Reliability, validity and relevance of needs assessment instruments for informal dementia caregivers: a psychometric systematic review

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ABSTRACT

Objective: The objective of this review was to identify reliable and/or valid needs assessment instruments for informal dementia caregivers that are relevant for clinical practice, research and informal caregivers.

Introduction: Informal dementia caregivers report important unmet needs at all stages of the disease. In addition, they often indicate that health care providers insufficiently attend and adapt to their multiple needs. A systematic and patient-centered assessment is needed to address this lack of knowledge and understanding. However, existing quantitative needs assessment questionnaires are limited in terms of psychometric testing. Qualitative measures are time-intensive and difficult to conduct on a large scale, with growing economic pressure. Information about the methodological quality and the characteristics of needs assessment instruments are crucial for clinicians and researchers to make informed decisions about the most reliable and valid tool for their specific purpose.

Inclusion criteria: This review considered studies on multidimensional needs assessment instruments for informal dementia caregivers living at home. Psychometric studies or other types of studies with sufficient data to evaluate methodological quality were included if they considered at least one outcome for reliability or validity.

Methods: Studies in English, French or German and published until February 2019 were searched in four databases: Embase, MEDLINE, CINAHL and PsycINFO. After screening the titles, abstracts or full texts for eligibility, the provisional included studies were assessed for methodological quality with a standardized tool for systematic reviews of measurement properties. After data extraction using a standardized tool, the quality of the measurement properties was rated and compared using predefined quality criteria.

Results: Eighteen articles covering 14 different needs assessment instruments were included in the review. Eleven publications focused on the development or the evaluation of an instrument. In addition, a development report, a manual and five studies, not aimed primarily at validation but containing sufficient information about the development or the evaluation of the used instruments, were included. The systematic evaluation of the instruments revealed that half of them had excellent content validity. In contrast, structural validity was rarely examined, and mostly with an insufficient sample size or a questionable analysis. None of the instruments had optimally tested and good internal consistency. Regarding reliability, test-retest agreement was rarely tested and inter-rater agreement was evaluated using controversial procedures. Comparing the different instruments reviewed, the “Partnering for better health – living with chronic illness: dementia” had the best psychometric evidence, and the “Questionnaire of consultation expectations” was also partly supported, while most other instruments presently had limited psychometric soundness.

Conclusions: Despite the good evidence for some psychometric properties, further developments in the field of needs assessment for informal dementia caregivers are needed, particularly regarding structural and construct validity, as well as test-retest reliability and sensitivity to change. To enhance conceptual clarity, the development of an underlying theoretical model of needs should be prioritized.

Keywords Dementia; informal caregivers; instrument; needs assessment; psychometrics

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The authors declare no conflict of interest.

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## Summary of Findings

### Reliability, validity and relevance of needs assessment instruments for informal dementia caregivers

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| Psychometric property | Summary or pooled results | Overall rating | Factors determining the quality of evidence | GRADE of the quality of evidence |
|------------------------|---------------------------|----------------|--------------------------------------------|----------------------------------|
| Content validity       | Seven studies with excellent content validity, but the identified domains were not consistent. | +/- inconsistent | - 1 inconsistency | Moderate |
| Internal consistency   | One study had an adequate dimensionality analysis but low alphas; all others had either their dimensionality evaluated with small samples, or no proper evaluation of their dimensionality, although some had high alphas. | - insufficient | - 1 risk of bias | Moderate |
| Structural validity    | Four studies evaluated the factor structure of their instrument: one with an excellent factor analysis meeting all criteria, two with an adequate factor analysis but samples sizes were too limited, and one with a factor analysis inadequately performed separately for each dimension. Identified factor structures varied from five to eight dimensions. | - insufficient | - 1 inconsistency | Moderate |
| Reliability – test-retest | Two studies used a satisfactory procedure to evaluate test-retest agreement, one obtained a good correlation, and the other had two correlations above .70 and three below .70. Results for measurement error were rated as good for one study. | +/- inconsistent | — | High |
| Reliability – inter-rater | Inter-rater agreement was evaluated with questionable procedures. | ? indeterminate | - 2 risk of bias | Low |
Introduction

Dementia is characterized by a progressive decline of cognitive and social functions. This limits the autonomy of those affected by dementia, and makes it difficult for them to cope with daily life. They become increasingly dependent on the care of others, particularly informal caregivers. Informal caregivers are individuals who regularly provide unpaid care, assistance and/or supervision to a close person with reduced autonomy. Studies show that informal dementia caregivers provide care of a higher intensity (more hours per day) and longer duration (over more years), compared to caregivers of people without dementia. In a survey of Alzheimer Europe, almost half of informal dementia caregivers spent more than 10 hours per day providing care. This is comparable to data from the United States (US), where informal dementia caregivers spent on average nine hours per day caregiving, and 31.1% of informal dementia caregivers provide care for two to three years, 18.5% for four to five years and 38.4% for six or more years. For 2015, the estimated economic value of unpaid care provided by informal dementia caregivers in the US was US$221.3 billion. In Switzerland, an example of a Western European country, informal caregivers contributed 80 million unpaid hours in 2016, a substantial increase compared to 64 million hours invested in 2012 and 52 million in 2010. The contribution of informal caregivers is expected to further increase due to the rising care needs of an aging population and the growing prevalence of multiple chronic conditions, in particular dementia. In addition, the number of available formal carers is not expected to increase accordingly. Informal caregivers thus play a key role, not only for the people with dementia but also for society in the sustainability of the health care system. Recognizing the valuable contribution of informal caregivers and providing them adequate support is therefore a core public health issue.
Caring for a person with dementia is a challenging experience, and the burden of informal dementia caregivers is higher compared to informal caregivers of persons with other chronic conditions. Straining continuous care, an unpredictable course, and neuropsychiatric symptoms of the person with dementia can cause high levels of stress, which often leads to physical, psychological, emotional, social and financial problems. In addition, family caregivers often have no experience in performing care, feel unprepared, and are lacking the required knowledge and support from health care providers to deliver appropriate care. Informal caregivers report feelings of tiredness, stress, helplessness, and loneliness, and show high prevalence of depression and anxiety. Due to the nature of dementia, informal caregivers also struggle with feelings of guilt, ambivalence, grief and loss. Identified physical problems can create an increased risk for vascular disease, impaired wound healing, decreased immunity, and reduced likelihood to engage in preventive health behavior. Poor physical and psychological health conditions not only impair the quality of life of informal dementia caregivers but also affect their ability to provide care to the person with dementia and to sustain their own social support network, which leads to social isolation. Burden and health deterioration of informal dementia caregivers are core predictors of early institutionalization and mistreatment of their care recipient.

Due to the challenges of caregiving and the associated burden, informal dementia caregivers report important unmet needs at all stages of the disease. Their needs cover very diverse areas, such as information about the illness and support resources; support for their own emotional concerns; support on how to communicate with the care recipient, the family or the service providers; practical support in daily care and respite; and financial support. Informal dementia caregivers often report that health care providers insufficiently attend and adapt to their multiple needs, and complain about care fragmentation and poor coordination, which ultimately increases their stress and underutilization of support services despite their needs. Underutilization of health care and other support resources contributes to the exhaustion of the informal dementia caregivers and precipitates institutionalization of their care recipient, thereby increasing health care costs. Informal caregivers do not always express spontaneously and directly how their needs can be met. Therefore, evaluating their needs in a systematic manner is crucial to supporting them in fulfilling their needs in a person-centered way in order to promote the quality of life of the caregiver and the affected person, as well as to maintain the caring situation at home.

Most studies on the needs of informal dementia caregivers have used qualitative study designs. Existing quantitative questionnaires have limitations: very few items for caregivers, poor validation, or lack of empirical evidence about needs dimensions (factor structure). This limits their use in both research and clinical practice. In addition, many of the assessment instruments, particularly semi-structured interviews, are time-intensive (e.g. assessment alone takes on average two hours or 90 minutes). Furthermore, most of the collected information in interviews is qualitative. As such it is usually extensive, and more time is needed to prepare the information to make it available for the caregiver or other service providers (e.g. transcriptions). In view of the growing economic pressure on the social system, and of the rising support needs associated with population aging, such resources are impossible to secure on a large scale.

A search in MEDLINE, CINAHL, the JBI Database of Systematic Reviews and Implementation Reports and the Cochrane Database of Systematic Reviews was performed in January 2017, and again in January and July 2018, to identify completed and in-progress systematic reviews on needs assessment instruments for informal dementia caregivers. Three systematic reviews examining dementia needs assessment instruments were identified. Two reviews focused on documenting the diverse instruments available, focusing on their content and methodological approach to measure the needs, with no specific interest in their psychometric properties. Novais et al. included all types of studies that conducted a needs assessment, and Bangerter et al. concentrated on quantitative findings. The third review (Mansfield et al.) critically examined the psychometric properties of a very limited number of instruments (N = 4) due to highly restrictive inclusion criteria; only peer-reviewed studies in English published until August 2015 were included. In addition, they targeted instruments where all items were directly asked about their needs.
Information about the methodological quality of measurement instruments are crucial for clinicians and researchers to make informed decisions about the best tool for their specific purpose. Therefore, the current review aimed to expand on the previous three reviews by focusing specifically on (at least partially) validated instruments and documenting in detail their psychometric properties. For this purpose, we followed the COSMIN (COnsensus-based Standards for the selection of health Measurement INstruments) guidelines for systematic reviews of measurement properties, which are recommended for psychometric reviews. In addition, the current review expanded on that of Mansfield et al. by including: i) instruments with diverse application methods (i.e. also professionally-assessed) or with items for both caregivers and persons with dementia, ii) studies published until February 2019 and in English, French and German, and iii) the CINAHL database as a relevant source for caregiver literature. The current review also includes recommendations for conducting further psychometric validations in the field of needs assessment among informal dementia caregivers. This review provides comprehensive and systematic information to guide further development of well-validated needs assessment instruments for use in research and clinical settings.

This review was conducted according to an a priori published protocol.

Inclusion criteria
The inclusion criteria were developed following COSMIN guidelines and JBI guidance. The COSMIN guidelines for systematic reviews of measurement properties recommend the following inclusion criteria: i) the instrument should aim to measure the construct of interest (types of intervention(s)/phenomena of interest), ii) the study sample should comprise the target population of interest (types of participants), iii) the study should concern the type of measurement instrument of interest (self-reported or professionally interviewed), iv) the aim of the study should be the development of a measurement instrument or the evaluation of one or more of its measurement properties (types of studies).

Participants
This review considered studies that included informal caregivers of persons with dementia living at home as the study sample or as a part of it. Where possible, only the corresponding data were included in the review. Informal caregivers were defined as individuals who regularly provide unpaid care, assistance and/or supervision to a close person with reduced autonomy, in this context with dementia. In regards to the person with dementia, there was no restriction regarding the type of dementia.

Instruments/construct
This review considered studies that report on needs assessment instruments for informal dementia caregivers. Needs was defined as “a condition that is important to the subject and that is not being satisfied in the subject’s present environment.” The application method of the instruments was either self-reported or professionally interviewed. Further inclusion criteria were added while identifying and screening the literature to complement those of the a priori protocol: Instruments needed to be multidimensional (e.g. include more dimensions than only information needs). Instruments with items for informal caregivers and people with dementia were included if they contained at least two dimensions for the caregivers. Measuring the needs of informal dementia caregivers had to be an explicit objective of the instrument or of specific dimensions.
Outcomes
This review considered studies that included the following outcomes for psychometric properties:

- Reliability (test-retest reliability, inter-rater reliability, internal consistency)
- Validity (content validity, construct validity, structural validity, sensitivity to change)

In order to characterize the instruments, we additionally documented the following instrument characteristics:

- Purpose (original intended use)
- Application method (self-reported, professionally interviewed)
- Administration burden (training for clinicians, time for completion)
- Number of items and domain structure.

As not every article provided data for all of the psychometric outcomes, articles that reported at least one outcome regarding reliability or validity were considered. Criterion-related validity was not considered in the results as there was no reasonable gold-standard available for the included instruments (in accordance with COSMIN guidelines). No data were found for the outcome sensitivity to change.

Types of studies
This review considered psychometric studies, namely, instrument development or instrument evaluation studies. Other types of studies (in which needs assessment instruments were used) were included to identify eligible instruments and their responsible authors. If no published or unpublished psychometric study was available, other types of studies (e.g. a survey) were only included if they provided sufficient information to evaluate the methodological validity of at least one psychometric property according to the COSMIN checklist.

Methods
This review was conducted according to an a priori published protocol and has been registered with PROSPERO (CRD42018090611).

Search strategy
The search strategy aimed to find both published and unpublished studies. A three-step search strategy was utilized in this review. An initial limited search of MEDLINE and CINAHL was undertaken followed by analysis of the text words contained in the title and abstract, and of the index terms used to describe the articles. A second search was undertaken in February 2019 across all included databases. The search strategy considered all identified keywords and index terms as well as search blocks for dementia and patient-reported outcome measures provided by the study group of the Biomedical Information of the Dutch Library Association (KNVI) and a search block combined with a filter for measurement properties, as suggested by and available from the COSMIN website. Thirdly, the reference lists of all selected full texts were searched for additional studies. A complementary search was performed in the included databases and gray literature using the names of the needs assessment instruments identified in the three foregoing steps, and authors were contacted to obtain possible additional gray literature relating to their instrument. Studies published in English, German and French were considered for inclusion in this review. There was no limitation regarding the publication time.

The databases searched included Embase, MEDLINE, CINAHL and PsycINFO. In contrast to the a priori protocol, Embase was searched as recommended by COSMIN, and CINAHL was searched instead of OVID Nursing.

The search for unpublished studies included Google Scholar, ProQuest Dissertations and Theses, ResearchGate (contact with relevant researchers), homepages with information about needs assessment/outcome tools (e.g. www.decideproject.co.uk) and homepages of dementia or caregiver associations or organizations (e.g. www.alz.org, www.alzheimer-europe.org, www.caregiver.org, www.rosalynnncarter.org).

Seven relevant researchers identified during the literature search were contacted by email to obtain unpublished psychometric studies or testing of the instrument, or to request further or lacking data. Two of them provided the instrument itself to complete the data extraction. In two cases, additional data were delivered to evaluate the methodological quality. One unpublished publication was found by contacting the responsible researcher. Three authors were not able to send additional information. The full search strategy is provided in Appendix I.
Study selection
Following the search, all identified citations were collated and uploaded into EndNote X7.8 (Clarivate Analytics, PA, USA) and duplicates removed. Titles and abstracts were then screened by two independent reviewers for assessment against the inclusion and exclusion criteria for the review. Potentially relevant publications were retrieved in full and their citation details imported into the standardized data extraction tool developed for this review. The full texts of selected citations were assessed in detail against the inclusion and exclusion criteria by two independent reviewers. Reasons for exclusion of full-text studies that did not meet the inclusion criteria are reported in Appendix II. Any disagreements that arose between the reviewers at each stage of the study selection process were either resolved through discussion, or with a third reviewer.

Assessment of methodological quality
Publications selected for retrieval were assessed by two independent reviewers for methodological validity prior to inclusion in the review using the COSMIN checklist. The COSMIN checklist is a standardized tool recommended for use in systematic reviews of measurement properties. This tool fulfills the specific requirements of a psychometric review and has already been successfully used in another JBI review protocol. The original checklist consists of 12 boxes with five to 18 items per box, and is a modular tool. We therefore used only the seven boxes evaluating relevant psychometric properties for our review, namely, internal consistency, reliability, measurement error, content validity, structural validity, hypotheses testing and responsiveness. The box for criterion-related validity was not included as there is no reasonable gold-standard available. The three boxes for additional methodological standards (item response theory models, interpretability and cross-cultural validity) were not used as they focus on more advanced properties. The last box on generalizability was used for data extraction of study characteristics. Each item of the checklist is assessed on a four-point response scale: excellent, good, fair and poor. However, some items have only two or three response options (e.g. only excellent, fair or poor). The methodological quality scores per box were obtained by taking the lowest rating across all items in the box (“worst score count”). The lowest score of any box presented the overall score of the reviewed study. Studies with poor scores in all boxes would have been excluded from the review as this would indicate inadequate methodological quality. In this review, none of the included studies were excluded due to poor scores in all boxes. The checklist used for assessing the methodological quality can be found in the a priori published protocol.

All disagreements that arose between the reviewers were resolved through discussion, or with a third reviewer.

Data extraction
Data were extracted from papers included in the review using the standardized data extraction tool developed for this review. This tool was inspired by the standardized data extraction tools in JBI System for the Unified Management, Assessment and Review of Information (JBI SUMARI; JBI, Adelaide, Australia), by selecting the relevant parts and adapting them to the specificity of a psychometric review. JBI SUMARI is a web application designed to support researchers and practitioners in the entire process of conducting a systematic review. The data extracted included specific details about: i) study characteristics, ii) instrument characteristics and iii) outcomes of significance for the review question and specific objectives. Although the comprehensiveness and accuracy of the provided data varied across studies, the following data were collected:

i) Study characteristics: citation details, aim of the study, study design and method, setting, population characteristics, definition of informal caregivers and needs.

ii) Instrument characteristics: name of the instrument, purpose, target population, application method, administrative burden, number of items and domain structure, range of scores, response options/format.

iii) Psychometric outcomes: reliability (test-retest reliability, inter-rater reliability, internal consistency), validity (content validity, construct validity, structural validity).

Data were extracted and double checked independently by two reviewers (first and second author). To minimize errors, the data extraction form was first pilot tested and a standardized form was used.
Disagreements between the reviewers were resolved by discussion, or with a third reviewer.

**Data synthesis**
The main aim of the data synthesis was to compare outcomes to provide recommendations on the most suitable instrument for research, clinical use and informal caregivers. The findings on instrument characteristics, reliability and validity were compared and described in narrative form, including tables to aid data presentation. The domain structure of the instruments and a summary of their content are presented in a table and in narrative form (see Appendix III and Table 1).

The quality criteria from Terwee et al. were used to judge the psychometric outcomes of the different instruments, namely, their content validity, internal consistency, construct validity, test-retest reliability (agreement), and inter-rater reliability (reliability). The quality of the instruments was assessed as positive, indeterminate, or negative, with a fourth category for “no information available and doubtful design or method.” The results of this appraisal are presented in a narrative form and in Appendix IV.

**Results**

**Study inclusion**
A total of 4909 records were identified through the systematic search in the four databases. Searches in gray literature and requests to relevant researchers for unpublished literature or publications about specific instruments, identified *a priori*, revealed seven additional publications. After removing duplicates, 3468 records remained. Another 3404 records were excluded for not meeting the inclusion criteria after screening their titles or abstracts, resulting in 64 full texts assessed for eligibility. After reading the full texts, 19 articles met the inclusion criteria and were assessed for quality. One publication was excluded due to insufficient data to evaluate the methodological quality. Eighteen articles covering 14 different needs assessment instruments were included in the review with, in four cases, two different publications describing the same instrument, as they provided complementary information. A flow chart of the study selection is presented in Figure 1. The excluded full texts and the reasons for exclusion according to the inclusion criteria or the critical appraisal are listed in Appendix II. Table 2 presents an overview of the included instruments, their acronyms and the authors of the included publications.

**Characteristics of included studies**
Eleven of the 18 included publications focused on the development or the evaluation of an instrument. In addition, a development report and a manual were included to assess the content validity of an instrument or to provide additional results for the psychometric testing. Five other studies, not primarily aimed at validation but containing sufficient information about the development or the evaluation of the used instruments, were also integrated in the review. The publication dates ranged from 1996 to 2019.

Nine of the development or evaluation samples included only caregivers of persons with dementia.

| Table 1: Summary of topics |
|---------------------------|
| **Need for information and education** |
| - Information about dementia and treatment |
| - Nursing skills/care tasks/dementia specific caring skills |
| - Information resources |
| - Formal help/services in the region/community resources |
| - Characteristics, access and availability of services |
| - Organizing care/type of provider |
| **Needs related to emotional support** |
| - Respite support |
| - Family time/shared activities |
| - Mental health counseling/psychiatric care |
| - Informal network/support from family and friends |
| - Support from society |
| - Relationship to person with dementia |
| - Counseling negative emotions |
| - Being a caregiver/assume caregiver role |
| - Sleep |
| **Need for other accessible and appropriate services** |
| - Medical care for caregiver |
| - General assistance or household chores |
| - Financial and legal support |
The remaining nine samples were mixed, including caregivers of persons with dementia residing in the community or living in institutions. Study samples differed in the relationship status, but caregivers were mostly spouses or offsprings of the persons...
with dementia. In all studies, the majority of caregivers were female. The mean age of the caregivers ranged from 51 to 68 years (with a standard deviation of 10 to 14.5 years), and the age range of people with dementia ranged from 76 to 85 years (with a standard deviation of 5.9 to 9.8 years). An overview of the characteristics of the different samples is presented in Appendix V.

Description of instruments

Five of the instruments were developed or tested in the US, five in the United Kingdom (UK), and one each in Austria, Singapore, Greece, Canada, and the Netherlands. Eleven instruments contained only items for caregivers (CADI, RAM, QCNE, UNM, EAC, QNP, PBH-LCI:D, CNCD, NAS, CNA-D, SIDECAR) while three also included items for persons with dementia (CARENAP, JHDCNA, Tayside). Three instruments were recommended for use in both clinical and research settings (CADI, PBH-LCI:D, RAM) while one was recommended only for research purposes (CNA-D), and four others only for clinical assessments (CARENAP, CNCD, EAC, Tayside). For the last six instruments, the intended context of use was not specified (JHDCNA, NAS, QCNE, QNP, SIDECAR, UNM). With the application method, seven instruments were self-administered (CNCD, NAS, PBH-LCI:D, QCNE, EAC, SIDECAR, UNM), three were used in professional interviews (CARENAP, CNA-D, JHDCNA), and two could be either self-administered or professionally interviewed (QNP, Tayside). For two instruments, the application method was not clearly stated, but the descriptions suggested for both a self-reported application method (CADI, RAM). For seven instruments, the administration time was described, ranging from five to 50 minutes (CADI, CARENAP, CNA-D, NAS, RAM, QCNE, QNP). Information about the required training for

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**Table 2: Overview of instruments and authors**

| Acronym | Instrument name | Authors of included studies |
|---------|-----------------|-----------------------------|
| CADI    | Carers Assessment of Difficulties Index | Charlesworth et al. (2007)⁴⁹ |
| CARENAP | The Care Needs Assessment Pack for Dementia | McWalter et al. (1996; 1998)⁵⁰,⁵¹ |
| CNA-D   | The Carers’ Needs Assessment for Dementia | Wancata et al. (2005)⁵²,⁵³ |
| CNCD    | Caregivers’ Needs Checklist for Dementia | Vaingankar et al. (2013; 2018)⁵⁴,⁵⁵ |
| JHDCNA  | The Johns Hopkins Dementia Care Needs Assessment | Hughes et al. (2014)⁵⁶ |
| NAS     | Needs Assessment Survey | Wackerbarth et al. (2002)⁵⁷ |
| PBH-LCI:D | Partnering for Better Health–Living with Chronic Illness: Dementia | Sadak et al. (2015)⁵⁸ |
| RAM     | Risk Appraisal Measure | Czaja et al. (2009)⁵⁹ |
| QCNE    | Questionnaire of Carers Needs Evaluation | Dimakopoulou et al. (2015)⁶⁰ |
| EAC     | Questionnaire on consultation expectations [Echelle d’attentes de consultation (EAC)] | Laprise et al. (2001)⁶¹ |
| QNP     | Questionnaire National Dementia Programme Survey Needs and problems of informal caregivers of persons with dementia | Peeters et al. (2010)⁶², Van der Poel and van Beek (2006)⁶³ |
| SIDECAR | Scales measuring the Impact of DEmentia on CARers | Oyebode et al. (2019)⁶⁴ |
| Tayside | Tayside Profile for Dementia Planning | Gordon et al. (1997)⁶⁵ |
| UNM     | Unmet Need Measure | Gaugler et al. (2004)⁶⁶ |
clinicians was mentioned for two instruments. For one of them, no specific abilities or prior knowledge were necessary (QNP), whereas the other instrument could only be administered by professionals with experience in assessments and interviewing (CARENAP). Response options for all instruments were either nominally or ordinally scaled. For nine instruments, a total (CADI, CNA-D, CNCD, JHDCNA, PBH-LCi:D, RAM, EAC, UNM) or a mean score (NAS) could be obtained, with higher scores indicating higher unmet needs in most of the instruments. Four instruments did not use a scoring system (CARENAP, QCNE, QNP, Tayside). No concrete information was provided for SIDECAR on this aspect. The instruments differed regarding the domain structure and number of items. The number of items for caregivers ranged from 12 to 70 items. Appendix III provides an overview of the domain structure, number of items and response options of the different instruments. Further detailed information about the characteristics of the instruments is presented in Appendix VI.

**Methodological quality**

For each study, we evaluated the methodological quality of their assessment of six different psychometric properties, namely, content validity, structural validity (i.e. the factor structure of the instrument), internal consistency, reliability (including both test-retest and inter-rater), measurement error and construct validity. Criterion validity was not considered as there is currently no gold standard, and responsiveness (i.e. sensitivity to change) was excluded as no study assessed it. Table 3 provides an overview of the quality (excellent, good, fair or poor) of the assessment for each study and each specific standardized question (Q1 and following) regarding the psychometric properties evaluated in this study. For CNA-D, two studies testing construct validity with different variables were available and therefore evaluated separately, thus generating in total 15 studies for methodological quality. In the other three cases where two publications described the same instrument, as they contained complementary information, we treated them as one study when assessing methodological quality.

**Content validity** was documented in 13 of the 15 studies. Seven of these 13 studies had excellent ratings for each of the five specific criteria considered (QCNE, CARENAP, QNP, PBH-LCi:D, CNCD, CNA-D, SIDECAR), and another study had good or excellent ratings (NAS). The five other studies all had at least one poor rating. Three of them failed to include informal caregivers in the item development process (i.e. Q2: RAM, UNM, EAC). Three did not assess if all items together comprehensively reflected the construct to be measured (i.e. Q4: CADI, Tayside, EAC). One did not assess if all items referred to relevant aspects of the construct to be measured (i.e. Q1: CADI). In summary, various examples of excellent methodological quality were available regarding content validity.

**Structural validity** is relevant for instruments aiming at measuring different domains of needs, which was clear for all studies. However, only four studies evaluated the factor structure of their instrument, one with a factor analysis meeting all the criteria (CADI), two with adequate factor analysis but samples of too limited size (PBH-LCi:D, CNCD), and a last one with a factor analysis inadequately performed separately for each dimension (QCNE). In summary, sample size issues and inappropriate statistical analysis limited the strength of the limited evidence regarding structural validity.

**Internal consistency** was reported in 10 of the 15 studies. Information on the percentage of missing data (Q2) and how they were handled (Q3) was provided in only two of these 10 studies (RAM, EAC). The sample size used to assess internal consistency was optimal (Q4, N=100 or more) for seven studies (CADI, RAM, QCNE, UNM, QNP, PBH-LCi:D, CNCD), good for the EAC (50<N<99), moderate for the CNA-D (30<N<49), and poor for CARENAP (N<30). As presented above, four studies tested the unidimensionality of their subscales before computing the Cronbach alphas (Q5), in two cases with an appropriate sample size (Q6: CADI, QCNE) and in two with far too small samples (PBH-LCi:D, CNCD), while six studies did not assess unidimensionality. Seven studies computed the internal consistency statistic for each subscale separately (Q7: CADI, QCNE, UNM, EAC, PBH-LCi:D, CNCD, CNA-D), while three computed a single alpha for all items irrespective of possible dimensions (RAM, CARENAP, QNP). Eight studies calculated an appropriate statistic (Q9 and/or Q10: CADI, RAM, QCNE, UNM, EAC, PBH-LCi:D, CNCD, CNA-D), while two provided only item-total correlations (CARENAP, QNP). In summary, most studies met a majority of criteria for methodological quality, with the
| Citation          | Box                      | Q1  | Q2  | Q3  | Q4  | Q5  | Q6  | Q7  | Q8  | Q9  | Q10 | Q11 | Q12 | Q13 | Q14 |
|-------------------|--------------------------|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|
| JHDCNA Hughes et al. (2014) | Construct validity       | exc | good | exc | good | exc | good | exc | fair | good | -   | -   | -   | -   | -   |
| CADI Charlesworth et al. (2007) | Content validity         | poor | exc  | fair | poor | exc | -   | -   | -   | -   | -   | -   | -   | -   | -   |
|                    | Structural validity      | yes  | exc  | exc  | exc  | exc | exc | exc | exc | exc | na  | -   | -   | -   | -   |
|                    | Internal consistency     | yes  | exc  | exc  | exc  | exc | exc | exc | exc | na  | na  | -   | -   | -   | -   |
| RAM Cjaza et al. (2009) | Content validity         | exc  | poor | exc  | exc  | exc | -   | -   | -   | -   | -   | -   | -   | -   | -   |
|                    | Internal consistency     | yes  | good | fair | exc  | poor | na  | poor | exc | exc | exc | exc | exc | -   | -   |
|                    | Construct validity       | good | fair | exc  | fair | good | exc | good | fair | exc | good | -   | -   | -   | -   |
| QCNE Dimakopoulou et al. (2015) | Content validity         | exc  | exc  | exc  | exc | exc | exc | exc | exc | -   | -   | -   | -   | -   | -   |
|                    | Structural validity      | yes  | good | fair | exc  | poor | na  | -   | -   | -   | -   | -   | -   | -   | -   |
|                    | Internal consistency     | yes  | good | fair | exc  | exc | exc | exc | exc | na  | na  | -   | -   | -   | -   |
| UNM Gaugler et al. (2004) | Content validity         | exc  | poor | exc  | exc  | exc | exc | exc | exc | exc | na  | -   | -   | -   | -   |
|                    | Internal consistency     | yes  | good | fair | exc  | poor | na  | exc | fair | na  | exc | exc | -   | -   | -   |
|                    | Construct validity       | good | fair | exc  | exc  | exc | exc | exc | exc | good | fair | fair | good | -   | -   |
| Tayside Gordon et al. (1997) | Content validity         | exc  | exc  | exc  | poor | fair | -   | -   | -   | -   | -   | -   | -   | -   | -   |
|                    | Reliability              | good | fair | good | exc | fair | poor | poor | poor | na  | exc | na  | na  | na  | na  |
| EAC Laprise et al. (2001) | Content validity         | exc  | poor | exc  | poor | fair | -   | -   | -   | -   | -   | -   | -   | -   | -   |
|                    | Internal consistency     | yes  | exc  | exc | good | poor | na  | exc | exc | exc | na  | na  | -   | -   | -   |
|                    | Reliability              | good | exc  | exc | fair | exc | exc | exc | exc | exc | good | fair | good | fair | exc |
|                    | Construct validity       | good | fair | good | exc | exc | exc | exc | exc | exc | good | n   | -   | -   | -   |
| CARENAP McWalter (1996, 1998) | Content validity         | exc  | exc  | exc  | exc  | exc | exc | exc | exc | exc | -   | -   | -   | -   | -   |
|                    | Internal consistency     | yes  | good | fair | poor | poor | na  | poor | poor | na  | exc | good | -   | -   | -   |
|                    | Reliability              | exc  | fair | poor | exc | exc | exc | exc | exc | poor | na  | exc | na  | na  | na  |
| QNP – Peeters et al. (2010) / Van der Poel and van Beek (2006) | Content validity         | exc  | exc  | exc  | exc | exc | exc | exc | exc | exc | -   | -   | -   | -   | -   |
|                    | Internal consistency     | yes  | good | fair | exc  | poor | na  | poor | poor | good | na  | -   | -   | -   | -   |
| PBH-LCED Sadak et al. (2015) | Content validity         | exc  | exc  | exc  | exc | exc | exc | exc | exc | exc | -   | -   | -   | -   | -   |
|                    | Structural validity      | yes  | good | poor | fair | exc | exc | exc | exc | -   | -   | -   | -   | -   | -   |
|                    | Internal consistency     | yes  | good | fair | exc | poor | exc | poor | exc | exc | na  | poor | -   | -   | -   |
|                    | Reliability              | exc  | good | exc  | good | exc | good | exc | exc | good | exc | exc | na  | na  | na  |
|                    | Measurement error        | good | good | exc  | exc | good | exc | exc | good | exc | -   | -   | -   | -   | -   |
|                    | Construct validity       | exc  | exc  | good | exc | good | exc | exc | good | exc | fair | -   | -   | -   | -   |
| CNCD Vaingankar et al. (2013, 2017) | Content validity         | exc  | exc  | exc  | exc | exc | exc | exc | exc | -   | -   | -   | -   | -   | -   |
|                    | Structural validity      | yes  | good | poor | fair | exc | exc | na  | -   | -   | -   | -   | -   | -   | -   |
|                    | Internal consistency     | yes  | good | fair | exc | poor | exc | poor | exc | exc | na  | na  | -   | -   | -   |
|                    | Construct validity       | good | fair | exc  | exc | good | exc | good | exc | exc | na  | exc | na  | na  | na  |
| NAS - Wackerbarth et al. (2002) | Content validity         | exc  | good | exc  | exc | exc | exc | exc | exc | exc | -   | -   | -   | -   | -   |
| CNA-D Wanacata et al. (2005) | Content validity         | exc  | exc  | exc  | exc | exc | exc | exc | exc | exc | -   | -   | -   | -   | -   |
|                    | Internal consistency     | yes  | good | fair | poor | na  | exc | fair | na  | exc | na  | -   | -   | -   | -   |
|                    | Reliability              | good | fair | exc  | exc | exc | exc | good | exc | good | exc | na  | exc | na  | na  |
|                    | Construct validity       | good | good | fair | good | exc | good | exc | good | exc | -   | -   | -   | -   | -   |
the inter-rater agreement was tested, based on a
by having two interviews conducted by two different
and CARENAP. For CNA-D, test-retest and inter-
CNA-D), while this was not the case for CNA-D
30.5 days. However, this was only the case for
the measurement occasions were clearly or assum-
items performed? Q9. Were there any important flaws
in the design or methods of the study? Q10. Were design and statistical methods adequate for the hypotheses to be tested?
Reliability: Q1. Was the percentage of missing items given? Q2. Was there an assessment of whether all items refer to relevant aspects of the construct to be measured? Q3. Was there an assessment of whether all items together comprehensively reflect the construct to be measured? Q4. Were there any important flaws in the design or methods of the study? Q10. Were design and statistical methods adequate for the hypotheses to be tested?
Convergent validity: Q1. Was the percentage of missing items given? Q2. Was there a description of how missing items were handled? Q3. Was the sample size included in the analysis adequate? Q4. Were at least two measurements available? Q5. Were the administrations independent? Q6. Was the time interval stated? Q7. Were patients stable in the interim period on the construct to be measured? Q8. Was the time interval appropriate? Q9. Were the test conditions similar for both measurements? Q10. Were there any important flaws in the design or methods of the study? Q11. Was there an adequate description of the comparator instrument(s)? Q12. Were the expected absolute or relative magnitude of correlations or mean differences included in the hypotheses? Q7. For convergent validity: Was an adequate description of the comparator instrument(s) adequately described? Q9. Were there any important flaws in the design or methods of the study? Q10. Were design and statistical methods adequate for the hypotheses to be tested?

Table 3. (Continued)

| Citation | Box | Q1 | Q2 | Q3 | Q4 | Q5 | Q6 | Q7 | Q8 | Q9 | Q10 | Q11 | Q12 | Q13 | Q14 |
|----------|-----|----|----|----|----|----|----|----|----|----|-----|-----|-----|-----|-----|
| CNA-D    | Construct validity | good | good | fair | exc | exc | good | good | exc | exc | -   | -   | -   | -   | -   |
| Kaiser et al. (2005) | SIDECAR | Oyebode et al. (2019) | Construct validity | exc | exc | exc | exc | exc | - | - | - | - | - | - | - |
and inter-rater assessment, as well as as irregular or too long time intervals, and modest sample sizes, undermined the limited efforts to evaluate reliability. Measurement error was only reported for one study (PBH-LCI:D), with no information on the percentage of missing data (Q1) and how they were handled (Q2). However, the study had a good sample size of N = 79 (Q3), and used a clear (Q6) and appropriate (Q8) time interval between the two measures (i.e. two weeks), suggesting that the informal dementia caregivers were stable over that period (Q7). It was also assumable that the measurements were independent (Q5) and test conditions were similar (Q9), and appropriate statistics were used (Q11). The very limited evidence about measurement error was thus of good quality.

Construct validity was documented in eight of the 15 studies. Only JHDCNA and PBH-LCI:D provided information on the percentage of missing data (Q1) and how they were handled (Q2). Six studies had optimal or good sample sizes (Q3, N > 50: JHDCNA, RAM, UNM, EAC, PBH-LCI:D, NAS), while CNA-D had a moderate sample in both studies with N = 45. In six studies, multiple hypotheses (PBH-LCI:D, CNCD, CNA-D) or a minimal number of hypotheses (JHDCNA, EAC, CNA-D) were formulated a priori (Q4), while RAM and UNM formulated only vague hypotheses. EAC was the sole study to specify the expected magnitude of the association (Q6). Seven studies described adequately all or most constructs measured by the comparator instruments (Q7), with only UNM delivering a poor description. In contrast, only JHDCNA and CNA-D provided comprehensive evidence of the measurement properties of the comparator instruments (Q8), with all of the others giving only partial information. Finally, all studies used clearly or assumably appropriate statistics, with the exception of PBH-LCI:D which used parametric statistics while the standard deviations for some variables suggested non-normal distributions. In summary, there was some quality empirical evidence on construct validity, although the psychometric properties of the comparator instruments were often insufficiently described.

**Review findings**

**Quality assessment of psychometric properties**

We will first summarize the evidence available for each type of psychometric feature, with the exclusion of criterion validity, which is irrelevant in the absence of a gold standard, and of responsiveness or sensitivity to change, which was never evaluated for the reviewed instruments. We will then briefly discuss the psychometric properties of each instrument. The relevant information is presented in Appendix IV.

Content validity was documented for 13 of the 14 reviewed instruments. As mentioned above, the procedure used to optimize content validity was evaluated satisfactory in seven of these 13 instruments (CARENAP, CNA-D, CNCD, NAS, PBH-LCI:D, SIDECAR, QNP). Most studies generated items based on a literature review and/or an expert consultation, and then reviewed these items in collaboration with experts and at least five informal dementia caregivers. However, the initial pool of the SIDEAR instrument was inductively developed on the basis of 42 interviews with caregivers. Caregivers, researchers and carer consultants were included in further steps of questionnaire development and testing of content validity. For the QNP, the final set of items was further submitted to additional informal dementia caregivers to optimize understandability. For three other instruments, a doubtful design was used (CADI, QCNE, Tayside; see Appendix IV for more details), and the three last instruments failed to include the target population in the process of item development (RAM, EAC, UNM).

Internal consistency was assessed for 10 of the 14 reviewed instruments. For three of these 10 instruments, the Cronbach alphas were computed for dimensions based on the results of a factor analysis. The latter supported a structure with five to eight dimensions (five for CNCD, with three to seven items; seven for PBH-LCI:D, with three to eight items; eight for the CADI, with two to seven items per dimension). However, for CADI, most alphas were below .70 despite an adequate factor analysis, and for CNCD and PBH-LCI:D, the factor analysis had largely insufficient sample sizes but all alphas were above .70. These results are therefore all partly problematic. For four other instruments, Cronbach alphas were provided for each dimension and comprised between .70 and .95, but for three of them no factor analysis was conducted (CNA-D, EAC and UNM), and for QCNE the factor analysis was inadequate; therefore this evidence should be considered inconclusive. Three other instruments reported only...
the alpha for the full scale without considering the dimensions, with low values in for CARENAP and RAM, and a high value for QNP.

**Reproducibility** was evaluated for four of the 14 reviewed instruments in terms of test-retest agreement (CNA-D, PBH-LCI:D, EAC, Tayside), for three in terms of inter-rater reliability (CARENAP, CNA-D, Tayside), and for PBH-LCI:D in terms of measurement error. CNA-D and PBH-LCI:D showed satisfactory test-retest agreement, as evaluated with a proper procedure, with correlations in the .70 range. For two other instruments where the procedure was questionable, the average Kappa was excellent for EAC but varied between poor and excellent for the different subscales of Tayside. As presented above, all the procedures used to evaluate the inter-rater reliability were problematic. For CARENAP, which compared the simultaneous evaluation of an interviewer and an observer, thereby increasing the likelihood of agreement, the Kappas were high. For Tayside, which compared the ratings of a professional to the self-report of the informal caregiver, thereby reducing the chances of agreement, the Kappas were very low. For CNA-D, which evaluated the test-retest and inter-rater reliability in combination, based on two interviews conducted by different persons two weeks apart, the mean Kappa was high. The results for measurement error for PBH-LCI:D were good. Sound evidence regarding reproducibility is therefore still scarce, and limited to test-retest agreement and measurement error.

**Validity** was consistently assessed based on construct validity in seven of the 14 reviewed instruments. For CNA-D, PBH-LCI:D, RAM and EAC, precise a priori hypotheses were formulated and at least 75% of the results were in accordance with them. In contrast, CNCD had less than 75% of its well-formulated hypotheses confirmed. Most hypotheses focused on associations between unmet needs and the caregiver’s objective (e.g. number of hours of care, problem behaviors or functional dependency of the care recipient) or subjective burden, depression or anxiety symptoms or psychological distress, amount of formal or social support received, self-care or quality of life. These postulated associations were either based on plausible links with other common relevant outcomes for informal dementia caregivers (e.g. depression and subjective burden for RAM; subjective burden for CNA-D; burden and psychological morbidity for CNCD), or on different theoretical models, namely, the patient activation model for PBH-LCI:D, and Caplan’s model of mental health consultation for EAC. For JHDCNA and UNM, the formulated hypotheses were too vague and numerous, seeming to pre-empt associations between all needs domains and all outcomes, which resulted in low rates of confirmation. They were based on plausible links with objective and subjective burden for JHDCNA, and Pearlin’s stress process model for UNM. There is thus some evidence for construct validity.

**Best-validated instruments**

Regarding individual instruments, the best-validated one is currently PBH-LCI:D by Sadak et al. PBH-LCI:D was developed using an appropriate procedure regarding content validity, included six domains confirmed in a factor analysis and with good internal consistency, demonstrated adequate test-retest stability after two weeks, and showed the expected correlations with other variables indicating construct validity. EAC (in French) by Laprise et al. had appropriate evidence of test-retest reliability and construct validity, although informal caregivers were not involved in the item development process and Cronbach alphas were computed without a dimensionality analysis. Four other instruments had adequate support for content validity, but insufficient evidence on all other psychometric properties (CARENAP, CNCD, QNP, NAS). The excellent content validity of SIDECA was provided a valid basis for further psychometric testing of this instrument, which seems to be currently underway. The CNA-D and RAM had good evidence for construct validity, but inconclusive evidence for all other properties. The last five instruments currently provide no convincing evidence on any psychometric property (CADI, JHDCNA, QCNE, Tayside, UNM).

**Content and structure of dimensions**

The topics assessed in the instruments can be divided into three thematic groups of needs: i) need for information and education, ii) needs related to emotional support, and iii) need for other accessible and appropriate services. All instruments contained at least one item in the first and second group. Ten instruments comprised items in the third group. The first group of needs was typically assessed with items about the need for education regarding care tasks, especially dementia-specific caring skills (included in
all instruments), items about information on local services or community resources for persons with dementia or caregivers (included in 13 instruments). Needs for information about dementia and its treatment were present in 10 instruments, and needs in relation to the characteristics, accessibility and availability of services were included in nine instruments. In the second thematic group, the most considered topics were counseling for negative emotions (nine instruments); support from the informal network such as family, friends or other caregivers (nine instruments) or society (eight instruments); and respite (eight instruments). Financial and legal support (seven items) was the most common topic in the third group of needs. Table 1 provides a summary of the topics assessed in the different instruments.

Discussion

This psychometric literature review identified 14 needs assessment instruments for informal dementia caregivers with empirically evaluated measurement properties. Their systematic evaluation, based on the COSMIN criteria, revealed that half of them had excellent content validity. In contrast, the structure validity was rarely examined, and factor analyses were in most cases of low quality because of insufficient sample sizes or questionable procedures. None of the instruments had an optimally tested and good internal consistency, as the sole one with an adequate dimensionality analysis had low alphas, and all others had either their dimensionality evaluated with small samples, or no proper evaluation of their dimensionality – although some had high alphas. Regarding reliability, test-retest agreement was rarely tested, and only two instruments used a satisfactory procedure and obtained good correlations. Inter-rater agreement was relevant only for professionally interviewed instruments and evaluated using inadequate procedures. Regarding validity, in the absence of a gold standard, no criterion validity could be assessed, but construct validity was evaluated in more than half of the instruments, with satisfactory procedures and results on four of them. Comparing the different instruments reviewed, PBH-LCID was rated as having the best psychometric evidence, and EAC was also partly supported, while most other instruments had limited or no proof of their psychometric soundness. SIDECAR showed very promising results regarding content validity. However, further psychometric testing is needed and seems to be currently underway.

This overview highlighted the importance of guidelines such as COSMIN in guiding the efforts in the development of instruments with optimal psychometric properties in a specific field. As we noted for needs assessment in informal dementia caregivers, despite international investments in instrument development, optimal standards were on many occasions not achieved due to problematic procedures and analyses, as well as largely insufficient sample sizes. Our review therefore provides essential information to inform future efforts in the development of such measures in order to achieve more robust psychometric results.

Regarding needs assessment in informal dementia caregivers, our review showed that there are presently several instruments with adequate content validity, developed in diverse countries (Singapore, Austria, the Netherlands, the UK and the US). This information provides an excellent starting point for further development. The priority should now be to identify the structure, in terms of the number of different domains required to cover the diverse needs of informal dementia caregivers. The preliminary factor-analytic evidence that we reviewed identified five to eight domains of needs, but overall the instruments reviewed comprised two to 18 different need domains. Currently, there is no established theoretical model to organize the diverse and complex needs of our population of interest. Pini et al. made a first attempt in this direction by developing a needs-led framework based on qualitative interviews to conceptualize the impact of caring on the lives of family caregivers. A robust theoretical framework might enhance the conceptual clarity around the assessment of these needs, and could be further informed by explorations of the factorial structure with sufficient sample sizes. Nevertheless, cultural differences in the experience of dementia caregiving could affect the content of needs assessment instruments. Such differences were difficult to evaluate at the stage of content validity. They should be examined at later stages of instrument validation by testing the measurement invariance.

Our review also identified different approaches that were applied to assess the construct validity of some of the reviewed instruments. These approaches involved testing the associations between needs and other related constructs, predominantly the informal
caregiver’s objective or subjective burden, psychological symptoms, the amount of support received, self-care or quality of life. The postulated associations were mostly based on plausible links with common outcomes for informal dementia caregivers, and sometimes on a theoretical model, with different models being used. The diversity of these procedures indicated that further conceptualization of the place of needs within relevant theoretical models could strengthen the nomological net and thereby support more solid examination of the construct validity. Another challenge in this area is to ensure that the content of the items of the different instruments are not overlapping, to prevent spurious correlations. This would typically be the case for needs assessment including items about burden or depression, for example, when their construct validity is tested with instruments measuring these constructs.

Regarding reliability, test-retest stability was scarcely assessed. And inter-rater agreement was always tested with questionable procedures, despite its importance as multiple professionals are normally involved with informal dementia caregivers. Substantial efforts need to be invested in these aspects, but we also have to keep in mind that their assessment is challenging in a fragile population such as informal dementia caregivers. Indeed, a short time interval is required as the situation of the person with dementia and the larger context can change quickly, thereby modifying the needs of the caregiver. Yet it is certainly difficult to obtain two assessments within one or two weeks from chronically stressed and often exhausted caregivers.

Finally, needs assessments were mainly used for evaluations at one specific point in time. Increasingly, the assessment of needs might be considered as an outcome to analyze the impact of interventions, or used to document the evolution of needs over time. The use of needs assessment as an outcome or longitudinal measure requires a satisfactory sensitivity to change, which was not assessed in the reviewed instruments, and warrants scientific attention in the future.

This review has both strengths and limitations. First, our use of a highly structured procedure based on the COSMIN criteria is a strength. However, COSMIN uses a very stringent evaluation of the reviewed instruments, as the final appreciation for each psychometric property is based on the lowest grade across all specific criteria for this property. Although the resulting synthesis could give the impression of a globally poor quality of evidence, we believe it is very helpful in the process of achieving high psychometric standards. Secondly, we were able to include instruments and articles in more diverse languages than previously published reviews, although our limitation on our language skills did not allow us to examine publications in any Asian or Arabic languages. Thirdly, despite our efforts, we could not always access gray literature, as some authors did not respond, persons in charge of the project were absent, or authors were in a commercial process preventing them to provide access to the manual. Our conclusions are therefore limited to available information.

**Conclusion**

**Recommendations for practice**

This review revealed several instruments for measuring the needs of informal dementia caregivers. However, the evidence for their use in the clinical or research setting is often limited. The two best-validated instruments are PBH-LCI:D by Sadak et al. and EAC by Laprise et al. While PBH-LCI:D is in English and intended for use in clinical and research settings, EAC is in French and recommended only for clinical use. Both of them include a scoring system that allows for comparison of results at different time points and between different informal caregivers. PBH-LCI:D and EAC contain items covering the most common topics found across all instruments, namely, the need for information and education, needs related to emotional support and the need for other accessible and appropriate services. Their self-administered application method suggests less effort in administration time for professionals. Nevertheless, there is no information regarding administration burden for both, which would be important in terms of usability, especially in the clinical setting. In regards to well-documented cultural differences in the experience of dementia caregiving, and the absence of empirical evidence on the measurement invariance of the available instruments across diverse cultural groups, caution should be taken in using them in cultural contexts different from those in which the instruments were developed.

**Recommendations for research**

Although we identified moderate to high evidence of strong psychometric properties for PBH-LCI:D and
EAC, this review highlights the need for further developments in the field of needs assessment in informal dementia caregivers, particularly in structural validity and construct validity, as well as test-retest reliability and sensitivity to change. The evaluation of both forms of validity would certainly benefit from a more robust theoretical framework about the core dimensions of needs in informal dementia caregivers, and the relationship between needs and other relevant outcomes for this population. We also need to identify appropriate procedures to assess test-retest reliability with minimal additional burden for informal dementia caregivers, and to evaluate sensitivity to change appropriately despite this involving a demanding procedure.

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43. Biomedical Information of the Dutch Library Association. COSMIN checklist. Qual Life Res 2012;21(4):651–7.
44. COSMIN checklist. Qual Life Res 2012;21(4):651–7.
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### Appendix I: Search strategy

#### Embase

**Search on 21 Feb 2019**

| Search | Query | Records retrieved |
|--------|-------|-------------------|
| #1     | ('needs'/exp OR needs:ab,ti) | 1340 |
| #2     | ('caregiver'/exp OR 'caregiver' OR 'family'/exp OR 'caregiver OR 'family':ab,ti) AND ('alzheimer disease'/exp OR 'senile dementia'/de OR 'presenile dementia'/exp OR 'multinifarct dementia'/exp OR 'diffeus lewy body disease'/exp OR 'frontotemporal dementia'/exp OR 'alzheimer' OR 'dementia' OR 'amnesia'/ti,ab)) |  |
### SYSTEMATIC REVIEW

S. Kipfer and S. Pihet

**MEDLINE via Pubmed**

Searched on 21 Feb 2019

| Search | Query | Records retrieved |
|---|---|---|
| #1 | (need[Title/Abstract]) | |
| #2 | (caregiver[tiab] OR informal carer[tiab] OR Family[Mesh] OR familial[tiab] OR caregivers[Mesh] OR relative[tiab] OR spouse[tiab] OR child[tiab] AND (Dementia[Mesh] OR Alzheimer Disease[Mesh] AND Alzheimer[tiab] OR dementia[tiab] OR amnésia[tiab]) | 1620 |
| #3 | (assess[tiab] OR evaluate[tiab] OR outcome[tiab] OR identif[tiab] OR index[tiab] OR indices[tiab] OR instrument[tiab] OR inventory[tiab] OR measure[tiab] OR questionnaire[tiab] OR profile[tiab] OR scale[tiab] OR score[tiab] OR status[tiab] OR survey[tiab] OR test[tiab]) | |
| #4 | #1 AND #2 AND #3 AND ((instrumentation[sh] OR methods[sh] OR Validation Studies[pt] OR Comparative Study[pt] OR Psychometrica[MeSH] OR psychometry[tiab] OR clinometer[tiab] OR clinimetric[tiab] OR outcome assessment AND (health care) AND MeSH OR outcome assessment[tiab] OR outcome measure[tiab] OR observer variation[MeSH] OR observer variation[tiab] OR Health Status Indicators[Mesh] OR reproducibility of results[MeSH] OR reproducibility[tiab] OR discriminant analysis[MeSH] OR reliable[tiab] OR valid[tiab] OR valid[tiab] OR coefficient of variation[tiab] OR coefficient[tiab] OR homogeneity[tiab] OR homogeneous[tiab] OR internal consistency[tiab] OR (cronbach[tiab] AND alpha[tiab] OR alpha[tiab])) OR (item[tiab] AND (correlation[tiab] OR selection[tiab] OR reduction[tiab]))) OR agreement[tiab] OR precision[tiab] OR imprecision[tiab] OR precise values[tiab] OR test-retest[tiab] OR test[tiab] AND retest[tiab]) | |
| #5 | #4 NOT (addresses[Publication Type] OR biography[Publication Type] OR case reports[Publication Type] OR comment[Publication Type] OR directory[Publication Type] OR editorial[Publication Type] ORfestschrift[Publication Type] OR interview[Publication Type] OR lectures[Publication Type] OR legal cases[Publication Type] OR legislation[Publication Type] OR letter[Publication Type] OR news[Publication Type] OR newspaper article[Publication Type] OR patient education handouts[Publication Type] OR popular works[Publication Type] OR congresses[Publication Type] OR consensus development conference[Publication Type] OR.nih[Publication Type] OR practice guideline[Publication Type] OR NOT (animals[MeSH Terms] OR NOT humans[MeSH Terms]) | |

Publication time: no limitation / Language: no limitation
### CINAHL via EBSCOhost

Searched on 21 Feb 2019

| Search | Query                                                                 | Records retrieved |
|--------|----------------------------------------------------------------------|-------------------|
| #1     | (AB need) OR (TI need)                                                | 816               |
| #2     | (TI carer* OR informal carer* OR famil* OR caregiv* OR relative* OR spouse* OR child*) OR AB (carer* OR informal carer* OR famil* OR caregiv* OR relative* OR spouse* OR child*) OR (MH “family”) OR (MH “caregivers”) AND (MH “Dementia-”) OR (MH “Dementia, Presenile-”) OR (MH “Dementia, Senile-”) OR (MH “Dementia, Vascular-”) OR (MH “Dementias, Multis-Infarct”) OR (MH “Frontotemporal Lobal Degeneration-”) OR (MH “Frontotemporal Dementia-”) OR TI (alzheimer* OR dementia* OR amening*) OR AB (alzheimer* OR dementia* OR amening*) | 1750788          |
| #3     | TI (assess* OR evaluate* OR outcome OR outcomes OR identify* OR index OR indices OR instrument OR instruments OR interview* OR inventory OR measure OR measures OR questionnaire OR questionnaires OR profile OR profiles OR scale OR scales OR score OR scores OR status OR survey OR surveys OR test*) OR AB (assess* OR evaluate* OR outcome OR outcomes OR identify* OR index OR indices OR instrument OR instruments OR interview* OR inventory OR measure OR measures OR questionnaire OR questionnaires OR profile OR profiles OR scale OR scales OR score OR scores OR status OR survey OR surveys OR test*) | 3013866          |
| #4     | #1 AND #2 AND #3 AND (MH “Psychometrics”) OR (TI psychometr* OR AB psychometr*) OR (TI clinometr* OR AB clinometr*) OR (MH “Outcome Assessment”) OR (TI outcome assessment OR AB outcome assessment) OR (TI outcome measure* OR AB outcome measure*) OR (MH “Health Status Indicators”) OR (MH “Reproducibility of Results”) OR (MH “Discriminant Analysis”) OR (TI reproducib* OR AB reproducib*) OR (TI relia* OR AB relia*) OR (TI unreliab* OR AB unreliab*) | 5837457          |
| #5     | #4 NOT (PT “addresses” OR PT “biography” OR PT “case reports” OR PT “comment” OR PT “directory” OR PT “editorial” OR PT “festschrift” OR PT “interview” OR PT “lectures” OR PT “legal cases” OR PT “legislation” OR PT “letter” OR PT “news” OR PT “newspaper article” OR PT “patient education handout” OR PT “popular works” OR PT “congresses” OR PT “consensus development conference” OR PT “consensus development conference, nh” OR PT “practice guideline”) NOT (MH “animals”) NOT (MH “humans”) | 12867290         |

Publication time: no limitation / Language: no limitation
## PsycINFO via OVID

### Searched on 3 Mar 2019

| Search | Query | Records retrieved |
|--------|-------|-------------------|
| #1     | ((need:ab. or need:ti. or needs:sh.) | 1133 |
| #2     | ((carer’ or informal carer’ or famili’ or caregiv’ or relative’ or spouse’ or child’).ab. or (carer’ or informal carer’ or famili’ or caregiv’ or relative’ or spouse’ or child’).ti. or caregivers:sh. or family:sh.) AND (exp Dementia/ or exp Dementia with lewy bodies/ or exp Presenile Dementia/ or exp Semantic Dementia/ or exp Senile Dementia/ or exp Vascular Dementia/ or exp Alzheimer’s Disease/ or (Alzheimer’ or dementia’ or ameni’).mp..ab. or (Alzheimer’ or dementia’ or ameni’).mp..t.) | |
| #3     | (assess’ OR evaluate’ OR outcome OR outcomes OR identif’ OR index OR indices OR instrument OR instruments OR interview’ OR inventory OR measure OR measures OR questionnaire OR questionnaires OR profile OR profiles OR scale OR scales OR score OR scores OR status OR survey OR surveys OR test’).ab. OR (assess’ OR evaluate’ OR outcome OR outcomes OR identif’ OR index OR indices OR instrument OR instruments OR interview’ OR inventory OR measure OR measures OR questionnaire OR questionnaires OR profile OR profiles OR scale OR scales OR score OR scores OR status OR survey OR surveys OR test’).ti. OR (exp outcome/ OR exp outcomes/ OR exp index/ OR exp instrument/ OR exp instruments/ OR exp inventory/ OR exp questionnaire/ OR exp questionnaires/ OR exp profile/ OR exp scale/ OR exp score/ OR exp survey/ OR exp surveys/) | |
| #4     | #1 AND #2 AND #3 AND (Validation Studies.ti. or Comparative Study.ab.) OR (exp Psychometrics/) OR (psychometr’).ti.ab. OR (clinimetr’).tw. OR (exp Psychometrics/ or exp Rating Scales/ or exp Test Constructions/ or exp Measurement/ or exp Test Validity/ or outcome assessment).ti.ab. OR (outcome measure’).tw. OR (observer variation).ti.ab. OR (Health Status Indicators).ti.ab. OR (reproducib’).ti.ab. OR (Discriminant Analysis’).ti.ab. OR (relabil’ or unrelabil’ or valid’ or coefficient or homogeneity or homogeneous or ‘internal consistency’).ti.ab. OR (cronbach’ and (alpha or alphas)).ti.ab. OR (item and (correlation’ or selection’ or reduction’)).ti.ab. OR (agreement or precision or imprecision or ‘precise values’ or ‘test-retest’).ti.ab. OR (test and retest).ti.ab. OR (relabil’ and (test or retest’)).ti.ab. OR (stability or interrater or inter-rater or intrarater or intrarater or intertester or inter-tester or intratester or intra-tester or interobserver or inter-observer or intraobserver or intraobserver or intertechnician or inter-technician or intratechnician or intra-technician or interexaminer or inter-examiner or intraexaminer or intra-examiner or interassay or interassay or intra-assay or intra-assay or interindividual or inter-individual or intraindividual or intra-individual or interparticipant or inter-participant or intra-participant or kappa or kappas or repeatab’).ti.ab. OR (replicab’ or repeated) and (measure or measures or findings or result or results or test or tests’).ti.ab. OR (generaliz’ or generalisa’ or concordance’).ti.ab. OR (intraclass and correlation’).ti.ab. OR (discriminative or known group or factor analysis or factor analyses or dimension’ or subscale’).ti.ab. OR (item discriminant or interscale correlation’ or error or errors or individual variability’).ti.ab. OR (valid’ or variability and (analysis or values’)).ti.ab. OR (uncertainty and (measurement or measuring’)).ti.ab. OR (standard error of measurement or sensitiv’ or responsiveness’).ti.ab. OR (((minimal or minimally or clinical or clinically) and (important or significant or detectable) and (change or difference’)).ti.ab. OR ((|small’ or (real or detectable) and (change or difference’)).ti.ab. OR (meaningful change’).ti.ab. OR (ceiling effect’)).ti.ab. OR (floor effect’).ti.ab. OR (item response model’).ti.ab. OR (IRT’).ti.ab. OR (rasch’).ti.ab. OR (Differential item functioning’).ti.ab. OR (DIF’).ti.ab. OR (computer testing’).ti.ab. OR (item bank’).ti.ab. OR (cross-cultural equivalence’)).ti.ab. | |
| #5     | #4 NOT (‘addresses’it or ‘biography’it or ‘case reports’it or ‘comment’it or ‘directory’it or ‘editorial’it or ‘testeschrift’it or ‘interview’it or ‘lectures’it or ‘legal cases’it or ‘legislation’it or ‘letter’it or ‘news’it or ‘newspaper article’it or ‘patient education handout’it or ‘popular works’it or ‘congresses’it or ‘consensus development conference’it or ‘consensus development conference’it or ‘nih’it or ‘practice guideline’it).mp. not (‘animals’/exp not ‘humans’/exp) [mp = title, abstract, heading word, table of contents, key concepts, original title, tests & measures] | |

Publication time: no limitation / Language: no limitation
### Appendix II: Excluded studies

**Articles ineligible following full-text review**

| Reference                                                                                                                                                                                                 | Reason for exclusion |
|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|----------------------|
| Reynolds T, Thornicroft G, Abas M, Woods B, Hoe J, Leese M, et al. Camberwell Assessment of Need for the Elderly (CANE). Development, validity and reliability. Br J Psychiatry 2000;176:444-452.                          | Criteria 1           |
| Nicolaou PL, Egan SJ, Gasson N, Kane RT. Identifying needs, burden, and distress of carers of people with frontotemporal dementia compared to Alzheimer’s disease. Dementia 2010;9(2):215-235.                      | Criteria 1, 4        |
| van der Roest HG, Meiland FJM, Comijs HC, Derksen E, Jansen APD, van Hout HPJ, et al. What do community-dwelling people with dementia need? A survey of those who are known to care and welfare services. Int Psychogeriatr 2009;21(5):949-65. | Criteria 1, 4        |
| Orrell M, Hancock GA, Liyanage KCG, Woods B, Challis D, Hoe J. The needs of people with dementia in care homes: the perspectives of users, staff and family caregivers. Int Psychogeriatr 2008;20(5):941-51.                  | Criteria 1, 2, 4     |
| Miranda-Castillo C, Woods B, Orrell M. The needs of people with dementia living at home from user, caregiver and professional perspectives: a crosssectional survey. BMC Health Serv Res 2013;13:43.                                                      | Criteria 1, 4        |
| Fernandes L, Goncalves-Pereira M, Leuschner A, Martins S, Sobral M, Azevedo LF, et al. Validation study of the Camberwell Assessment of Need for the Elderly (CANE) in Portugal. Int Psychogeriatr 2009;21(1):94-102.             | Criteria 1           |
| Ducharme F, Kergoat M-J, Coulombe R, Lévesque L, Antoine P, Pasquier F. Unmet support needs of early-onset dementia family caregivers: a mixed design study. BMC Nurs 2014;13(1):49.                                              | Criteria 2, 4        |
| Ducharme F, Lévesque L, Caron C, Hanson E, Magnusson L, Nolan J, et al. Testing of a tool to assess the support needs of family caregivers of an elderly relative at home: A tool with potential for application in French-speaking Europe. [Mise à l’essai d’un outil d’évaluation des besoins de soutien des proches-aidants d’un parent âgé à domicile : Un outil ayant un potentiel d’application en Europe francophone]. Rech soins infirm 2010;101(2), 67-80. French. | Criteria 2           |
| Hanson E, Nolan J, Magnusson L, Sennemark E, Johansson L, Nolan M. COAT: The Carers Outcome Agreement Tool: a new approach to working with family carers. Getting Research into Practice (GRiP) Report No 1. Project Report. Sheffield: University of Sheffield; 2006. | Criteria 2, 4        |
| Toye C, Lester L, Popescu A, McInerney F, Andrews S, Robinson AL. Dementia Knowledge Assessment Tool Version Two: development of a tool to inform preparation for care planning and delivery in families and care staff. Dementia 2014;13(2):248-56. | Criteria 1           |
| Scholzel-Dorenbos CJ, Arons AM, Wammes JJ, Rikkert MG, Krabbe PF. Validation study of the prototype of a disease-specific index measure for health-related quality of life in dementia. Health Qual Life Outcomes 2012;10:118                      | Criteria 1           |
### Reference

| Reference | Reason for exclusion |
|-----------|----------------------|
| Smith SC, Lamping DL, Banerjee S, Harwood R Foley B, Smith P, et al. Measurement of health-related quality of life for people with dementia: development of a new instrument (DEMQOL) and an evaluation of current methodology. Health Technol Assess 2005;9(10):1-93 | Criteria 1 |
| Cooper C, Katona C, Livingston G. Validity and reliability of the brief COPE in carers of people with dementia: the LASER-AD Study. J Nerv Ment Dis 2008:196(11):838-43. | Criteria 1 |
| Stephan A, Mayer H, Renom Guiteras A, Meyer G. Validity, reliability, and feasibility of the German version of the Caregiver Reaction Assessment scale (G-CRA): a validation study. Int Psychogeriatr 2013;25(10):1621-8. | Criteria 1 |
| Chow TW, Pio FJ, Rockwood K. An international needs assessment of caregivers for frontotemporal dementia. Can J Neurol Sci 2011;38(5):753-7. | Criteria 1,4 |
| Ewing G, Brundle C, Payne S, Grande G. The carer support needs assessment tool (CSNAT) for use in palliative and end-of-life care at home: a validation study. J Pain Symptom Manage 2013;46(3):395-405. | Criteria 2 |
| Leong J, Madjar I, Fiveash B. Needs of family carers of elderly people with dementia living in the Community. Australas J Ageing 2001;20:133-138. | Criteria 1,4 |
| George LK, Fillenbaum GG. OARS methodology. A decade of experience in geriatric assessment. J Am Geriatr Soc 1985;33(9):607-15. | Criteria 1 |
| Francis GM, Munjas BA. Needs of family caregivers and persons with Alzheimer’s disease. Am J Alzheimers Dis 1992;7(4): 23–31. | Criteria 1 |
| Jennings LA, Reuben DB, Everton LC, Serrano KS, Ercoli L, Grill J, et al. Unmet needs of caregivers of individuals referred to a dementia care program. J Am Geriatr Soc 2015;63(2):282-9. | Criteria 1,4 |
| Fortinsky RH, Kercher K, Burant, CJ. Measurement and correlates of family caregiver self-efficacy for managing dementia. Aging Ment Health 2002;6(2):153-60. | Criteria 1 |
| Borson S, Scanlan JM, Sadak T, Lessig M, Vitaliano P. Dementia Services Mini-Screen: a simple method to identify patients and caregivers in need of enhanced dementia care services. Am J Geriatr Psychiatry 2014;22(8):746-55. | Criteria 1 |
| Roud H, Keeling S, Sainsbury R. Using the COPE assessment tool with informal carers of people with dementia in New Zealand. N Z Med J 2006;119(1237):U2053. | Criteria 1 |
| Mizoguchi T, Iijima S, Niino N, Orimo H. Reliability and validity of a Japanese version of the cost of care index. Nihon Ronen Igakkai Zasshi 1995;32(6):403-9. | Criteria 1,2 |
| Pérez-Fuentesa M, Gámez Linaresa JJ, Ruiz Fernández DM, Molero Juradoa M. Inventory of overburden in alzheimer’s patient family caregivers with no specialized training. Int J Clin Health Psychol 2017;17(1):56-64. | Criteria 1 |
| Givens JL, Jones RN, Mazor KM, Prigerson HG, Mitchell SL. Development and psychometric properties of the family distress in advanced dementia scale. J Am Med Dir Assoc 2015;16(9):775-80. | Criteria 2 |
### Reference

| Reference                                                                 | Reason for exclusion |
|--------------------------------------------------------------------------|----------------------|
| Rosness TA, Haugen PK, Gausdal M, Gjora L, Engedal K. Carers of patients with early-onset dementia, their burden and needs: a pilot study using a new questionnaire - Care-EOD. Int J Geriatr Psychiatry 2012;27(10):1095-6. | Criteria 4           |
| Keady J, Nolan M. Behavioural and Instrumental Stressors in Dementia (BISID): refocussing the assessment of caregiver need in dementia. J Psychiatr Ment Health Nurs 1996;3(3):163-72. | Criteria 1           |
| Sato S, Kazui H, Shimizu Y, Yoshida T, Yoshiyama K, et al. Usefulness of carer-held records to support informal caregivers of patients with dementia who live at home. Psychogeriatrics 2018;18(3):166-174. | Criteria 1,4         |
| Cova I, Travi N, Maggiore L, Cucumo V, Mariani C, et al. What are the caregivers’ needs on dementia care? An integrated qualitative and quantitative assessment. Neurol Sci 2018;39(6):1085-1091. | Criteria 4           |
| Aoun, SM, Toye C, Slatyer S, Robinson A, Beattie E. A person-centred approach to family carer needs assessment and support in dementia community care in Western Australia. Health Soc Care Community 2018;26(4):e578-e586. | Criteria 4           |
| Bass DM, Judge KS, Snow AL, Wilson NL, Morgan R, Looman WJ, et al. Caregiver outcomes of partners in dementia care: effect of a care coordination program for veterans with dementia and their family members and friends. J Am Geriatr Soc 2013;61(8):1377-86. | Criteria 4           |
| Bass DM, Judge KS, Snow AL, Wilson NL, Looman WJ, McCarthy C, et al. Negative caregiving effects among caregivers of veterans with dementia. Am J Geriatr Psychiatry 2012;20(3):239-47. | Criteria 4           |
| Bass DM, Judge KS, Snow AL, Wilson NL, Morgan RO, Maslow K, et al. A controlled trial of partners in dementia care: veteran outcomes after six and twelve month. Alzheimers Res Ther 2014;6(1):9. | Criteria 4           |
| Amieva H, Rullier L, Bouisson J, Dartigues J-F, Dubois O, Salamon R. Needs and expectations of Alzheimer’s disease family caregivers. Rev Epidemiol Sante Publique 2012;60(3):231-8. | Criteria 4           |
| Zegelin A, Schnepf W, Riesner C, Herder K. Brief report: Review of the applicability, practicability and modalities of implementing the CarenapD assessment for the German outpatient dementia sector [Kurzbericht: Überprüfung der Anwendbarkeit, Praktikabilität und Modalitäten der Implementierung des Assessments CarenapD für den deutschen ambulanten Sektor bei Demenz] Hannover: Pflegebibliothek - Wittener Schriften; Schlütersche Verlagsgesellschaft; 2010. German. | Criteria 4           |
| Riesner C. The role of family carers of people with dementia in evaluating needs, using the Carenap study as an example [Die Rolle pflegender Angehöriger von Menschen mit Demenz in der Bedarfsbestimmung am Beispiel der Carenap-Studie.] Pflege 2014;27(4):243-255. German. | Criteria 4           |
| Zwaanswijk M, Peeters JM, van Beek AP, Meerveld JH, Francke AL. Informal caregivers of people with dementia: problems, needs and support in the initial stage and in subsequent stages of dementia: a questionnaire survey. Open Nurs J 2013;7:6-13. | Criteria 4           |
### Inclusion criteria recommended by the COSMIN guidelines for systematic reviews of measurement properties:

1. The instrument should aim to measure the construct of interest (types of intervention(s)/phenomena of interest).
2. The study sample should concern the target population of interest (types of participants).
3. The study should concern the type of measurement instrument of interest (self-reported or professionally interviewed).
4. The aim of the study should be the development of a measurement instrument or the evaluation of one or more of its measurement properties (types of studies).

#### Article excluded on critical appraisal

| Reference | Reason for exclusion |
|-----------|----------------------|
| Hodgson C, Higginson I, Jefferys P. Carers’ checklist: an outcome measure for people with dementia and their carers. London: The mental health foundation; 1998. | Not enough information for critical appraisal; development report not available. |
## Appendix III: Domain structure, number of items, and response options

| Instrument acronym | Domain structure/sections (number of items) | response options/format | Range of scores |
|--------------------|---------------------------------------------|-------------------------|-----------------|
| **CADI**<br>Charlesworth et al. (2007)⁵⁷ | 8 dimensions (30 items):<br>1. Carer’s reaction to caring (7 items)<br>2. Degree of physical help (4 items)<br>3. Carer–PwD relationship (4 items)<br>4. Restrictions on social life (4 items)<br>5. Professional support (3 items)<br>6. Family support (3 items)<br>7. Interpersonal demands (3 items)<br>8. Financial consequences (2 items) | The frequency of experiences was rated on a three-point scale<br>1. ‘never applies’<br>2. ‘sometimes applies’<br>3. ‘always applies’ | Total range from 30 to 90 |
| **CARENAP**<br>McWalter (1996/1998)⁵⁰,⁵¹ | Instrument contains 4 sections:<br>1. Basic and referral information<br>2. Assessments for the PwD (7 dimensions): health and mobility, self-care and toileting, social interaction, thinking and memory, behaviour and mental state, house-care, and community living<br>3. Assessments for the carer (5 dimensions):<br>   - Health<br>   - Daily difficulties<br>   - Support<br>   - Rest<br>   - Feelings<br>4. Personal history | Response option for carers: yes or no<br>Response options for PwD: 'No Need' - they have no difficulty or they manage with no help at all; 'Met Need' - they have difficulties but they manage with the help they are getting; 'Unmet Need' - they need help or more help than they currently get<br>Types of help for unmet needs: social stimulation/activity, prompting/supervision, doing tasks for the person, aids and adaptations, specialist assessment, counseling for the person, behavior management, carer advice/training, don’t know | No scoring |
| **CNA-D**<br>Wancata et al. (2005)⁵² / Kaiser et al. (2005)⁵³ | 18 problem areas and 2 - 6 possible interventions per problem area<br>1. Lack of information about dementia<br>2. Lack of information about treatment<br>3. Lack of information about services<br>4. Financial burden<br>5. Legal issues<br>6. Disappointment caused by the illness, concerns about the patient’s future<br>7. Communication problems and conflicts with the patient<br>8. Burdened by behavioral problems of the patients<br>9. Problems caused by crises<br>10. Not enough time for oneself (including caring for the relative becomes sick)<br>11. Social isolation, conflicts within the family<br>12. Burden caused by dangerous situations<br>13. Fear of stigmatization and discrimination<br>14. Feelings of guilt, being blamed<br>15. Missing nursing skills<br>16. Difficulties concerning household tasks<br>17. Burned out or overstrained by care<br>18. Physical or psychiatric illness of the carer | 1. Problem areas (three-point scale): no, mild, moderate or serious problem.<br>2. Interventions (five-point scale): No need - intervention not needed and not received, Overprovision - intervention not needed, but received, Unmet need - intervention needed, but not received, Partially met need - intervention needed and received, but insufficiently, Met need - intervention needed and sufficiently received | No scoring |
| **CNCD**<br>Vaingankar et al. (2013, 2017)⁵⁴,⁵⁵ | 5 dimensions (26 items):<br>1. Information (7 items)<br>2. Access to services (3 items)<br>3. Availability of services (6 items)<br>4. Other formal support services (4 items)<br>5. Informal support (6 items) | Caregivers agreement rated on a 5-point response scale: I did not need ‘X’ / Strongly agree / Agree / Disagree / Strongly disagree | Response scale combined to a 3-point scale for analysis: I did not need ‘X’ = 0 / Strongly agree/ Agree = 1 // Disagree/ Strongly disagree = 2),<br>Range: 0 - 52<br>Total score obtained by adding the item scores, with higher scores indicative of higher unmet need |
| Instrument acronym | Domain structure/sections (number of items) | response options/format | Range of scores |
|-------------------|--------------------------------------------|-------------------------|-----------------|
| **JHDCNA**
Hughes et al. (2014) | 4 caregiver domains (12 items):
   - p. Caregiver Education (3)
   - q. Resource Referrals (5)
   - r. Caregiver Mental Health Care (4)
   - s. Caregiver General Medical / Health Care (3)
15 care recipient domains (77 items):
   - a. Dementia Evaluation / Diagnosis (6)
   - b. Treatment of Cognitive Symptoms (2)
   - c. Treatment of Neuropsychiatric Symptoms (5)
   - d. Behavior Management (3)
   - e. Medication Management
   - f. Medication Administration (3)
   - g. General Medical / Health Care (8)
   - h. Allied Health & Nursing Care (5)
   - i. Safety (9)
   - j. Assistance with Daily Activities (10)
   - k. Meaningful Activities (6)
   - l. Legal Issues / Advance Care Planning (6)
   - m. Assistance with Health Insurance (5)
   - n. Patient Education (1)
   - o. Caregiver Availability (3) | 1) Clinicians rated items as being needed or not and if needed:
2) whether the need was “unmet” if: (a) not addressed and potentially beneficial interventions are available or (b) addressed but potential benefits of available interventions have not yet been achieved. A “fully met” need = addressed and potential benefits of available interventions achieved to the extent possible for the individual. | Unmet caregiver needs derived from the JHDCNA included: dementia education (1, 0), skills counseling (1, 0), community resource availability (1, 0), emotional support (1, 0), respite support (1, 0), mental health counseling/psychiatric care (1, 0), primary medical care (1, 0), specialist medical care (1, 0), and other professional medical care (1, 0). Unmet caregiver need for Community Resource Referrals services included: the Alzheimer’s Association, eldercare attorney, Office on Aging/social services, geriatric care management, and adult protective services. The total count of unmet needs was the sum of the five items of this domain |
| **NAS**
Wackerbarth et al. (2002) | Instrument contains 3 sections:
   1. Information needs (20 items):
      - a. General information about dementia and providing care
      - b. Diagnosis and treatment
      - c. Legal and financial issues
   2. Support needs (19 items):
      - a. Supporting the care receiver
      - b. Interacting with and supporting others
      - c. Finding support for myself
   3. Information about the caregiving experience, socio-demographic information about the respondent and care receiver, and the caregiving duties (activities of daily living) | Section 1 and 2: Respondents rate the need in terms of importance on a scale from 0 to 3: 0 = not important, 1 = slightly important, 2 = important, 3 = essential. Contains also the response option “doesn’t apply to me” | No total score calculated (only means) |
| **PBH-LCED**
Sadak et al. (2015) | 6 domains (32 items)
   - 1. Ability to recognize and to anticipate day-to-day symptoms and challenges with multiple dimensions of patient health (8 items)
   - 2. Ability to manage sudden changes in the dimensions of patient’s health, to engage health services and to practice self-care (8 items)
   - 3. Patient’s medications (5 items)
   - 4. Ability to manage day-to-day symptoms and challenges with the dimensions of patient’s health (4 items)
   - 5. Ability to recognize sudden/worrisome changes in the dimensions of patient’s health (4 items)
   - 6. Ability to advocate for patient in health care situations (3 items) | 5-level Likert type response scale ranging from (1) ‘Disagree Completely’ to (4) ‘Agree Completely,’ with an additional response option (0) ‘Not my responsibility’ | Possible total scores can range from 0 to 128, with higher scores indicating higher activation |
| Instrument acronym | Domain structure/sections (number of items) | response options/format | Range of scores |
|--------------------|---------------------------------------------|-------------------------|-----------------|
| RAM                | 6 Domains (16 items) and suggested intervention strategies for each domain | Different dichotomous or ordinal response options according to the question | Domains 1, 2, 3, 4, 6: higher score indicates higher risk. Domain 5: lower score indicates less satisfaction with support and higher risk. High risk: summed all high risk answers for the upper limit of 38–40. 27 is sum of 1/2 high risk and 1/2 medium risk answers. Moderate risk: 1/2 moderate answers and 1/2 high-risk answers. Low risk: Upper limit of 11 is sum of all moderate risk answers. |
| QCNE               | Instrument contains four sections (19 items): | 19 closed (yes or no response) and two open ended questions | No scoring, no summary score |
| EAC                | 4 domains (28 items): | 5-level Likert type response scale ranging from (1) no expectations at all to (5) a lot of expectations (5): not at all, a little, moderately, many, a lot of expectations | Total score: 28–140 |
| QNP                | Instrument contains three sections: | Response option part 1 (perceived actual problems): ‘yes’, ‘actually yes’, ‘actually no’, ‘no’, or ‘not applicable’. Response option part 2 (importance of problems): ‘of paramount importance’, ‘very important’, ‘important’ or ‘not so important’. Response option part 3 (needs for specific types of professional support): list of suggestions for support, directly linked to a particular problem area | No scoring |
| Instrument acronym | Domain structure/sections (number of items) | response options/format | Range of scores |
|-------------------|---------------------------------------------|-------------------------|----------------|
| SIDECAR           | 70 items (without explicit division into domains) See Oyebode et al. (2018)** for all items | Binary response format (Agree/Disagree) | No information found |
| Tayside Gordon et al. (1997)** | Instrument contains three sections (31 items): 1. Needs (14 items): a. Mobility (1 item) b. Personal care (2 items) c. Domestic tasks (1 item) d. Behavior (2 items) e. Solitude (2 items) f. Material needs (3 items) g. Informal carer problems (3 items: coping, practical problems & emotional upset) 2. Current care inputs (6 items): Informal carer involvement (2 items), Service receipt (2 items), Respite (1 item) and Service refusal by sufferer (1 item) 3. Demographics and background (11 items): Sufferer demographics (3 items), Evidence of cognitive impairment (2 items), Informal carer demographics (5 items), Informal support to carers (1 item) | Level of commitment for the domains: mobility, personal care, domestic tasks and behaviour needs rated as: * Independent = Attention needed less than once a week * Long interval = Attention needed not more than once a day * Short interval = Attention needed more than once a day at prearranged times * Critical interval = Attention needed at short (< 2h) and/or unpredictable intervals Other response options: frequencies, quantities, yes/no or other dichotomous response options. | Key concept is the use of a descriptive profile; no summative score or categorization |
| UNM Gaugler et al. (2004)** | 7 domains (34 items): 1. Help with ADL tasks (5 items) 2. Help with IADL tasks (6 items) 3. Dementia symptoms (2 items) 4. Timing of care (3 items) 5. Formal support (8 items) 6. Information (4 items) 7. Confidante/family support (6 items) | Questions for each item: “Do you need more help with/help providing ____?” Response option: yes or no | The “yes” responses for each domain were summed to create unmet need scores |

**CG: Caregiver, PwD: Person with dementia, ADL: Activities of daily living, IADL: Instrumental activities of daily living, CADI: Carers Assessment of Difficulties Index, CARENAP: The Care Needs Assessment Pack age for Dementia, CNA-D: The Carers’ Needs Assessment for Dementia, CNCD: Caregivers’ Needs Checklist for Dementia, JHDCNA: The Johns Hopkins Dementia Care Needs Assessment, NAS: Needs assessment survey, PBH-LCD: Partnering for Better Health – Living with Chronic Illness: Dementia, RAM: Risk Appraisal Measure, QCNE: Questionnaire of carers needs evaluation, EAC: Questionnaire consultation expectations, QNP: Questionnaire National Dementia Programme Survey Needs and problems of informal caregivers of persons with dementia, SIDECAR: Scales measuring the Impact of DEmentia on CARers, Tayside: Tayside Profile for Dementia Planning, UNM: Unmet need measure.
### Appendix IV: Quality criteria for psychometric outcomes

| Instrument acronym/author (year) | Content validity | Internal consistency | Reproducibility | Construct validity |
|----------------------------------|------------------|----------------------|-----------------|-------------------|
| CADI Charlesworth et al. (2007)⁴⁹ | ? Target population was questioned for additional problems but, despite those mentioned, no changes were made to the items | Despite adequate factor analysis, 5 out of 8 Cronbach’s alphas are below 0.70 | 0 | 0 |
| CARENAP McWalter (1996/1998)⁵⁰,⁵¹ | + | Cronbach’s alpha assessed for the full scale only AND is 0.56 | 0 | 0 |
| CNA-D Wancata et al. (2005)⁵² / Kaiser et al. (2005)⁵³ | + | Cronbach’s alpha(s) calculated per dimension AND between 0.70 and 0.95; BUT no factor analysis performed | ? Evaluated in combination with inter-rater: Two interviews conducted by different persons two weeks apart; Mean Kappa 0.84 | + |
| CNCD Vainisikar et al. (2013, 2017)⁵⁴,⁵⁵ | + | Factor analysis performed but insufficient sample size (109 for 26 items); Cronbach’s alpha(s) calculated per dimension AND between 0.70 and 0.95 | 0 | 0 |
| JHDCNA Hughes et al. (2014)⁵⁶ | 0 | No information on procedure | 0 | 0 |
| NAS Wackerbarth et al. (2002)⁵⁷ | + | | 0 | 0 |
| PBH-LCSD Sadak et al. (2015)⁵⁸ | + | Factor analysis performed but insufficient sample size (130 for 32 items); Cronbach’s alpha(s) calculated per dimension AND between 0.70 and 0.95; | + Pearson correlations of 0.76 | + |
| RAM Czaja et al. (2009)⁵⁹ | - Informal caregivers were not involved in item development | - Cronbach’s alpha assessed for the full scale only AND is 0.65 | 0 | 0 |
| QCNE Dimakopoulou et al. (2015)⁶⁰ | ? Description of the concepts is too vague (e.g. wishes, problems with services) | Factor analysis inadequately performed (separately for each dimension); Cronbach’s alpha(s) calculated per dimension AND between 0.70 and 0.95 | 0 | 0 |

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**Note:** The table above summarizes the quality criteria for psychometric outcomes of various instruments. The criteria include content validity, internal consistency, agreement (test-retest reliability), reliability (inter-rater reliability), and construct validity. Each instrument is evaluated based on its adherence to these criteria, with a summary of the findings provided for each.
### SYSTEMATIC REVIEW

| Instrument acronym/ author (year) | Content validity | Internal consistency | Reproducibility | Construct validity |
|---|---|---|---|---|
| EAC Laprise et al. (2001)\(^{61}\) | - Informal caregivers were not involved in item development | ? Cronbach’s alpha(s) calculated per dimension AND between 0.70 and 0.95; BUT not factor analysis performed | + Pearson correlations of 0.63 for the total scale and 0.61 to 0.75 for the 4 subscales | 0 | + |
| QNP Peeters et al. (2010)\(^{62}/\) Van der Poel and van Beek (2006)\(^{63}\) | + | ? Cronbach’s alpha assessed for the full scale only | 0 | 0 | 0 |
| SIDECAR Oyebode et al. (2019)\(^{64}\) | + | 0 | 0 | 0 | 0 |
| Tayside Gordon et al. (1997)\(^{65}\) | ? Lack of details on the procedure | 0 | ? Time interval is too variable; Weighted Kappa of 0.41 to 0.89 for the 4 subscales | ? Time interval is too variable and ratings by worker and carer are not comparable; Weighted Kappa of -0.17 to 0.14 for the 4 subscales | 0 |
| UNM Gaugler et al. (2004)\(^{66}\) | - Informal caregivers were not involved in item development | ? 5 out of 6 Cronbach’s alphas are adequate BUT factor analysis not performed | 0 | 0 | ? Hypotheses are too vague and numerous, so that only 33% were confirmed |

\(+\): positive rating, \(?\): indeterminate rating due to doubtful design or method, \(-\): negative rating, \(\_\): no information available

CADNI: Carers Assessment of Difficulties Index, CARENAP: The Care Needs Assessment Pack for Dementia, CNA-D: The Carers’ Needs Assessment for Dementia, CNCD: Caregivers’ Needs Checklist for Dementia, JHDCNA: The Johns Hopkins Dementia Care Needs Assessment, NAS: Needs assessment survey, PBH-LCI-D: Partnering for Better Health – Living with Chronic Illness: Dementia, RAM: Risk Appraisal Measure, QCNE: Questionnaire of carers needs evaluation, EAC: Questionnaire consultation expectations, QNP: Questionnaire National Dementia Programme Survey Needs and problems of informal caregivers of persons with dementia, SIDECAR: Scales measuring the Impact of Dementia on CARers, Tayside: Tayside Profile for Dementia Planning, UNM: Unmet need measure
## Appendix V: Characteristics of included studies

| Acronym/instrument name | Authors/study design | Validation sample size | Mean age (SD or R)/age groups (n) | Distribution of sex, relationship status | Supplementary characteristics | Setting(s)/country |
|-------------------------|----------------------|------------------------|----------------------------------|------------------------------------------|-------------------------------|-------------------|
| CADI                    | Charlesworth et al. (2007) | Internal consistency & structural validity: CG: n = 232 Content analysis: 671 questionnaires | Internal consistency & structural validity: CG: M = 68 years (SD = 11.5) Content analysis: no information | Internal consistency & structural validity: CG: 65% female, 66% spouse, 26% children Content analysis: no information | —                             | Community setting |
|                         | Psychometric study    |                        |                                  |                                           |                               | UK                |
| CARENAP                 | McWalter et al. (1996, 1998) | Inter-rater reliability: CG: n = 27 Professionals: n = 28 Internal consistency & inter-rater reliability: CG: n = 27 Professionals: n = 28 Content validity: PwD: M = 82.7 years (SD = 8.34) Professionals: not reported | Internal consistency & inter-rater reliability: CG: n = 75.9 years (SD = 6.3) Professionals: not reported Content validity: PwD: M = 78.7 years (SD = 7.4) Professionals: not reported | Internal consistency & inter-rater reliability: CG: 63% female, 35% daughters, 30% wives, 26% husbands, 7% brothers, 4% son PwD: 70.6% female Content validity: PwD: 62% female, 25% male, 12% unidentified | —                             | Community setting |
|                         | The Care Needs Assessment Pack for Dementia |                        |                                  |                                           |                               | Community setting |
|                         | Same samples in both studies |                        |                                  |                                           |                               | Scotland          |
|                         | Psychometric study / Manual |                        |                                  |                                           |                               |                   |
| CNA-D                   | Wancata et al. (2005) | Content validity: CG: n = 40 Professionals: n = 40 Concurrent validity, test-retest, inter-rater reliability & internal consistency: CG: n = 45 | Content validity: CG: M = 51.1 years (SD = 13.4) Professionals: M = 77.5 years (SD = 9.3) Concurrent validity, test-retest, inter-rater reliability & internal consistency: CG: M = 60.9 years (SD = 11.9) Professionals: M = 77.5 years (SD = 9.3) Content validity: PwD: M = 82.7 years (SD = 8.34) Professionals: not reported | Content validity: CG: M = 62.5% female, 39.5% offspring Concurrent validity, test-retest, inter-rater reliability & Internal consistency: CG: 73% female, 32% offspring and 46% partners of PwD PwD: 38% female | Content validity: Professionals: 28.2% medical doctors, 25.6% nurses, 12.8% psychologists, 12.8% social workers, 7.7% occupational therapists Duration prof. experience: M = 13.9 years (SD = 9.2) Concurrent validity, test-retest, inter-rater reliability & internal consistency: PwD: 72% Alzheimer’s, 13% vascular, 15% not diagnosed exactly, Housing situation: PwD: 59% home-dwelling, 49% cohabiting with CG, 40% in nursing facility | Community setting |
|                         | (2005) |                        |                                  |                                           |                               | Austria           |
|                         | Same sample for concurrent validity in both studies | Psychometric study |                        |                                  |                                           |                   |
|                         | Kaiser et al. (2005) | |                        |                                  |                                           | Community setting |
|                         | | | | | | Nepal            |
|                         | | | | | |                   |
| CNCD                    | Vaingankar et al. (2013) | Internal consistency, structural validity & hypothesis testing: CG: n = 109 | Internal consistency, structural validity & hypothesis testing: CG: M = 55.9 years (SD = 10) PwD: M = 84.9 years (SD = 7.5) Content validity: CG: M = 52.9 years (SD not reported) PwD: R = 54-91 years (M, SD not reported) | Internal consistency, structural validity & hypothesis testing: CG: 62% female, 12% spouse, 75% child, 13% other relative/friend PwD: 68.8% female Content validity: CG: 60% female 60% children, 20% spouses, 20% other relative | Internal consistency, structural validity & hypothesis testing: CG Education: 7.4% None/Minimal, 19.5% complete primary, 43.5% complete secondary, 29.8% completed tertiary CG Employment: 45.8% paid full-time, 12.1% paid part-time, 5.6% unemployed, 0.9% student, 15.9% housewife/husband, 19.6% retired = 19.6% Content validity: PwD duration of diagnosis: R = 1 - 23 years | Community setting |
|                         | (2018) for internal consistency, structural validity & hypothesis testing | |                        |                                  |                                           | Singapore         |
|                         | Psychometric study | | | | |                   |
|                         | Vaingankar et al. (2016) | Internal consistency, structural validity & hypothesis testing: CG: n = 63 | Internal consistency, structural validity & hypothesis testing: CG: M = 55.9 years (SD = 10) PwD: M = 84.9 years (SD = 7.5) Content validity: CG: M = 52.9 years (SD not reported) PwD: R = 54-91 years (M, SD not reported) | Internal consistency, structural validity & hypothesis testing: CG: 62% female, 12% spouse, 75% child, 13% other relative/friend PwD: 68.8% female Content validity: CG: 60% female 60% children, 20% spouses, 20% other relative | Internal consistency, structural validity & hypothesis testing: CG Education: 7.4% None/Minimal, 19.5% complete primary, 43.5% complete secondary, 29.8% completed tertiary CG Employment: 45.8% paid full-time, 12.1% paid part-time, 5.6% unemployed, 0.9% student, 15.9% housewife/husband, 19.6% retired = 19.6% Content validity: PwD duration of diagnosis: R = 1 - 23 years | Community setting |
|                         | (2013) | | | | | Singapore         |
| Acronym/instrument name | Authors/study design | Validation sample size | Mean age (SD or R)/age groups (n) | Distribution of sex, relationship status | Supplementary characteristics | Setting(s)/country |
|-------------------------|----------------------|------------------------|----------------------------------|------------------------------------------|------------------------------|-------------------|
| JHDCNA                  | Hughes et al. (2014)⁵⁴ | Cross-sectional study  | Hypothesis testing: CG: n = 246 PwD: n = 254 | Hypothesis testing: CG: M = 66.13 years (SD 13.3) PwD: M = 83.6 years (SD 5.9) | Hypothesis testing: CG: 74.8% female 59.8% non-spouse PwD: 75.2% female | Community setting Baltimore, US |
| NAS                     | Wackerbarth et al. (2002)⁵⁷ | Cross-sectional survey | Content validity: 15 interviews with CG | No information for sample of content validity (only survey) | No information for sample of content validity (only survey) | Community setting Kentucky, US |
| PBH-LCID                | Sadak et al. (2015)¹⁴ | Psychometric study     | Internal consistency, Test-retest reliability, Measurement error, STRUCTURAL VALIDITY, Hypothesis testing, Responsiveness: CG: n = 130 / PwD: n = 130 | Content validity: no information | Content validity: no information | Community setting Washington/Minnesota, US |
|                         |                      |                        |                               |                                           |                              |                   |

Note: (Continued)
### SYSTEMATIC REVIEW

S. Kipfer and S. Pihet

| Acronym/instrument name | Authors/study design | Validation sample size | Mean age (SD or R)/age groups (n) | Distribution of sex, relationship status | Supplementary characteristics | Setting(s)/country |
|-------------------------|----------------------|------------------------|----------------------------------|------------------------------------------|-------------------------------|-------------------|
| RAM                     | Czaja et al. (2009)²⁰ | Internal consistency, hypothesis testing: CG: n = 642 / PwD: n = 642 | CG: M = 60.6 years (SD = 13.3)  PwD: Hispanic: M = 78.2 years (13.9), White/Caucasian: M = 78.8 years (SD = 12.3), Black/African American: M = 78 (SD = 13.2) | CG: 80% female Hispanic: 39.2% spouse  White/Caucasian: 57.5% spouse  Black/African American: 29.9% spouse  PwD: 58% female | CG: 33% Hispanic, 34% White, 33% Black/African American  PwD: MMSE: severe cognitive deficits: Hispanic: M = 11.4, White/ Caucasian M = 14.4, Black/African American: M = 11.5  PwD ADL: Limitations: substantial functional impairments: Hispanic: M = 3.5, White/Caucasian: M = 3.3, Black/African American: M = 3.4  IADL: Limitations: Hispanic: M = 6.8, White/Caucasian M = 6.9, Black/African American: M = 6.8 | Community setting 5 geographical areas in the US |
| QCNE                    | Dimakopoulou et al. (2015)²⁰ | Internal consistency & structural validity: n = 248 | CG: M = 63.16 years (SD = 14.54) | Internal consistency & structural validity: CG: M = 78.2% female, 48.6% child, 41.1% spouse, 10.5% other | Years spent in caring: male CG M = 5.29 (SD = 2.92), female CG: M = 4.23 (SD = 2.4)  PwD: 69.3% Cohabiting, 30.6% living alone  CG Education: 16.9% 1–6 years, 35.5% 7–12 years, 41.1% 13–16 years, 6.5% ≥ 17 years | Community setting Greece |
| EAC                     | Laprise et al. (2001)²⁰ | Internal consistency, test-retest reliability, hypothesis testing: CG: n = 78 | CG: 56.4% between 28–58 years, 43.6% + 60 years | Internal consistency & structural validity: CG: M = 66.6% female, 23% spouse, 77% child | Care receiver: 59% dementia, 16.6% cardiovascular diseases, 6.4% cancer  37% living in an institution, 62.8% community setting | Hospital & Nursing home setting Canada (French-speaking part) |
| QNP                     | Peeters et al.(2010)² / Van der Poel and van Beek (2006)² | Content validity: 6 psychologists ou travailleurs sociaux & 18 CG  Internal consistency: CG: n = 984 | CG: M = 62.8 years (SD = 13.3)  PwD: 78.2 years (SD = 8.8) | Internal consistency: CG: M = 71.6% female, 50.8% spouses, 38.2% children  PwD: 58% female | Internal consistency: CG: Frequency of care: 54.8% daily, 18.3% 3–6 times/week, 21.9% 1–2 times/week, Duration of caregiving: 23% more than 5years, 20% 2–3 years, 18% 1–2 years  PwD: 69% homedwelling (with spouse or child), 31% living in institution, duration of dementia symptoms: 27% more than 5 years, 20% 2–3 years 16%, 1–2 years, 15% 4–5 years  Content validity: no information | Community setting, Nursing homes Netherlands |
| Acronym/instrument name          | Authors/study design                   | Validation sample size | Mean age (SD or R)/age groups (n) | Distribution of sex, relationship status | Supplementary characteristics | Setting(s)/country         |
|---------------------------------|---------------------------------------|------------------------|----------------------------------|------------------------------------------|-------------------------------|----------------------------|
| SIDECAR                         | Oyebode et al. (2019)                  | Content validity:      | Content validity:                | Content validity:                        | Content validity:             | Community setting          |
|                                 |                                       | CG: n = 22             | CG (n = 22) age groups (years):  | CG (n = 22): 77.3% female               |                               | UK                        |
|                                 |                                       | In addition six       | 23–65: n = 12                    | 63–79: n = 8                             |                               |                            |
|                                 |                                       | researchers, two       |                                 | ≥ 80: n = 2                              |                               |                            |
|                                 |                                       | CG and a charita-     |                                 |                                          |                               |                            |
|                                 |                                       | ble sector carers’    |                                 |                                          |                               |                            |
|                                 |                                       | organization were     |                                 |                                          |                               |                            |
|                                 |                                       | included in an initial |                                 |                                          |                               |                            |
|                                 |                                       | check of content      |                                 |                                          |                               |                            |
|                                 |                                       | validity.              |                                 |                                          |                               |                            |
|                                 |                                       |                       |                                 |                                          |                               |                            |
| Tayside                         | Gordon et al. (1997)                   | Inter-rater & test-    | No information                   | No information                         | No information                | Community, residential and |
|                                 |                                       | retest validity:       |                                 |                                          |                               | hospital setting            |
|                                 |                                       | n = 1122 assessments   |                                 |                                          |                               |                            |
|                                 |                                       | of 805 individuals    |                                 |                                          |                               | Scotland                   |
|                                 |                                       |                       |                                 |                                          |                               |                            |
| UNM                             | Gaugler et al. (2004)                  | Internal consistency & | Internal consistency & hypothesis | Internal consistency & hypothesis testing:| Internal consistency & hypothesis testing: | Community and institutional |
|                                 |                                       | hypothesis testing:   | hypothesis testing:             | CG community setting:                  | CG community setting:         | setting                    |
|                                 |                                       | CG: n = 344 (community | CG: n = 344 (community setting) | M = 62.23 years (SD = 13.48)            | M = 48.09 years (SD = 31.17)  | Kentucky, US               |
|                                 |                                       | setting)               |                                  | PwD community setting:                  |                              |                            |
|                                 |                                       | PwD: n = 344 (community |                                  | M = 76.36 years (SD = 9.09)             |                              |                            |
|                                 |                                       | setting)               |                                  |                                         |                              |                            |
|                                 |                                       | Total sample:          |                                  |                                         |                              |                            |
|                                 |                                       | n = 694 (Community     |                                  |                                         |                              |                            |
|                                 |                                       | sample = 344, Nursing  |                                  |                                         |                              |                            |
|                                 |                                       | home = 137, bereaved = |                                  |                                         |                              |                            |
|                                 |                                       | 216)                  |                                  |                                         |                              |                            |
|                                 |                                       | Content validity:      |                                  |                                         |                              |                            |
|                                 |                                       | consultations with    |                                  |                                         |                              |                            |
|                                 |                                       | clinical and research  |                                  |                                         |                              |                            |
|                                 |                                       | experts in dementia   |                                  |                                         |                              |                            |
|                                 |                                       | caregiving            |                                  |                                         |                              |                            |

ADL: Activities of daily living; CADD: Cornell Scale for Depression in Dementia; CG: Caregiver; GAD: General Anxiety Disorder Assessment; IADL: Instrumental activities of daily living; M: Mean; MMSE: Mini-Mental State Examination; NPI-Q: Neuropsychiatric Interview-Questionnaire; PGDRS: Psychogeriatric Dependency Rating Scale; PHQ9: Patient Health Questionnaire; PwD: Person with dementia; SD: Standard deviation; SF-12: Short-Form Health Survey; UK: United Kingdom; US: United States; ZBI: Zarit Burden Inventory.
| Instrument acronym | Purpose (research/clinical) | Target population & setting | Application method | Respondent/administrative burden | Languages/country in which the instrument was evaluated |
|-------------------|-----------------------------|-----------------------------|-------------------|---------------------------------|----------------------------------------------------------|
| CADI              | Clinical & research use     | CG of PwD living in the community | Not clearly stated, the authors repeatedly use the term interview, but they never say who conducted the interview, and the presentation format of the questionnaire is suitable for self-administration. | 25 minutes to administer | English United Kingdom |
| CARENAP           | Clinical use                | CG and PwD living in the community | Professionally interviewed - information gathered from different sources & over a period of two weeks (2–3 visits) | PwD Assessment: 60 minutes to administer CG Assessment: 30 Minutes to administer. For professionals with experience in assessment/interviewing and PwD. | English Scotland |
| CNA-D             | Research use (service planning) | CG of PwD living in the community or in nursing facilities | Professionally interviewed | 50 minutes to administer | German Austria |
| CNCD              | Clinical use                | CG of PwD living in the community | Self-administered | No information | English, Chinese, Malay, Tamil Singapore |
| EAC               | Clinical use                | CG of older persons living in the community or in an institution | Self-administered | No information | French Canada |
| JHDCNA            | Not specified, was used in a cross-sectional study | CG and PwD living in the community | Clinician rated / professionally interviewed | No information found | English United States |
| NAS               | Not specified, was used in a survey | CG of PwD living in the community or in an institution | Self-administered | 30 minutes | English United States |
| PBH-LCID          | Clinical & research use     | CG of PwD living in the community | Self-administered | No information found | English United States |
| RAM               | Clinical & research use     | CG of PwD living in the community | Not clearly stated, the authors often highlight the ease of administration, suggesting self-report. | 5–7 minutes to assess key areas of potential risk | Spanish English United States |
| QCNE              | Not specified (used in a cross-sectional survey) | Informal dementia caregivers | Self-administered | 20 minutes to complete | Greek |
| QNP               | Not specified (was used in a national survey) | CG of PwD living in the community or in an institution | Self-administered or professionally interviewed | 30 minutes Interviewer does not need interview abilities or knowledge | Dutch Netherlands |
| SIDECAR           | Not specified               | CG of wilting in the community | Self-administered | No information found | English United Kingdom |
| Tayside           | Clinical use                | CG and PwD living in the community, residential or hospital setting | Self-administered or professionally interviewed | No information found | English Scotland |
| UNM               | Not specified (was used in a cross-sectional survey) | CG of PwD living in the community, residential or hospital setting | Self-administered | No information found | English United States |

CADI: Carers Assessment of Difficulties Index (Charlesworth et al., 2007)^[49]; CARENAP: The Care Needs Assessment Pack for Dementia (McWalter et al., 1996;1998)^[50-51]; CG: Caregiver; CNA-D: The Caregivers’ Needs Assessment for Dementia (Wancata et al., 2005; Kaiser et al., 2005)^[52-53]; CNCD: Caregivers’ Needs Checklist for Dementia (Vaingankar 2013; 2017)^[54-55]; EAC: Questionnaire consultation expectations (Laprise et al. (2001)^[61]; JHDCNA: The Johns Hopkins Dementia Care Needs Assessment (Hughes et al., 2014)^[56]; NAS: Needs assessment survey (Wackerbarth et al, 2002)^[57]; PBH-LCID: Partnering for Better Health – Living with Chronic Illness: Dementia (Sadak et al, 2015)^[58]; PwD: Person with dementia; QCNE: Questionnaire of carers needs evaluation (Dimakopoulou et al., 2015)^[60]; QNP: Questionnaire National Dementia Programme Survey Needs and problems of informal caregivers of persons with dementia (Peeters et al., 2010); Van der Poel and van Beek, 2006)^[62-63]; RAM: Risk Appraisal Measure (Czaja et al., 2009)^[59]; SIDECAR: Scales measuring the Impact of Dementia on CARers (Oyebode et al., 2019)^[64]; Tayside: Tayside Profile for Dementia Planning (Gordon et al. 1997)^[65]; UNM: Unmet need measure (Gaugler et al. 2004)^[66].