Towards a middle-range theory of ‘Stability of home-based care arrangements for people living with dementia’ (SoCA-Dem): findings from a meta-study on mixed research

Kerstin Köhler ,1,2 Jan Dreyer ,1,2 Iris Hochgraeb ,1,2 Milena von Kutzleben ,3 Christiane Pinkert ,1,2 Martina Roes ,1,2 Bernhard Holle ,1,2

ABSTRACT

Background Most people with dementia and their informal carers live at home and strive to create a stable care situation for as long as possible. This preference of dyads is consistent with the global policy of ageing in place. Therefore, we aimed to develop a middle-range theory of stability guided by two research questions: How is stability of home-based care arrangements for people living with dementia constituted? What are the essential factors influencing stability?

Methods Within the ‘Stability of home-based care arrangements for people living with dementia’ project (SoCA project) at the German Center for Neurodegenerative Diseases (DZNE), we conducted a meta-study on mixed research. The analytical steps of meta-data analysis, meta-method and meta-theory are merged in an integrative synthesis. Eligible publications were identified through systematic database searches (MEDLINE, CINAHL and PsycINFO; last searched on 3 January 2017), backward/forward citation tracking and snowballing. All publications were screened against predefined inclusion criteria and evaluated through a quality appraisal. The analytical approach was thematic synthesis.

Results 99 publications were included. The middle-range theory conceptualises stability as a complex phenomenon comprising three components including eight concepts that are dynamically inter-related. The conceptual model visualises: (1) the trajectory of the dementia care arrangement, which involves a cyclic process of change and balancing over time; (2) the characteristics of the care arrangement, including needs, the carer role, the dyadic relationship and resources; and (3) the context, which is determined by society and culture and the respective healthcare system. The relevance of each concept in relation to stability changes over time. The forming of each concept is actively shaped by the informal carer.

Discussion This middle-range theory provides a thorough understanding of the stability of home-based care arrangements for people living with dementia and can be used to guide future research and practice.

Strengths and limitations of this study

- We developed a middle-range theory (SoCA-Dem theory) that provides an innovative conceptualisation of the phenomenon ‘Stability of home-based care arrangements for people living with dementia’.
- The SoCA-Dem theory can guide future research and policy development and can also be used to improve health and social care practice.
- Currently, we elaborate the interplay of the concepts that constitute ‘stability’ on a descriptive level: causal relationships need to be explored in subsequent empirical research.
- To better understand the complexity of home-based care arrangements, the predominant perspective of the informal carer within the SoCA-Dem theory should be complemented by the perspective of the person living with dementia and the diverse informal and professional supporters.
- This meta-study is based primarily on research from Europe and North America, which might limit the applicability of the SoCA-Dem theory in societies with different cultural contexts.

INTRODUCTION

Worldwide, most people with dementia live at home. It is the explicit wish of most people with dementia to stay in their familiar environments, and most informal carers wish to support the person with dementia at home for as long as possible. Driven by this motivation, informal carers are a vital resource for people living with dementia at home and play a crucial role in supporting dementia care in...
the community. In current research, policy and practice, these preferences are mirrored in programmatic agendas, such as ‘ageing in place’, and reflected by an increasing focus on ‘social health’. These agendas emphasise the capability of people with dementia, their families and their social networks and aim to create prerequisites to enable dyads to live well with dementia in the places of their choice.

This dyadic perspective can be complemented by the perspective of the social security and healthcare system. In light of demographic changes and the constantly growing prevalence of dementia worldwide, societies are challenged to provide adequate health and social care for an increasing number of ageing individuals and, thus, need to appropriately address limited financial resources. As dementia is among the strongest predictors of cost-intensive institutional care, the support of home-based care has been discussed as an option to reduce expenses.

The stability of home-based care arrangements is an important goal for people living with dementia and their informal carers and reflected by current social security and healthcare policy. Accordingly, the term ‘stability’ is used as a buzz word in research and policy contexts, but a thorough definition of stability is usually not provided. Often, stability and instability are recognised as opposing phenomena and are operationalised in a binary way. Thus, the transition of a person with dementia from home to an institution is often understood as a demarcation line between stability and instability. The time until institutionalisation greatly varies, and numerous studies have uncovered the predictors and risk factors of institutionalisation, but these studies did not focus on the process that finally leads to the termination of home-based care. Other studies have illuminated the interactions between individuals involved in home-based care, but these interactions were not clearly linked to the phenomenon of stability or instability. In our understanding, continuing to live at home does not necessarily indicate a stable situation, and in the worst case, the needs of the involved persons can be neglected, leading to an exhausting situation for the person with dementia or the informal carer. In contrast, living in a nursing home can be perceived as a satisfying and stable living situation. At first glance, the prevalent binary conceptualisation of stability versus instability based on the criterion of institutionalisation is appealing, but at a second glance, this conceptualisation does not capture the complexity of the phenomenon.

The significance of the interactions between all persons involved in a home-based dementia care arrangement and the central importance of the needs of these persons are highlighted in a published working definition of stability (see figure 1). Here, stability is defined as a dynamic process that unfolds over the trajectory of dementia care that is intentionally shaped by the actors and can be achieved only if the needs of the person with dementia and the informal carer are addressed. This definition is an essential part of the foundation of this meta-study. It reflects the work conducted by Corbin and Strauss regarding managing chronic illness, Nolan and colleagues regarding family care and von Kutzleben regarding informal dementia care trajectories, all of which conceptualise care as a process of vibrant interactions between individuals over time.

The working definition and this meta-study are integral parts of the ‘Stability of home-based care arrangements for people living with dementia’ project (SoCA project) at the German Center for Neurodegenerative Diseases (DZNE), site Witten, Germany. The SoCA project strives to develop a sound theory-based understanding of the complex phenomenon of stability and further explore the newly developed middle-range theory by conducting subsequent empirical studies using qualitative and quantitative research methodologies to finally identify and develop stability promoting interventions and care structures.

**AIM**

The aim of this study was to develop a middle-range theory of the ‘Stability of home-based care arrangements for people living with dementia’. The theory generation process was driven by the following research questions: How is stability of home-based care arrangements for people living with dementia constituted? What are the essential factors influencing stability?

**METHODOLOGY**

**Study design**

We aimed to generate theory from published research findings. For this purpose, we performed a meta-study according to the approach described by Paterson and colleagues and adapted the original meta-study methodology for the integration of mixed research by including qualitative and quantitative, mixed-method studies, as well as systematic reviews. During the theory development process, we considered the recommendations by Liehr and Smith, In and Risjord regarding the creation of middle-range theories. In the subsequent sections, we provide a very brief description of our methodology. The
details of the methodology were published in the respective study protocol.27 This meta-study is registered at the International Prospective Register of Systematic Reviews (PROSPERO registration number CRD42016041727). The reporting of this publication follows the recommendations in the ENTREQ statement.28

**Literature search**

Following the framework described by Boell and Cecez-Kecmanovic,29 30 we searched for literature in open-ended ‘hermeneutic loops’, including the repeated steps of searching, sorting, selecting, acquiring, reading, identifying and refining. In parts, the cyclic literature search process was performed simultaneously with the ongoing data analysis: the final goal of the whole theory building process was to sufficiently encircle our phenomenon of interest by reaching conceptual saturation.31

The following search techniques were combined: berry picking,32 database searches, backward and forward citation tracking and snowballing. The database searches were performed between 24 September 2015 and 16 November 2015 in MEDLINE, CINAHL and PsycINFO and updated again on 3 January 2017 (referred to as ‘updated database searching’ in figure 2). Since the phenomenon of stability is rarely investigated directly, we used our previously published definition of stability18 to draw from it nine separate search strands and nine syntaxes. The complete syntaxes of the database searches are shown in online supplemental appendix A. During the iterative cyclic process of searching and analysing, we included further publications identified through citation tracking and snowballing.

**Screening process**

The titles and abstracts of all identified studies were screened by four authors (KK, JD, IH and MvK) in alternating teams of two authors. Full-text screening was performed by one author, who discussed his or her decision with at least one colleague. If disagreement occurred, consensus was reached by discussion. EndNote X7 was used to manage the screening process.

**Eligibility criteria**

The original meta-study methodology was designed for the synthesis of qualitative research only. In contrast, we decided to include all types of empirical research and (systematic) reviews. The rationale for this decision has been described elsewhere.27 The inclusion criteria are shown in online supplemental appendix B. We included
publications written in English, German or French; no time limits were set.

Quality appraisal
We applied the quality appraisal recommended by Dixon-Woods and colleagues,33 which includes five criteria to identify studies with major methodological deficits (online supplemental appendix B). The decision to include or exclude a publication was made based on this quality appraisal and the publication’s contribution to the proposed middle-range theory. Each paper was assessed by one researcher. If he or she was in doubt of the methodological quality, he or she discussed the paper with at least one other researcher until consensus was reached.

Analysis and data collection process
A meta-study combines the following three analytical steps, leading to a final meta-synthesis: meta-theory, meta-method and meta-data analyses. During our iterative analytical process, these steps were considered in an integrated process rather than individually. The insights gained from one analytical step immediately shaped the subsequent analyses. All analyses were performed using the Professional Research Software for Qualitative, Quantitative and Mixed Methods Research (MAXQDA 2018). The coding of the data was performed in teams of at least two researchers. As Thomas and Harden,34 we translated the widespread method of thematic analysis for the use in systematic reviews, we chose this analysis approach.

Meta-theory
Meta-theory involves a critical exploration of the theoretical frameworks that guided the included research.24 For each publication, we analysed the stated research gaps, aim(s), research question(s), explicitly stated theoretical frameworks and/or underlying concepts.

Meta-method
Meta-method analyses consider how the application of research methodologies influences the findings and shapes the current understanding of the phenomenon under research.24 For this purpose, we analysed the study design, methodology, sample, sampling, setting, data collection, data analysis and standardised measures used in each publication.

Meta-data analysis
A meta-data analysis is a comparative analysis of the findings of all included studies; this analysis contributes to the creation of an integrated body of knowledge.24 In our extended meta-study approach that included mixed research, we ‘qualitised’ the quantitative data. Our aim was to understand which concepts applied in quantitative studies are associated with ‘stability’ and how. Following Pope and colleagues35 and Sandelowski and colleagues,36 our main approach for ‘qualitising’ quantitative data was to extract relevant concepts from the texts, tables and figures of included quantitative studies, mixed-methods studies and systematic reviews and to compare these concepts across all studies. Concepts applied in quantitative studies often gave indications of an association with (in-)stability, which was examined in depth in qualitative studies. For details on ‘qualitising’, see also Dreyer and colleagues.27 We deduced a preliminary coding scheme based on our working definition.16 Following Thomas and Harden,34 we extended this coding scheme and analysed the findings of the included studies in three stages. We started the ‘line-by-line coding’ with a small introductory text set (figure 2). Gradually, the code system grew by induction, and the new codes were applied for each publication that was added over time while going back and forth in iterative loops of searching and analysing literature. In several coding rounds, the codes were translated to and compared across all publications of the meta-study sample. We organised these codes into ‘descriptive themes’, and we created major ‘analytical themes’ that formed the components and concepts of the proposed middle-range theory. Finally, no additional codes emerged from the added studies, and conceptual saturation was reached.31

Meta-synthesis
In the meta-synthesis, we elaborated on the relationship between the analytical themes. While we engaged more deeply with the material, at a certain point, the emerging middle-range theory no longer substantially changed. To ensure that the meta-synthesis was complete, we sampled a selection of publications from our sample and reviewed these publications again to determine whether the statements were all consistent with the status quo of the middle-range theory or any statements that might change the theory were overlooked. Our middle-range theory is visualised in a conceptual model that displays the interplay among the components and concepts in an organised form.

Patient and public involvement
Patients or the public were not involved in this meta-study.

RESULTS
Sample description
We included 99 publications in our meta-study. The detailed process of searching and selecting eligible publications is documented in the flow diagram in figure 2. During the screening process, 5 publications were excluded during the quality appraisal. During the analysis phase, the whole ‘interventions’ search strand was excluded from the sample because the content of the retrieved intervention reviews and studies did not contribute to the intended theory generation. A statement explaining the reasons for this decision is provided in online supplemental appendix C. Detailed information of each publication is provided in online supplemental appendix D, and a complete bibliography of the included papers is displayed in online supplemental appendix E.
The included publications were authored by researchers in North America (the USA: n=31 and Canada: n=12) and Europe (Germany: n=12, the UK: n=11, the Netherlands: n=8 and other European countries: n=12); members of international consortia (n=8); and researchers in Asia or Australia (n=5); however, no publications authored by researchers in Africa or South America were included. The publications cover a wide range of scientific fields, including geriatrics, geriatric psychiatry, gerontology, dementia, nursing, palliative care, social sciences and so on. More than half of the included publications (n=56) were published from 2010 onwards, 26 were published between 2000 and 2010, 13 were published in the 1990s and 4 were published in the 1980s.

50 included studies used quantitative methodologies, and 31 studies used qualitative methodologies. Furthermore, we included 7 mixed-methods studies and 11 systematic reviews (3 synthesised qualitative studies and 9 synthesised quantitative studies). 54 publications reported studies involving cross-sectional designs, and 34 publications reported studies involving longitudinal designs.

Only 21 publications explicitly stated the theoretical framework guiding their research process. Thereby, stress theories and stress models were chosen the most frequently (n=10). In 78 publications, no explicit theoretical framework was indicated, but the dominance of a stress/burden perspective was also apparent in these publications. For instance, 71 publications in our sample refer to the work conducted by Zarit et al., Pearlin et al. and/or Aneshensel et al., who are researchers that advanced the understanding of stress and burden in caregiving. In the discussion section of our article, this conceptual bias and its impact on the development of our middle-range theory are reflected.

Components and concepts of the middle-range theory and their interplay

In this meta-study, we developed a middle-range theory of the ‘Stability of home-based care arrangements for people living with dementia’ (called the SoCA-Dem theory) that emphasises the perspective of the informal carer. The current focus on the informal carer perspective in this middle-range theory is driven by our own previous work concerning informal dementia care trajectories, which identified stability as a guiding principle of informal carers. Therefore, we created a search strategy in which the presence of an informal carer in the home-based care arrangement was mandatory. As a consequence, most identified studies placed the informal carer at the centre of the research. Nevertheless, we are keenly aware that the perspectives of all actors (including the person living with dementia) in care arrangements are meaningful and shape the respective care arrangement, and we reflect on this conceptual bias in the discussion section of this publication.

We illustrate the middle-range theory in a conceptual model. The model visualises three components comprising eight concepts that altogether constitute stability of home-based care arrangements for people living with dementia from the perspective of the informal carer (see figure 3).

- Component 1 – trajectory: according to our understanding and consistent with Corbin, a chronic illness trajectory refers to the course of an illness/condition over time and the actions that shape that course. The arrow at the bottom visualises the time course of the trajectory of a home-based care arrangement that covers the time span from the onset of the care arrangement to its termination. Above the arrow, a cyclic process displays the events and actions that shape the trajectory. This cyclic process comprises the two concepts of change and balancing and reruns in many loops over the trajectory.
- Component 2 – characteristics: four concepts, namely, needs, care role, dyadic relationship and resources, and the forming of these concepts characterise the stability of a home-based care arrangement.
- Component 3 – context: the trajectory of a home-based care arrangement and its characteristics are influenced by the context, which includes the two concepts society and culture and the healthcare system.

The components and concepts of this middle-range theory are related in a dynamic interplay and cannot be interpreted in isolation from each other. Each component and concept is entwined with the other components and concepts, and all components and concepts influence and are influenced by the other components and concepts.

At any time in the trajectory, change may occur and trigger an action (balancing) of the informal carer. This balancing influences whether the needs of the person with dementia and the informal carer are successfully met or remain unmet. Balancing influences the informal carer’s experience of the care role, the development of the dyadic relationship and the management of resources. In contrast, met or unmet needs, the specific experiences of the informal carer, the development of the dyadic relationship and the management of resources may enable or hinder the informal carer from balancing in such a way that supports the stability of the care arrangement. In addition, this interplay between change and balancing over time and the characteristics of the care arrangement is influenced by the societal and cultural background and the respective healthcare system.

Thus, the forming of each concept is variable and change happens over the course of time, here over the care trajectory; each concept can assume a forming that promotes the stability of the home-based care arrangement or a forming that may challenge the stability. In addition, a specific concept in a specific form may have a minor impact on stability at the beginning of the trajectory but a major influence during the late phases and vice versa. At any time, there can be stable and unstable phases. These mechanisms of the dynamic interplay between the components and concepts of the SoCA-Dem theory are further clarified in the subsequent paragraphs, which
introduce the concepts one by one. To ensure the readability of the results section, we decided to cite only one key reference for each argument. Online supplemental appendix D presents information (for each of the included publications) on the question if a publication contributed to our understanding of a specific concept of the developed middle-range theory.

**Change**

**Conceptualisation**

To a great extent, the trajectories of home-based dementia care arrangements are driven by change. One major change is the progression of dementia. Dementia is a neurodegenerative syndrome associated with a progressive deterioration of cognitive and physical functioning that increasingly challenges the person with dementia in managing his or her daily life independently. Consequently, the person with dementia often relies on the support of family members, friends, neighbours and professional service providers. The change caused by dementia introduces various additional challenges to the home-based care arrangement. In general, gradual changes can mostly be well balanced, whereas unexpected changes challenge the stability of the home-based care arrangement.

**Analyses and synthesis**

Table 1 provides an overview of the fields in which change occurs.

If change occurs, the informal carer appraises the change and decides whether and how he or she needs to balance the changed situation. The appraisal of change is highly individual as follows: some informal carers interpret the first cognitive symptoms of the person with dementia as a part of normal ageing, while others identify these changes as pathological and seek a diagnosis. During this phase, some dyads experience a period of uncertainty while coping with ‘diagnostic shock’, whereas others feel relieved by the final diagnosis. This appraisal of the impact and meaning of a specific change by the informal carer is as important as the change per se.

In the context of this middle-range theory, change is the most crucial if it challenges the stability of the home-based care arrangement and provokes crises that cannot be balanced. MacNeil Vroomen and colleagues built a conceptual model of stability of home-based care arrangements for people living with dementia.

![Conceptual model of stability of home-based care arrangements for people living with dementia.](http://bmjopen.bmj.com/)

**Figure 3** Conceptual model of stability of home-based care arrangements for people living with dementia.
theory of crises in dementia care. Their conceptualisation of the dementia crisis process and the repeating alternation between imbalance and homeostasis is consistent with our understanding of stability.

Change can trigger positive or negative consequences. If the informal carer can accept the (unavoidable) changes in the person with dementia and the related changes in his or her own life and decide to acknowledge the positive aspects of his or her situation, this may result in supportive changes in his or her attitude and ultimately facilitate the finding of meaning.

Change can be slow, developmental and predictable or rapid, situational and unexpected. Different forms of dementia or rather late or early onset of dementia, can result in different changes. In general, changes that can be foreseen (such as a progressive cognitive decline) are easier to balance than changes that are unexpected (such as a fall). Gradual changes can be monitored without time pressure, whereas in the face of a rapid change, informal carers must act spontaneously. Some authors have concluded that serious and unexpected changes rather than constant changes are the initial triggers leading the informal carer to start thinking about the termination of home-based care. Liken reported that 80% of the informal carers in her sample moved their relatives with dementia to institutions due to unexpected events.

### Table 1 Overview of the fields of change addressed in the meta-study sample

<table>
<thead>
<tr>
<th>Change related to:</th>
<th>Fields of change</th>
</tr>
</thead>
<tbody>
<tr>
<td>Person with dementia</td>
<td>Cognition, behaviour, (Instrumental) Activities of Daily Living (ADL/IADL) functioning, physical health, mental health and age</td>
</tr>
<tr>
<td>Informal carer</td>
<td>Physical health, mental health, ADL/IADL functioning and age</td>
</tr>
<tr>
<td>Needs</td>
<td>Needs of the person with dementia and needs of the informal carer</td>
</tr>
<tr>
<td>Carer role</td>
<td>Self-conception and external conception of the carer role and having/assuming other roles</td>
</tr>
<tr>
<td>Dyadic relationship</td>
<td>Roles in the dyad, reciprocity and relationship quality</td>
</tr>
<tr>
<td>Resources</td>
<td>Practical skills, mental resources, financial resources and arrangement of informal/formal networks</td>
</tr>
<tr>
<td>Society and culture</td>
<td>Policy strategy, societal discourse of dementia, demographics and spatial environment</td>
</tr>
<tr>
<td>Healthcare system</td>
<td>Policy strategy, availability and financing of health and social care services</td>
</tr>
</tbody>
</table>

### Balancing

#### Conceptualisation

Change in a current care situation triggers an action by the informal carer. The most prominent action strategy in the proposed middle-range theory is balancing. Informal carers strive to balance the situation to maintain the stability of their care arrangements. Balancing can either target internal processes (eg, changing mindset) or practical actions. Imbalance occurs if there is a mismatch between the acute challenge caused by a change and the informal carer’s repertoire to adapt to this challenge. If the informal carer loses control and fails to balance the situation, the home-based care arrangement often moves into a crisis, followed by a period of instability.

#### Analyses and synthesis

We identified balancing as a process that is often divided into four steps. First, the informal carer appraises a change and reflects on whether he or she should adapt to the change. Thus, the informal carer weighs the risk of taking no action against the perceived benefit of adaptation. Second, the informal carer decides how to adapt to maintain the stability of the situation. Third, the informal carer acts with the intention to balance the care situation. Fourth, the informal carer evaluates whether his or her balancing was adequate to maintain the stability of the care arrangement.

The notion of different types of informal carers who perform balancing in different ways was very prominent in our sample. In some studies, a distinction was made between informal carers who proactively plan ahead for the future and informal carers who delay decisions and actions until they obviously need to react in a crisis. Regarding the maintenance of stability of the home-based care arrangement, a proactive balancing strategy seems to promote stability, whereas a reactive attitude challenges stability.

The following examples illustrate that the fields of actions targeted by the balancing of an informal carer are diverse; informal carers balance the needs of the person with dementia with their own needs to care for their physical and emotional well-being. Regarding the carer role, informal carers balance their mindsets; they accept the fact that their family members have dementia and choose to have a positive attitude, modify their beliefs and expectations and attempt to find peace of mind. Furthermore, informal carers balance the duties they face in their carer roles with their duties in other roles, such as being a parent or having a professional occupation. Informal carers balance the reciprocity in their dyadic relationships and the interactions within their informal and formal support networks and strive to establish collaborative partnerships.

In this context, informal carers value their support networks as an essential source of the support needed to continue in their carer roles. In addition, they balance the challenge of caring for a person with dementia by developing expertise...
in their knowledge regarding dementia and their practical skills.43

Related to our understanding of a trajectory as a cyclic process of change and balancing over time, some publications in our sample contextualised their research with theories concerning chronic illness trajectories42 and conceptualised dementia care as occurring in phases over time.62 Some studies shed light on informal carers’ balancing throughout the whole care trajectory.63 However, overall, our sample strongly focused on the decision making of informal carers regarding the institutionalisation of the person with dementia.64 Therefore, in extent research, there is an emphasis on balancing during the phase of the trajectory in which instability is impending or already definite.

Needs
Conceptualisation
The needs of the person living with dementia and the informal carer occupy a central position in our conceptual model because a stable care arrangement can be created and maintained only if these needs are met. Over the trajectory of a home-based care arrangement, needs change, and continuous balancing is necessary. Conflicts due to the competing needs of one person or the divergent needs of both persons in the dyad can emerge and must be resolved to the greatest extent possible.

Analyses and synthesis
A prevailing assumption among the studies in the sample was that the stability of a care arrangement is at risk if the needs of the person with dementia or his or her informal carer are unmet.65

Regarding the person with dementia, the commonly stated needs were health-related needs or needs related to limitations in the activities of daily living.66 These needs were discussed as a consequence of the progression of dementia or ageing and multimorbidity in general. Regarding the informal carer, the identified needs were often related to the consequences of giving care, the informal carers’ well-being,67 or lack of resources.68 Needs related to the lack of resources often included needs for professional support services or educational programmes.50

Furthermore, during the analysis process, we identified numerous needs of the person with dementia and the informal carer that were related to one of the concepts of the characteristics of the care arrangement, such as the need to find meaning in the demanding carer role;47 the need to preserve reciprocity in the dyadic relationship46 and the need for resources in the form of a supporting network of family, friends, neighbours and professionals62 or specific services (such as care).69 According to these findings, needs represent a concept with a very large conceptual overlap with the other concepts and major influence; whether needs are met or unmet shapes the forming of the other concepts which, in turn, either promotes or challenges stability. Therefore, our conceptual model positions needs as the central concept.

As dementia progresses, the informal carer becomes increasingly responsible for anticipating the needs of the person with dementia. Considering the needs of the person with dementia and his or her own needs is not an easy task for the informal carer as these needs can be conflicting.52 Furthermore, the anticipated needs of the person with dementia could also conflict with each other, such as the need for autonomy versus the need for safety.70 Additionally, the informal carer could have conflicting needs, such as the need to be a dedicated carer and the need to care for himself or herself. Therefore, these conflicts in needs could be distinguished as interneed conflicts (ie, the persons in the dyad have conflicting needs) and intraneed conflicts (ie, the needs of one person conflict with each other).

Carer role
Conceptualisation
The way an informal carer forms his or her carer role has a crucial impact on the stability of home-based care arrangements. Becoming and being an informal carer can be perceived as a positive mission, an undesired obligation or a mélange of both positive and negative experiences. Regarding the stability of home-based care arrangements, how the informal carer assumes the carer role and manages to balance this role with other social roles seem to be important.

Analyses and synthesis
At the very beginning of the trajectory, assuming the carer role is a gradual and unconscious process.42 Subsequently, informal carers realise their actions and identify as carers.41 Adapting to the new role is often challenging43; for instance, because informal carers assume new responsibilities71 and feel unprepared,60 adapting to the new role can be particularly challenging if the beginning of the care trajectory is unexpected.72

Assuming the carer role is usually motivated by the dyadic relationship between the informal carer and the person with dementia.70 In general, there are societal expectations to care for family members. These expectations vary across countries and cultures and are reflected in the respective healthcare systems.73

Informal carers have different feelings regarding their roles. For some informal carers, caring is a reciprocal act as follows41: adult children often understand caring as an opportunity to give back to their parents47 and spouses often view caring as fulfilling their marriage vows.58 Being intrinsically motivated to provide care supports stability.55 A positive appraisal of giving care increases the motivation of the informal carer78 and the likelihood that he or she will continue caring at home.74 Intrinsically motivated informal carers take pride in their roles and often experience personal growth.47 For other informal carers,
assuming the carer role is considered a moral obligation. Such carers feel as if they have no choice either because no one else is available or others expect them to provide care. This situation can lead to feelings of entrapment and overload in the carer role. Informal carers who experience few if any positive aspects and feel burdened by their duties have a lower commitment to continue caring at home and tend to develop a desire to institutionalise the person being cared for in the future and actually end the home-based care arrangement sooner. Attitudes towards the carer role can change over the trajectory of the care arrangement, which was described in the studies in the sample as a modification of beliefs or personal expectations, a cognitive shift or a result of adapting and learning.

A consequence of assuming the carer role may be role conflicts with other obligations in the life of the informal carer. These obligations can be a professional occupation, the informal carer’s family life (especially among carers who have children) or other social interactions with friends or neighbours. In particular, adult children are concerned with conflicting obligations. If the informal carer does not successfully balance his or her social roles, the stability of the care arrangement could be endangered.

**Dyadic relationship Conceptualisation**
Caring occurs within relationships, and most informal carers perceive caring as a natural continuation of the dyadic relationship with the person living with dementia. Therefore, the structure of the dyadic relationship (ie, kinship relation, family status and living situation) and the quality of the dyadic relationship (ie, primarily the renegotiation of roles and the maintenance of reciprocity) impact the stability of the home-based care arrangement. The direction of the effect of the relationship quality on stability may reverse over the trajectory.

**Analyses and synthesis**
Most studies in our sample stated that close kinship relationships promote the stability of a home-based care arrangement, whereas distanced kinship relationships increase the likelihood of a prompt cessation of home-based care. In general, spouses tend to continue to care at home the longest, children tend to end home-based care sooner than spouses and other informal carers tend to institutionalise sooner than children. Regarding the family status of the person with dementia, there is consensus that being married or living in a partnership is associated with the expense of the informal carer’s physical and financial health. The availability of at least one informal carer seems to be indispensable. Living in the same household as the informal carer is supportive of the maintenance of a stable situation at home, whereas living alone is often a challenge. These structural aspects of the dyadic relationship do not directly influence stability but rather function as indicators of the other concepts included in our middle-range theory. The kinship relationship, family status and living situation influence the quality of the dyadic relationship (eg, spousal affection vs filial piety), the self-construction of the carer role (eg, motives to provide care) and the availability of informal and formal resources (eg, having adult children).

A prevailing assumption among the studies in the sample was that a good past or present relationship quality supports the continuation of home-based care, whereas a low past or present relationship quality challenges stability. Over the trajectory, the forming of the dyadic relationship is influenced by changes in the person with dementia. In many chronic illness trajectories, dyadic relationships tend to develop an asymmetric structure, with the carer being dominant. These changes are especially meaningful in a dementia care trajectory as they usually lead to a shift in former roles and are often accompanied by a strong decline in the reciprocity of the dyadic relationship.

In the dementia care trajectory, premorbid roles within the dyadic relationship are questioned and adapted to the current situation. Sometimes, a swap of premorbid (often traditional and gender-related) roles occurs, for example, carer husbands perform household tasks, carer wives manage financial issues and children assume parent-like roles. Many informal carers perceive this process of adapting long-standing roles in their dyadic relationships as a challenging transition that may be accompanied by a decline in the quality of the dyadic relationship, which is often conceptualised as shrinking reciprocity.

In reciprocal relationships, informal carers have a high willingness to continue care at home. If informal carers appreciate the opportunity to reward the love and care they received in the past, they may be more likely to continue. In contrast, informal carers start to complain about a decline in the reciprocity of their dyadic relationships, they lost a major source for continuing care. These informal carers report changes in their feelings towards their relatives with dementia and a decreased level of affection, emotional distance and estrangement and a perception of having fallen out of love and ultimately describe the relationship with the relative with dementia as lost.

Over the trajectory of a care arrangement, the direction of the effect of the dyadic relationship on the stability of the home-based care arrangement may change. At the onset and during the early phases of the trajectory, it is assumed that the closer the kinship relationship is and the higher the quality of the dyadic relationship is, the higher the chance that the home-based care arrangement will remain stable. However, at some point, the direction of this effect may reverse. In the long term, the intent to keep the person with dementia at home for as long as possible is often associated with the expense of the informal carer’s physical and emotional health.
mental well-being or the health and safety of the person with dementia. If the informal carer does not anticipate the critical phase when he or she will no longer be able to balance the care situation adequately, he or she might miss the opportunity to ensure a smooth and well-planned termination of home-based care.

Resources
Conceptualisation
Resources are pivotal for informal carers to balance the consequences of constant change and create and maintain a stable care arrangement. The more resources available in general, the more available resources fit the dyad’s needs, and the smarter these resources are mobilised, the better the stability of the care arrangement. The availability and use of resources change over the trajectory and can be differentiated as intrapersonal resources and interpersonal resources.

Analyses and synthesis
In many publications, various resources were studied to examine their impacts on stability. If informal carers judge their resources as inadequate to balance the situation, the stability of the care arrangement could be endangered. In the category of intrapersonal resources, mental resources (eg, resilience and related concepts, such as mastery, a sense of competence and self-efficacy) were the most common resources researched. Skills, such as hands-on care skills or communication skills, were the second most studied group of resources researched in the intrapersonal category. Furthermore, many studies focused on the education, income levels and the physical health of the informal carer.

In the category of interpersonal resources, the studies in our sample investigated social support (eg, family members, friends or neighbours) and formal support (eg, respite care, community nurses or meal services). Regarding interpersonal resources, notably, the overall amount of potential social or formal support is often less important than the quality of the relationships and interactions between the persons involved. Meaningful interactions and communication between the informal carer and his or her supporters is highly important for the creation of stable care arrangements.

All resources are dynamic and may change during the trajectory of a home-based care arrangement. For instance, the informal carer may experience a decline in support from his or her social network and an increase in social isolation if family members and friends withdraw. In contrast, meaningful new relationships, for example, with the members of a support group, could grow over the trajectory. The possible positive development of resources, such as increased resilience or the learning of new skills, were discussed as gains of caring, while a decline in resources, such as deteriorating health or decreasing financial resources, were discussed as burdens that may endanger the stability of home-based care.

One study in our sample adopted an explicit strength-based perspective as its theoretical foundation and highlighted the meaning of resources in the context of positive outcomes of caregiving. Other studies explicitly referred to a stress model, for example, the stress process model or the stress-coping model. In these stress models, resources play an important role in coping with challenging life events. Studies adopting a theoretical perspective focusing on social inequality in the allocation of resources were missing in our sample, which could be interpreted as the lack of a societal perspective and emphasis on the individual/dyadic perspective.

Society and culture
Conceptualisation
Home-based dementia care arrangements are influenced by the society and culture in which the persons with dementia and their informal carers live. Therefore, the prevailing values of a specific culture shape the stability-related context.

Analyses and synthesis
Regarding cultural values, we included studies addressing the societal expectations of different cultures in different geographical regions, for example, China and Europe, and research addressing the societal expectations of different ethnic groups in one country, for example, Chinese people in Australia or Latinos in the USA. In some cultures and ethnic groups, prevailing traditional values lead people to prefer family care and refuse to involve professional support services. In such contexts, the included studies described ‘caring as a fulfillment of cultural values’ or a ‘powerful cultural tradition of family care’. In a German study, von Kutzleben and colleagues identified the country of origin of the person living with dementia as a significant influencing factor as follows: informal carers who cared for a person with dementia who was native born. Mausbach and colleagues and Sun and colleagues reported that in the USA, Latinos and African-Americans tend to institutionalise their relatives later than their Caucasian counterparts.

In cultures bound to traditional values, (mostly female) family members are expected to assume the informal carer role. On the one hand, these informal carers often feel valued by their social networks and tend to have positive attitudes towards their carer role. On the other hand, informal carers are frequently placed under pressure (by the care recipient him/herself or other family members) to fulfil the expected traditional roles regardless of the personal cost or consequences. Particularly if there are divergent notions of familial obligation between different family members and/or the informal carer refuses to accept the assigned traditional role, role conflicts and family disagreement emerge with possible consequences on the stability of the care arrangement.
Regarding the cultural/societal context, distinctive changes in the near future are expected. The profound demographic changes, increasing number of elderly people, reduction in family size, geographic spread of family members and workplace participation (of women) could result in the older generation relying on fewer potential family caregivers and a smaller amount of informal resources than before. In relation to cultures still bound to traditional values, ‘modern’ families may no longer be able to maintain the traditional roles or may consciously shift from the traditional values held by older generations.

Healthcare system

Conceptualisation

The national policy agenda, the associated promotion of specific formal support services and the availability and financing of such services affect the informal carer’s options to use or reject the use of formal services. In addition, the national policy context and the respective healthcare system are shaped by the perceived cultural and societal values. This context influences the stability of home-based care arrangements.

Analyses and synthesis

Dramé and colleagues identified the following universally accepted principles in all member countries of the Organisation for Economic Cooperation and Development: remaining at home for as long as possible, delaying institutionalisation, supporting informal carers and coordinating services at a local level whenever possible. However, how these principles are incorporated into national policy agendas and implemented in healthcare insurance systems and care practice widely differ across different cultures and countries worldwide. Some national policy agendas rely on extensive family participation (eg, Asia, eastern and southern Europe, Germany and the USA), while in other agendas, the additional integration of formal support services (including institutional care) is highly accepted (eg, Australia, Western Europe and Scandinavia). According to these diverse policy aims, how financial resources are governed and allocated varies, which often influences the decision of informal carers to continue or end home care.

In our sample, there was a consensus that most home-based care arrangements require some type of formal support services at some point during the trajectory. Therefore, the use of formal support services may either function as a protective factor for stability because the unmet needs of the care arrangements are addressed or accelerate the termination of home-based care because positive experiences with the increasing integration of formal support services pave the way for the final decision to institutionalise the person with dementia. Informal carers particularly value education and training, case management, home care services, respite services, support groups, hospice services and financial aid to support the maintenance of a stable situation at home.

The setup of different healthcare systems and how formal services are promoted and funded seriously affect the availability and use of such services. Informal carers state a need for adequate financial aid. In the US context, Ceponi-Martin and colleagues reported that a higher percentage of spending on in-home care services and a higher number of home health agencies decreased the risk of nursing home placement. In the Italian context, Sansoni and colleagues argued that the coverage of nursing home costs by the state could increase the rate of nursing home placement. Families who cannot afford to pay for a nursing home avoid placement and tend to continue care at home, but if the costs for institutional care are covered by the state, families with a lower income tend to institutionalise their relatives earlier, while families with a higher income more often continue home-based care as they can pay for community-based services out of pocket.

In summary, the existence and availability of formal support services and adequate financial resources (either covered by a healthcare system or private means) allow informal carers to decide which formal support services support the maintenance of stability in their current situations.

DISCUSSION

In this meta-study, we developed a middle-range theory of the ‘Stability of home-based care arrangements for people living with dementia’ (SoCA-Dem theory) that emphasises the perspective of the informal carer. We conceptualised stability as a complex phenomenon comprising the following three components: (1) the trajectory of the home-based care arrangement comprising a cyclic process of change and balancing over time; (2) the characteristics of the care arrangement, including the concepts needs, carer role, dyadic relationship and resources; and (3) the context, including the concepts of society and culture and the healthcare system. Most concepts of the proposed middle-range theory have been well studied individually but have not been examined from an integrative perspective. The strength of this study and additional value of our new middle-range theory are that this study links the concepts to the phenomenon of stability and sheds light on the dynamics linking the components and concepts.

The following three ongoing debates in the scientific community became obvious in our meta-theory and meta-method analyses, influenced our conceptualisation of stability and allowed us to recognise some limitations: (1) the debate concerning predominant research focusing on dyads (mostly from the perspective of the informal carer) versus a more heterogeneous understanding of care arrangements; (2) the debate concerning the predominant stress/burden perspective in research focusing on informal care versus a framing of informal care that also considers positive aspects; and (3) the debate concerning the predominance of retrospective and cross-sectional...
studies versus the need for prospective and longitudinal studies.

1. As a consequence of the predominant focus of most studies in our sample on informal carers, our middle-range theory primarily reflects the perspective of one actor in the dyadic care arrangement and highlights the pivotal role of the informal carer in the creation and maintenance of stability. This emphasis on the informal carer can be interpreted as a limitation that, to some extent, may have been triggered by our search strategy in which the involvement of an informal carer in the care arrangement was chosen as an inclusion criterion. Although we were actually interested in the perspectives of all actors in dyadic care arrangements, the included publications nearly exclusively adopted the perspective of the informal carer and failed to provide insight into the perspectives of the person with dementia and other actors in the care arrangement. There is a vivid debate regarding this prevalent shortcoming in research concerning informal dementia care. For example, Ceci and colleagues\textsuperscript{106} criticised that the heterogeneity of care arrangements is often overlooked. We do not question that the person with dementia has a voice and plays an active role in the creation and maintenance of his or her care arrangement\textsuperscript{105}; we are aware that there is a growing population of people with dementia who live alone and need to cope without the help of an informal carer\textsuperscript{106}; and we know that most home-based care arrangements rely on the support of formal service providers and have a triadic structure.\textsuperscript{107} Home-based care arrangements are often diverse ‘convoys of care’ and will likely become even more diverse in the future due to demographic shifts and changes in gender roles and family structures. In subsequent research, this middle-range theory should be extended to include diverse actors’ perspectives regarding stability, ideally through participative research approaches.\textsuperscript{106} 110 However, even in its present form, the SoCA-Dem theory is open to a heterogeneous conceptualisation of care arrangements as it already positions diverse informal and formal support networks as pivotal resources for the dyadic care arrangement. In addition, we assume that this middle-range theory may also be valid in the context of diverse chronic diseases. Currently, we hypothesise that our middle-range theory can be applied in future research focusing on more diverse care arrangements and other chronic diseases, but the components and concepts of the theory might change in their meaning and forming.

2. Many publications included in our sample highlighted the progression of dementia as the main driver of change leading to a growing care dependency of the person with dementia and increasing stress and burden for the informal carer. This assumption is mirrored by a high number of publications in our sample that (explicitly or implicitly) used a stress/burden framework as their theoretical perspective and therefore relied on a dominant but increasingly questioned research paradigm.\textsuperscript{111} Informal carer burden has often been identified as a main outcome to be studied and targeted in intervention studies, but it has been criticised that this approach usually leads to a mismatch between the simple research logic of ‘identified problem – targeted outcome’ and the individual and much more complex real-life experiences of dyads living at home.\textsuperscript{112} In-depth qualitative research focusing on the structure and underlying meanings of informal caring has been found to be very valuable in this respect. For example, von Kutzleben\textsuperscript{21} revealed that informal carers follow their own logics and strategies in response to change. Informal caring proved to be the expression of an individual informal care concept, which can be defined as the intrapersonal manifestation of motives, aims and personal boundaries. Considering these individual informal care concepts helps enhance our understanding of how informal carers perceive burden and that the reduction in burden is not necessarily the most important priority of all carers. The authors of the publications included in our meta-study had very diverse understandings of the phenomena of stress and burden. This conceptual ambiguity has also been critically discussed in the literature,\textsuperscript{113} and in our case, it complicated a coherent interpretation of the study results regarding burden and its influence on stability. Although the stress/burden perspective was dominant in our sample, in some included studies, the nascent paradigm shift to a more multifaceted understanding of informal dementia care, including positive experiences, was apparent.\textsuperscript{5} The SoCA-Dem theory emphasises the coexistence of both positive and negative experiences; for meaningful interpretation, burden and positive experiences always need to be contextualised within the complex interplay of all theory components and concepts.

3. Several up-to-date phase models\textsuperscript{21} 114 115 emphasise constant change in the everyday lives of dyads living at home, followed by ongoing adaptation in a dynamic non-linear process. Our views are consistent with those described by von Kutzleben,\textsuperscript{21} who highlighted the importance of distinguishing between different phases of the dementia care trajectory as in each phase, a different type of change may affect stability in a specific way. According to this study, the middle stages of the trajectory during which the person with dementia is still quite actively involved in everyday life but already has difficulties mastering the activities of daily living independently might be perceived as particularly challenging with regard to the maintenance of stability. In some studies in our sample, the research was contextualised using theories concerning chronic illness trajectories\textsuperscript{119} and transitions,\textsuperscript{116} and 34 longitudinal studies showed that researchers valued following dyads over a certain time span. However, there are methodological challenges to assessing the development of trajectories, even in longitudinal research. The definite beginning of a trajectory is difficult to determine. Consequently,
the studies in our sample largely included care arrangements in various and undefined phases within their individual trajectories and did not acknowledge this issue as a limitation. Furthermore, most studies focused on rather late phases of the trajectory, often on the phase during which a transition to an institution occurred. To generate deeper knowledge of stability in all phases of the trajectory, as recommended by Kralik and colleagues,117 there is a need for future longitudinal studies that provide valid assessments of the current stage of the trajectory of the individual care arrangement, ideally include care arrangements at the very beginning of their trajectories and follow such trajectories over a long period. In this type of longitudinal study, the focus should be on the specific forming of each concept of our middle-range theory during each phase of the trajectory and the influence of each concept on the creation of stability.

Limitations
This meta-study has limitations. Although we critically reflected and discussed its results with peer researchers, our theoretical perspective as well as our previous empirical research influenced the development of the SoCA-Dem theory. Theoretically, we conceptualised dementia care as a trajectory that is shaped by the interaction of individuals over time.19 20 Empirically, the study by von Kutzleben21 identified the maintenance of stability as a leading principle of informal carers and shed light on this specific perspective. Furthermore, our working definition18 guided the deduction of the search strands and the preliminary coding scheme for this meta-study. Though the concepts of the new theory were created in an inductive and reflexive approach in the analysis phase, further research is needed to reduce existing bias within the SoCA-Dem theory. According to Roy,178 the circular relationship of theory to research to practice is particularly evident in middle-range theories. Their use in research and practice provides feedback for further theory development.25 Currently, we elaborate the interplay of the theory concepts on a descriptive level. We provide a middle-range theory that helps to organise and interpret the relationships of relevant concepts and that is rather constitutive than causal.26 To enhance the explanatory power of the SoCA-Dem theory, the causal relationships between the concepts need clarification beyond the current level. Another limitation concerns the dominance of research from Europe and North America in our sample. The possibility to reflect social and cultural differences was incorporated in the concept of society and culture as well as in the concept of healthcare system. Nevertheless, our SoCA-Dem theory might be influenced by a western perspective and therefore its applicability in other contexts needs to be explored. Finally, an additional limitation is grounded in the large number of included studies. To analyse and synthesise all these studies was time-consuming. For this reason, we did not perform the theoretical sampling as extensively as we planned according to the study protocol,27 and the time between the last database search and the publication of this meta-study is rather long.

Conclusion
This middle-range theory conceptualises the phenomenon of the stability of home-based care arrangements for people living with dementia. By adopting the perspective of the informal carer, the SoCA-Dem theory contributes to an enhanced understanding of the dynamic processes that contribute to stable or unstable phases within the often long-lasting dementia care trajectory.

Elaborating on the perspective of the person with dementia is a necessary important goal for future research focusing on ‘stability’ and will enhance our understanding of the identified components and concepts. In addition to the aspired inclusion of the perspective of the person with dementia and other actors involved in home-based care arrangements, we highlight the need to further clarify the conceptual overlap and interplay of the theoretical components and concepts and the need to better understand the development of stability in all phases of the dementia care trajectory. All this will contribute to an ongoing improvement of the explanatory power of the SoCA-Dem theory and to its usefulness for scientists and healthcare practitioners. In the SoCA project, we address the current knowledge gaps in subsequent empirical research applying qualitative and quantitative methods. We aim to be able to distinguish between stable and unstable care arrangements and identify or develop interventions. Already, our middle-range theory can provide guidance for healthcare providers and policy makers: the concepts of the SoCA-Dem theory define topics that need to be considered to support the stability of home-based care arrangements. Support should be adapted to the individual forming of the concepts in the respective care arrangement. Effective interventions have to take into account the specific needs of the person living with dementia and the informal carer at a particular point of time.

Finally, we encourage the scientific community, policy makers and healthcare providers to use, reflect on and refine the SoCA-Dem theory and contribute to the ongoing discourse regarding stability to better enable dyads to live in the place of their choice currently and in the future.

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Contributors
KK, JD, IH, MvK and BH substantially contributed to the conception of this meta-study, including the development of the research questions, search strategies, eligibility criteria, data extraction schemes and analytic procedures. KK, JD and IH performed the data analysis and synthesis process. The proposed middle-range theory was repeatedly discussed, refined and finalised as a team by all authors. KK, JD and IH contributed to the draft of this manuscript. All authors read, provided feedback and approved the final manuscript.

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ORCID ids

Kerstin Köhler http://orcid.org/0000-0002-8681-565X
Jan Dreyer http://orcid.org/0000-0002-9400-4787
Iris Hochgraeber http://orcid.org/0000-0003-1425-555X
Milla von Kutzleben http://orcid.org/0000-0002-6443-2593
Christianne Pinkert http://orcid.org/0000-0002-2692-1671
Martina Roes http://orcid.org/0000-0003-4531-8584
Bernhard Holle http://orcid.org/0000-0003-2549-7765

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Appendix ‘Towards a middle-range theory of ‘Stability of home-based care arrangements for people living with dementia’ (SoCA-Dem): Findings from a meta-study on mixed research’

Kerstin Köhler1,2, Jan Dreyer1,2, Iris Hochgraeber1,2, Milena von Kutzleben3,1, Christiane Pinkert1, Martina Roes1,2, Bernhard Holle1,2

1 German Center for Neurodegenerative Diseases (DZNE), Witten, Germany
2 Witten/Herdecke University, Faculty of Health, School of Nursing Science, Witten, Germany
3 Carl von Ossietzky Universität Oldenburg, Faculty of Medicine and Health Sciences, Department of Health Services Research, Oldenburg, Germany

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Appendix A: Systematic database searches

We searched the databases MEDLINE, CINAHL and PsycINFO with nine separate search strands:

1. search strand stability
2. search strand (self-)caring actions
3. search strand needs & demands
4. search strand crises
5. search strand positive aspects
6. search strand end-of-life-care
7. search strand institutionalization
8. search strand burden
9. search strand interventions

All search syntaxes consist of similar domains. The first domain addresses the specific topic of the search strand. The next three domains define the population (the person with dementia, the informal carer) and the setting (community setting). The last domain defines the outcomes (stability OR institutionalization). Depending on the respective search strand, these domains were used in different ways to either increase or decrease the sensitivity of the search. Thereby, at least the syntax for the specific search strand and the search domain for dementia were used for every search strand.

Exemplary, the search syntaxes for all search strands for the database search in MEDLINE via PubMed are displayed. No filters were applied.

1. Search strand ‘Stability’

2. **Search strand ‘(Self-)caring action’**


3. **Search strand ‘Needs and demands’**


4. **Search strand ‘Crises’**


5. **Search strand ‘Positive aspects’**


6. **Search strand ‘End-of-life care’**


7. **Search strand ‘Institutionalisation’**

8. **Search strand 'Burden'**


9. **Search strand 'Interventions'**

Appendix B: Inclusion criteria and quality appraisal

To be included in the meta-study, the studies had to fulfil the following inclusion criteria:

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<th>MANDATORY</th>
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<td>• study focuses on care arrangements for people living with dementia</td>
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<td>• involvement of informal carer(s) in care arrangement</td>
</tr>
<tr>
<td>• community setting (or institutional setting if study is retrospective or</td>
</tr>
<tr>
<td>reflects the transition process)</td>
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<tr>
<td>• written in English, German, or French</td>
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AND

EITHER...

<table>
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<th>... OR ...</th>
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<tr>
<td>Study defines or researches stability of home-based care arrangements for</td>
</tr>
<tr>
<td>people living with dementia.</td>
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</table>

OR...

| Study defines or analyses one of the following concepts or phenomena       |
| included in the working definition of stability:                          |
| • (self-)caring actions                                                   |
| • needs/demands of people living with dementia and their carer(s)         |
| • dynamics in care arrangements                                           |
| • crises                                                                  |
| • successful caring routines                                             |
| • transition to an institutional setting                                 |

... AND ...

| Study makes a statement whether the person living with dementia can stay at |
| home (longer/better) or not.                                               |

The decision to include or exclude a publication was made in light of the contribution of the publication to the synthesis. Nevertheless, to avoid the inclusion of ‘fatally flawed’ publications, the following questions suggested by Dixon-Woods and colleagues (2005) were applied to judge the methodological quality of the publications:

- Are the aims and objectives of the research clearly stated?
- Is the research design clearly specified and appropriate for the aims and objectives of the research?
- Do the researchers provide a clear account of the process by which their findings were produced?
- Do the researchers display enough data to support their interpretations and conclusions?
- Is the method of analysis appropriate and adequately explicated?

Reference:

Appendix C: Statement of the reasons for the exclusion of the publications from the 'interventions' search strand

In the literature search one of our nine search strands was the 'interventions' strand. Our motive to operate this search strand was to include intervention studies or reviews that used the 'odds of institutionalisation' and/or 'time to institutionalisation' (in this case interpreted as a manifestation of stability/instability) as outcomes. We expected that these intervention studies and reviews might explicate in detail why and how the respective intervention was supposed to and has succeeded or failed to prolong a stable situation at home. Actually, the retrieved intervention reviews (Brodaty, Green, & Koschera, 2003; Olazaran et al., 2010; Pimouguet, Lavaud, Dartigues, & Helmer, 2010; Pinquart & Sörensen, 2006; Reilly et al., 2015; Smits et al., 2007; Spijker et al., 2008; Zabalegui et al., 2014) suggest, that (in particular multicomponent) interventions may decrease the odds of institutionalisation and increase the time that dyads spent at home. But both, intervention studies and reviews, remained vague in their statements with relation to our target phenomenon ‘stability’ in general and specifically to its constitutive concepts. Therefore, they did not contribute meaningful to the creation of the midrange theory, and in deviation to the study protocol we decided during the analysis process to remove the whole search strand from the meta-study sample. This experience with the 'interventions' search strand points to an ongoing debate on an emerging reorientation of intervention research in the field of home-based and informal dementia care that should to be considered in future research (e.g. Ceci, Symonds Brown, & Judge, 2018; Dawson, Bowes, Kelly, Velze, & Ward, 2015).

Literature:


Appendix D: Overview of the meta-study sample

The following table gives an overview of the n=99 publications that we included in this meta-study. For each included publication we list the author(s), year of publication, country, search strand, aim, study design, sample/setting, main findings, and contribution to theory concepts. In addition to the study design, we indicate for quantitative studies if they have a longitudinal or a cross-sectional approach. For all other studies we only indicate if they have a longitudinal approach. In the column 'contribution to theory concept(s)' we indicate to which concept(s) of our middle-range theory the respective publication contributed. The concepts of the theory are: change, balancing, needs, carer role, dyadic relationship, resources, society and culture, health care system. Thereby, concepts that are addressed very prominent in a publication are displayed in bold type; further addressed concepts are displayed in regular type. Following the principles of conceptual saturation within the coding process, we did not code all information on each concept in each publication. For this reason, it is probable, that a publication includes information with regard to a theory concept, but the concept is not listed in the respective column.

### Abbreviations

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<th>Abbreviation</th>
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<tr>
<td>AChEI</td>
<td>Acetylcholinesterase inhibitor</td>
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<td>AD</td>
<td>Alzheimer’s disease</td>
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<tr>
<td>ADL</td>
<td>Activity of daily living</td>
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<tr>
<td>DLB</td>
<td>Dementia with Lewy bodies</td>
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<tr>
<td>FC</td>
<td>Formal carer</td>
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<tr>
<td>IC</td>
<td>Informal carer</td>
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<tr>
<td>GP</td>
<td>General Practitioner</td>
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<tr>
<td>HRQoL</td>
<td>Health Related Quality of Life</td>
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<tr>
<td>LOD</td>
<td>Late onset dementia</td>
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<tr>
<td>MMSE</td>
<td>Mini Mental State Examination</td>
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<tr>
<td>NH</td>
<td>Nursing home</td>
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<td>NHP</td>
<td>Nursing home placement</td>
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<tr>
<td>NPI</td>
<td>Neuropsychiatric Inventory</td>
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<tr>
<td>PwAD</td>
<td>Person with Alzheimer’s disease</td>
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<tr>
<td>PwD</td>
<td>Person with dementia</td>
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<tr>
<td>YOD</td>
<td>Young onset dementia</td>
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### Table

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<thead>
<tr>
<th>Author (Year) (Country)</th>
<th>Search Strand</th>
<th>Aim</th>
<th>Study Design</th>
<th>Sample/Setting</th>
<th>Main Findings</th>
<th>Contribution to theory concept(s)</th>
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</thead>
<tbody>
<tr>
<td>Afram et al. (2014) (International Consortium: Estonia, Germany, Finland, France, Netherlands, UK, Spain, Sweden)</td>
<td>Berries</td>
<td>to explore reasons for institutionalisation of PwDs according to ICs as well as variation in reasons between countries</td>
<td>mixed methods</td>
<td>786 ICs (PwD institutionalised)</td>
<td>Mainly PwD-related reasons were stated (neuropsychiatric symptoms, care dependency, cognition). Neuropsychiatric symptoms were among the most often mentioned reasons in most countries. Besides PwD-related reasons, IC burden and the inability to care were stated. Countries differ significantly in reasons according to ICs. Spouses more often stated reasons related to themselves compared with child ICs.</td>
<td>change, carer role, resources, society and culture, health care system</td>
</tr>
<tr>
<td>Afram et al. (2015) (Netherlands)</td>
<td>Needs &amp; Demands</td>
<td>to provide insights into problems and needs of ICs of PwDs during care transition from home-based care to institutional long-term care</td>
<td>systematic review: qualitative synthesis</td>
<td>13 publications</td>
<td>Regarding the needs and problems of ICs during the care-transition period, the most stated topics were emotional concerns, knowledge/information, and support. The care-transition period should be considered as a continuum, as similar needs and problems were identified prior and after admission to long-term-care.</td>
<td>needs, carer role, resources, health care system</td>
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<td>Andrén and Elmståhl (2008) (Sweden)</td>
<td>Institutionalisation</td>
<td>to determine the effectiveness of a psychosocial intervention for ICs in delaying institutionalisation of PwDs</td>
<td>quantitative: experimental (longitudinal)</td>
<td>intervention group: 153 ICs, control group: 155 ICs</td>
<td>There were significant delays (6 months) in institutionalisation for PwDs with adult children as ICs. Factors associated with institutionalisation are the ICs’ influence on intervention, IC gender and the PwD severity of dementia.</td>
<td>change, dyadic relationship</td>
</tr>
<tr>
<td>Badrakalimuthu et al. (2014) (UK)</td>
<td>End-of-life-care</td>
<td>to undertake a systematic review and narrative synthesis of the literature concerning place of death of PwDs, and the PwDs, ICs and health-care-providers preferences for place of death of PwDs</td>
<td>systematic review: narrative review</td>
<td>6 publications</td>
<td>Studies of death certificate data show that PwDs die more commonly in NHs than in other locations. In contrast, prospective studies show that death is more common in own residence or hospital. Older age, male gender, availability of hospital were associated with hospital death, availability of NH beds was associated with death in NH and enrolment in hospice was associated with death in own residence. Little is known about patient, IC and health professional preferences or the extent to which PwDs discuss their preferences with their families and healthcare providers. Preferences of proxy decision-makers are met more often than those of the PwDs.</td>
<td>needs, resources, health care system</td>
</tr>
<tr>
<td>Bakker et al. (2013) (Netherlands)</td>
<td>Institutionalisation</td>
<td>to investigate time from symptom onset to institutionalisation and predicting factors for PwDs with YOD compared to PwDs with LOD</td>
<td>quantitative: observational (longitudinal)</td>
<td>308 dyads, 226 dyads YOD, 102 dyads LOD (PwD living at home at baseline)</td>
<td>Time from symptom onset to institutionalisation was nearly 9 years for PwDs with YOD and 4 years for PwDs with LOD. Time to institutionalisation was significantly predicted by apathy in the YOD group and by IC’s competence in caring for the PwDs in both groups.</td>
<td>change, carer role, dyadic relationship, resources</td>
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<tr>
<td>Caldwell et al. (2014) (Australia)</td>
<td>Institutionalisation</td>
<td>to investigate the decision-making process for placing a PwD on a waiting list for a NH; why offers of a place in a NH are accepted or declined; the influence of cultural factors in people living in Australia from a Chinese background compared with those from an English-speaking background</td>
<td>qualitative: unspecific design</td>
<td>27 ICs (PwD institutionalised or on a NH waiting list)</td>
<td>ICs were at different stages of decision-making when they applied for a waiting list (ready for placement, just in case, urgent need for placement). IC’s decisions were influenced by their emotions and expectations of NHs. The decision-making process was similar for both cultural groups, but Chinese ICs spoke more about their sense of duty, the need for a Chinese specific facility, and declining a place because of family disagreement.</td>
<td>change, balancing, carer role, dyadic relationship, society and culture, health care system</td>
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<tr>
<td>Caron et al. (2006) (Canada)</td>
<td>Institutionalisation</td>
<td>to explore the decision-making process with regards to institutionalisation of a PwD, from the perspective of the IC; to develop a theoretical model</td>
<td>qualitative: Grounded Theory</td>
<td>14 ICs for 8 PwDs (PwD institutionalised)</td>
<td>A theoretical model of the decision-making process was developed and three factors that were central to the process were identified: IC’s perceptions of their ability to provide care, IC’s evaluations of the PwD to make care decisions, and the evolving influence of contextual factors and interactions with healthcare professionals.</td>
<td>change, balancing, needs, carer role, resources</td>
</tr>
<tr>
<td>Cepoiu-Martin et al. (2016) (Canada)</td>
<td>Institutionalisation</td>
<td>to summarise the effects of various individual, IC and system-related factors on the risk of institutionalisation of PwDs residing in the community</td>
<td>systematic review: narrative review and meta-analysis</td>
<td>review part: 59 publications, meta-analysis: 37 publications</td>
<td>White race, dementia severity and older age increased the risk of institutionalisation. Married PwDs living with their carer had a lower risk. Behavioural and psychological symptoms of dementia, the degree of functional impairment, and carer burden had a consistent effect on the risk of institutionalisation.</td>
<td>change, dyadic relationship, resources, society and culture, health care system</td>
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<tr>
<td>Chang et al. (2010) (USA, data collection in Taiwan)</td>
<td>Institutionalisation</td>
<td>to understand the process and difficulties that Chinese ICs experience when making a NH placement decision for a PwD</td>
<td>qualitative: Grounded Theory</td>
<td>30 Chinese ICs in Taiwan (PwD institutionalised)</td>
<td>A stage-based model describes how ICs go through the decision-making process (initiating the placement decision, assessing and weighing the decision, finalising the decision, evaluating the decision), how they overcome difficulties to reach decisions, and how the decision-making process is related to the concept of Chinese filial piety, which is changing in modern Taiwanese society.</td>
<td>change, balancing, carer role, resources, society and culture, health care system</td>
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<tr>
<td>Chang et al. (2011) (USA, data collection in Taiwan)</td>
<td>Institutionalisation</td>
<td>to describe factors influencing decisional conflict among Chinese ICs regarding NH placement of PwDs</td>
<td>qualitative: unspecified design</td>
<td>30 Chinese ICs in Taiwan (PwD institutionalised)</td>
<td>Factors influencing IC’s decisional conflict included the Chinese value of filial piety, limited financial resources and information, placement willingness of the PwD, family disagreement, distrust of NH care quality, and limited NH availability. Factors influencing IC’s decisional conflict post placement included disappointment with NH care quality and self-blame for the placement decision.</td>
<td>change, balancing, carer role, resources, society and culture, health care system</td>
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<tr>
<td>Coe et al. (1999) (Canada)</td>
<td>Needs &amp; Demands</td>
<td>to explore male ICs perceptions of formal support</td>
<td>qualitative: Grounded Theory (longitudinal)</td>
<td>24 male ICs (PwD living at home or institutionalised)</td>
<td>The male ICs experienced a process of making concessions for care in four stages: resisting, giving in, opening the door, making the match. Personal barriers deterred the ICs from seeking help, and once the ICs engaged formal help, the influence of characteristics of the health care system and FCs had both enabling and disabling effects. For ICs who institutionalised, the phase of making the match was followed by redesigning their IC role.</td>
<td>balancing, needs, carer role</td>
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<tr>
<td>Cohen et al. (1994) (Canada)</td>
<td>Positive aspects</td>
<td>to review different concepts and measures of positive aspects of caring, and to describe the use of a simple measure of 'enjoyable aspects of caregiving'</td>
<td>quantitative: observational (longitudinal)</td>
<td>196 dyads (PwD living at home at baseline)</td>
<td>The measure of 'enjoyable aspects of caregiving' attained significant test-retest reliability and significantly correlated with IC burden and health. It also predicted IC desire to institutionalise but not actual institutionalisation. The paper calls for greater use of measures related to positive aspects and a better understanding of how these measures relate to one another and to outcomes.</td>
<td>carer role, dyadic relationship, resources</td>
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<tr>
<td>Cox (1996) (USA)</td>
<td>Institutionalisation</td>
<td>to identify what factors influence the IC hospital discharge decision and the IC satisfaction with the discharge plan, and to analyse if influences/satisfaction differ between discharges to home or to NH</td>
<td>quantitative: observational (cross-sectional)</td>
<td>172 ICs (PwD in hospital)</td>
<td>Social workers are important influencers in the decision to discharge from hospital to home or to NH. IC satisfaction depends on the amount of involvement in the discharge process.</td>
<td>change, balancing, needs, resources, health care system</td>
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<tr>
<td>Daly et al. (2012) (Ireland)</td>
<td>Berries</td>
<td>to develop a theory to explain the social processes employed by ICs to manage alterations to interactions within their social worlds</td>
<td>qualitative: Grounded Theory</td>
<td>20 ICs, 1 PwD (PwD living at home), 10 FCs from the health, social care and disability sectors</td>
<td>IC’s main concern was identified as ‘Living on the fringes’, which was stimulated by dementia-related stigma and living a different life. The theory of ‘Sustaining Place’ explains the social pattern of actions employed by ICs to manage this problem on behalf of themselves and the PwD.</td>
<td>change, balancing, needs, society and culture</td>
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<td>Davis et al. (2014) (USA)</td>
<td>(Self-)caring actions</td>
<td>to examine the conceptualisation of care problems by AD and by Parkinson’s disease spouse ICs and the efficacy of their problem management strategies, to propose a typology of IC management styles that can provide a contextual frame for understanding their experiences of burden</td>
<td>qualitative: unspecified design</td>
<td>12 ICs of a partner with: 6 AD and 6 Parkinson’s disease (living at home)</td>
<td>Three types of IC spouses are identified: adapters, strugglers, and case managers. The findings suggest that IC burden might be influenced more by the IC management style than by the demands of the care situation.</td>
<td>balancing, carer role, resources</td>
</tr>
<tr>
<td>De Vugt et al. (2005) (Netherlands)</td>
<td>Burden</td>
<td>to examine the impact of specific behavioural disorders and IC’s emotional reaction on institutionalisation</td>
<td>quantitative: observational (longitudinal)</td>
<td>119 dyads (PwD living at home)</td>
<td>IC distress related to PwD behaviour was a significant predictor of institutionalisation, while behaviour did not predict NHP. The study did not find a differential impact of specific aspects of PwD behaviour. Child ICs, especially daughters, were associated with shorter time to institutionalisation compared to spouse ICs.</td>
<td>change, dyadic relationship</td>
</tr>
<tr>
<td>Donovan (2006) (USA)</td>
<td>Positive aspects</td>
<td>to identify and understand the positive aspects of the lived experiences of ICs</td>
<td>qualitative: phenomenological approach</td>
<td>15 spousal ICs (PwD living at home)</td>
<td>The study revealed more positive than negative aspects of caring. ICs indicated positive aspects of caring through their behaviour and communication. A mélange of different actions facilitated the ICs to gain a feeling of competence and satisfaction in their carer role.</td>
<td>balancing, carer role, dyadic relationship, resources</td>
</tr>
<tr>
<td>Donovan et al. (2010) (USA)</td>
<td>Positive aspects</td>
<td>to give an in-depth description of how uplifted ICs think about care and conduct care</td>
<td>qualitative: phenomenological approach</td>
<td>15 spousal ICs (PwD living at home)</td>
<td>The study identified two primary themes of IC thinking and acting: (1) engaging in positive behaviours and (2) making adjustments in attitudes.</td>
<td>balancing, carer role, dyadic relationship, resources</td>
</tr>
<tr>
<td>Dramé et al. (2012) (France)</td>
<td>Institutionalisation</td>
<td>to identify factors predictive of institutionalisation over a period of 1 year among PwDs</td>
<td>quantitative: observational (longitudinal)</td>
<td>425 PwDs (PwD living at home at baseline)</td>
<td>There were 4 factors increasing the risk of institutionalisation: age 85 or older, inability to use the toilet, balance disorders and living alone. 3 factors decreased the risk: inability to transfer, increased number of children and increased initial MMSE score.</td>
<td>change, society and culture, health care system</td>
</tr>
<tr>
<td>Ducharme et al. (2012) (Canada)</td>
<td>Institutionalisation</td>
<td>to develop an explanatory model of the IC decision-making process regarding NHP of a PwD</td>
<td>qualitative: Grounded Theory (longitudinal)</td>
<td>18 ICs (7 PwD living at home, 11 PwD living in a retirement residence)</td>
<td>A theoretical model illustrates that the institutionalisation decision-making process is activated when the IC or a health care professional introduces the possibility of institutionalisation. The IC’s assessment of the pros and cons of institutionalisation is modulated primarily by interactions with the formal and informal social network.</td>
<td>change, balancing, resources, health care system</td>
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<tr>
<td>Ebly et al. (1999) (Canada)</td>
<td>Institutionalisation</td>
<td>to describe differences in short-term institutionalisation rates for PwDs who live alone as compared to PwDs residing with an IC</td>
<td>quantitative: observational (longitudinal)</td>
<td>317 dyads (PwD living at home at baseline)</td>
<td>One third of PwDs lived alone. Thinking about institutionalisation (intent) was predictive for actual short-term (2-year) institutionalisation and appeared to be influenced by living arrangements. ICs of PwDs living alone provided less hands-on assistance, experienced less burden, and were less likely to be depressed than those living with the PwD, but were more likely to have considered institutionalisation, driven by concerns about safety and support.</td>
<td>change, carer role, dyadic relationship</td>
</tr>
<tr>
<td>Feldman et al. (2009) (International Consortium: Belgium, Canada, Finland, France, Germany, UK)</td>
<td>Institutionalisation</td>
<td>to evaluate patient and treatment (galantamine and other AChEIs) factors associated with the time until institutionalisation</td>
<td>quantitative: experimental (longitudinal)</td>
<td>548 PwDs (PwD living at home at baseline)</td>
<td>Long-term treatment with galantamine or other AChEIs appears to be associated with a significant delay in the time to institutionalisation in PwDs. For each year of treatment the risk within a given period was reduced by 31% (galantamine) and 29% (other AChEI).</td>
<td>change, resources, health care system</td>
</tr>
<tr>
<td>Forbes et al. (2012) (Canada)</td>
<td>Needs &amp; Demands</td>
<td>to enable FCs, ICs, and PwDs (living in rural areas) to use dementia care information more effectively by examining their information needs, how these change over time, and how they access, assess, and apply the knowledge</td>
<td>qualitative: unspecific design (longitudinal)</td>
<td>5 PwDs (living at home), 14 ICs, 14 FCs</td>
<td>6 stages of the dementia care journey were described. ICs identified the need for different types of knowledge during each stage. Barriers to knowledge exchange included: lack of rural community-based services; ICs reluctant to seek help and had limited energy; and lack of integration of dementia-related services and supports. Facilitators included: ICs with healthcare experience who were actively seeking information; development of trusting relationships between FCs, ICs, and PwDs; and formal mechanisms for exchanging information within and across rural community-based organisations.</td>
<td>balancing, needs</td>
</tr>
<tr>
<td>Frewer-Graumann (2014) (Germany)</td>
<td>(Self-)caring actions</td>
<td>to explore the construction of informal home-based care arrangements from the perspective of ICs</td>
<td>qualitative: Grounded Theory</td>
<td>14 dyads (PwD living at home)</td>
<td>Home-based care arrangements for PwDs are constructed by ICs in very different ways. Care arrangements are influenced by the progression of dementia, family dynamics, coping strategies, inner familiar relationships, professional occupation of ICs and secondary and tertiary social networks.</td>
<td>change, balancing, carer role, dyadic relationship, resources</td>
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<tr>
<td>Gallagher et al. (2011) (Ireland)</td>
<td>Institutionalisation</td>
<td>to determine whether modifiable IC attributes are significantly associated with desire to institutionalise; and whether these variables may be independent predictors in the context of other established PwD and IC-related variables</td>
<td>quantitative: observational (cross-sectional)</td>
<td>102 dyads (PwD living at home)</td>
<td>IC desire to institutionalise was significantly associated with a number of modifiable variables including IC coping style, self-efficacy, depression, burden, and the presence of an unmet service need. In a multivariate analysis, IC burden, IC depression, and non-spousal status were the only significant independent predictors for desire to institutionalise.</td>
<td>needs, dyadic relationship, resources</td>
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<tr>
<td>Gaugler et al. (2005a) (USA)</td>
<td>Institutionalisation</td>
<td>to determine how behaviour problems that occur early in the caring influence time to NHP and change into burden and depression over time</td>
<td>quantitative: observational (longitudinal)</td>
<td>4761 dyads (PwD living at home at baseline)</td>
<td>ICs who have to manage frequent behaviour problems early, were more likely to institutionalise. ICs who were faced with severe, early behaviour problems reported greater increases in burden and depression over the 3 years study.</td>
<td>change, carer role</td>
</tr>
<tr>
<td>Gaugler et al. (2005b) (USA)</td>
<td>Needs &amp; Demands</td>
<td>to determine how unmet needs for activity of daily living tasks influenced NHP, death, or loss to follow-up in dementia</td>
<td>quantitative: observational (longitudinal)</td>
<td>5831 dyads (PwD living at home)</td>
<td>Greater unmet needs were predictive for institutionalisation, death, and loss to follow-up. Unmet needs may be useful in identifying PwDs at risk of institutionalisation and death.</td>
<td>needs</td>
</tr>
<tr>
<td>Gaugler et al. (2007) (USA)</td>
<td>Positive aspects</td>
<td>to examine the concept of resilience in dementia caring, and determine how resilience is potentially predictive of key exit transitions from at-home dementia care</td>
<td>quantitative: observational (longitudinal)</td>
<td>1979 ICs (PwD living at home at baseline)</td>
<td>The present study utilized data from ICs to ascertain whether resilience influences transitions from PwD caring, such as institutionalisation, care recipient death, or loss to follow-up. It was revealed that high baseline resilience (low burden, high care demands) was associated with less frequent institutionalisation and loss to follow-up as well as more frequent care recipient mortality.</td>
<td>resources</td>
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<tr>
<td>Gaugler et al. (2001) (USA)</td>
<td>Institutionalisation</td>
<td>to determine factors that predict the problems and help ICs perceive while institutionalising the PwD</td>
<td>quantitative: observational (longitudinal)</td>
<td>185 ICs (PwD living at home at baseline)</td>
<td>The personal and social resources of ICs prior to NHP (e.g., mastery, family conflict, socioemotional support, duration of care, education, and physical health) were predictors of difficulties during searching for a NH, perceived family help, and perceived professional help.</td>
<td>resources</td>
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<tr>
<td>Gaugler et al. (2011) (USA)</td>
<td>Burden</td>
<td>to determine whether IC burden is a mediator between the effects of behavioural disturbance and institutionalisation</td>
<td>quantitative: experimental (longitudinal)</td>
<td>5831 ICs (PwD living at home at baseline)</td>
<td>IC burden fully mediated the relationship between four behavioural disturbances and institutionalisation.</td>
<td>change</td>
</tr>
<tr>
<td>Gaugler et al. (2009) (USA)</td>
<td>Institutionalisation</td>
<td>to identify factors that predict NHP of PwDs</td>
<td>systematic review: meta-analysis</td>
<td>80 publications</td>
<td>Identification of numerous factors (with regard to PwDs and their ICs) that predict institutionalisation. PwD and IC factors should be considered when assessing the risk of institutionalisation.</td>
<td>change, carer role, dyadic relationship, resources</td>
</tr>
<tr>
<td>Gilhooly (1986) (UK)</td>
<td>Institutionalisation</td>
<td>to shed light on features of caring which are associated with IC’s preference for institutional care</td>
<td>quantitative: observational (cross-sectional)</td>
<td>48 ICs (PwD living at home), 2 groups: co-resident and non-resident</td>
<td>The dyadic relationship is an important determinant of motivation to continue care at home. In addition, variables associated with institutionalisation included age of ICs, IC’s contacts with friends, satisfaction with help from relatives, another dependent relative, and the employment of ICs. Help from services and IC’s well-being were not correlated with preference for institutionalisation.</td>
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<td>Glass (2016) (USA)</td>
<td>End-of-life-care</td>
<td>to document and examine the experiences of end-of-life care of PwDs and ICs and the role that environment and hospice play in these experiences</td>
<td>qualitative: phenomenological approach</td>
<td>4 ICs (PwD living at home or institutionalised)</td>
<td>The cases reveal a range of end-of-life experiences, suggesting that there is not just one ‘good’ path. The extent of care needed, the responsiveness of the individual, the health of the ICs, and the residence and support situations can all intersect in a variety of ways that make no single scenario the answer for all.</td>
<td>balancing, needs, resources</td>
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<tr>
<td>Gort et al. (2007) (UK)</td>
<td>Burden</td>
<td>to analyse the Zarit Burden Scale’s ability to identify signs of IC collapse</td>
<td>quantitative: observational (cross-sectional)</td>
<td>66 ICs (PwD living at home)</td>
<td>The Zarit Burden Scale was useful to identify IC burden and to predict collapse of IC.</td>
<td>change</td>
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<tr>
<td>Groen-van de Ven et al. (2017) (Netherlands)</td>
<td>(Self-)caring actions</td>
<td>to examine the decisions made and the related key events in the trajectories of care networks including PwDs, their FCs and ICs</td>
<td>qualitative: unspecific design (longitudinal)</td>
<td>113 respondents altogether (23 PwDs, 44 ICs, and 46 FCs) (PwD living at home or institutionalised)</td>
<td>Four decision themes were constructed: managing daily life, arranging support, community living, and preparing for the future; eight key events delineate decision trajectories of PwD; decisions and key events differ between PwD living alone and living with IC.</td>
<td>balancing</td>
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<tr>
<td>Hagen (2001) (Canada)</td>
<td>Institutionalisation</td>
<td>to explore factors influencing ICs decision-making process related to placing a PwD in a NH</td>
<td>qualitative: unspecific design (longitudinal)</td>
<td>5 ICs (PwD living at home)</td>
<td>The study develops a theoretical model of factors influencing institutionalisation-related decision-making, including independence, perceived presence of others, fear of loneliness, negative NH attitudes, sense of existential self, and guilt.</td>
<td>carer role, dyadic relationship, resources</td>
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<tr>
<td>Hill et al. (2017) (UK)</td>
<td>End-of-life-care</td>
<td>to identify and describe the aspects of end-of-life care for PwDs that are most important to them and their carers</td>
<td>mixed methods</td>
<td>14 PwDs, 21 ICs (PwD living at home), 22 bereaved ICs</td>
<td>Different views on what is important about end-of-life care were identified and grouped into four viewpoints: family involvement, managing the present, pragmatic expectations and autonomy and independence.</td>
<td>balancing, needs, dyadic relationship</td>
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<tr>
<td>Hirschfeld (2003) (Israel, data collection in the USA)</td>
<td>Positive aspects</td>
<td>to present the concept of mutuality, which emerged as the crucial factor in continuing home care vs. institutionalisation</td>
<td>mixed methods</td>
<td>30 dyads (PwD living at home)</td>
<td>Mutuality emerged as the major parameter for families managing life with senile brain disease and influenced the decision to institutionalise an impaired family member.</td>
<td>balancing, dyadic relationship</td>
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<tr>
<td>Janssen et al. (2017) (International Consortium: Germany, Ireland, Italy, Netherlands, Norway, Portugal, Sweden, UK)</td>
<td>Stability</td>
<td>to identify different profiles of ICs and to investigate differences between the identified carer profiles in quality of life, in depression and in perseverance time</td>
<td>quantitative: observational (cross-sectional)</td>
<td>453 dyads (PwD living at home)</td>
<td>Five profiles of ICs were identified. These profiles differ statistically significant with regard to quality of life, depressive symptoms and perseverance time.</td>
<td>balancing, carer role, resources</td>
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<td>Author (Year) (Country)</td>
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<td>Kajiwara et al. (2015)  (Japan)</td>
<td>Positive aspects</td>
<td>to identify factors that influence the continuation of in home care of PwDs</td>
<td>quantitative: observational (cross-sectional)</td>
<td>354 ICs, 343 PwDs (PwD living at home)</td>
<td>IC’s current feelings about caring and the continuation of caring of PwDs with worsening symptoms were affected by a positive appraisal of caring and by the burden experienced by the ICs. The continuation of caring was indirectly affected by the PwDs cognitive impairment and by the behavioural and psychological symptoms of dementia score.</td>
<td>change, carer role</td>
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<tr>
<td>Kellett (1999) (Australia)</td>
<td>Institutionalisation</td>
<td>to explore the experience of family breakdown and NHP</td>
<td>qualitative: phenomenological approach</td>
<td>14 ICs (PwD institutionalised)</td>
<td>The analysed IC stories of institutionalisation uncovered 5 shared meanings: experiencing a loss of control; being disempowered; feeling guilt, sadness and relief simultaneously; possessing a sense of failure; and having to make a forced and negative choice.</td>
<td>balancing, carer role</td>
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<tr>
<td>Kraijo et al. (2014) (Netherlands)</td>
<td>Stability</td>
<td>to introduce the concept ‘perseverance time’ and to validate it in a sample of ICs</td>
<td>quantitative: observational (cross-sectional)</td>
<td>223 ICs (PwD living at home)</td>
<td>The concept of ‘perseverance time’ (Pt) is a valid measure for the IC ability and willingness to continue home care. Health of PwD, IC living apart from the patient, and male gender of IC were positively associated with Pt; need for supervision, intensity of informal care provision, and reductions in working hours and hobbies in order to be able to provide care were negatively associated.</td>
<td>change</td>
</tr>
<tr>
<td>Kraijo et al. (2015) (Netherlands)</td>
<td>Institutionalisation</td>
<td>to focus the final decision by ICs to have their PwDs placed in a NH and to question whether the admission took place at the right time in the IC’s point of view</td>
<td>mixed methods</td>
<td>14 spousal ICs (PwD institutionalised)</td>
<td>The placement decision had to be made in phases over time: placement of the PwD on a waiting list, and actual placement in a NH (stage-based model). The second and final decision often had to be taken under time pressure. The following classification emerged: Placement was at the right time, too early, too late or out of control.</td>
<td>balancing, dyadic relationship, resources</td>
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<tr>
<td>Kwon et al. (2012) (South Korea)</td>
<td>Institutionalisation</td>
<td>to explore and describe the process of NHP decision-making and adaptation among adult child carers of PwDs in Korea</td>
<td>qualitative: Grounded Theory</td>
<td>16 ICs (adult children, PwD institutionalised)</td>
<td>The core category in the process of decision making and adaptation to institutionalisation among ICs was accepting the inevitable and reorienting to changes. The developed stage-based model consisted of four phases: realizing a dead end, seeking a way out, accepting the inevitable decision, and reorienting to changes. Though ICs were confronted with various challenges, they managed to accept the decision and learned to reorient themselves to the changed care situation.</td>
<td>balancing, carer role, society and culture</td>
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<td>Ledgerd et al. (2016) (UK)</td>
<td>Crises</td>
<td>to identify the main causes of crisis and interventions to treat or prevent crisis in PwDs based on different stakeholder perspectives</td>
<td>quantitative: observational (cross-sectional)</td>
<td>719 participants: 20 academics, 562 health care sector staff, 54 ICs, 23 social care sector staff, 16 emergency service staff, 12 voluntary sector staff, 4 PwDs, 28 others</td>
<td>Wandering, falls and infection were highly rated as risk factors for crises across all stakeholder groups. Consumers rated aggression as less important, but severity of memory impairment as much more important than the other groups did. Education and support for ICs, and home care staff were highly valued for preventing crises. Well-trained home care staff, communication equipment, emergency contacts and access to respite were highly valued for managing crises.</td>
<td>change, resources, health care system</td>
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<tr>
<td>Lethin et al. (2016) (Sweden)</td>
<td>(Self-)caring actions</td>
<td>to investigate IC’s experiences of formal care when caring for a PwD through the stages of the dementia</td>
<td>qualitative: unspecific design</td>
<td>23 ICs (PwD living at home or PwD institutionalised)</td>
<td>ICs require collaboration with FCs to get support adjusted to individual’s needs, specific to stages of dementia. Three main turning points are experienced: the dementia diagnosis; when ICs realise increased need for FCs to continue caring at home; when PwD is institutionalised, ICs experience FCs reactive to needs, proactive reaction is required.</td>
<td>balancing, needs, carer role, health care system</td>
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<tr>
<td>Lewis (2015) (USA)</td>
<td>End-of-life-care</td>
<td>to discover a substantive theory that identifies the main problems that ICs of PwDs face at the end of life and the basic social processes by which they resolve that problem</td>
<td>qualitative: Grounded Theory</td>
<td>83 ICs (PwD passed away or living at home or institutionalised)</td>
<td>Carers attempt to resolve the challenge of caring at the end of life through a 5-stage basic social psychological process of rediscovering including missing the past, sacrificing self, yearning for escape, reclaiming identity, and finding joy. Health care professionals can support ICs through this journey by validating, preparing them for future stages, and encouraging natural coping strategies identified in this process.</td>
<td>balancing, carer role, dyadic relationship</td>
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<tr>
<td>Liken (2001a) (USA)</td>
<td>Crises</td>
<td>to examine crises experienced by ICs</td>
<td>qualitative: unspecific design</td>
<td>20 ICs (PwD institutionalised within the past 6 months)</td>
<td>All elements of a crisis were evident among ICs in the process of institutionalisation of the PwD. Dangerous behaviours resulting from cognitive decline led to IC fear and anxiety, which served as crisis precipitants. Perceived lack of family support in conjunction with physical and psychological exhaustion were crisis mediators. These factors, along with the IC’s failed attempts to keep the PwD at home, caused the crisis process moving toward the outcome of institutionalisation.</td>
<td>change, balancing</td>
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<td>Liken (2001b) (USA)</td>
<td>Institutionalisation</td>
<td>to examine underlying values that precipitated the NHP</td>
<td>qualitative: unspecific design</td>
<td>23 ICs (PwD institutionalised)</td>
<td>Three major themes describe values underpinning the critical incident surrounding the move of the PwD to an assisted living facility. Critical incidents centred on two major factors: the PwDs safety and the ICs emotional well-being. Conflicting values underpinning the critical incident were the PwDs autonomy versus the ICs responsibility, obligation to PwD versus obligation to self, and logic versus emotion.</td>
<td>change, balancing, needs, carer role</td>
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<tr>
<td>Lindgren (1993) (USA)</td>
<td>(Self-)caring actions</td>
<td>to conceptualise the experiences of ICs over time and to portray IC careers in the context of dementia</td>
<td>qualitative: unspecific design</td>
<td>10 ICs (PwD living at home)</td>
<td>In the IC career a process of learning, adaptation to role demands and restructuring of the IC's life evolves, and unique stages of the IC career are formed. The Encounter Stage is characterized by the need for rapid adjustment to major changes, for information about illness characteristics, and instruction in providing quality care. The Enduring Phase is the long-term, heavy-duty caring phase where supportive interventions are needed. During the Exit Stage, institutionalisation is best done when time can be taken to choose a place. Finding the right home and financial issues are two problems that need to be solved.</td>
<td>change, balancing, carer role, dyadic relationship, resources, health care system</td>
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<tr>
<td>Lloyd et al. (2016) (UK)</td>
<td>Positive aspects</td>
<td>to critically evaluate the empirical findings of qualitative studies that have explored positive aspects of caring in dementia</td>
<td>systematic review: qualitative synthesis</td>
<td>14 publications</td>
<td>Multiple positive dimensions of caring and several factors that were important in supporting their positive caring experience are identified. Conceptualisations of positive aspects of caring (PAC): role satisfaction, emotional rewards, personal growth, competence and mastery, faith and spiritual growth, relationship gains, sense of duty and reciprocity. ICs perceptions of the process of PAC: acceptance, choosing positive caregiving attitude, commitment to relationship, creating opportunities, drawing strength from faith/past challenges/others.</td>
<td>change, balancing, needs, carer role, dyadic relationship</td>
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<tr>
<td>Lord et al. (2016) (UK)</td>
<td>Institutionalisation</td>
<td>to understand the NHP decision-making process, the needs and difficulties that PwDs and ICs encounter and how these may be overcome</td>
<td>qualitative: unspecific design</td>
<td>7 PwDs, 13 ICs (PwD living at home, 1 PwD institutionalised)</td>
<td>Decision-makers balanced the competing priorities of remaining somewhere familiar, family’s wish they remain at home, reduction of risk and effects on ICs and PwDs physical health. The PwDs frequently presented their lack of autonomy as decisions about institutionalisation were made. Family consultation usually helped ICs but sometimes exacerbated tensions. Direct professional support was appreciated where it was available.</td>
<td>balancing, needs</td>
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<tr>
<td>Luppa et al. (2008) (Germany)</td>
<td>Berries</td>
<td>to review publications that analyse predictors of institutionalisation of PwDs</td>
<td>systematic review: narrative review</td>
<td>42 publications</td>
<td>Predictors of institutionalisation were classified according to a provided conceptual framework in the categories sociodemographic and relationship characteristics of change, needs, dyadic relationship,</td>
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<td>Luppa et al. (2012) (Germany)</td>
<td>Berries</td>
<td>to investigate risk factors for institutionalisation for PwDs, considering characteristics at the time of the diagnosis.</td>
<td>quantitative: observational (longitudinal)</td>
<td>254 PwDs (living at home at baseline)</td>
<td>30% of PwDs were institutionalised over the study course (6 years). Mean time until NHP was 4.1 years. Significant characteristics of NHP were marital status (being single or widowed), higher severity of cognitive impairment and mobility impairment.</td>
<td>change, dyadic relationship</td>
</tr>
<tr>
<td>MacNeil Vroomen et al. (2013) (Netherlands)</td>
<td>Berries</td>
<td>to review definitions of crises, and to create a standardised definition that can be utilised for research, policy and clinical practice</td>
<td>systematic review: qualitative synthesis</td>
<td>27 publications</td>
<td>Crisis in dementia is defined as a process where a stressor causes an imbalance requiring an immediate decision to be made which leads to a desired outcome and therefore a resolution of the crisis. If the crisis is not resolved, the cycle continues.</td>
<td>change, balancing, resources, health care system</td>
</tr>
<tr>
<td>Markiewicz et al. (1997) (Canada)</td>
<td>(Self-)caring actions</td>
<td>to examine the influence of IC’s attachment styles and personality traits on their experiences of caring for a PwD</td>
<td>quantitative: observational (longitudinal)</td>
<td>113 dyads (PwD living at home)</td>
<td>The attachment style can contribute to explain aspects of IC’s experiences. Those who chose to institutionalise the PwD were higher on the avoidance factor than those choosing to maintain the PwD in the community.</td>
<td>carer role, dyadic relationship</td>
</tr>
<tr>
<td>Mausbach et al. (2004) (USA)</td>
<td>Institutionalisation</td>
<td>to compare institutionalisation rates of PwDs cared for by Latina and Caucasian female ICs, and to explore IC and PwD related predictors</td>
<td>quantitative: observational (longitudinal)</td>
<td>264 female ICs (154 female Caucasians, 110 female Latinas) (PwD living at home at baseline)</td>
<td>Hispanic Americans delayed institutionalisation longer than their Caucasian counterparts. PwD characteristics or IC demographics did not explain these results, although Hispanic Americans who identified greater benefits or more-positive aspects in caring at baseline were less likely to institutionalise the PwD than those who reported fewer benefits.</td>
<td>carer role, society and culture</td>
</tr>
<tr>
<td>McLennon et al. (2010) (USA)</td>
<td>Institutionalisation</td>
<td>to identify common themes from the statements of ICs who decided to institutionalise their relative with AD or Parkinson’s disease</td>
<td>qualitative: unspecific design (longitudinal)</td>
<td>9 ICs of PwD, 2 ICs of persons with Parkinson’s disease (PwD living at home at baseline)</td>
<td>2 categories identified from the ICs stories were ‘anticipating the inevitable’ and ‘reaching the limit’ (stage-based model). 3 to 4 months before institutionalisation, ICs discussed knowing that they would not be able to continue caring at home. The most frequent reasons for institutionalisation were serious health events. There were more institutionalisations in the AD participant group than in the Parkinson’s disease group.</td>
<td>change, needs</td>
</tr>
<tr>
<td>Mitchell et al. (2004) (USA)</td>
<td>End-of-life-care</td>
<td>to examine and contrast the end-of-life experience of PwDs who died within one year of admission to either a NH or a community-based home care service</td>
<td>quantitative: observational (cross-sectional)</td>
<td>2730 PwDs institutionalised, 290 PwDs living at home</td>
<td>NH residents dying with advanced dementia were older, had greater functional impairment, and more behaviour problems compared to home care clients. Few subjects in the NH and home care cohorts were perceived to have less than 6 months to live. Only few individuals were referred to hospice. Hospitalizations were frequent. Pain and shortness of breath were common in both settings.</td>
<td>change</td>
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<td>Morycz (1985) (USA)</td>
<td>Institutionalisation</td>
<td>to explore whether the strain experienced by ICs of PwDs will be strongly related to the desire to institutionalise</td>
<td>quantitative: observational (longitudinal)</td>
<td>80 dyads (PwD living at home)</td>
<td>Results rely to the self-developed theoretical model: The desire to institutionalise a PwD was found to be greater when the IC’s experiences increased strain or burden, when PwD is widowed, when there is more physical labour involved in caring tasks, and when PwD lives alone. Intensity of family strain (or felt stress) can best be predicted by the availability of social support to the IC: Less support implies more strain. For male IC groups and for all black ICs, strain does not predict desire to institutionalise.</td>
<td>dyadic relationship, resources</td>
</tr>
<tr>
<td>Okura et al. (2011)</td>
<td>Institutionalisation</td>
<td>to examine the association between neuropsychiatric symptoms and risk of institutionalisation and death</td>
<td>quantitative: observational (longitudinal)</td>
<td>537 PwDs living at home at baseline, 109 PwDs institutionalised</td>
<td>The presence of one or more neuropsychiatric symptoms was not associated with higher risk of institutionalisation or death during the 5-year study period. Although when assessing each symptom individually, depression, delusions, and agitation were each associated with a higher risk of institutionalisation. IC distress mediated the association between delusions, agitation and institutionalisation. Depression and hallucinations were associated with significantly higher mortality.</td>
<td>change</td>
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<tr>
<td>Oliva y Hausmann et al. (2012) (Germany)</td>
<td>Institutionalisation</td>
<td>to explore the influence of IC needs, IC burden and role conflicts on the risk of institutionalisation of PwD</td>
<td>quantitative: observational (longitudinal)</td>
<td>373 ICs (PwD living at home at baseline)</td>
<td>A greater distance between households, increased need in housekeeping, higher income, increased IC burden and low self-evaluation of caring-quality are related to an increase in chance for institutionalisation. A constant need for supervision and lack of social appreciation are associated with a smaller chance.</td>
<td>change, dyadic relationship, health care system</td>
</tr>
<tr>
<td>Peacock et al. (2010)</td>
<td>Positive aspects</td>
<td>to better understand family care by applying a strength-based perspective</td>
<td>qualitative: unspecific design</td>
<td>39 ICs (PwD living at home alone, PwD living with IC, PwD institutionalised)</td>
<td>ICs can view their role as an opportunity to give back, to discover personal strengths, and to become closer to the PwD. The results suggest that identifying and mobilising the IC’s strengths can be an effective strategy for supporting ICs in their role.</td>
<td>carer role, resources</td>
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<tr>
<td>Pimouguet et al. (2016) (France)</td>
<td>Institutionalisation</td>
<td>to investigate the influence of early specialist referral for PwDs on the risk of institutionalisation and functional decline</td>
<td>quantitative: observational (longitudinal)</td>
<td>179 PwDs (for institutionalisation analysis), 243 PwDs (for ADL analysis), (PwD living at home at baseline)</td>
<td>One third of the incident PwDs had consulted a specialist for cognitive problems early in the disease course. PwDs who had consulted a specialist early presented a higher rate of being institutionalised than those who did not. Early specialist referral was not associated with functional decline.</td>
<td>resources</td>
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<td>Pinquart and Sörensen (2006) (Germany and USA)</td>
<td>Berries</td>
<td>to identify average effects of interventions found on IC burden, depression, indicators of positive subjective well-being, ability/knowledge of ICs, symptoms of ICs, and the risk of institutionalisation</td>
<td>systematic review: meta-analysis</td>
<td>127 publications</td>
<td>Interventions for ICs of PwDs had small but significant effects on burden, depression, ability/knowledge, subjective well-being and symptoms of PwDs. Only structured multicomponent interventions reduced the risk of institutionalisation.</td>
<td>resources</td>
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<td>Pinzon et al. (2013) (Germany)</td>
<td>End-of-life-care</td>
<td>to describe the preferred and actual places of death, symptom prevalence, and quality of care of PwDs</td>
<td>quantitative: observational (cross-sectional)</td>
<td>310 ICs of PwDs and 931 ICs of people without dementia, who died at home or in an institution</td>
<td>Almost half of the PwDs died at home. Compared to other countries this number is high. Most PwDs and their ICs preferred death at home to death anywhere else. PwDs living with at least one relative were more likely to die at home. The ICs were critical of the quality of care on standard hospital wards, citing the limited temporal availability of staff and limited emotional support.</td>
<td>dyadic relationship, society and culture, health care system</td>
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<tr>
<td>Porter et al. (2016) (USA)</td>
<td>Institutionalisation</td>
<td>to investigate the association between total NPI score, the NPI-4 (an agitation/aggression sub-scale) and individual domains of the NPI and NHP</td>
<td>quantitative: observational (longitudinal)</td>
<td>641 ICs: 352 cases (PwD institutionalised in the first 6 months of study), 289 controls (PwD living at home)</td>
<td>An increase in the NPI score implied an increase in odds of institutionalisation, also being married or being a male IC predicted institutionalisation. PwDs who entered a NH were significantly more likely to have behavioural and psychological symptoms of dementia related to agitation/aggression one month prior to institutionalisation than PwDs who stayed at home.</td>
<td>change</td>
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<tr>
<td>Quinn et al. (2015) (UK)</td>
<td>Needs &amp; Demands</td>
<td>to investigate how meaning, motivation and relationship dynamics influence the IC’s subjective experience of caring</td>
<td>qualitative: phenomenological approach</td>
<td>12 ICs (10 PwD living at home, 2 PwD institutionalised)</td>
<td>6 key themes emerged, which were encompassed under an overarching theme of ‘balancing needs’. This theme describes the IC’s struggle to balance their own needs against those of the PwD. Dilemmas emerged from the desire to try to preserve the ICs relationship with the PwD, while recognising that this relationship was changing. The IC’s relationship with the PwD influenced the IC’s motivations to provide care and gave caring meaning. Although ICs were motivated to provide care the battle to balance needs meant that ICs recognized there might come a time when they may need to cease caring.</td>
<td>balancing, needs, carer role, dyadic relationship, resources</td>
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<tr>
<td>Redfern et al. (2002) (UK)</td>
<td>(Self-)caring actions</td>
<td>to advance the understanding of care for a PwD at home with an emphasis on routines, control and care goals</td>
<td>qualitative: unspecific design</td>
<td>20 dyads (PwD living at home)</td>
<td>Accomplishing care is a major challenge for PwDs and their ICs. In most cases development of effective coping strategies, handling competing goals, controlling the situation and maintaining a preferred sense of self are challenges in dementia care that have to be mastered.</td>
<td>balancing, dyadic relationship</td>
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<tr>
<td>Romero-Moreno et al. (2014) (Spain)</td>
<td>Institutionalisation</td>
<td>to analyse the potential of both frequency of and satisfaction with leisure to explain IC’s distress, risk of institutionalisation, IC’s stressors and resources variables</td>
<td>quantitative: observational (cross-sectional)</td>
<td>275 ICs (PwD living at home)</td>
<td>ICs with a high satisfaction with and high frequency of leisure showed a more positive profile on resource variables, health outcomes and lower levels of risk of institutionalisation compared with the other groups. ICs with low satisfaction with and low frequency of leisure used significantly less adaptive emotional regulation strategies and showed worse consequences on health outcomes.</td>
<td>carer role</td>
</tr>
<tr>
<td>Rongve et al. (2014) (Norway)</td>
<td>Institutionalisation</td>
<td>to study time until and predictors for NHP for persons with mild dementia and to compare differences in time to NHP between PwDs with DLB and PwADs</td>
<td>quantitative: observational (longitudinal)</td>
<td>189 PwDs (PwD living at home at baseline)</td>
<td>Patients diagnosed with DLB had nearly 2 years shorter time to institutionalisation than those diagnosed with AD. Predictors of shorter time to institutionalisation were a DLB diagnosis, the use of antipsychotic medication, more advanced age, longer duration of dementia symptoms prior to diagnosis, living alone, higher reported carer distress, and more neuropsychiatric symptoms. The use of AChEIs was associated with halved risk of institutionalisation. In the DLB group, the use of antipsychotics was associated with increased risk of institutionalisation.</td>
<td>change, resources</td>
</tr>
<tr>
<td>Runte (2018) (Germany)</td>
<td>Institutionalisation</td>
<td>to analyse a cohort of PwDs in respect of institutionalisation over a follow-up period of 4.5 years and to investigate factors associated with the institutionalisation of PwDs, in general and stratified by sex</td>
<td>quantitative: observational (longitudinal)</td>
<td>652 PwDs (PwD living at home at baseline)</td>
<td>Risk of institutionalisation is higher in women than in men and when cared for by a care service in comparison to an IC. Inhibiting factors are care level and positive evaluation of caring by ICs. Stratified analysis by sex revealed that the risk of institutionalisation in men is influenced by their relationship to their IC, in women by duration of care at baseline.</td>
<td>change, dyadic relationship, resources, health care system</td>
</tr>
<tr>
<td>Saks et al. (2015) (International Consortium: Estonia, Germany, Spain, Finland, Sweden, France, UK)</td>
<td>(Self-)caring actions</td>
<td>to investigate the extent of variability in decisions of individuals and multidisciplinary groups from different European countries about the most appropriate setting for PwDs</td>
<td>quantitative: observational (cross-sectional)</td>
<td>161 experts in dementia care</td>
<td>There is considerable variation in individual’s recommendations for the most appropriate setting for PwDs. Group-based decisions were more consistent, but country-specific patterns remained.</td>
<td>dyadic relationship, resources, society and culture, health care system</td>
</tr>
<tr>
<td>Sansoni et al. (2013) (Italy)</td>
<td>Institutionalisation</td>
<td>to review literature about factors that influence the institutionalisation of PwDs</td>
<td>unsystematic review</td>
<td>17 publications</td>
<td>Factors like IC burden, hours involved in care, stress, social isolation, gender, relationship to the PwD, availability of support resources, and IC characteristics influence the decision for institutionalisation.</td>
<td>change, society and culture, health care system</td>
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<tr>
<td>Author (Year) (Country)</td>
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<td>Study Design</td>
<td>Sample/Setting</td>
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<tr>
<td>Schacke et al. (1998) (Germany)</td>
<td>Stability</td>
<td>to analyse the meaning of various dimensions of IC burden for the IC's well-being and the stability of the care situation</td>
<td>quantitative: observational (cross-sectional)</td>
<td>78 ICs (PwD living at home)</td>
<td>Stability of home care is associated with perceived inadequacy of IC resources. Lower resources determine a higher probability of institutionalisation within the next 12 months. Other dimensions of IC stress/burden were not significant. An explicit distinction is made between IC burden which directly refers to the situation and the more general concept of well-being which is potentially, but not necessarily, affected by the caring situation. IC's life satisfaction, self-esteem, and their subjective perception of the stability of the care situation were regarded as indicators for potential outcomes of caring.</td>
<td>resources</td>
</tr>
<tr>
<td>Schäufele et al. (2005) (Germany)</td>
<td>Stability</td>
<td>to identify care gaps and possible undesirable development and to determine factors that stabilise or endanger the care situation</td>
<td>quantitative: observational (cross-sectional)</td>
<td>151 PwDs, 155 persons without dementia and their IC</td>
<td>Most care arrangements are judged as stable by the IC. Subjective burden, inadequate support and male gender of IC are associated with instable care arrangements.</td>
<td>change, resources</td>
</tr>
<tr>
<td>Schirra-Weirich and Wiegelmann (2017) (Germany)</td>
<td>Stability</td>
<td>to develop a typology of PwD and IC dyads to better address their specific needs in a case management approach</td>
<td>quantitative: observational (longitudinal)</td>
<td>40 dyads (PwD living at home)</td>
<td>Dyads are divided into three types based on generational distinctions of care concepts. Case management is a mental support for intra-generational care settings and a functional support for inter-generational care settings. Case management based on a typology seems to be an appropriate support for ICs and PwDs.</td>
<td>balancing, career role</td>
</tr>
<tr>
<td>Schoenmakers et al. (2009) (Belgium)</td>
<td>Institutionalisation</td>
<td>to investigate the differences between characteristics of community dwelling PwDs and non-demented elderly and their ICs at the moment of definitive institutionalisation</td>
<td>quantitative: observational (cross-sectional)</td>
<td>143 GPs (reporting data of 300 PwDs and 507 non demented controls, already institutionalised)</td>
<td>PwDs with concomitant diseases were mainly placed because of unmanageable behavioural disturbances. Strikingly, these PwDs were more often confronted with a delay in definitive institutionalisation due to their high care dependence. Although IC burden was an important motivation, it did not seem to be a motive in the final institutionalisation. Behavioural disturbances are independently of IC burden, FC support, or a spousal relationship the main direct reason for institutionalisation.</td>
<td>change, resources</td>
</tr>
<tr>
<td>Schölzel-Dorenbos et al. (2010) (Netherlands)</td>
<td>Needs &amp; Demands</td>
<td>to make an inventory of needs assessment instruments, to explore the interaction between unmet needs and HRQoL and to relate these to the conceptual model of Maslow’s hierarchy of needs in order to design systematic review: qualitative synthesis</td>
<td>2 needs assessment instruments: CANE and CNA-D</td>
<td>PwDs identified significantly fewer needs than ICs. The most important needs, that also determine a large part of HRQoL, are: need for information, support with regard to symptoms of dementia, social contact and company, and health monitoring and safety. Focusing unmet needs can improve HRQoL.</td>
<td>needs</td>
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<td>Shega et al. (2008) (USA)</td>
<td>End-of-life-care</td>
<td>to evaluate the impact that hospice enrolment has on the terminal care of PwDs, and to describe the symptom burden these patients face at the end of life</td>
<td>quantitative: observational (cross-sectional)</td>
<td>135 ICs (PwD living at home or in a hospice)</td>
<td>The results suggest that dying PwDs who are enrolled in hospice programs are more likely to die in their locations of choice and their ICs are more likely to perceive that care is excellent or good than those who do not obtain hospice services. No effects on pain or other symptoms could be demonstrated in this analysis.</td>
<td>resources</td>
</tr>
<tr>
<td>Shim (2011) (USA)</td>
<td>Positive aspects</td>
<td>to explore how ICs of PwDs are able to have positive care experiences, how the relationship between the ICs and the PwADs changes over time, and how this relationship differs between dementia ICs who report positive care experiences and those who do not</td>
<td>mixed methods</td>
<td>PwD and people with Parkinson’s disease (living at home). Study 1: n= 91 IC/PwAD dyads, Study 2: n= 21 spousal ICs of PwAD, Study 3: n= 12 ICs of PwAD</td>
<td>The relational dynamics between ICs and PwDs are important and can be the source of a positive care experience. For the ICs their current attitude toward the relationship dynamic with the PwD is what seemed to bring the most meaning. By finding caregiving meaningful, ICs can obtain peace and pleasure in the currently lived experience. IC attitudes toward the relationship with PwDs and the PwD’s degenerative decline influence the quality of the care experience.</td>
<td>change, balancing, dyadic relationship</td>
</tr>
<tr>
<td>Spitznagel et al. (2006) (USA)</td>
<td>Berries</td>
<td>to understand predictors of desire to institutionalise in ICs of PwDs</td>
<td>quantitative: observational (cross-sectional)</td>
<td>72 ICs (PwD living at home)</td>
<td>ICs with a desire to institutionalise had significantly higher burden, greater dementia knowledge, more family dysfunction, and decreased social support compared with no-desire ICs. Findings emphasise the importance of IC and family relationship variables in desire to institutionalise, suggesting potentially modifiable targets for IC interventions. Dementia knowledge was associated with higher desire to institutionalise, suggesting that educational programs alone may not be helpful to delay institutionalisation.</td>
<td>change, dyadic relationship, resources</td>
</tr>
<tr>
<td>Stephan et al. (2015) (International Consortium: Estonia, Germany, Finland, France, Netherlands, UK, Spain, Sweden)</td>
<td>Institutionalisation</td>
<td>to explore the perceptions of ICs and FCs regarding reasons for the institutionalisation of PwDs in eight European countries</td>
<td>mixed methods</td>
<td>1160 ICs (PwD living at home) and their health care practitioners (number not reported)</td>
<td>Agreement about reasons for institutionalisation between ICs and FCs was low-to-moderate. Discrepancy was high for reasons related to IC burden. Furthermore differences between countries emerged.</td>
<td>change, resources, society and culture, health care system</td>
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<tr>
<td>Stiens et al. (2006) (Germany)</td>
<td>Institutionalisation</td>
<td>to investigate associations of filial maturity with PwD institutionalisation and IC burden by using the Louvain Filial Maturity Scale</td>
<td>quantitative: observational (longitudinal)</td>
<td>61 filial ICs (PwD living at home at baseline)</td>
<td>Filial maturity had no influence on the IC’s feelings of burden, but higher ‘parental consideration’ resulted in lower IC burden. Adult children with more ‘filial obligation’ continued to care for their parents more often, even when experiencing great burden and stress. Institutionalisation was caused mainly by PwDs growing needs and increasing behavioural problems.</td>
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<tr>
<td>Sun et al. (2013) (USA)</td>
<td>Institutionalisation</td>
<td>to understand the role of quality of care in mediating between caring stressors and IC desire to institutionalise a PwD</td>
<td>quantitative: observational (cross-sectional)</td>
<td>612 racially diverse ICs (PwD living at home)</td>
<td>Potentially harmful behaviours of the ICs were positively related to IC’s desire to institutionalise their care recipients. The mediation effect of exemplary care was significant in the African American group only.</td>
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<tr>
<td>Toot et al. (2013) (UK)</td>
<td>Berries</td>
<td>to identify which factors may lead to crisis for PwDs and ICs and identify interventions these individuals believe could help in crisis</td>
<td>qualitative: unspecific design</td>
<td>18 PwDs (living at home), 15 ICs, 19 FCs</td>
<td>Causes of crises for PwDs were risks and hazards in their home, whereas for ICs carer stress and their own mental health problems were predominant. FCs focused on problems with service organisation and coordination leading to crisis. PwDs valued support from family and friends, access to mobile phones and home adaptations to reduce risks. ICs valued assistive technology and home adaptation. Both ICs and FCs valued carer training and education, care plans and well-coordinated care.</td>
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<tr>
<td>Toot et al. (2017) (UK)</td>
<td>Institutionalisation</td>
<td>to understand the factors that influence the decision for PwDs to move into NHs, with particular emphasis on factors that might be responsive to intervention</td>
<td>systematic review: narrative review and meta-analysis</td>
<td>26 publications, meta-analysis: 15 publications</td>
<td>Poorer cognition and behavioural and psychological symptoms were associated with an increased risk of institutionalisation. Impairments in ADLs were identified as a significant risk. There was an association between IC burden and risk of institutionalisation, but findings with regard to IC depression varied, as did physical health associations, with some studies showing an increased risk of institutionalisation following hip fracture, reduced mobility, and multiple comorbidities.</td>
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<tr>
<td>Treloar et al. (2009) (UK)</td>
<td>End-of-life-care</td>
<td>to undertake detailed interviews of ICs who had supported PwDs at home, with the aim of identifying the major factors which make such care feasible</td>
<td>mixed methods</td>
<td>14 bereaved ICs (PwD living at home or institutionalised)</td>
<td>The study revealed blockages to accessing support as a result of poor understanding of the needs of advanced dementia care as well as organisational prejudice. Home based palliative care of PwDs can be achieved with very positive outcomes.</td>
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**Contribution to Concepts**

- carer role
- needs, society and culture
- change, needs, carer role, resources, health care system
- change, needs, carer role, resources
- change, needs, resources
- change, needs, resources
<table>
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<tr>
<th>Author (Year) (Country)</th>
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<tbody>
<tr>
<td>Tucker et al. (2016) (International Consortium: UK, Estonia, Netherlands, Spain, Sweden, Finland, Germany, France)</td>
<td>Institutionalisation</td>
<td>to explore the extent to which client characteristics, decision-maker attributes, and country influence judgments of institutional long-term care appropriateness for PwDs</td>
<td>quantitative: observational (cross-sectional)</td>
<td>161 experts in dementia care</td>
<td>Client characteristics, decision-maker attributes, and country seemed to play a role in influencing FC’s perceptions of the appropriateness of NHs for PwDs. Decision-makers were more likely to recommend NHs for PwDs who required help with mobility or had multiple care needs, and appeared to give more weight to IC’s than PwD’s wishes. Community-based social workers were less likely than other FC groups to favour NHP. Experts in Finland, Germany, and the UK were less likely to recommend NHP than experts in France, the Netherlands, and Estonia. Experts in Sweden and Spain took an intermediate position.</td>
<td>change, resources, society and culture</td>
</tr>
<tr>
<td>Verbeek et al. (2015) (International Consortium: Estonia, Germany, Finland, France, Netherlands, UK, Spain, Sweden)</td>
<td>Institutionalisation</td>
<td>to explore inter-country variations associated with institutionalisation of PwDs</td>
<td>quantitative: observational (longitudinal)</td>
<td>2014 dyads (PwD living at home or recently institutionalised)</td>
<td>Inter-country differences were found in recently institutionalised PwD’s characteristics. Most consistent factors associated with institutionalisation were ADL and IC burden, although country differences may be more prominent for this factor.</td>
<td>change, dyadic relationship, society and culture</td>
</tr>
<tr>
<td>Volicer et al. (2003) (USA)</td>
<td>End-of-life-care</td>
<td>to evaluate the quality of care in home settings and compare it with care provided in institutions</td>
<td>quantitative: observational (cross-sectional)</td>
<td>154 ICs (PwD living at home or institutionalised (various institutions))</td>
<td>Psychiatric symptoms and increased IC burden were the most common reason for institutionalisation, and psychiatric care was associated with longer stay at home. Presence of advance directives decreased hospital stay and increased the likelihood of dying in a NH. Care recipients dying at home had fewer symptoms and less discomfort than care recipients dying in other settings. These results indicate that quality end-of-life care can be provided at home and is facilitated by hospice programs, effective pain control, and psychiatric care.</td>
<td>change, resources</td>
</tr>
<tr>
<td>von Kutzleben et al. (2015) (Germany)</td>
<td>Berries</td>
<td>to assess home-based care arrangements, to illustrate utilisation of formal services, and to identify factors associated with perceived stability of the care situation from the IC’s perspective</td>
<td>quantitative: observational (cross-sectional)</td>
<td>84 dyads (PwD living at home)</td>
<td>Most frequently used formal services were home care nursing services, day care, and respite care. Companion home visit, home care nursing services and day care were used over the longest periods of time. IC’s perceived stability of care situation was high, and was associated with the country of origin and sex of PwD.</td>
<td>dyadic relationship, society and culture, health care system</td>
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<td>Wackerbarth (1999a) (USA)</td>
<td>Berries</td>
<td>to understand the decision processes of ICs and to develop a model of their decision processes</td>
<td>qualitative: unspecific design</td>
<td>28 ICs (PwD living at home or institutionalised)</td>
<td>The proposed model charts the caring experience over time and documents decisions made to maintain a stable care situation. It has three main components: tolerance line, status points, and decision events. ICs are divided into 5 types: ICs who plan ahead, take it one day at a time, have difficulty implementing decision, receive decision support from FCs and ICs who feel that decisions are made by others or by the disease itself.</td>
<td>change, balancing, needs</td>
</tr>
<tr>
<td>Wackerbarth (1999b) (USA)</td>
<td>(Self-)caring actions</td>
<td>to describe the complexities of the overall decision making situation, to characterise a specific care decision, and to identify decision support needs</td>
<td>mixed methods</td>
<td>80 ICs (PwD living at home or institutionalised)</td>
<td>Most prevalent decisions involved institutionalisation, communicating the diagnosis and the driving privilege. Most challenging decisions involved providing care and relocation. The most prevalent question involved the timing of actions.</td>
<td>balancing</td>
</tr>
<tr>
<td>Wattmo et al. (2011) (Sweden)</td>
<td>Institutionalisation</td>
<td>to identify risk factors for early institutionalisation in AD, focusing on the impact of longitudinal change in cognition, ADL, service utilisation, and AChEIs</td>
<td>quantitative: observational (longitudinal)</td>
<td>880 PwDs (PwD living at home at baseline)</td>
<td>Factors that precipitated institutionalisation were: lower cognitive and functional abilities at baseline, faster rate of decline in Instrumental Activities of Daily Living Scale, female gender, living alone, substantial increase in adult day care, and a lower mean dose of AChEI. The rate of functional but not cognitive decline was a strong risk factor for institutionalisation.</td>
<td>change</td>
</tr>
<tr>
<td>Willoughby et al. (1991) (Canada)</td>
<td>(Self-)caring actions</td>
<td>to understand the process of caring from the perspective of ICs of PwDs</td>
<td>qualitative: Grounded Theory</td>
<td>10 ICs (PwD institutionalised)</td>
<td>The finding yield a five-stage model of gaining and relinquishing control of caring (emerging recognition, taking control, losing control, adjusting to the psychiatric institution, moving on). Shifts in the understanding of the process delineated each stage.</td>
<td>change, balancing, carer role, dyadic relationship</td>
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<tr>
<td>Winslow et al. (1999) (USA)</td>
<td>Burden</td>
<td>to describe the longitudinal pattern of IC burden and to compare IC burden of wives who continue in home-based care and those who institutionalised the PwD</td>
<td>quantitative: observational (longitudinal)</td>
<td>376 ICs (PwD living at home) and 523 ICs (PwD institutionalised)</td>
<td>Spouse ICs who provided home-based care to their husbands experienced an increase in burden over time. There appeared to be a threshold of burden when institutionalisation occurred. Immediately after placement the IC appeared to experience a reduction in burden.</td>
<td>change, needs</td>
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<tr>
<td>Wuest et al. (1994) (Canada)</td>
<td>(Self-)caring actions</td>
<td>to explore the reciprocal process of becoming strangers in Canadian ICs and PwDs and their interaction on a continuum from intimacy to alienation</td>
<td>qualitative: Grounded Theory</td>
<td>15 dyads (PwD living at home)</td>
<td>The continuum from intimacy to alienation comprises three dimensions: dawning, holding on and letting go. ICs struggle on a daily basis with their commitment to the person they knew and their increasing detachment from the stranger the PwD has become.</td>
<td>change, balancing, carer role, dyadic relationship</td>
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<td>Zarit et al. (1986) (USA)</td>
<td>Burden</td>
<td>to investigate changes over time for ICs of PwDs, with a focus on factors associated with institutionalisation, to determine if differences between husbands and wives found at the initial testing remained stable over time</td>
<td>quantitative: observational (longitudinal)</td>
<td>64 ICs (PwD living at home)</td>
<td>Institutionalisation was more strongly associated with subjective factors, particularly IC burden, than with objective indicators of the severity of dementia. Although wives initially reported more burden than husbands, no differences were found at the follow-up.</td>
<td>change</td>
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</table>
Appendix E: Bibliography of the meta-study sample

All 99 publications included in our meta-study (including those, that are not cited in the publication itself) are listed below.


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