# Abstract

Gender is a neglected dimension in public discourse related to people with dementia. Those living with this condition are typically portrayed in policies and strategies in gender neutral terms as ‘people with dementia’ and ‘family carers’ as if gender does not matter, when clearly it does. The purpose of this scoping review was to take stock of knowledge about gender differences in relation to dementia care to inform policy and future research. The work is grounded in a feminist perspective to citizenship, as this provide a lens with which to expose and examine gendered assumptions within dementia studies. A search of four databases, including CINAHL, Web of Science, Medline, and Cochrane was conducted using systematic techniques between May and July 2014. A repeat search was conducted in February 2015. We found a significant amount of valuable research concerned with gender differences in relation to dementia care published from 1990 to 2014; the majority of which, lacks a feminist citizenship perspective. Moreover, a disproportionate number of studies focused solely on caregivers rather than citizens with dementia. As such, questions about gender equality are not being raised and the voices of men and women with dementia are silent. Thus we argue for increased gender-sensitivity in policy making and recommend that social scientists inject a feminist citizenship perspective into their work.

*Key words*: citizenship; feminist theory; gender, dementia care; health policy

**What is known about this topic**

* Gender is a neglected dimension in public discourse about dementia
* Gender is a key factor structuring the lives of those living with dementia
* There are gender differences in the caregiving experience

**What this paper adds**

* A critical review of the research literature on gender differences in dementia care from a feminist citizenship perspective
* Insights into the gaps in the research literature
* Recommendations as to how policy makers can incorporate gender into national plans

# Background

There is an urgent need to improve the delivery of health and social in the community for people living with dementia. The number of people diagnosed with dementia in 2015 worldwide is estimated at 46.8 million, reaching over 74 million by 2030 and 131 million in 2050 (Alzheimer’s Disease International, 2015a). The establishment of a World Dementia Council, appointment of a World Dementia Envoy following the G8 dementia summit in London in December 2013, and fact that the WHO hosted the first ever Ministerial Conference on dementia in March 2015, demonstrates the growing importance worldwide of improving care services for the millions of people living with dementia.

Generally speaking, the majority of people affected by dementia are women. It is estimated that 61% of people with dementia are women and 39% are men. This is likely to be because women live longer than men, as age is the biggest known risk factor for the condition (Alzheimer’s Research UK, 2015). However, different dementias sub-types may have different prevalent rates and so life expectancy is not the only factor to consider. Furthermore, as with other major health conditions, it is predominantly women who take on the role of caring for someone with dementia; studies suggest that in most countries up to three quarters of family caregivers are women (Bamford & Watson, 2012). For these reasons, dementia has recently come to the attention of feminists (Wilkinson, 2015); in that, it represents a struggle for resources and an unequal division of labour (Freedman, 2001).

Because age is the single biggest risk factor, the majority of people living with dementia are also older, aged sixty-five years and above. Thus, it is important to consider gender in the context of aging when discussing this topic. Since the late 1990s, scholars have been arguing for more critical feminism in gerontology - that is, focusing on how patriarchal practices and assumptions affect people in later life (see, for example, Ray & Fine, 1999). This work has paved the way for increased critical attention on older woman. For example, there has been research into older women’s contribution to civic society (McHugh, 2012), and work on the role of older women as ‘transmitters of values’ (Queniart & Charpentier, 2013). However, little attention has been paid to older women with dementia within this area of feminist inquiry, as a dementia diagnosis tends to exclude people from mainstream ageing studies. This is a significant oversight, as we need a critical feminism for people with dementia too.

Gender clearly matters in the context of illness, impairment, and caregiving (Annandale and Hunt, 2000). It is especially important in respect of people with a disability like dementia where ‘forms and impacts of disablism are always refracted in some way through the prism of gendered locations and gender relations’ (Thomas, 1999: 28). For example, the loss of a handbag is thought to symbolise the loss of independence for a woman with dementia (Buse & Twigg, 2014); and a man with dementia in a care home may feel socially excluded because there are so few other men to relate to (Bartlett, 2007). Thus, gender, as well as age, will inevitably be a factor structuring the lives of those living with dementia.

Researchers have begun to focus on gender differences in relation to the lived experience of dementia and significant findings are emerging. For instance, one qualitative study has found that male carers are less likely to facilitate the autonomy of their partners than their female counterparts (Boyle, 2013: 240). While it is unclear whether this situation is due to dementia, or more of a reflection of the spousal relationship through the life course, such results are significant in terms of understanding the restrictions placed on a woman’s agency. Other research shows that husbands and wives respond differently to signs of dementia in their spouse; with men being slower to recognise the symptoms than women, and women quicker to recognise subtle changes but failing to take action quickly (Hayes, Zimmerman, and Boylstein, 2010). Such work underscores the importance of incorporating gender into the design, delivery and evaluation of dementia care policies (International Longevity Centre, 2011). Moreover, it shows that research on gender in the context of dementia care is growing and producing results that warrant the attention of policy makers.

Given what we know about gender, it is curious why the topic is not more prominent in dementia care research and policy. Influential texts use gender neutral terms such as ‘people with dementia’ and ‘family carers’ rather than distinguishing between men and women (e.g. Kitwood, 1997; Sabat, 2005). Even literature reviews on self-identity (Caddell & Clare, 2011) and relationships (Ablitt, Jones, & Muers, 2009) - topics deeply connected with gender – there is no reference to gender, either in relation to the studies reviewed or the review itself. It would seem that the dementia care literature is gender blind.

Gender is neglected in national dementia strategies too. In 2015, seventeen countries, namely, Australia, Denmark, England, Finland, France, Israel, Republic of Korea, Luxembourg, Malta, Netherlands, Norway, Northern Ireland, Switzerland, Taiwan, Scotland, USA and Wales had developed a national dementia strategy (http://www.alz.co.uk/alzheimer-plans). Other than referring to female sex as a risk factor for dementia, no distinction is made in these national plans to men and women. The incidence and prevalence of dementia is not even considered from a gendered perspective, as numbers of *people* with dementia are given, rather than numbers of men and women. Given the strategic importance of these plans, which provide the language and set the agenda for the development of healthcare services, it is both curious and of concern that national dementia strategies throughout the world should neglect gender.

There are signs of gender moving into the dementia care landscape, with the publication of new reports about women and dementia. For example, Alzheimer’s disease International recently produced a report on the extent to which dementia affects women across the globe (e.g. ADI, 2015b). Also, in the UK the Joseph Rowntree Foundation has published a piece called Dementia Through Women’s Eyes, as part of its Dementia Without Walls programme (Savitch, Abbott & Parker, 2015). Although not overtly political or feminist in tone, such work does show a growing awareness of gender. However, as feminist gerontologists have pointed out, the risk with focusing on women is that men become invisible and the relational aspects of gender, privilege, and oppression are overlooked (Calasanti, 2004). It is important, then, to consider gender as an influential force in everybody’s lives.

In this review we therefore take stock of existing research on gender differences in the context of dementia care to inform policy and reflect on future research directions. The review is grounded in a feminist perspective to citizenship, as this provide a lens with which to expose and examine gendered assumptions implicit within dementia care studies. Such a lens also encourages us to think beyond care practices to broader issues of knowledge production and power relations. All of which are important if we want to promote the citizenship of people living with dementia, as it draws critical attention to the problem of binary thinking – e.g. men and women are different (Lister, 2003) and how the processes of research affects the results of that research (Letherby, 2003). We recognise that there is no single or unified approach to feminism, and so for us the project is about analysing how gender differences are investigated and handled by researchers to determine the extent to which the field of dementia studies is working to achieve a social (gender) justice - gender differences being cultural, as opposed to sex differences, which are biological. Essentially, this article seeks to examine the relationships between feminist theory, citizenship, and dementia care.

The scoping review includes the vast number of studies on gender differences among male and female caregivers and the small but growing area of research related to the gendered nature of the lived experience of dementia. We highlight the lack of a feminist citizenship perspective in dementia studies and argue for greater gender sensitivity in policy and research arenas related to dementia care.

# Method

A scoping review of published studies on gender differences in the context of dementia care was conducted between May 2014 and February 2015. Scoping reviews aim to provide a comprehensive coverage of the literature on a given topic area and are primarily concerned with the breadth and depth of existing knowledge rather than its quality (Arksey & O’Malley, 2005). The threefold aim of this scoping review was to (1) summarise and discuss research findings on gender differences in the context of dementia care and through a feminist citizenship lens, (2) identify research gaps in the existing literature, to (3) inform policy and future research. Criteria for inclusion in the scoping review were that the study was: published in a peer review journal; addressed gender differences in the context of dementia care (including Alzheimer’s disease; Lewy body dementia, vascular dementia, and ‘senile dementia’); and written in English. The criteria for exclusion in the review were: non-research articles (including thought pieces, meeting abstracts, and letters to the editor), reporting on gender as a risk factor for dementia and biological studies on sex differences. No date restrictions were imposed. We wanted to locate and review all relevant studies to identify any trends in research and to avoid making recommendations for future research if similar work had been carried out previously.

We searched for primary studies on gender differences in the context of dementia care. This included studies that reported exclusively on gender differences in relation to the caregiving role, and studies that differentiated between men and women’s experience of living with dementia. We took the decision to include a wide range of studies because we felt it was important to gain a comprehensive picture of the research arena in order to summarise what is currently known about gender differences in the context of dementia care. Initially, we searched the Cochrane Library using the terms ‘dementia’ and ‘gender’ and found no relevant reviews. A controlled vocabulary and key word search was then conducted using the following databases: CINAHL, Medline, The Allied and Complimentary Medicine Databases, Web of Science, and International Bibliography of the Social Sciences. The controlled search terms were ‘dementia, including Alzheimer’s disease; Lewy body dementia, vascular dementia’, ‘pre-senile’ and ‘senile dementia’; gender, gender roles, gender differences, caregivers. The key word search terms were ‘gender’, ‘men’, ‘women’ and ‘dementia’. Relevant additional studies identified from lateral searching of references were also reviewed. The literature search was carried out in May 2014. A total of 23 publications satisfied the inclusion criteria and were selected for a full-text review. Reference lists of the selected articles were visually scanned to ascertain whether any key studies had been missed. Ten additional publications were found in this way. The search was repeated in February 2015 (with the date limit 2014-2015) and three further studies were found, bringing the total amount of papers to review to 35. Retrieved articles were stored and shared between authors using the digital reference manager Mendeley. A template for extracting data and reviewing studies was developed and agreed by the authors. It contained the following categories (a) full details of citation (b) knowledge established (c) theoretical perspective (d) study design and methods (e) sample size and profile (f) methodological strengths and limitations (g) key findings related to citizenship and gender (h) recommendations for research.

# Findings

Publication dates were from 1990 to 2015. Studies included in this review were conducted in the United States, Canada, the United Kingdom, Finland, Sweden, Norway, Spain, Germany, Cyprus, and Japan. An early finding was that researchers have been sensitive to gender differences in relation to dementia care since the early 1990s. Several studies published at this time discuss gender differences in the context of dementia care. The studies reviewed used either qualitative methods, predominantly interviews (n=17), quantitative methods, predominantly large-scale surveys (n=15), or mixed methods (n=2). Four studies were literature reviews. The majority of the studies focused solely on gender differences in relation to the caregiving role, primarily husbands. Eight studies focused on gender differences in relation to the experience of living dementia. We identified two main themes in the studies reviewed by taking into account both the sample and whose perspective researchers had taken; these were (1) gender differences among caregivers and (2) the gendered nature of the lived experience (Table 1).

The 35 papers are presented in Table 1.

**Table 1**

Studies on gender difference in context of dementia care

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| Study  | Sample | Study design  | Key findings | Comments  |
| *male caregivers*  |  |  |  |
| Baker, Robertson, & Connelly (2010)  | Husbands (n=70) currently caring for their wives with dementia | Survey methodology; explores gender identity using questionnaires  | Male caregivers with traditional beliefs about masculinity are more likely to say that (a) they are not feeling burdened, (b) they feel uncertain about caring and (c) they are more likely to articulate positive aspects to being a spousal carer than men with less traditional beliefs about masculinity (p.326). | Male perspective is privileged; women’s perspective is absent. |
| Kramer (1997) | Husbands (n=74) caring for their wives with dementia. | Cross-sectional analysis using structured interviews. | Social resources and good health help men feel better about caregiving; highly educated men were least likely to feel good about caregiving.  | Some discussion of structural inequalities in the workplace; women’s voices are absent  |
| Kramer (2000) | Husband caregivers (n=74) (same sample as study above). 60% participated in this follow-up study  | Longitudinal study using interview method; scales were used to measures stressors and well-being, including depression | Men adapt to caregiving role over time.  | Women’s voices are absent |
| Kirsi & Jylha (2000) | Sample of 159 caregivers: 58 wives, 47 daughters, 36 other female carers (sister, friend, daughter in law), 3 sons and 15 husbands.  | A discourse analysis of 15 stories written by husbands about their wives with dementia | There is a tension between being a caregiver and a man | No discussion of how this is produced or what might be done to change it.  |
| Kirsi, Hervonen, & Jylhä (2004) | Written accounts (N=44): 11 husbands/33 wives and other female carers.Interviews (N=36): 13 husbands/23 wives and other female carers. | Qualitative study conducted in Finland ; 13 focused interviews with husbands who had given care to their wives with dementia | Men’s experiences of caregiving are diverse | Questions raised about how gender is reproduced; voices of women are absent, but the authors recognize this is problematic.  |
| Knutsen & Råholm, (2009) | Nine men caring for their wives with dementia took part in interviews | A phenomenological study conducted in Norway | Caregiving is a daily struggle; men do not like not being able to fulfil themselves and loss of control.  | Taking on household responsibilities and caring role is assumed to be a burden for men; Voices of women are absent.  |
| Neufeld (1998) | Male caregivers (N=22) (husbands, sons, brothers) were interviewed four times. One focus group with 7 interviewees | Qualitative component of a larger study conducted in Canada which involved female caregivers  | Most men gave care on the basis of obligation, which was accompanied by negative or mixed feelings.  | Provides an example of men’s citizenship within the private sphere  |
| Russell (2001) | Men (n=14) currently caring for their wives with dementia | Qualitative study using interviews conducted in USA near New York.  | Men are capable, nurturing and innovative carers  | The need for gender-sensitive services is highlighted and female perspective is considered  |
| Brown, Chen, Mitchell, & Province (2007) | Men (n=9) caring for their wives with dementia at home for an average of 6 years  | Qualitative grounded theory study  | Help-seeking by older husband caregivers is complex and gender-specific (p.352) | Voices of women are absent  |
| Baker & Robertson (2008) | Reviewed 93 articles  | Systematic review of literature to assess the status of knowledge about men’s coping with the role of dementia caregiver | Male caregivers are still poorly understood compared to female caregivers | The emphasis on coping is individualistic; hence, wider socio-political relations are overlooked.  |
| Robinson, Bottorff, Pesut, Oliffe, & Tomlinson (2014) | Reviewed 30 articles  | Scoping review of articles published between 2007 and 2012 | None of the studies use a gender framework to inform study design or analysis.  | A gender lens is needed to critique society and develop an understanding of what needs to be done and changed  |
| *female caregivers* |  |  |  |
| Eriksson, Sandberg, & Hellström (2013) | Women (n=12) caring for their husbands with dementia  | Qualitative study; used interview method to explore meanings of home, their partner’s condition and everyday life | The commitment and responsibilities that women experience in their day-to-day caring must be acknow-ledged (p.159).  | Goal here is to raise gender awareness rather than achieve gender justice.  |
| Perry (2002). | Wives (n=20) all but one of whom lived at home with their husband with dementia  | Qualitative study conducted in Canada  | Wives took over their husbands’ roles and responsibilities  | Provides an example of women’s citizenship within the private sphere |
| Strang (2001) | Women caregivers (n=8) were interviewed twice  | Qualitative study conducted in Western Canada using repeat interviews  | The women were committed to the welfare of their family members  | Draws explicitly on a feminist perspective but no discussion of women’s citizenship or agency.  |
| *Caregivers per se* |  |  |  |
| Winter, Gitlin & Dennis (2011) | Male (35) and female (202) caregivers (n=237) of community-dwelling people with dementia.  | Used baseline data from participants enrolled in a randomised study of non-pharmacological intervention to manage behavioural symptoms  | Stronger desire to institutionalise was associated with lower quality of relationship for male but not female caregivers  | Needs of male caregivers are privileged; the citizenship of women, including those who are institutionalised, is not discussed.  |
| Romero-Moreno et al. (2013) | Caregivers (n=351) (58.97% daughters, 10.54% sons, 19.66% wives and 10.38% husbands).  | Questionnaire based study conducted in Spain  | Daughters with higher levels of guilt who do not engage in leisure activities may be especially vulnerable to psychological distress  | No discussion of why daughters should feel guilty; or how feelings of guilt might play a role in women’s oppression.  |
| Chappell, Dujela & Smith (2014) | Caregivers (n=873) as part of a larger research program on drug therapy for people with AD | Quantitative study using structured interviews conducted in British Columbia, Canada  | Gender as social positions/categories can work together and in opposition to shape experiences of caregivers´ burden | Draws explicitly on an inter-sectionality framework but only in terms of gender and relationships.  |
| Gibbons et al., (2014) | Spouses/carers (n=65) of people with dementia - 45 (69.2 female) and 20 (30.8) male | Used survey methodology and measures to clarify differences in the experiences of wives and husbands in respect of burden and health. Conducted in Canada  | Female gender is a marker that places caregivers at increased risk of high burden and less support  | Hints at social injustices but not explicit about it.  |
| Hooker, Manoogian-O’Dell, Monahan, Frazier, & Shifren (2000) | Spouse caregivers (n=175) of people with AD (88 – 59% women) and Parkinson’s Disease (87 - 64% women)  | Quantitative study using structured interviews/questionnaires conducted in USA | AD caregiving wives tended to report greater depression and were significantly more stressed and anxious than were AD caregiving husbands (p571).  | Gender lens is not apparent but calls for future research to investigate how gender differences are (re) produced  |
| Schiffczyk, Jonas, Lahmeyer, Müller, & Riepe (2011) | People with dementia (n=212) and their proxies – primary proxy was the spouse in 180 of 194 dyads (93%) | Quantitative study using questionnaires conducted in Germany | Found a higher score of depression in female proxies than men..  | A gender-specific analysis is taken but it might have skewed results, as you will not see the effects of other variables (e.g. socio-economic status) if the sample is split according to gender. |
| Hayes, J. Zimmerman, M and Boylstein (2010) | Men (n=13) and women (n=15), all of whom cared for their spouses with dementia, took part in intensive qualitative interviews | An interview study conducted in the US | Husbands/men were slower than the wives/women to recognize the symptoms of Alzheimer’s disease and related disorders (ADRDs) within their spouse. | The gender relations within the context of marriage are presented as quite stereotype. The authors do not problematize this.  |
| Miller & Kaufman, (1996) | Spouse caregivers (n=215). Male and female, African-American and white | Qualitative study conducted in USA using a grounded theory approach and interview method.  | Men were less likely to refer to caregiving in terms of emotional work | Participants were asked three open-ended questions about gender; such directedness is unusual in dementia care research |
| Papastavrou et al , (2009) | Primary caregivers (n=172)  | Cross sectional descriptive study where all data were obtained from questionnaires. Conducted in Cyprus.  | Women use emotional coping strategies (such as day dreaming and wishful thinking) more often than men, who used more practical strategies.  | Voices of people with dementia are absent.  |
|  Pöysti et al. (2012) | Dyads of wife-husband couples. Mean (n=335).  | Quantitative study conducted in Finland; Participants were from two intervention trials  | Male caregivers of people with dementia experienced lower burden than female caregivers despite care recipients more severe disease  | Differences between men and women highlighted; Persons with dementia positioned as having no agency |
| Boyle (2013a) | 21 dyads of wife/husband couples in which one partner had a diagnosis of dementia, took part in the study which examined the gendered nature of support | Qualitative study conducted in the north of England | Gender inequality in relationships persists even when women develop dementia.  | A gender perspective is implicit and recommended to shift focus from women’s capacity to an analysis of interdependency  |
| Boyle (2013b) | Same study as above | Same study as above  | Gender influenced whether people with dementia were given the support necessary to exercise their capacity. |
| Borden & Berlin (1990) | Spousal caregivers (n=61) of older adults with dementia  | completed a questionnaire and more than half participated in a follow-up interview | Women are more prone to psychological distress than men in the context of the caregiving experience | Gendered assumptions are made about the way men and women approach illness and are affected by it.  |
| Corcoran (1992) | Scope of review is unclear.  | Literature review of gender differences in dementia caregiving and consequences of each approach for dementia management plans  | Gender differences may influence spousal caregivers definition and resolution of care management problems  | The need for gender-sensitive practice is acknowledged. |
| *men and women with dementia*  |  |  |
| Ono, Tamai, Takeuchi, & Tamai, (2010) | Hospitalised patients (n=325) with dementia; 294 were divided into men and women.  | Longitudinal study carried out over 7 year period  | Most women did not return to their homes  | Neither the reasons for this, nor what needs to be done to promote equal health outcomes are explored  |
| Buse & Twigg (2014) | People with dementia (n=32): nine men and twenty-three women took part in study  | Qualitative study conducted across three care homes in Kent and with people with dementia living in their own homes | The loss of a handbag can signify the loss of independence for women with dementia | Men’s perspective is absent.  |

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| --- | --- | --- | --- | --- |
| Ward, Campbell & Keady (2014) | People with dementia (7 women and 5 men)  | Qualitative interviews using ‘appearance biographies’ method  | Men and women reveal themselves through their appearance and the processes of managing it  | Findings on gender are not fully utilised. |
| Lindesay & Skea (1997) | Older people with dementia (n=36) (19 men and 17 women) and 37 care staff (30 women and 7 men) were observed interacting over 6 weeks | Non-participant, time-sampling observation and coding of interactions between staff and residents in single setting in the UK | Residents and staff are more likely to interact with male residents than female residents  | Reference is made to feminist theory but it is not used to analyse findings  |
| O’Connor, Phinney, & Hulko (2010) | One aboriginal woman with dementia living with a same-sex partner  | Individual case study, which generated data over a one year period through in-depth interviews, participant observation and access to health records. Conducted in Canada  | Gender as social positions/categories can work together and in opposition to shape the dementia experience | Intersectionality is considered in relation to poststructuralist and social constructionist concept of subjectivity, subject position and discourse |
|  |  |  |  |  |
| Forbes et al., (2008) | Sample of 467 males = 247; females = 220  | Cross-sectional survey design using data from the 2003 Canadian Health Survey Cycle 2.1  | Although women reported receiving more services and rated the availability of community services higher than their male counterparts, they also reported greater unmet home care needs than the men  | A gender perspective was used throughout the study  |
| Baron, Ulstein, & Werheid (2015) | Data from 21 European and North American countries were pooled. | Reviewed 73 high-quality RCTs on psychosocial interventions (PSI) outcomes in AD and MCI with respect to gender distribution and gender differences.  | Three key findings. (1) Only 45 (62%) of the RCTs reported gender ratios. In these studies women were under-represented. (2) only 19% of the RCTs analysed and reported on gender effects. In these studies women were under-represented. (3) In 40% of RCTs reporting gender analyses, gender effects on treatment outcomes were reported.  | The study raises questions about equal opportunities for women to get involved in research.  |
| Zuidema, Jonghe, Verhey, & Koopmans (2009) | People with dementia (n=1319) were recruited from 26 nursing homes across the Netherlands | Cross sectional cohort study conducted within a 4-week period  | Physically aggressive behaviour more common in men, whereas female patients demonstrated more verbally agitated behaviour  | Behaviours are attributed to the condition rather than quality of care or wider societal factors  |

## Gender differences in relation to caregiving

Twenty eight of the thirty-five studies reviewed studies focused on gender differences in relation to caregiving. This is clearly an established and growing field of enquiry, with one literature review dating back to the early 1990s (Corcoran, 1992) and another study published last year (Gibbons et al., 2014). With the exception of the study by Gibbons, et al, (2014) and another recent study reported in two papers (Boyle, 2013a,b), the perspective of the person with dementia (or care recipient) is completely absent from these studies. These findings will be reported in relation to three subthemes: caregiving from a male perspective; caregiving from a female perspective, and differences between male and female caregivers.

### Caregiving from a male perspective

Eleven of the studies focused on caregiving from a male perspective, usually the husband. Six studies were qualitative in design, three were quantitative and two were reviews of the literature on men caring for wives with dementia (Baker & Robertson, 2008, Robinson et al., 2014). These studies present valuable findings in respect of an older man’s experience of caring for his wife with dementia.

Some studies provide evidence of the interplay between caregiving and masculinity. This research reveals tensions between being a caregiver (a feminised activity) and being a man. For example, one study found that caring responsibilities can make it difficult for a man to maintain contact with their male friends (Kirsi & Jylha, 2000; Kirsi, Hervonen, & Jylhä, 2004). In another study, it was found that ‘gender role conflict’ is more significant that ‘gender identity’ as ‘some men may connote care as a feminised activity and may struggle to construe themselves as in the caring role. This may explain why some men are reluctant to access services’ (Baker, Robertson, & Connelly, 2010: 326).

In contrast, an early study suggests that some men with ‘traditional views about masculinity may view caregiving as an opportunity to express gratitude for the care their wives provided as a spouse and mother’ (Borden & Berlin, 1990: 609). More recently it has been found that some husbands may want to take control to feel good about the caregiving role, especially if they are highly-educated (Kramer, 1997, Kramer, 2000). Some evidence is contradictory. In one qualitative study it was found that men may experience isolation and invisibility because other family members do not see or recognise what they are doing (Russell, 2001). Whereas another study found that male caregivers are likely to be supported by other family members, usually women (Brown et al., 2007). In sum, male participants report wide variations in their experiences of caregiving.

### Caregiving from a female perspective

Three of the studies reviewed focused on caregiving from a female perspective. These were all qualitative in design. Research conducted by Perry (2002) uncovers a ‘process of interpreting caring’, beginning when the wife recognizes the husband is changing, moving to a phase when the wife takes over the husband’s roles and responsibilities (p.307). The researcher found that the women construct a new daily life through the process, as new identities and roles take root for the husband. Similarly, another study found that women view their caregiving role and responsibilities as paramount to other duties and concerns in everyday life, including care for themselves and need for support (Eriksson et al., 2013). This was echoed in the third study which found that women are so motivated by family obligations and ethics of care that they hesitate to access services; hence, the authors call for a ‘female/care burden-liberation’ to solve the problem (Strang, 2011). These studies show a sensitivity to gender dynamics and social power structures.

### Differences between male and female caregivers

Twelve of the studies focused on gender differences in the caregiving role and included both men and women in their samples. Five studies were qualitative in design, five were quantitative, one was a mixed method design and one was a review of spousal differences in dementia care plans (Corcoran, 1992). Most of these studies are about the ‘burden’ of caring and report on coping strategies and the adverse effects of the caregiving role; critically in these studies ‘(f) emale gender is a marker that places (female carers) at increased risk of high burden’ (Gibbons et al, 2015: 6). For example, in one of the earliest studies reviewed, gender was found to be a significant ‘correlate of psychological well-being, in that men reported lower levels of psychological distress than women’ (Borden & Berlin, 1990: 608). Similarly, researchers found that caregiver wives of people with AD reported more depression, anxiety and stress than caregiver husbands (Hooker et al., 2000). The same finding was reported from a study conducted in Greece where gender differences in overall burden depression, with females experiencing higher levels than men (Papastavrou, Tsangari, Kalokerinou, Papacostas and Sourtzi, 2009).

Some studies examine the relationship between burden and severity of impairment. One study found that ‘male caregivers of people with dementia experienced lower burden than female caregivers despite care recipients more severe disease (Pöysti et al., 2012). Similar findings were reported by Norwegian researchers who examined 27 ‘text pieces’ from a survey of adults providing care to people with dementia in their own homes and found that men more than women, say that they are satisfied with formal care and the collaboration with the rest of the family (Aasgaard 2007).

One quantitative study explored the relationship between feelings of guilt and depression in the context of dementia care-givers (Romero-Moreno et al., 2013). This study found that ‘daughters with higher levels of guilt who do not engage in leisure activities may be especially vulnerable to suffering psychological distress’ (p1). Thus providing useful evidence on how caregiving affects younger women.

Only one study addressed the issue of power dynamics between couples. This was reported in two papers by Boyle (2013a, 2013b). Findings from this study show that men may take over certain household tasks (such as cooking) when their wife develops dementia in order to gain a sense of control (Boyle, 2013a). Furthermore, as previously noted, male carers are less likely to facilitate the autonomy of their partners than their female counterparts (Boyle, 2013: 240). As the researcher explains, this shows that gender inequality in relationships persists even when women develop dementia.

## Gender differences in the lived experience

Nine of the thirty-eight studies reviewed focused on gender differences in relation to the lived experience of dementia. Findings from these studies will be reported in relation to three subheadings: effects of gender on health outcomes, effects of gender on participation, and effects of gender on use of services.

### Effects of gender on health outcomes

One study conducted in Japan focused on the effects of gender on health outcomes. The researchers explored health outcomes after discharge from a hospital ward, and found that the length of hospital stay was longer for women than men, and women were more likely than men to be institutionalised (rather than return home) on discharge from the hospital (Ono,Tamai,Takeuchi & Tamai, 2010).

### Effects of gender on social participation

Two studies examined the effects of gender on social participation. One of these examined men and women’s participation in interactions in care homes, while the other was concerned with participation in clinical trials. In the study of interaction contacts between staff and residents in a long-term care facility, it was found that men initiated significantly more interactions with staff than female residents did. Women did not initiate any interaction with male staff. Female staff initiated more interaction with residents than male staff; and finally, both male and female staff initiated higher proportions of interaction with male residents than with female residents (Lindesay & & Skea, 1997). In the other study, it was found that gender can affect levels of participation in clinical trials (Baron et al, 2015). Three key findings are reported here (1) Only 45 (62%) of the RCTs reported gender ratios. In these studies women were underrepresented (2) only 19% of the RCTs analysed and reported on gender effects. In these studies women were underrepresented (3). In 40% of RCTs reporting gender analyses, gender effects on treatment outcomes were reported. While these are very differently designed studies carried out over twenty years apart, they both show how gender can affect a person’s opportunity to participate in social life.

### Effects of gender on service use

Three of the studies reviewed examined the effects of gender on service use. One mixed method study conducted in Canada examined gender differences in relation to the use and availability of home and community-based services for people with dementia (Forbes et al., 2008). In this study, women with dementia reported greater difficulty moving about the home and were more likely to receive personal care and respite care from family members. Men were more likely to report sub-optimal functional health and to suffer the effects of a stroke (p.44). Another interesting finding was that women with dementia visited the dentist and eye specialist less frequently than men. The study also found that men with severe dementia and comorbidities were living in their homes longer than women with dementia. This implies, the authors suggest, that although men with dementia are in poorer health, their caregivers (usually their wives and/or other family members) are willing and able to care for them longer than those who are caring for women with dementia. As a result, caring for men with dementia (usually wives) may be particularly vulnerable to negative health outcomes (Forbes, et, al, 2008: 52).

In two separate Canadian studies, intersectionality theory is used to explore the position of one younger aboriginal woman with dementia (O’Connor, Phinney, & Hulko, 2010) and relations between caregivers and relatives with dementia (Chappell, Dujela and Smith, 2014) in respect of using support services. Using intersectionality theory meant that these researchers explored gender as one of several factors affecting a person’s experience of dementia and use of services. For example, in the individual case study, the women described ‘feeling out of place’ in the Alzheimer’s support group because others were much older than her and she was the only person with childcare concerns (O’Connor, Phinney, & Hulko, 2010: 36). Likewise, the results from the other study revealed that daughters experience the highest burden but also the highest esteem, suggesting that caregiving role is less salient for their self-identities, perhaps because of their younger age (Chappell, Dujela and Smith, 2014). The findings of both these studies highlight the complexity of women (and men’s lives) and the importance of taking into account other differences, such as age, ethnicity, and sexual orientation, as well as gender, when researching the lived experience of dementia.

# Limitations

We only included studies written in English and therefore acknowledge a publication bias. The review has not included work on gender published in Japanese and cited by (Ono et al., 2010), or the research on gender differences in respect of caregiving published in Korean (Lee, Cho and Kim, 2009). It is possible that other international studies have been excluded. The review did not account for study quality.

# Discussion and conclusion

The findings of this review show that men and women respond differently to the caregiving role and that gender effects the lived experience of dementia. The reviewed studies provide valuable evidence of how gender differences matter in the context of dementia care, and findings reinforce the point that others have made about factoring in such differences when policies and strategies are developed (Boyle, 2013; Brown, et. al. 2007; Dao & Woo, 2014; Forbes et al., 2008). In particular, it is recommended that ‘policy needs to be more gendered in order to promote the equality of women with dementia (Boyle, 2013a: p347). Much more emphasis is needed on influencing the policy agenda to ensure it is gender-sensitive. A point researchers have been making for over a decade (Russel 2001).

Integrating a feminist perspective into health policy is a global priority; it is widely recognised that ‘gender-based differences in access to and control over resources, in power and decision-making, and in roles and responsibilities have implications for women’s and men’s health status, health-seeking behaviours and access to health services (World Health Organisation, 2010: 14). Studies in this review have clearly established the fact that in regards to dementia care ‘female gender is a marker that places caregivers at increased risk of higher burden and less support’ (Gibbons, et. al. 2014: 16). Moreover, findings show that people with dementia are not being included or represented in studies; their experiences are not valued and so they have fewer opportunities than caregivers to say what it is like to live with this illness and receive care. Applying a feminist citizenship perspective to dementia care research in the future may help to explain why such inequalities persist, and provide a framework with which to work towards social equality and justice.

We found that research on gender differences in relation to dementia care lacks a feminist perspective – that is, few researchers start from the premise that society tends to be unequal, hierarchical and based on masculinised values and that this needs to change (Letherby, 2003). Instead, masculinised tools and measures which subjugate participants are commonly used, especially in caregiving studies (Schiffczyk et al., 2011; Romero-Moreno et al., 2013), and findings related to gender are not fully utilised or extended to discussions about what needs to change structurally and politically to end oppression. A case in point is that women’s socialisation for caregiving is a long-standing feminist issue, yet several of the studies reviewed on this topic did not problematize the position that women found themselves in (e.g. Ono et al, 2010). Moreover, the disproportionate number of studies on male caregivers identified in this review, raises questions about whether the traditional view that women are ‘natural carers’ is being challenged or reproduced in dementia care research. Richer interpretations are therefore needed that link the personal experience of dementia with wider inequalities and social locations, including gender.

It is worth emphasising here that gender does not equal women: men are at risk of oppression too. But not from being a man (Calasanti, 2004). The studies reviewed here show how one social identity can intersect with another to create disadvantage. So, for example, in the same way as the voices of women with dementia were silent in most of the caregiving studies, the voices of men with dementia were silenced too. Men were thus disadvantaged by their health status rather than gender. Additionally, some studies found variations in peoples’ experiences of caregiving based on education (Kramer, 1997), ethnicity (O’Connor, Phinney & Hulko, 2010), and age (Romero-Moreno et al., 2013) as well as gender. Such evidence highlights the need for a feminist citizenship lens in the field of dementia studies; people living with dementia will then be recast as political agents and oppressive power dynamics can be challenged (Lister, 2003).

Our aim in this work has been to take stock of knowledge about gender differences in relation to dementia care to inform policy and future research. We conclude that while there is a growing body of work concerned with gender differences in the context of dementia care, most of it fails to address the marginalisation of people living with dementia. Furthermore, the field tends to focus on the individual experiences of men and women rather than wider gender relations. As feminist disability scholars point out, ‘the experiences of individuals tell us not simply about the particular, the micro-environments, in which individuals live out their lives, but also about the general, the macro-environments which make up the broader social context of these lives’ (Thomas, 1999: 84). Fundamental questions remain therefore as to how policy makers can act now to address inequalities and promote citizenship. Moreover, how should health and social care professionals incorporate gender awareness into the support they provide (Alzheimer’s Disease International, 2015). Such questions will remain unanswered unless social scientists inject a critical gender perspective into their work, and use findings about gender differences to influence the policy arena.

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