors: Nil Impact: Nil	<ul style="list-style-type: none"> • All were outpatients • Convenience sampling
Vaingankar <i>et al.</i> , ⁽⁴⁵⁾ 2012	Examine the predictors of perceived burden among relatives of people with chronic physical or mental illness	Close relative of person with any chronic physical or mental illness	Family Burden Module	2,458	Prevalence: 30.9% perceived high burden resulting from their relative's health condition. Predictive factors: 1. Multivariate analysis showed the following were more likely to perceive burden: • Women, OR 1.58 (1.17-2.14) • People who could open up to family or friends, OR 1.65 (1.1-2.48) • People with dysthymia, OR 4.91 (1.11-21.77) 2. Malays were less likely to perceive burden, OR 0.68 (0.52-0.89) Impact: Nil	<ul style="list-style-type: none"> • Data collected from a national household survey with good response rate (75.9%) • Broad definition of 'family caregiver' used in study • Larger number of predictors analysed, but significant predictors only explained 10% of variance; authors attributed this to lack of information on other important factors

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Table 1. Contd...

Study, year	Study aim	Population	Caregiver measure	Study size	Findings	Remarks
Lim <i>et al.</i> , ⁽⁴⁶⁾ 2011	Examine the factors associated with caregiving outcomes among caregivers of PWD	Family caregivers of PWD attending dementia clinic or recruited through email	ZBI	107	Prevalence: Nil Predictive factors: 1. Significant predictors of burden were: • Behavioural problems • Dementia severity • Use of behavioural disengagement and criticism as coping strategies 2. Use of behavioural disengagement and criticism consistently predicted burden above and beyond behavioural problems, even after statistically adjusting for caregiver characteristics 3. Sociodemographic characteristics of caregivers (age, sex, race, relationship with recipient, marital status, employment, education, caregiving duration, co-residency, help from FDW) had no associations with burden 4. High mean score of burden (34.2) associated with caregiving for PWD. Impact: Nil	<ul style="list-style-type: none"> Multiple comparisons were done without adjustment of significance levels (attributed to exploratory nature of study to capitalise on chance findings and avoid overlooking any significant effect) Predictors of burden accounted for 48% of the variance in this study
Siddiqui <i>et al.</i> , ⁽⁴⁷⁾ 2010	Evaluate the prevalence of stress in caregivers of elders with osteoporotic hip fractures, and determine the contributory factors	Primary caregivers of patients admitted for osteoporotic hip fracture	CSI	76	Prevalence: Proportion of caregivers who were significantly stressed: • 1 wk from admission: 59.2% • 6 mth from admission: 50% Predictive factors: 1. Financial strain was the only significant cause after multivariate analysis, OR 5.04 (1.46-17.38) 2. Mental status and functional status of care recipients were not predictors of caregiver stress. Impact: Nil	<ul style="list-style-type: none"> Majority of patients were Chinese (86.8%) and had high Charlson Comorbidity Index score (63.2%) 31.5% of the caregivers had maids
Mehta <i>et al.</i> , ⁽⁴⁸⁾ 2005	Examine the factors affecting stress among family caregivers of older persons	Principal family caregivers of homebound elderly patients	CSS (modified from CSI, locally validated)	61	Prevalence: Nil Predictive factors: 1. Increased caregiver stress was associated with ADL and instrumental ADL dependency of care recipient, and performance of physical tasks 2. Female caregivers had higher levels of stress, but difference was not statistically significant 3. Age of caregiver had an extremely low correlation with stress Impact: Nil	<ul style="list-style-type: none"> Purposive, non-random sampling All patients were Chinese with low income and educational level, as data came from a voluntary welfare agency (Hua Mei Mobile Clinic)
Tan <i>et al.</i> , ⁽⁴⁹⁾ 2005	Assess the impact of neuropsychiatric symptoms on distress in family and professional caregivers	Caregivers of patients with dementia	NPI-D and NPI-NH	85	Prevalence: Nil Predictive factors: 1. Neuropsychiatric symptoms were positively correlated with caregiver distress 2. Family caregivers were significantly more distressed than professional caregivers over the delusion, agitation, depression and aberrant motor domains Impact: Nil	This study assessed both family and professional caregivers

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Table 1. Contd...

Study, year	Study aim	Population	Caregiver measure	Study size	Findings	Remarks
Siew <i>et al.</i> , ^[50] 2003	Examine caregiver stress associated with caring for ADL-dependent elderly in the community	Caregivers of community dwelling ADL-dependent elderly	ZBI	41	Prevalence: 53.7% reported little or no burden, 39% reported mild to moderate burden, 7.3% reported moderate to severe burden, none reported severe burden. Predictive factors: Nil Impact: Nil	Elderly in the study were not severely ADL-dependent
Heok <i>et al.</i> , ^[51] 1997	Investigate the factors associated with emotional well-being of caregivers of PWD	Family caregivers of elderly Chinese patients with dementia presenting to memory clinic	GHQ-28	50	Predictive factors: 1. GHQ scores correlated significantly with duration of care, depression and total behavioural score on multiple regression analysis 2. No significant association with gender, age and employment of caregivers; type and severity of dementia Impact: Nil	Potential selection bias in using hospital patients

ADL: activities of daily living, AMA: American Medical Association, BAS: Burden Assessment Scale, BPSD: behavioural and psychological symptoms of dementia, CGA: Comprehensive Geriatric Assessment, CQOLC: Caregiver Quality of Life Index-Cancer, CRA: Caregiver Reaction Assessment, CSI: Caregiver Strain Index, CSS: Caregiver Stress Scale, ECOG: Eastern Cooperative Oncology Group, EuroQol-VAS: EuroQol-Visual Analog Scale, FBIS: Family Burden Interview Schedule, FDW: foreign domestic worker, GHQ: General Health Questionnaire, Hb: haemoglobin, LTC: long-term care, NPI-D: Neuropsychiatric Inventory Caregiver Distress Scale, NPI-NH: Neuropsychiatric Inventory Caregiver Nursing Home Version, NPI-Q: Neuropsychiatric Inventory Questionnaire, OR: odds ratio, PWD: person with dementia, QoL: quality of life, SRH: self-rated health, ZBI: Zarit Burden Interview

with increased burden in one study,^[38] but this finding was not replicated in another study.^[46] Some studies found an effect of ethnicity: Malay caregivers showed less burden and negative impact,^[42,45] while Chinese caregivers had greater impairment on self-esteem compared to their Malay and Indian counterparts.^[37] Not all studies, however, reported ethnicity as a significant factor.^[38,46]

Socioeconomic factors also played a role. Caregivers who were unemployed or had lower income,^[37] stayed in one- or two-room public housing,^[42] or experienced financial strain^[47] had worse outcomes. However, employment was not found to be a significant predictive factor in some studies.^[46,51] Having the support of a foreign domestic worker was found to moderate the impact of caregiving for elderly requiring assistance in at least one activity of daily living and was associated with better caregiving outcomes.^[43] This mitigating effect of external help was, however, not seen in all studies.^[38,46] The coping strategies used by caregivers of persons with dementia also played a role; for instance, the use of behavioural disengagement or criticism as a coping strategy was linked to increased burden.^[46]

Four of the included studies measured the associated outcomes of caregiver burden. Increased caregiver burden was associated with reduced quality of life and increased depressive symptoms.^[35,40,42] It also disrupted caregivers' work and schedules.^[35,37] Two of these studies found that caregivers were more likely to have poorer self-rated health and more outpatient visits.^[37,42]

DISCUSSION

Population studied and prevalence of caregiver burden

Studies on caregiver burden in Singapore have largely concentrated on those caring for the elderly or demented. Caregivers of patients with other medical conditions or demographic characteristics (e.g. paediatric patients with genetic disorders) are under-represented and would, therefore, be prime candidates for further research endeavours. This will likely give a more complete picture of caregiver burden in Singapore.

The prevalence of caregiver burden in the included studies was reported to be 23.0%–59.2%. This is similar to the prevalence reported by other studies outside Singapore that used similar measurement scales.^[52-55] With Singapore's ageing population and more elderly developing multiple and complex health conditions, the prevalence would be expected to increase even further.^[25] Health policymakers may be better informed to allocate appropriate resources to develop integrated and targeted health intervention programmes in the community to address this pressing issue.

Measures of caregiver burden

ZBI is the most widely used among the included studies and has been validated in Singapore.^[56] The use of the ZBI as a

measure has certain advantages. Firstly, it has been extensively studied and validated in a wide variety of settings, with well-documented psychometric properties. Secondly, its wide use allows healthcare providers to standardise reporting of caregiver burden and easily compare results obtained across different studies. However, its use as a unidimensional measure to produce a summary score may overlook the uniqueness and rich data from the individual dimensions.^[31]

One observation is that studies in Singapore have uniformly adopted the use of quantitative measures to evaluate caregiver burden. The advantages of quantitative measures are their value for statistical analysis, as well as the ease of adaptation for use in clinical settings due to their directness and practicality. However, it has been argued that caregivers may conceptualise burden in ways that differ from descriptions used in measures, leading to a potential failure of the instruments to capture the entire breadth of burden.^[31] Future mixed method approaches may enhance the comprehensiveness of research on this topic.

Predictive factors

The predictive ability of sociodemographic factors has been mixed in this review, which is consistent with other international studies which found that caregiver outcomes may not directly correlate with caregiver sociodemographic factors.^[23,57-59] For example, female caregivers were shown to be at higher risk in some studies,^[60,61] but not in others.^[62,63] Some studies showed Malay caregivers experienced less burden and negative impact,^[42,45] whereas other studies reported ethnicity as not a significant factor.^[38,46] The real-life difficulty in predicting caregiver burden based on sociodemographic factors was also reflected in one study done on a broad population, which used national household survey data with a large sample size and good response rate but only found a limited number of predictors.^[45] Furthermore, these predictors only explained 10% of the variance observed, which the authors attributed to a lack of information on other significant factors related to burden. This suggests that there are likely to be hidden predictors and confounders that are beyond the usual variables captured, which warrants more explorative qualitative studies. Further studies may also explore the complex interplay of the role of culture, social values and sociodemographic factors. Coping strategies appeared to influence the perceived burden in Singapore caregivers.^[46] This is consistent with other studies which reported that strategies focused on problem-solving may be more effective in reducing caregiver burden than strategies that are focused on emotions.^[64,65] This is of potential value, as caregiver coping strategy is a modifiable variable that can be pre-empted and targeted in tailored health intervention programmes for them. A future research direction may be an interventional trial to assess the effectiveness of coping strategy programmes to reduce caregiver burden.

Pertaining to care recipient factors, caregiver burden increased with caregiving demands, physical dependency and duration of

time spent on care.^[35,36,38,48] Similar findings about the impact of physical dependency have also been shown in other studies.^[66] In addition, behavioural and neuropsychiatric symptoms, as well as the presence and severity of dementia were found to increase caregiver burden.^[38,41,46,51] This is also consistent with findings in the existing literature that behavioural disturbances are often the most prominent predictive factor of burden.^[67,68]

Impact of caregiver burden

The included studies evaluated the negative impact of caregiver burden on caregivers. This is important, as caregivers can be forgotten in the doctor-patient relationship.^[69-71] Engaging the caregivers and addressing their needs are crucial, as caregivers play an essential role in keeping patients well in the community.^[64,68,72] Further research efforts may also focus on how other caregiving constructs, such as caregiver grief, can have a negative impact on adverse caregiver outcomes.^[73,74] On the other hand, it is also imperative to evaluate the effect of caregiver burden on care recipients (patients); however, none of the included studies had done so. Given the measured prevalence of caregiver burden in Singapore, it would be of great concern if caregiver burden translates into worse clinical outcomes for patients and increased healthcare costs. Future research efforts may focus on whether caregiver burden is associated with worse patient outcomes such as mortality, quality of life and healthcare expenditure.

Strengths and limitations

This article is an attempt to review the existing medical literature on caregiver burden in Singapore – a developed, multiracial and multicultural city-state. Our findings highlight important challenges in studying this highly subjective issue with cultural sensitivity. This article further builds on the current literature on caregiver burden by providing unique experiences in a multicultural setting in Singapore, which can inform future studies that intend to examine caregiver burden in other settings with non-homogeneous cultural backgrounds. One limitation of this review is that the focus of studies done in Singapore may limit its generalisability. Secondly, the risk of bias was assessed by one author. The search strategy may be further optimised by including more comprehensive search terms (such as ‘carer’) and other databases such as PsychInfo and CINAHL. Furthermore, this review did not have a registered protocol prior to the commencement of the study, which is another of its limitation. Finally, the broad topic of this review may limit its applicability to specific patient populations.

CONCLUSION

There is a significant prevalence of caregiver burden in Singapore. The ZBI is the most commonly used tool to measure caregiver burden in Singapore. In particular, caregivers of people with functional, cognitive and behavioural impairments appear to be at increased risk of burden. In our local context,

caregiver burden is associated with various adverse caregiver outcomes, such as reduced quality of life and increased depressive symptoms. Given the limited number of studies in Singapore, future research efforts may include: studies that focus on under-represented populations beyond the elderly with dementia; mixed method studies to uncover more predictive factors for caregiver burden, especially cultural and sociodemographic factors in the Malay population; and studies that examine the impact of caregiver burden on care recipients.

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Conflicts of interest

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