Factors of Resilience in Informal Caregivers of People with Dementia from Integrative International Data Analysis

Karlijn J. Joling  Gill Windle  Rose-Marie Dröes  Franka Meiland  Hein P.J. van Hout  Janet MacNeil Vroomen  Peter M. van de Ven  Esmé Moniz-Cook  Bob Woods

Bangor University, Bangor, UK; VU University Medical Centre, Amsterdam, The Netherlands

Key Words
Informal caregiving · Resilience · Dementia · Adversity · Data harmonization

Abstract

Background/Aims: Although caring for a person with dementia can be stressful, some caregivers appear to experience few negative consequences to their well-being. This study aimed to examine what proportion of caregivers demonstrates resilience under different challenging circumstances and to identify factors related to their resilience. Methods: Baseline data from 4 studies from the Netherlands and UK among informal caregivers of people with dementia were harmonized and integrated. Caregiver resilience was defined as high levels of psychological well-being despite different types of high caregiving demands. Multivariate regression analyses identified factors significantly related to caregiver resilience. Results: The integrated data set included 15 harmonized variables with data from 1,048 caregivers facing a high care demand. The prevalence of resilience varied between 35 and 43%, depending on the demand for high care. Being a male caregiver, caring for a female, living apart from your relative, and low caregiver burden were positively related to caregiver resilience. Conclusion: Caregivers have the capacity to demonstrate resilience despite significant challenges. This study demonstrates how harmonization of data from multiple existing studies can be used to increase power and explore the consistency of findings. This contributes to a better understanding of which factors are likely to facilitate caregiver resilience and offers insights for developing services.
Introduction

Dementia is recognized globally as a major public health priority. Across the world, the majority of the care for persons with dementia (PwD) is provided by family members [1]. These caregivers clearly play a socially important and economically valuable role within society, but the challenge of informal care provision also presents a considerable risk for negative psychosocial consequences to the caregiver, often associated with the chronic stress involved with caregiving. Compared to non-caregivers, caregivers have an increased risk of depressive symptoms and physical health problems [2, 3].

Despite this, there are groups of caregivers who, even in the face of considerable caregiving demands, appear to manage relatively well [4]. This can potentially be interpreted as a sign of resilience. Little is known about resilient caregivers, who have fulfilled their care task relatively successfully, with few negative consequences for themselves. Understanding how some caregivers are able to function well and remain resilient when caring is an important step in rebalancing the current focus on stress and poor outcomes. This new approach to rethinking dementia caregiving could contribute to developing new programmes of services for those supporting PwD.

Resilience in the Context of Dementia Caregiving

Resilience is generally understood as a positive outcome despite exposure to an adversity or risk [5], and focuses on strengths rather than weaknesses. Inspired by ecological systems theory [6], Windle and Bennett [7] developed a theoretical resilience framework for caregivers. This recognizes that caregivers will draw on individual resources, but also interact with their environment by drawing on community and societal resources which may facilitate or hinder resilience. The absence of resources may lead to poor outcomes or further caring challenges. Considering this framework, resilience can be described as “the process of negotiating, managing and adapting to significant sources of stress or trauma. Assets and resources within the individual, their life and environment facilitate this capacity for adaptation and “bouncing back” in the face of adversity” [8, p. 163]. Whilst there is currently no standardized measure of caregiver resilience, this definition might facilitate outcome measurement, since it outlines the key elements of resilience: the encounter with adversity, the ability to resist and adapt to the adversity (through a range of assets and resources), and a positive outcome in the face of such challenges.

The few studies that attempt to understand resilience in this context provide an indication of how resilience could be measured and which factors are likely to influence resilience. Recently, Cherry et al. [9] synthesized the literature to identify factors and resources associated with good outcomes in the context of dementia care and concluded that carers’ resilience is supported by social and cultural factors, properties of the caring relationship, and psychological characteristics. Applying the caregiver resilience framework, a qualitative study identified from interviews with spousal dementia caregivers that staying positive, using downward comparisons with others, actively seeking knowledge, strong social relationships, and use of respite care facilitated their resilience [10]. Hindering factors included having a negative outlook, perceived loss of social relationships, and feeling isolated. Earlier quantitative studies which measured resilience with self-report instruments identified resilience as an important predictor of change in burden, life satisfaction, and perceived stress [11] and found that social support moderated this relationship [12]. These resilience measures mainly reflect inner psychological attributes such as self-esteem and mastery. Although these can be considered indicators of resilience, such measures may fail to capture the dynamic nature of resilience, as they do not describe some of its broader social determinants. Just one study attempted to identify resilience in dementia caregivers by examining a combination of
a positive outcome and exposure to adversity [13]. High caregiver resilience was defined as reports of low burden in the face of frequent demands that can occur directly in the context of dementia care [behavioural problems, dependency in activities of daily living (ADLs) and instrumental ADLs in the PwD]. High resilience was present in 45% of the 1,979 caregivers. Caring for a female, providing more care, caring for a longer duration, and greater utilization of (in)formal resources were positively associated with high resilience, whilst caregiver ethnicity or race, caregiver instrumental ADL dependencies, cohabitation, greater caregiver education and income, and greater cognitive impairment of the care recipient were negatively associated with high resilience in univariate analyses [13]. Inner psychological attributes were not examined, yet these are likely to play an important role in understanding the process of resilience.

The Present Study

Reflecting the caregiver resilience framework and drawing on the work of Gaugler et al. [13], we conceptualize resilience as a relatively high level of psychological well-being in the caregiver (‘a positive outcome’) despite various substantial demands that occur directly in the context of dementia care (‘adversities’), as visualized in figure 1. The first aim of this study is to examine what proportion of caregivers demonstrates high resilience under these different challenging caregiving demands. Secondly, we explore to what extent a variety of
internal and external resources are associated with caregiver resilience. To address these aims, this study harmonizes and integrates data from 4 studies conducted in the Netherlands and the UK.

Combining data sets from individual studies creates the opportunity to study a larger, more heterogeneous group of dementia caregivers. To be able to study resilience, a large sample size is particularly important, as it first requires the selection of the group of persons who are exposed to a significant risk or challenge. Also, combining data sets allows exploration of the consistency and generalizability of results from single studies. It responds to calls for increased data sharing and using limited resources more efficiently to answer new research questions [14]. Although the process of pooling original individual data across studies is a complicated and challenging task, it provides advantages over techniques synthesizing the relevant summary statistics, such as meta-analysis [14], and may be a way forward to advance the current state of knowledge about dementia caregiving. The present study aims to contribute to the existing literature on caregiver resilience by considering multiple samples and exploring the challenges and possibilities of harmonizing data across studies in this field. Accordingly, we first illustrate how the data were harmonized and subsequently address the two research aims.

**Materials and Methods**

**Data Sources**

Data were derived from 4 longitudinal studies conducted in the Netherlands and the UK selected as they included PwD and their primary informal caregivers living in the community at baseline and assessed key variables to measure resilience and its associated factors. The studies were the following:

- The REMCARE Study: a randomized controlled trial evaluating the (cost-)effectiveness of joint reminiscence groups for PwD and their caregivers as compared with usual care. A total of 488 dyads were recruited in the UK through NHS memory clinics and community mental health teams for older people [15].
- The FamCare Study: including a cohort of 157 informal caregivers of PwD referred to NHS community mental health teams for people with challenging behaviour in the UK (www.challengedemcare.com).
- The COMPAS Study: examined the (cost-)effectiveness of case management among community-dwelling PwD and their informal caregivers. A total of 521 dyads from regions with and without case management across the Netherlands were included [16].
- The Meeting Centres Support Programme (MCSP) studies: 2 studies evaluating the effects of the Dutch community-based MCSP compared with regular psychogeriatric day care in nursing homes. A total of 149 dyads of PwD and their informal caregivers participating in the day care programmes were included in the evaluation studies [17, 18].

**Study Sample**

For the current study, the baseline data from the study participants included in the original samples were used. This resulted in a total of 1,315 dementia caregivers in the combined data set. As the presence of significant adversity is a condition to be able to demonstrate resilience, the caregivers who faced substantial caregiving demands were selected from this data set to address the research aims (fig. 2).

**Definitions of Caregiver Resilience**

Caregiver resilience was defined as relatively high reported levels of psychological well-being ('positive outcome') while being exposed to various types of stressors that represent high demands on caregivers ('adversities'): caring for someone with more severe dementia, limitations in basic self-care, behavioural problems, and providing a substantial amount of care. Consequently, the caregivers who faced the high caregiving demand of interest were coded as highly resilient if they reported good psychological well-being, and as low resilient if their well-being was poor. In this way, a resilience definition was constructed for each of the 4 high caregiving demands we selected. Since it is often a combination of stressors that places a heavy burden on caregivers, we subsequently constructed a fifth resilience definition that distinguished caregivers facing more than 1 of the high caregiving demands from caregivers with no or 1 high caregiving demand.
Data Harmonization

To obtain a common set of variables that could be used to assess the research aims, where feasible we followed guidelines for integrative data analysis (IDA). IDA offers methods for integrating original individual data from multiple studies for secondary analysis [14, 19]. First, the scales and items in each of the studies were carefully reviewed. The exact wording of the relevant variables were examined by the first author and discussed with G.W. and B.W. to determine whether the variables and categories had the same face value across studies. To create the harmonized variable, categories of the original items were transformed and relabelled in each study depending on the wording and ordering. If the same item(s) was/were used, but on a different point scale, we converted scores to a common scale. Scales for care burden in each study were standardized. The harmonization process is described under Measures. Online supplementary table 1 (for all online suppl. material, see www.karger.com/doi/10.1159/000449131) gives a detailed overview of the study-specific measures together with the harmonized variables and categories.

Measures

Psychological Well-Being (‘Positive Outcome’)

Psychological well-being (‘positive outcome’) in the caregiver was measured with the General Health Questionnaire (GHQ), 12-item (FamCare and COMPAS) and 28-item (REMCARE and MCSP studies) versions [20]. The GHQ is a widely used and well-validated self-report instrument for symptoms of psychological distress. The person is asked to assess changes in his/her mood, feelings, and behaviours in the last 4 weeks. Higher scores indicate more distress. Goldberg et al. [21] compared the validity of the GHQ-12 and GHQ-28 and showed that the shorter GHQ is remarkably robust and works as well as the longer instrument to detect a case (the areas under the curve were 0.88 and 0.87, respectively). For both versions, we used the recommended GHQ scoring method thresholds (5/6 for the GHQ-28 and 1/2 for the GHQ-12) to detect psychiatric morbidity in order to distinguish between caregivers having good and those having poor psychological well-being [21].

High Caregiving Demands (‘Adversities’)

More Severe Dementia. The Clinical Dementia Rating (CDR) [22], Global Deterioration Scale (GDS) [23], and Mini-Mental State Examination (MMSE) [24] were used. These measures were harmonized into a dichotomous variable with mild versus moderate or severe dementia as categories following common classifications and ranges from the literature. Perneeczky et al. [25] showed that the MMSE classification has a substantial agreement with the CDR stages for the categories of mild, moderate, and severe dementia. Furthermore, Choi et al. [26] demonstrated high correlations between the GDS, CDR, and MMSE, reflecting a good concurrent validity. The mean MMSE scores in the GDS and CDR groups in this study were in line with the classification we used.

Fig. 2. Flow diagram of the study sample.
**Limitations in Basic Self-Care.** We defined considerable limitations in daily functioning of PwD as having problems with performing the basic self-care tasks. In the MCSP studies, the item measuring ‘help with dressing’ from the Assessment Scale for Elderly Patients (ASEP) was used (no vs. some or full help needed) [27]. For the other studies, the self-care item of the EQ-5D was used (no vs. some problems or unable to wash or dress themselves) [28]. Both measures were based on caregiver proxy reports. These measures were harmonized into a dichotomous variable with no limitations versus some or severe limitations as its categories.

**Behavioural/Mood Problems.** To assess this high caregiving demand, we focused on the presence of significant mood symptoms in the PwD, as these were measured in all studies. REMCARE and the first MCSP study included the Cornell Scale for Depression in Dementia (CSDD) [29], and we used the cut-point of ≥8, which indicates symptomatology consistent with clinically significant depression [30], to define significant mood problems in the PwD. In the FamCare and COMPAS and the second MCSP study, the Neuropsychiatric Inventory (NPI) was used to measure behavioural problems [31]. The NPI measures the frequency and severity of neuropsychiatric symptoms in 12 domains. The severity and frequency of each domain are scored and multiplied to calculate the domain composite score (range 0–12) with a score of ≥4 indicating the presence of clinically relevant symptoms, and a score of ≥9 indicating severe symptoms [e.g. 32, 33]. We used the NPI ratings of the combined 5 mood domains depression/dysphoria, anxiety, irritability/lability, sleep, and appetite (NPI-M) [34, 35] and defined the presence of significant behavioural problems as having 2 or more clinically relevant or at least 1 severe mood symptom. Both the CSDD and the NPI are valid, reliable instruments that are widely used to measure ‘behavioural’ disturbance in PwD.

**Providing a Substantial Amount of Care.** This was assessed in the REMCARE and FamCare studies by asking the caregiver how many hours he/she spent each week caring or performing care tasks for their relative. In the COMPAS study, caregivers were asked to report the hours they spent in the last week on specific tasks (household activities, personal care, transport, or visits outside the home) in which they supported the PwD or which they had taken over. The MCSP studies measured the number of days a week, and hours per day, a caregiver spent on average caring for the PwD. In each study, we converted these reports to the number of hours spent caring per week. Based on the estimate that a dementia caregiver spends around 40 h per week providing care duties on average, we used a threshold of ≥40 care hours to identify caregivers with a substantial caring role in order to create a dichotomous variable harmonized over the studies [36, 37].

**Potential Associated Factors of Resilience**

Based on previous studies on caregiver resilience [9, 10, 12, 13] and an evidence review on resilience in later life [38], a variety of individual and external resources were considered to be potentially associated with resilience, as visualized in figure 1.

**Sociodemographic Characteristics**

In all studies the caregivers were asked their own and their relative’s gender, age or date of birth, and employment status. The response categories for employment status differed slightly, and therefore we dichotomized this into (self-)employed or not employed to create a harmonized variable. Caregivers’ educational level was assessed in the COMPAS and MCSP studies with similar categories to record the caregiver’s highest completed education. We dichotomized these categories into lower or elementary versus secondary or higher education. The REMCARE and FamCare studies asked at what age the person left full time education. For these 2 studies, we used a threshold of 16 years to distinguish between caregivers with lower and higher education in the harmonized variable.

**Characteristics of the Care Context**

**Type and Quality of the Relationship.** All studies assessed the type of relationship between the caregiver and the PwD and whether they were living together. From these questions we constructed the harmonized variables ‘cohabiting’ (yes/no) and ‘relationship to the PwD’ (spouse/other). To examine the quality of the caregiver-patient relationship, the single item ‘I wish that my relative and I had a better relationship’ of the Short Sense of Competence Questionnaire (SSCQ) was used in the COMPAS, FamCare, and MCSP studies [39]. The answer categories ranged from 1 ‘agree very strongly’ to 5 ‘strongly disagree’, but in the MCSP studies the neutral option was not used. To obtain equivalent scores, we converted the item scores in the other 3 studies to a 4-point score. In the REMCARE study, the Quality of the Patient-Caregiver Relationship questionnaire (QCPR) was available [40]. We selected the QCPR item ‘There is a big distance in the relationship..."
between my relative and myself’, which we assumed to be the most similar in meaning to the SSCQ item. The item was measured on a 5-point Likert scale and converted to a 4-point score to create the harmonized variable.

Care Burden. This was assessed in all studies, but different scales were used. In the FamCare and COMPAS studies and the second MCSP study, the NPI Caregiver Distress Scale (NPI-D) was available [31]. The NPI-D asks caregivers to rate the emotional distress they experienced in relationship to 12 neuropsychiatric symptoms on a 6-point scale (0 ‘not at all’ to 5 ‘extremely distressing’). In the REMCARE study, the Relative Stress Scale (RSS) was used, a validated measure of general caregiver stress [41]. The 15 items are rated on a 5-point scale ranging from ‘never’ to ‘always’ or from ‘not at all’ to ‘considerably’. The first MCSP study used the 13-item Caregiver Strain Index (CSI) [42]. Each item asks if a stressor or an example of a situation that depicted the stressor was present (yes/no). On all scales, higher scores indicate a higher care burden. To standardize these scales, the total scores were converted into Z-scores in each study.

Type of Dementia and Time since Symptoms. These were assessed in the COMPAS and MCSP studies. For these studies, we dichotomized dementia type into a diagnosis of Alzheimer disease versus other dementias or not specified. Time since symptoms was assessed by asking the caregiver when the first symptoms of dementia in their relative had started, and was expressed in years. Harmonization was not reached in all studies, as these 2 variables were not assessed in the REMCARE and FamCare studies.

Social and Community Resources

Use of Services. All studies asked the caregiver if their relative received home care and day care services. In the REMCARE and FamCare studies, the number of visits (home care) or days (day care) over the past 3 months was recorded, while the COMPAS study recorded the number of hours per week during this period. The MCSP studies assessed if home care services were used (yes/no) at the time of baseline assessment. All participants included in the MCSP studies were participating in some type of day care, because of the study aim. To harmonize the use of home and day care, we dichotomized the variables into yes/no. The REMCARE and FamCare studies also recorded the number of visits from a case manager in the past 3 months. In the COMPAS study this was assessed as part of the inclusion criteria. Because this service was not measured in the MCSP studies, harmonization for the primary combined analysis was not possible.

Social Support. This was measured in all studies, but with different approaches. The MCSP studies used the Social Support List (SSL-12), a valid 12-item scale assessing the level of social support by means of social interactions with members of the primary social network [43]. The other 3 studies used a single question with slightly different wording to assess whether regular help from family or friends was present (yes/no). Harmonization between these 2 approaches examining family/social support was not possible, and these variables were therefore analysed separately in an ancillary analysis.

Feelings of Loneliness. These were only measured in the COMPAS and MCSP studies, with the De Jong-Gierveld Loneliness Scale (score 0–11), with higher scores indicating more loneliness [44]. Therefore, harmonization of this variable was not feasible across all samples.

Caregivers’ Inner Psychological Attributes

Various psychological attributes of the caregiver were represented in several of the studies. Sense of competence in providing care was assessed in all studies, except REMCARE, with the SSCQ [39]. The SSCQ includes 7 items scored on a 5-point Likert scale. Because the MCSP studies used the 4-point item version, we applied the dichotomized scoring method to calculate total scores, which recodes the answer categories of ‘agree very strongly/agree/neutral’ into 0 and ‘disagree/strongly disagree’ into 1. Higher total scores (0–7) indicate a better sense of competence.

Other inner psychological characteristics of the caregivers were only available in 1 of the studies:
- Use of coping styles was measured in the MCSP studies with the Jalowiec Coping Scale [45].
- Sense of mastery was assessed in the COMPAS study with the Pearlin Mastery Scale [46].
- Feelings of guilt were assessed in the FamCare study with the Guilt Scale [47].

Consequently, harmonization of these characteristics in the combined data set of all samples was not possible.

Statistical Analyses

Frequencies and descriptive statistics were used to describe sample characteristics and calculate the prevalence of caregiver resilience (aim 1). To address the second aim, first, candidate variables with a significant association with resilience in univariate logistic regression (p < 0.05) were selected for the multivariate analysis for each of the resilience definitions. To detect multicollinearity, we checked for strong correlations...
between the selected variables (Spearman’s ρ > 0.7). Second, the selected variables were entered in a multivariate logistic regression model and removed stepwise until all variables showed a statistically significant association with caregiver resilience (p < 0.05). Finally, we performed 2 ancillary analyses. First, we investigated if findings were consistent across studies. Differences in prevalence rates of resilience between study samples were tested with logistic regression analyses, with study membership included in the model as an independent categorical variable. When the overall effect of study membership was significant, pairwise comparisons between studies were conducted. To determine whether differences were statistically significant, the significance level was set at p < 0.0083 (0.05/6 comparisons for each resilience definition) after a Bonferroni correction for multiple comparisons. Next, we examined to what extent the identified factors associated with resilience in the primary analyses were consistent across studies. To do this, we examined whether the interaction between the variable and study membership was significant when included in the final multivariate model of the primary analysis. In the second analysis, we explored if the variables that could not be harmonized across all studies were of significance. For this purpose, we added the variable to the final multivariate model from the primary analysis and used the χ² goodness-of-fit test to compare the 2 models in the sample(s) of caregivers for which the variable was available.

Study membership was included as a fixed factor (categorical variable) in all analyses using data from multiple studies to control for between-study heterogeneity in hypothesis testing (fixed-effect IDA [19]). Before creating the harmonized variable for severity of dementia, we used maximum likelihood imputation as implemented by the EM algorithm in SPSS to impute missing MMSE total scores in the COMPAS study. In order to replace missing MMSE scores by their most likely values while also taking into account the mechanism that generated the missingness, statistically significant predictors of the MMSE score and predictors of its missingness were included in the maximum likelihood estimation procedure to obtain the required imputed values. All analyses were performed using SPSS version 20.

Results

Study Sample

From the combined data set of 1,315 caregivers, 1,048 faced a high caregiving demand of interest and their data were used for analysis in the present study (fig. 2). The characteristics of these caregivers and their relatives with dementia are described in table 1.

Table 1. Description of the study samples

<table>
<thead>
<tr>
<th></th>
<th>REMCARE (n = 379)</th>
<th>FamCare (n = 141)</th>
<th>COMPAS (n = 405)</th>
<th>MCSP studies (n = 123)</th>
<th>Total (n = 1,048)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Caregiver</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>characteristics</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean age (SD), years</td>
<td>69.1 (11.7)</td>
<td>66.0 (13.2)</td>
<td>64.3 (12.5)</td>
<td>64.4 (12.5)</td>
<td>66.3 (12.5)</td>
</tr>
<tr>
<td>Female gender, n (%)</td>
<td>262 (69.1)</td>
<td>102 (72.3)</td>
<td>273 (67.4)</td>
<td>92 (74.8)</td>
<td>729 (69.6)</td>
</tr>
<tr>
<td>Education, n (%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Elementary/lower</td>
<td>167 (44.1)</td>
<td>60 (42.6)</td>
<td>61 (15.1)</td>
<td>51 (41.5)</td>
<td>339 (32.3)</td>
</tr>
<tr>
<td>Secondary or higher</td>
<td>205 (54.1)</td>
<td>81 (57.4)</td>
<td>339 (83.7)</td>
<td>69 (56.1)</td>
<td>694 (66.2)</td>
</tr>
<tr>
<td>(Self-)employed, n (%)</td>
<td>63 (16.6)</td>
<td>35 (24.8)</td>
<td>160 (39.5)</td>
<td>24 (19.5)</td>
<td>282 (26.9)</td>
</tr>
<tr>
<td>Cohabiting, n (%)</td>
<td>311 (82.1)</td>
<td>102 (72.3)</td>
<td>214 (52.8)</td>
<td>86 (69.9)</td>
<td>713 (68.0)</td>
</tr>
<tr>
<td>Spousal relationship, n (%)</td>
<td>261 (68.9)</td>
<td>74 (52.5)</td>
<td>204 (50.4)</td>
<td>84 (68.3)</td>
<td>623 (59.4)</td>
</tr>
<tr>
<td><strong>Care recipient (PwD)</strong> characteristics</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean age (SD), years</td>
<td>77.9 (7.3)</td>
<td>80.7 (7.6)</td>
<td>80.3 (7.8)</td>
<td>75.4 (7.8)</td>
<td>78.9 (7.8)</td>
</tr>
<tr>
<td>Female gender, n (%)</td>
<td>184 (48.5)</td>
<td>84 (59.6)</td>
<td>232 (57.3)</td>
<td>57 (46.3)</td>
<td>557 (53.1)</td>
</tr>
<tr>
<td>Type of dementia, n (%)</td>
<td>n/a</td>
<td>n/a</td>
<td>193 (51.6)</td>
<td>57 (46.3)</td>
<td>250 (47.3)</td>
</tr>
<tr>
<td>Alzheimer disease</td>
<td>n/a</td>
<td>n/a</td>
<td>193 (51.6)</td>
<td>57 (46.3)</td>
<td>250 (47.3)</td>
</tr>
<tr>
<td>Other/not specified</td>
<td>209 (47.7)</td>
<td>55 (44.7)</td>
<td>264 (65.0)</td>
<td>55 (44.7)</td>
<td>264 (50.0)</td>
</tr>
<tr>
<td>Mean years since first symptoms (SD)</td>
<td>n/a</td>
<td>n/a</td>
<td>4.5 (2.9)</td>
<td>4.2 (3.2)</td>
<td>4.4 (2.9)</td>
</tr>
</tbody>
</table>

SD = Standard deviation; PwD = person with dementia; n/a = not available. 1 Included COMPAS and MCSP study participants.
Table 2. Prevalence of high resilience, defined as good psychological well-being (GHQ score below the clinically relevant cut-point) in the face of different high caregiving demands, in the combined sample and in each study separately

<table>
<thead>
<tr>
<th>High resilience in the face of:</th>
<th>REMCARE</th>
<th>FamCare</th>
<th>COMPAS</th>
<th>MCSP studies</th>
<th>Total sample</th>
</tr>
</thead>
<tbody>
<tr>
<td>Good psychological well-being</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>More severe dementia in the PwD</td>
<td>66 (51.2)</td>
<td>14 (24.1)</td>
<td>135 (43.7)</td>
<td>22 (44.0)</td>
<td>237 (43.4)</td>
</tr>
<tr>
<td>Limitations in basic self-care in the PwD</td>
<td>118 (52.9)</td>
<td>19 (19.2)</td>
<td>62 (41.3)</td>
<td>32 (42.1)</td>
<td>231 (42.2)</td>
</tr>
<tr>
<td>Behavioural/mood problems in the PwD</td>
<td>94 (45.2)</td>
<td>13 (17.1)</td>
<td>52 (30.6)</td>
<td>21 (36.2)</td>
<td>180 (35.2)</td>
</tr>
<tr>
<td>Providing a substantial amount of care</td>
<td>106 (53.0)</td>
<td>27 (31.0)</td>
<td>22 (28.2)</td>
<td>23 (30.7)</td>
<td>178 (40.5)</td>
</tr>
<tr>
<td>More than 1 high caregiving demand</td>
<td>121 (51.3)</td>
<td>20 (20.0)</td>
<td>74 (34.4)</td>
<td>29 (37.2)</td>
<td>244 (38.8)</td>
</tr>
</tbody>
</table>

Figures are numbers, with percentages in parentheses. PwD = Person with dementia.

Prevalence of Resilience: Good Psychological Well-Being despite High Caregiving Demands

Table 2 shows that caregivers were most often highly resilient when caring for a person with more severe dementia (43%), followed by caring for someone with limitations in basic self-care (42%) and providing a high amount of care (41%). The prevalence of resilience was lowest when dealing with behavioural/mood problems in the PwD (35%). About 39% demonstrated high resilience when facing more than 1 high caregiving demand.

Factors Associated with Caregiver Resilience

From the list of factors potentially associated with resilience, 15 of the 24 variables were harmonized across all samples and included in the combined data set. Table 3 illustrates the factors that remained significant in the final multivariate models. A high care burden was negatively associated with high resilience for all definitions. Also, a negative association was found between cohabiting with the PwD and high resilience for all definitions, except when the high caregiving demand referred to ‘providing a substantial amount of care’. Gender of the PwD (being female) or caregiver (being male) was positively associated with resilience under all challenging circumstances, except when ‘caring for a person with more severe dementia’.

Ancillary Analyses

Consistency of Findings across Studies

The first ancillary analysis showed statistically significant differences between studies in the prevalence of resilience for all definitions. Online supplementary table 2 presents the results of the pairwise comparisons between studies. In summary, applying the Bonferroni correction for multiple comparisons, the prevalence of resilience was significantly higher in the REMCARE study than in the other samples for most of the resilience definitions. The FamCare sample included the lowest number of highly resilient caregivers compared to the other samples for most definitions.

Most of the variables identified to be significantly related to high resilience in the primary multivariate analyses were consistent across studies. We looked for interaction effects with study membership in the multivariate regression models and found only a significant interaction with cohabiting status for the resilience definition ‘good psychological well-being in the face of more than 1 high caregiving demand’ (cohabiting status × study interaction: Wald $\chi^2 = 10.16$, d.f. = 3, $p = 0.02$). Only in the COMPAS sample, cohabiting with the PwD showed a significant negative association with high resilience (adjusted OR = 0.21, Wald $\chi^2 = 20.94$, d.f. = 1, $p < 0.001$, 95% CI 0.11–0.41).
Significance of Variables That Could Not Be Harmonized across all Studies

For 9 variables, harmonization was not feasible across all samples (online suppl. table 1). The second ancillary analysis examined if these variables were of significance for caregiver resilience. The $\chi^2$ goodness-of-fit tests indicated that a better sense of competence, greater mastery, and fewer feelings of loneliness in the caregiver were of significance for all resilience

<table>
<thead>
<tr>
<th>Table 3. Results from the univariate and multivariate regression analyses</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>a</strong> Examining the associated factors of resilience in the combined sample</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td><strong>Sociodemographics</strong></td>
</tr>
<tr>
<td>Age of caregiver</td>
</tr>
<tr>
<td>Male gender of caregiver</td>
</tr>
<tr>
<td>Higher education of caregiver</td>
</tr>
<tr>
<td>Caregiver (self-)employed</td>
</tr>
<tr>
<td>Age of PwD</td>
</tr>
<tr>
<td>Female gender of PwD</td>
</tr>
<tr>
<td><strong>Context of care</strong></td>
</tr>
<tr>
<td>Cohabiting</td>
</tr>
<tr>
<td>Spousal relationship</td>
</tr>
<tr>
<td>High relationship quality (1–4)</td>
</tr>
<tr>
<td>Higher care burden (Z-score)</td>
</tr>
<tr>
<td><strong>Community resources</strong></td>
</tr>
<tr>
<td>Use of home care</td>
</tr>
<tr>
<td>Use of day care</td>
</tr>
<tr>
<td><strong>b</strong> Continued for the resilience definitions with the high caregiving demands ‘substantial time spent caring’ and ‘more than 1 high care demand’</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td><strong>Sociodemographics</strong></td>
</tr>
<tr>
<td>Age of caregiver</td>
</tr>
<tr>
<td>Male gender of caregiver</td>
</tr>
<tr>
<td>Higher education of caregiver</td>
</tr>
<tr>
<td>Caregiver (self-)employed</td>
</tr>
<tr>
<td>Age of PwD</td>
</tr>
<tr>
<td>Female gender of PwD</td>
</tr>
<tr>
<td><strong>Context of care</strong></td>
</tr>
<tr>
<td>Cohabiting</td>
</tr>
<tr>
<td>Spousal relationship</td>
</tr>
<tr>
<td>High relationship quality (1–4)</td>
</tr>
<tr>
<td>Higher care burden (Z-score)</td>
</tr>
<tr>
<td><strong>Community resources</strong></td>
</tr>
<tr>
<td>Use of home care</td>
</tr>
<tr>
<td>Use of day care</td>
</tr>
</tbody>
</table>

Resilience was measured with 5 definitions defined as good or poor psychological well-being in the face of high caregiving demands. Dependent variable: 1 = high resilience; 0 = low resilience. OR = Odds ratio; PwD = person with dementia. * p < 0.05, entered in the multivariate model. Higher scores are better.
definitions when adding each of these variables to the final multivariate model derived from the primary analysis and using the data of the caregivers for which the variable was available (Table 4). Furthermore, fewer feelings of guilt and higher levels of social support (SSL score) were of significance in the face of some challenging circumstances. No added value was found for the type of dementia, years since first symptoms, coping styles, receiving support from a case manager, and regular help from family and friends.
Discussion

Main Findings and Interpretation

This study contributes new insights into resilience in dementia caregiving, utilizing data from 4 studies across 2 countries. Diverse approaches to measuring resilience showed that a considerable number of caregivers are able to have good psychological well-being, despite different challenges of caring. The prevalence of resilience was comparable when high caregiving demands were present in multiple care domains. High resilience was least prevalent when behavioural/mood problems in their relative were reported and associated with gender of the caregiver and PwD as well as characteristics of the context of care (cohabiting and care burden). These insights could help to identify those at high risk and provide important orientation in priority setting for intervention by services. The exploratory ancillary analyses suggested the importance of social relationships and some inner psychological attributes of the caregiver (mastery and sense of competence). As some of these are potentially modifiable, this provides indications for practitioners to focus their support – e.g. by providing interventions aimed at improving caregivers’ mastery and competence, particularly in managing challenging behaviour and (thereby) decreasing carer burden.

Comparison with Other Studies

Our estimates are quite similar to the rate of 45% reported by Gaugler et al. [13], who examined resilience, defined as lower or higher perceived burden in the face of frequent care demands, in a large sample of dementia caregivers. Several studies on resilience among older adults (not specifically caregivers) have been conducted. High scores on resilience scales have been reported and the suggestion made that these levels may remain steady or increase as older adults gain from the challenges they cope with successfully [48, 49]. Netuveli et al. [50] conducted one of the few longitudinal studies on resilience in older people, reporting a prevalence rate of 14.5%. In this study, resilience was measured as a GHQ-12 score that increased after exposure to an adversity (functional limitation, bereavement or marital separation, or poverty) and returned to its pre-exposure level in the following year. Our considerably higher estimates could be due to the different types of adversity or the cross-sectional measurement of resilience in our study.

The regression analyses of the combined sample indicated the importance of cohabitation, gender, and care burden. In line with the study by Gaugler et al. [13], we found that living apart from the PwD was related to high resilience. Cohabitation was not a significant factor among caregivers providing a high amount of care, suggesting that being heavily involved rather than living together threatens caregiver resilience. Whilst Gaugler found a positive (univariate) association between being a female caregiver and resilience, our results indicated that male caregivers and persons caring for a female were more often highly resilient. This difference in findings might be due to the fact that Gaugler defined high resilience as low burden levels despite high care demands, while we used good psychological well-being as the positive outcome to identify highly resilient caregivers. When experiencing a stressful situation, females might be more likely to respond with depressive or anxious symptoms (captured by the GHQ). Care burden was significantly related to resilience under all challenging circumstances. Previously, Chappell and Dujela [11] found that resilience is an important predictor of change in burden. It is also known that a high caregiver burden presents a risk for negative consequences to caregivers’ health [51–53]. Our study among caregivers in adversity indicates that despite relatively equal substantial care demands, the perceived burden of care varies and is a key factor for demonstrating resilience.

Prior studies among dementia caregivers provided evidence for the relevance of social supportive resources [10, 12]. Our ancillary analysis of some of the individual studies also
suggested that perceived low levels of social support and feelings of loneliness are related to low resilience. In contrast, this relationship was not found for the variable 'help from family and friends'. As this variable was a dichotomized measure, we hypothesize that it may not have been as sensitive as the continuous social support and loneliness measures. In addition, these analyses indicated the positive impact of inner psychological attributes such as mastery and sense of competence, and, to a lesser extent, the absence of guilt feelings. This corresponds with earlier findings from qualitative interviews reporting the importance of staying positive [10]. Previously, resilience was found to be positively related to a problem-focused coping style and the use of distancing, by not letting situations get to oneself and looking on the bright side, while escape-avoidance coping appeared to have a negative impact on resilience [54]. As males generally adopt more problem-focused coping styles and females use more emotion-focused styles when facing stress [e.g. 55], this might be one of the explanations why more males than females were highly resilient in our study. We were only able to explore the importance of coping styles in a small sample of caregivers. Possibly, this analysis might have lacked statistical power to detect a significant impact.

Strengths and Limitations

As far as we are aware, this is the first study addressing resilience simultaneously in multiple samples of dementia caregivers across nations. This enabled us to use a large, diverse sample and consider the heterogeneity of caregivers, contributing to a better understanding of the role of assets and resources related to resilience independent of specific study contexts. The use of existing data sets allowed us to add to the very limited work so far on resilience in dementia caregiving in an efficient way, but it also had its restrictions. A major challenge involved the variations across the data sets with regard to the measures that were used. Therefore, we needed to be pragmatic in measuring resilience. For example, we included only mood symptoms when examining behavioural problems in the PwD as adversity, because these symptoms were assessed in all studies. Although depression is one of the most common disturbances in PwD [e.g. 56] and puts a heavy strain on caregivers, there are also other important behavioural symptoms which are likely to increase the demand of care. Besides, the 2 scales we used might have differed in their ability to detect mood symptoms in dementia. A study which examined the effect of sertraline on depression in patients with Alzheimer disease suggested that the CSDD was more likely to detect differences in depression over time than the NPI-M [35]. This might be due to differences between items, but also because the CSDD combines patient observations with discussions with the caregiver to assess depression, whilst the NPI-M uses a caregiver interview only. It has previously been suggested that caregivers view depressed PwD as less depressed than do clinicians [57]. In our study, the prevalence of significant mood problems in the samples that used the NPI-M was indeed somewhat higher than in the samples using the CSDD – except in the FamCare sample, which showed the highest percentage of PwD with mood problems, as this study recruited people with a referral for challenging behaviour (data not shown).

We found variability between samples in resilience rates. This could be explained by differences in participants’ characteristics and sampling criteria. For example, the FamCare study included caregivers of PwD who were referred for displaying challenging behaviour. FamCare caregivers were most frequently facing a high caregiving demand, and the resilience rates were lowest under all the challenging circumstances. This could indicate that this sample involved a more severely afflicted patient group compared with the other studies. Compared with the other samples, REMCARE caregivers demonstrated high resilience most frequently for all definitions. They might have experienced better psychological well-being because they cared for relatively young persons in a mild disease stage. Although we cannot be sure to what extent variations in designs and methods across studies have impacted our
results, the use of several samples also provided the opportunity to observe these differences between caregivers. Besides, the ancillary analyses indicated that almost all factors that were identified to be significantly related to resilience in the primary analysis were consistent across studies. Only the association between cohabitation and resilience in the face of more than 1 high caregiving demand differed between studies. In the COMPAS sample, cohabiting with the PwD showed a significant negative association with high resilience, while this association was not significant in the other samples. The fact that a relatively large proportion of COMPAS caregivers were living apart from the PwD compared to the other samples might have influenced this association.

Since the studies were not originally designed to examine resilience, other factors that might be related to resilience, such as biological and physical characteristics or the presence of other major stressful life events, could have been missed. Some potential influencing factors, such as caregiving duration as well as psychological and social resources, would be useful to investigate, but they were not available for all samples and thus excluded from the primary combined analysis. The ancillary analysis using (some of) the individual samples indicated that inner psychological attributes and strong social relationships are important factors to examine in future research.

Although all study samples had longitudinal data available, we only used baseline data, as different follow-up times and the exposure to interventions during some of the studies would likely have influenced the results. Consequently, we could not demonstrate if resilience was present over time or identify which factors were key to achieve resilience. Therefore, future longitudinal research is needed to develop an understanding of how caregivers, despite significant challenges, are able to continue to function well in the long term and to determine the causal relationship to potential resources.

We tried to minimize problems of combining data sets and consider heterogeneity in studies by following IDA methods as much as possible. For example, we used fixed-effect IDA to control for between-study heterogeneity. Also, we aimed to optimally harmonize variables by examining the face validity of items, making transformations if these were highly overlapping across studies and standardizing scales if different measures were used for the same domain. This approach corresponds with previous efforts to harmonize data in a pragmatic way [58–61]. Even though limitations of our study are related to restrictions of secondary data analysis, this could also be seen as one of the strengths. Until now, research on dementia caregiving has predominantly focused on analysing data from single studies, with often relatively small, homogeneous samples. Also, existing population-based studies often do not provide enough information to study this target group thoroughly, especially around the types of caregiver challenges specifically arising from someone living with dementia. For example, in the UK, the Cognitive Function and Aging Studies (CFAS) do not ask about the type and context of care [62]. Besides, although dementia is a public health priority, the number of PwD (and their caregivers) still forms only a small percentage of the overall population. Despite its challenges, attempts to combine original data from secondary data sources may create new opportunities to test hypotheses and can provide advantages over focusing on one sample or time-consuming and expensive collection of primary data.

**Conclusion and Implications**

Examining this topic in multiple data sets simultaneously provides a starting point for future research. Our conceptual model reflects initial considerations of what might enable resilience for dementia caregivers, and our analyses sought to test this process. In summary, the findings indicate that although caregiving for a relative with dementia can be extremely
challenging, a group of caregivers appear to be able to manage relatively well. Resilience was observed across samples in the face of various types of high care demands. The finding that high resilience was associated with several factors confirms the conclusion of previous studies [10, 13] and indicates the multidimensional nature of resilience. Future studies should investigate the potential of different services to improve caregiver resilience. A standardized resilience measure in the context of caregiving would allow more accurate measurement across populations and may facilitate tailoring of interventions. In the face of the growing numbers of people living with dementia, ensuring those who support them can remain healthy is a policy priority. There is a challenge to apply research findings to the development and implementation of interventions which could, when successfully implemented, promote resilience in caregivers and ensure that providing care for PwD remains manageable in the future.

Acknowledgements

This work was supported by the Dutch Alzheimer’s Association (Alzheimer Nederland; grant No. WE.15-2014-05). The funder had no further role in the study design, in the collection, analysis, and interpretation of the data, in the writing of the report, and in the decision to submit the paper for publication.

Disclosure Statement

The authors declare that they have no competing interests.

References


