**Material Citizenship: An ethnographic study exploring object-person relations in the context of people with dementia in care homes**

**Authors:**

Kellyn Lee, Ph.D. School of Health Sciences, University of Southampton, Southampton, UK

Ruth Bartlett, Ph.D. Associate Professor, School of Health Sciences, University of Southampton, Southampton, UK

**Abstract**

Materiality has become an increasingly important topic in sociological studies of healthcare. How objects support the identity of people with dementia in care homes is an emerging area. Whilst early research has tended to focus on sentimental or cherished items (such as photographs or keepsakes) the present study focused on functional objects (such as curling tongs or a hairdryer) as a mechanism to actualise citizenship. This article presents findings from an ethnographic study into the everyday experiences of people with dementia living in a residential care home in southern England. Drawing on a framework analysis of observations of daily life, object-elicitation interviews with residents, in-depth interviews with staff and relatives and documentary research, the findings demonstrate that object-relations are a critical but overlooked site for citizenship. Residents are rarely involved in decision-making relating to their personal possessions, lack control over objects and are often discouraged from material interactions important to the maintenance and cultivation of identity. We introduce a new concept ‘material citizenship’ to advance thinking and practice in this area and argue that it is valuable for care practices to combine a material citizenship approach with existing care practices, thus elevating the importance of object-person relations in dementia care.

**Keywords: Dementia, care homes, materialities, citizenship, objects,**

**Introduction**

Materiality is an increasingly important topic in sociological studies of health and social care. The idea of a ‘materialities of care’ is used to make visible the often unnoticed, mundane aspects of a care setting, such as the use of topical creams in a care home (Nettleton, Buse and Martin, 2018: 243). Research on the ‘materialities of care’ focuses on the role or function of actual physical objects in the caring process. Objects are considered to be a significant mediator in a networked relation of care (Araujo, et al, 2020: 1). Take for example, how an ordinary piece of furniture (like a communal seat) mediates care by providing people with the physical space to share or connect with each other in a particular way (Brownlie and Spandler, 2018). Our interest in materiality is in how access to and use of functional objects might afford citizenship, particularly for persons living with dementia in care homes. Our definition of functional objects is ‘any inanimate item which a person can use to carry out a task, not necessarily to completion or to any perceived standard, which maintains and supports his/her identity’.

Concern for the citizenship of persons with dementia is growing, as evidenced by global policy work on the human rights and meaning of shared responsibility for people living with this condition (see, for example, Shakespeare, Zeilig and Mittler, 2019). To date, research on citizenship has focused on human relationships rather than object-person relations. In this article, we use a material lens to examine access to, and use of functional objects by persons with dementia living in a care home setting, and coin the phrase ‘material citizenship’ to advance thinking and practice in this area.

**Aligning citizenship, materiality and people with dementia in care homes.**

Citizenship is an increasingly important concept for enhancing the status, rights and opportunities afforded to people with dementia (Baldwin, 2008; Bartlett and O’Connor, 2010; Bartlett, 2014; Bartlett and O’Connor, 2007; Brannelly, 2011; Österholm and Hydén, 2016). Various studies have used the idea of citizenship to draw attention to power relations and discriminatory practices in care settings (Bartlett and O’Connor, 2010). For example, Brannelly (2011) investigated the [lack of] participation by people with dementia in decisions about their care, in terms of citizenship. In another study, communication barriers between social workers and persons with dementia were analysed as ‘citizenship as practice’ (Österholm and Hydén, 2014). The notion of citizenship is useful for bringing to the fore peoples’ relationships to the ‘larger structures of rule and belonging, which are often but not exclusively nation states’ (De Koning, Jaffe, and Koster 2015: 121). For people with dementia, such structures are invariably embedded in care systems and practices as the condition adversely affects a person’s capacity to take responsibility for themselves.

To date, social citizenship has lacked both a material lens and a practical application for successful implementation in a care context. We understand social citizenship to mean a ‘relationship, practice or status, in which a person with dementia is entitled to experience freedom from discrimination (and despair), and to have opportunities to grow and participate in life to the fullest extent possible' (Bartlett and O’Conner, 2010: 36). While the idea is applicable to any setting, research on citizenship has tended to focus on those living in their own homes and how it might actually manifest in a care home has yet to be examined. In this article we argue that the access to and use of functional objects, can be a useful mechanism to enable social citizenship.

Whilst dementia varies from person to person a common and noticeable difference is the way in which interactions with everyday objects can become challenging and cumbersome. Activities such as ironing a shirt or fastening a blouse, once easy and taken-for-granted will often require noticeable effort and concentration (Phinney and Chelsa, 2003). Other activities, such as making a drink, or cooking a meal can become difficult, maybe even impossible without help. This is because such tasks involve a high degree of executive functioning – i.e. using various objects in a specific sequential order, which is challenging for someone with a severe cognitive impairment like dementia. Nevertheless, removing access to the objects required to complete everyday tasks can deny a person citizenship, as it effectively refutes a person’s capacity for self-expression.

The importance of objects for the expression of identity and agency is well documented, and central to object-relations research (Belk, 1988; Csikszentmihalyi and Rochberg-Halton, 1981; Mccracken, 1987; Rubenstein, 1987). Recent studies have examined the link in various contexts and in relation to particular objects. For example, one study which explored the materialities of care in relation to people with dementia in care homes, found handbags to be an important object for female residents; the ‘rummaging in handbags’ sought to create boundaries of privacy and facilitated a sense of identity (Buse and Twigg, 2014: 11). Another study observed how people living in a dementia care unit reconfigured materialities to make a space for themselves and/or other residents (Cleeve, 2020). Other work, which does not include people with dementia, but is nonetheless relevant found that interactions with material culture (such as hand-held vacuum cleaners) help to facilitate interaction and meaningful relationships (Lovatt, 2018, 2020). These findings highlight the significance of functional objects in the context of a care home. Moreover, they confirm how achieving a sense of home can be made easier if a person has access to resources that maintain and cultivate identity (Molony, 2010; Cooney, 2012).

Whilst objects are often overlooked in dementia care research or viewed as secondary to person-person relationships or variables to control (Ceci, Symonds Brown and Purkis, 2019) they can be useful mechanisms offering insights into relationships supporting choice, control and agency (Driessen, 2019; Araujo *et al.*, 2020). For example, using a dining table to maintain a sense of dignity, control and agency (Kroger and Adair, 2008); a television to feel connected to the outside world (Nord, 2013); a mobile phone to maintain a spousal relationship (Ursin and Lotherington, 2018). The present study focused on functional objects (such as curling tongs or a hairdryer) that enable people to take part in activities of daily life.

The freedom to choose the life one wishes to have, which may include using objects, is an important aspect of citizenship (Pfister, 2012). It is especially relevant for people with a disability, as opportunities to live a meaningful life are often curtailed by others (Lid, 2015); through materialist practices. Take for example, Ursin and Lotherington’s (2018) report of a young man with dementia living in a nursing home who used his mobile phone each day to speak with his wife. On two occasions, staff found the mobile phone in the laundry, on bringing a third replacement mobile phone, staff informed his wife that the mobile phone would be kept in the staff room. Furthermore, she was asked not to call him on the mobile phone as staff could not hear it ring. As a result, the man’s chance to communicate with his wife on his own terms was lost, indicating a link between the materialities of care and citizenship, which we aim to examine further in this research.

In this article, we use a material lens to examine access to, and use of functional objects by people with dementia living in a care home. Rather than focusing on how such practices mediate care; we examine how functional objects might afford social citizenship. For a care home resident, the practice of citizenship might be as mundane as curling their hair with tongs, the point is that the action of using this functional object enables the resident to enact identity, exert some control and take some responsibility for themselves.. Increasingly, scholars are calling for more recognition of the capacity of people with dementia to take control, and regard ‘citizenship-in-and-as practice’ as a potentially useful framework for realising that (Nedlund, Bartlett and Clarke, 2019). The aim of this paper is to add to that small body of work by focusing on access to, and the use of functional objects in the context of people with dementia in care homes.

**Methodology**

The findings discussed in this article form part of data collected through doctoral research completed in 2019. The study was given ethical approval by the NHS Health Research Authority Social Care Research Ethics Committee (HRA Social Care REC) (Reference number: 17/IEC08/0003). The primary objective of the study was to understand how objects manifest in the everyday lives of people with dementia living in care homes. A secondary objective was to examine whether social citizenship was a useful lens to examine this. An ethnographic approach enabled observations of object-person relations in everyday life in the care home for residents, staff and relatives. It would provide insight into how functional objects are used and by whom. Whilst the study was conducted in two care homes in southern England[[1]](#footnote-1), this article draws on the findings of one ‘Southcote Manor’[[2]](#footnote-2).

Southcote Manor is a two-storey local authority residential care home offering care to 30 older adults including people with dementia. Each floor has corridors, with a mix of residents’ rooms, communal areas and bathrooms. Staff offices are located on the ground floor and residents move between floors when accompanied by staff. All bedrooms are of similar design but varied in size. Most rooms had identical furniture such as a bedside table, wardrobe, sink and commode. Some residents had brought in a small item of furniture from home, such as a chair, but this was often discouraged due to limited space and health and safety concerns.

A four-week hanging-out period (Bernard, 1995) commenced at Southcote Manor in July 2017. This provided an opportunity to build rapport prior to data collection and answer any questions about the study. Time was spent in communal areas, talking with residents, staff and relatives, taking part in organised activities, helping staff at mealtimes and eating with residents. Most residents and relatives appeared to accept the presence of a researcher, some residents mentioned enjoying having a new person to talk to. Some staff found it difficult, at times leaving the room when observations were taking place, and on two occasions challenging her presence. The hanging-out period allowed time to gain some familiarity of the building, memorise door codes, and observe care home life. Data collection was carried out by the first author between the months of August and November 2017.

A convenience sampling technique was used to recruit three groups of participants to the study (1) residents with dementia (2) staff employed for a minimum of six months, and (3) relatives who regularly visited the care home. It was important to elicit these differing perspectives to understand how decisions are made about functional objects. The registered manager of Southcote Manor acted as a gatekeeper and distributed participant information sheets to residents, staff, and relatives. The first author met with potential participants to discuss the study and answer any questions before deciding whether to take part.

Thirty-nine participants took part in the study; 15 residents with dementia (11 female and 4 male, average age 88); 16 staff (12 female and 4 male, average age 46) and eight relatives (6 female and 2 male, average age 60). Of the 39 participants that took part in the study, 38 were of white British origin, one member of staff was Romanian.

Data were collected over a 16-week period and included participant observations, object-elicitation interviews (with residents), in-depth interviews (with staff and relatives) and documentary research methods. Participant observations took place in the communal areas of the care home, three days per week, four hours per day, between the hours of 8.00am and 8.00pm. This resulted in 144 hours of participant observations. Participant observations and fieldnotes were written to capture informal conversations and object-person interactions such as, who used the remote control for the television, who carried out domestic chores, and whether residents had choice over where they sat.

Object-elicitation interviews were conducted with four residents, using the objects in a person’s room to guide the interview; asking questions such as; ‘shall we have a look at some of the things in your room?’, ‘how often do you use this?’ and ‘who decided what was brought in?’. Nineteen in-depth interviews were conducted: eleven with staff, and eight with relatives. Most in-depth interviews with staff and relatives took place in the staff office or staff lounge, one interview took place in a relative’s home. Within the interviews we explored which objects were important to the interviewee and how it might feel to have those objects taken away. We also explored who was involved in deciding which belongings residents could have and reasons why some objects were viewed as acceptable for care home life while others were not. Documents such as: quality standards documents, individual resident care plans and an inventory log were analysed to understand how objects were considered within care practices.

*Consent procedures*

Written consent was sought and provided by all participants except those who were assessed as lacking capacity to take part in research. For those participants, a personal consultee (relative, friend, spouse or independent advocate) provided either written or verbal agreement. People with dementia assessed as lacking mental capacity were included in participant observations when a personal consultee provided a favourable opinion. In line with HRA Social Care REC recommendations, only residents assessed as having capacity to take part in research were eligible to take part in object-elicitation interviews. In addition to this, a process consent method was used at every participant observation; this involved approaching each participant at every data collection point, making clear that research was taking place, what it involved and asking whether the participant was willing to take part.

*Data Analysis*

A framework analysis approach was used, using Nvivo 11, to enable structured and factual questions to be asked of the data (Ritchie and Spencer, 1994; Srivastava and Thomson, 2009). A prime concern was to describe and interpret what was happening within a particular context, how objects were used in practice and their inclusion in policy and practice documents.

The first step of familiarisation provided an opportunity to gain an overview of the data provided by the three participant groups. Transcripts and fieldnotes were read and audio-recordings listened to whilst making notes about their content. The second stage of the analytical process was to organise the data in a meaningful and manageable way to aid future exploration and the retrieval and examination of data later-on. The formation of framework categories informed by a priori concerns and emergent issues allowed the identification of categories that best fit the research questions. This is also meant the framework included the interests of the researcher and issues pertinent to participants (Richie and Spencer, 1994).

A set of preliminary codes were produced in step one pulling together all data sets, providing overall themes about what was said within interviews and what was observed in practice. The next step was to ask specific questions related to the research question such as: (1) How are decisions made relating to personal possessions and who is involved in decision-making? (2) Who is responsible for bringing personal possessions into the care home, and how are future objects acquired? (3) How are functional objects used in care homes and who uses them? (4) Are measures taken to control objects in the care home and if so what measures and by whom?

Codes created through interview and observational data were merged and overlapping codes were collapsed. These were reorganised into four categories: (1) decision-making relating to the ownership and use of functional objects, (2) access to functional objects to participate in everyday life, (3) safety or the perceived risk of having or using functional objects, and (4) relationships with objects, ownership and use of functional objects.

The third stage of the analytical process involved systematically applying each interview transcript and fieldnote to each framework category and comparing these to practice and policy documentary sources. This revealed how decisions were made, by whom and whether these decisions were influenced by policy guidance.

In stage four of the analytical process a framework matrix was generated. This provided a simple summarised version of the interview and observation data, (see table 1 for matrix staff interview data).

**Table 1. Matrix staff interview data**

|  |  |
| --- | --- |
| Safety | If they said they wanted to do their hair, the carers are quite happy here to let them, they would stay there, supervising them while they did their hair anyway. They do get that option anyway. They can’t just take it down. |
| Decision-making | No, usually it’s the families. We have done it for a couple of residents, when we’ve gone back to their home for them to get some belongings, but usually the family members have taken over and unfortunately some of them dump stuff before even letting the residents look at what’s left and things like that. |
| Taking action | Say I went into the dining room and I feel a little bit low, I could do with someone bringing out maybe one of my old knitted jumpers or something, or a picture. I personally would hope that a carer would listen to me, and then go and grab that item for me, so I could look at it or cuddle it. |
| Control | I know some have come in with mobile phones, so if they wanted to phone their families, but a lot of the families don’t want them to have telephones. |

The final stage of analysis was to use the matrices to pull together the key characteristics of the data and map and interpret the data as a whole. This provided an understanding of how decisions were made about objects, and who was involved, who had access to objects and how were they used and provided reasons why some residents had control over objects whilst others did not. It was also useful in illuminating contradictions and inconsistencies within and between groups and data sets.

**Findings**

Applying a material lens to care home life provided an opportunity to explore what citizenship looked like for those who took part in this study. By focussing on objects and viewing care home life through a material lens, we were able to determine whether citizenship was supported in care practices for people with dementia.

The study presents two key findings. First, we found a ‘*lack of* *opportunities to return home to collect personal possessions’.* Second, we found a ‘*lack of* c*ontrol over personal possessions and functional objects’*. From this, we developed the framework of ‘material citizenship’.

*Lack of opportunities to return home to collect personal possessions*

This study found that residents who moved directly from hospital to the care home lacked the opportunity to return home to collect their belongings. Furthermore, similar to other researchers (Innes, Kelly and Dincarslan, 2011; Lindley and Wallace, 2015) residents were often not involved in deciding which belongings they wanted to take and lacked control over what happened to their items. Take for example, 94-year-old Karen. Karen was admitted to hospital following a fall at home; it was there the decision was made for her to move directly to a care home, she was not given an opportunity to return home to collect her possessions. Karen had lived at Southcote Manor for three months when the study began, she would talk about her experience of moving into the care home at times with anger and other times with hopelessness at her lack of agency. She spoke of how decisions were made for her and how little control she had over her life and her possessions. Karen mentioned that she would have organised her own removals had she had control over her finances. Instead, she had to wait for her nephew to find the time and ‘hope’ that he would bring the items she had requested. Karen felt disempowered by the process of moving into the care home and the lack of opportunities to organise her possessions.

Staff were asked whether residents living in the care home had opportunities to return home to collect belongings:

*“No, usually it’s the families. We have done it for a couple of residents, when we’ve gone back to their home for them to get some belongings, but usually the family members have taken over and unfortunately some of them dump stuff before even letting the residents look at what’s left and things like that.”(Abby, staff, interview).*

This excerpt shows that whilst relatives usually take on the task of dealing with a person’s possessions, staff had on occasions returned to a person’s home to collect belongings. This was not common practice and residents did not accompany staff when doing so. Staff mentioned being the ‘go-between’ for residents, taking requests for items they wanted, contacting relatives to pass on this request but having no control over the outcome. Confounding our view that once a person becomes a ‘resident’ they lose control of their possessions.

Previous research shows that rather than viewing objects as a way in which to engage in normal social interactions, objects can become problematic (England and Dyck, 2011). The present study found the downsizing process was difficult for all concerned. An example of this is shown in a son’s explanation of why he excluded his mother when organising her belongings:

*“[If]I hadn’t done it that way, and my mother had been there, I most probably would have still been there today, trying to clear it all out because it would have been, oh no I can’t, I don’t want to get rid of that; where are you going to put it; there is no room to move a five-bedroom house…. into a one-bedroom home” (Finlay, relative, interview).*

For Finlay, the task of sorting his mother’s belongings was onerous; Finlay lived in the south of England, his mother’s house was in the north of England; he worked full-time and had two young children. Supporting a person to organise and collect belongings can be a complex process, fraught with difficulties. It is therefore unsurprising that relatives and staff may want to avoid this to reduce possible complications.

Other members of staff echoed these difficulties. For example, Elaine the registered manager believed that enabling a resident to return home presented a risk, the resident might refuse to return to the care home:

*“There’s always that risk that they won’t then, they want to stay and they don’t want to come back. But I mean I would do that; that would be me. If I went home and then had to pick bits, I wouldn’t want to leave again. That’s my home. Rather than putting them through that and the trauma and making them upset.” (Elaine, staff, interview).*

Although this is a plausible assumption, particularly as a person may not have chosen to live in the care home in the first instance, given that none of the participants in this study had an opportunity to return home, there is no evidence to support this claim. Nevertheless, it means that people have very little control or choice over personal possessions. Not affording people the chance to return home is a practice that denies new residents the right to access their personal possessions, we suggest, indicates a lack of material citizenship.

In addition to the aforementioned complexities of enabling a person to return home, staff and relatives reported short timeframes as a barrier to collecting personal possessions:

*“You’ve only got a short space sometimes to get it. Because either they’re in rented accommodation and if they’re not going to be able to go back there someone’s going to clear it, or they lived on their own and then it’s going to be packed up and sold, or they lived with someone else, and that’s not quite so bad, you know. But it can be difficult because there’s like a timespan when you can get these things. And sometimes they might think of it after it’s gone” (Elaine, staff, interview).*

Although Elaine was aware that access to personal possessions was a problem and timescales could be tight, there was no strategy in place to tackle this. Karen described how she had waited for her nephew to bring in her items:

*“He [nephew] thought he’d bring them to me I think, but time went on and he couldn’t seem to get here. And I never saw them [belongings] again.* *(Karen, resident, interview).*

This emphasises an imposed dependency; Karen was positioned as a passive recipient rather than an active citizen, she was unable to arrange delivery of her possessions, instead she had to wait for her nephew to bring the items to her. Karen mentions that she never saw some of her belongings again, demonstrating how little control she has.

*Lack of control over personal possessions and functional objects*

Through comparisons across codes, both within and between participant groups, and comparing the three data sources (interviews, participant observations and documentary resources) the responsibility for making decisions and the access people had to functional objects oscillated between relatives and staff. For example, staff would take responsibility for the removal of a resident’s possessions as and when they felt it was necessary:

*“the other day a resident was asking me for some of her Veet hair removal cream, she wanted to put some on her chin, but I saw it in the room there, it was on top of the chest of drawers, it was very reachable, but you know I had to grab it and put in my pocket, because I thought this is possibly something that she shouldn’t have, because it is meant to be on top of the wardrobe, so it’s well out of reach, but it wasn’t. So I said to her, she didn’t realise I’d put in my pocket, so I said to her I’ll just go down to the office and see if they’ve got it there. I went down to the office and said to Elaine , look like she’s got this Veet, I don’t know whether to give it to her, because it’s got a label on it that says ‘out of reach’, and Elaine says, no don’t give her that, just leave it down here in the office, and we’ll sort it. So I just ended up having to go back to her room and say, look it’s not here, I can’t find it, I think they’ve taken down to the office, and you know for safety reasons” (Michael, staff, interview).*

In this case, the object was removed by the member of staff without informing her. This study found no evidence of risk assessments taking place concerning functional objects. Instead, staff made subjective decisions on an ad hoc basis. Given the lack of guidance in policy and practice guides, and the focus on safety, this is perhaps unsurprising.

The local authority care home quality standards, included as a documentary source, provided guidance encouraging certain objects into the care home. The document listed items such as toiletries, soft furnishings, photographs and a small item of furniture. Clothing that should be able to withstand a 60-degree wash, be a size bigger than usual and have an elasticated waistband to assist staff with dressing and undressing. Objects were viewed and controlled through a service lens rather than as items important for enabling citizenship, or as a mechanism for gaining a deeper understanding of a person’s historical biography and present identities (Buse and Twigg, 2015). The guidance showed the tension around these spaces, as both a ‘home’ and as a regulated workplace.

Due to ethical considerations, it was not easy to ascertain how decisions were made about clothing. No observations were allowed in residents’ rooms. However, on one occasion, Karen was asked if she planned to attend the organised music session; she said she didn’t want to go in the clothes she was wearing. She had on a red cardigan that she did not like but ‘they’ insisted she put it on, and a floral skirt she said she would not have chosen to wear. Karen independently changed her clothes regularly throughout the day, with staff commenting ‘oh you’ve got changed again’. This suggests she had some freedom to access her clothes and change them as and when she pleased.

The degree of control residents had over functional objects, related to the level of support from staff. For example, whilst Karen was able to use her wooden walking stick covered in Scandinavian badges and carry her daily newspaper around the home, staff did not allow her curling tongs. Karen often commented on how she would like to style her hair:

*“Oh I would love to have something to do my hair, I like it curled, I want my curling tongs that is what would do it.” (Observation)*

Karen asked for curling tongs, but staff viewed this item as risky; they thought she might burn herself, or another resident as they frequently walked into each other’s rooms. Instead, staff provided Karen with plastic curlers, which she found difficult to use. This shows how perceptions of risk can act as a barrier to citizenship in a care home context (Buse and Twigg, 2014).

An expression of material citizenship was observed in the seating arrangements in the communal areas. Residents would ‘claim’ lounge chairs, that belonged to the home for themselves. Bill, a member of staff, explained that a lack of chairs in the downstairs lounge had resulted in ‘the ladies’ attempting to leave the table early in order to get ‘their’ seat. The following observation recorded this:

*Emily leaves her seat, her meal unfinished. Bill tells Emily he knows what she is up to, Emily responds with a laugh and makes her way to ‘her’ chair. (Observation)*

Objects such as remote controls for televisions, videos, CD’s and music players were present in communal spaces. Some staff asked residents which tv channel or music they preferred, but only staff were observed using the objects during fieldwork. A room located on the ground floor had an abundance of objects, such as an organ, puzzles, books and craft items but these were only accessible to staff. A few residents were observed taking cups to the sink or looking around the kitchenette area designed for their use, but staff generally encouraged residents to stay seated. Household chores were mainly carried out by staff with the exception of one relatively young resident with dementia – 69 year old Pippa. Pippa was afforded the responsibility of washing up, cleaning down, setting tables for mealtimes, dusting handrails and folding napkins:

*‘Pippa finishes her sandwich she then gets up from the table and takes her plate over to the sink. Pippa begins to wash up. Pippa picks up a cloth and begins to wash down the kitchen sides. Bill works around Pippa. Pippa is rinsing cups in the water. Bill works with Pippa giving directions so that he can pack the dishwasher.’ (Observation)*

Carrying out domestic chores was important to Pippa. She often became agitated if staff cleared away plates or began tidying the kitchen area. Pippa clearly struggled with executive functioning, often not knowing what to do with objects, nevertheless, staff used techniques such as guiding Pippa’s movements or showing her where to put place objects, akin to ‘scaffolding’(Österholm and Hydén, 2014). Staff supported her in carrying out chores considered risky such as, cleaning tables with detergent, washing up pots and cutlery, filling the dishwasher and making cups of coffee. When asked why Pippa was supported in these daily activities the response from staff was ‘because she’s young’. This shows another layer of complexity in object-person relations - that is, it is not simply about cognitive ability and risk but the perception of age and what that represents.

**Discussion**

In this article, we suggest that residents living with dementia at Southcote Manor lacked control over functional objects. Many residents were not involved in decision-making relating to the belongings they brought from home or were unaware of what had happened to them. Like other researchers, we found that people with dementia are routinely excluded from the decision-making process (Fetherstonhaugh *et al.*, 2016). In the context of this study, functional objects manifest in risk averse practices. Whilst risk management is necessary in care provision and can be a positive process, it can also disempower and be restrictive if done so discriminatorily. Examining decision-making and the use of objects through a social citizenship lens is useful in drawing out care practices that do or do not afford people with dementia the same opportunities as others. For example, Karen was denied access to her curling tongs, because staff considered it risky.  Whereas Pippa was encouraged to carry out domestic chores with items considered risky. A strength of material citizenship is that access to objects can enable people to live the life that they want to live and can be facilitated by care home staff. This may be more applicable to people with milder degrees of cognitive impairment however, objects can also function as an external memory support (Hyden, 2014) which could also be useful in care practice.

Whilst other studies have referred to residents’ interactions with their material surroundings, as providing the ability to continue routine practices such as organising their surroundings, cleaning and hosting (Nord, 2013; Lovatt, 2018), this study only observed one person enabled to carry out these tasks. Instead, staff were often observed discouraging interactions from taking place. By including people with dementia and applying a material lens to care home life we were able to draw out subtle injustices in care practice. Injustices that were not intentional, but nevertheless present. There was also a lack of guidance in the management of possessions, or an understanding of the importance of object-person relations in policy and practice documentation. By emphasising the importance of the materialities of care, applying a material lens to care practice, we were able to show a continuous reduction of control and independence for people with dementia living in a care home.

Other health sociologists have argued that by paying more attention to materiality, ‘everyday routines could become important occasions for care’ (Cleeve, Borell and Rosenberg, 2019; 126). Our analysis confirms this and takes it further by showing how material practices, specifically access to, and use of functional objects, has significance for social citizenship. For instance, disposing of a person’s belongings when they move into a care home is an example of a ‘material practice that is made mundane and inconsequential’ (Latimer, 2018: 379). Importantly, from our perspective, it shows a disregard for a person’s need for familiar objects to participate in life – i.e. social citizenship. The value of material citizenship is that it focuses on the nuances of a person’s day-to-day routines and rituals and the functional objects they need to keep these going. Material citizenship is thus defined as:

*‘the right to be included in decision-making relating to personal possessions and the right to have opportunities to use functional objects to perform everyday tasks.’*

The ability to make decisions and have those decisions upheld is central to self-determination (Fetherstonhaugh, Tarzia and Nay, 2013). Material citizenship aims to promote this by focusing on a person’s control over and rights and relationships in respect of functional objects, including for example, being able to participate in everyday life practices such as household chores. Additionally, the concept aims to help elevate an understanding of the significance of functional objects and the role they can play in care assessments and care planning. This is particularly relevant for people living with dementia in care homes, as policies and procedures are often designed to meet physical care needs rather than promote social citizenship.

The idea of material citizenship aligns and builds upon knowledge related to the materialities of care and social citizenship providing a pragmatic way of operationalising a person’s rights and responsibilities. Material citizenship provides a framework with which to educate, implement and evaluate the presence of social citizenship in a care home environment. By focusing on how decisions are made about belongings, and the access people have to functional objects, a balance can be achieved which provide assessments of proportional and reasonable risk of physical harm and citizenship. This would require conversations to take place between residents, relatives, and staff to explore and formally agree a plan of how a person can be supported to live the life they would like to live, including the objects needed to achieve this. For those with more advanced dementia, who may have difficulties with verbal communication, objects could be utilised as a method to explore how a person with dementia wants to live their life. Embedding material citizenship in organisational documentation is a potentially useful way of helping staff consider the role of functional objects in care. That said, larger-scale studies are needed, which involve more diverse groups of people with dementia, to understand fully the value and limitations of material citizenship as a conceptual framework for care.

As with all studies, there are several limitations. First, the data presented here were collected from a single study site in southern England and therefore no account is taken of variability between care homes, regionally, nationally or globally. Second, object-elicitation interviews were perceived as problematic by the ethics committee, and so people assessed as lacking mental capacity (i.e. a large proportion of the care home population) had to be excluded from taking part in the study. Third, in line with HRA recommendations, it was not possible to observe how functional objects were used during personal care and in resident’s room. As such, the findings only provide a partial picture of people’s access to and use of functional objects in care homes.

**Conclusion**

The growing number of people with dementia moving into a care home and increasing pressure with regard to a person’s rights raises urgent questions about peoples’ access to and use of functional objects. We have argued for the importance of bringing a material lens to sociological and psychological discussions of citizenship and care, specifically in the context of persons with dementia in residential care homes. Findings from this study suggest that functional objects are overlooked in the context of care homes and there is a lack of guidance for care workers. Thus, residents depend upon the whim of individual staff members, as to whether they can access a particular object, rather than through an organised and structured care planning and risk management process. Understanding how functional objects manifest in care practices made visible the lack of social citizenship afforded to people with dementia in this setting. We suggest it would be valuable for care practices to combine a material citizenship approach with existing care practices. This would elevate the importance of object-person relations, allowing staff and relatives to recognise that functional objects can help a person maintain identity and influence how they are perceived.

Finally, the concept of ‘material citizenship’ usefully extends our understanding social citizenship by foregrounding the role of objects in peoples’ lives. Specifically, it shows how functional objects are not only an extension of the self (Belk, 1988); but also, making decisions about and using functional objects enable people to take control and practice everyday citizenship, such as taking care of oneself and others. As such the concept, once it has been more widely applied and assessed in the dementia field, may be useful in other areas of health and social care where object-person relations are as important, but often overlooked, including hospitals and care homes for people for learning disabilities.

*The data that support the findings of this study are available on request from the corresponding author. The data are not publicly available due to privacy or ethical restrictions.*

**Acknowledgements**

Our sincere appreciation goes to all of those who took part in the study, and to the Alzheimer’s Society for funding this study

**Correspondence**

Faculty of Environmental and Life Sciences, School of Health Sciences, University of Southampton, Southampton, UK. Kellyn.lee@soton.ac.uk

**References**

Araujo, M. T., I. S. C. Velloso, C. Ceci, and M. E. Purkis. 2020. "The significance of overlooked objects: Materiality and care at home for people with dementia." Nursing Inquiry 27 (1). doi: 10.1111/nin.12306.

Araujo, M. T. *et al.* (2020) ‘The significance of overlooked objects: Materiality and care at home for people with dementia’, *Nursing Inquiry*, 27(1), pp. 1–8. doi: 10.1111/nin.12306.

Baldwin, C. (2008) ‘Narrative(,) citizenship and dementia: The personal and the political’, *Journal of Aging Studies*, 22(3), pp. 222–228. doi: 10.1016/j.jaging.2007.04.002.

Bartlett, R. (2014) ‘Citizenship in action: the lived experiences of citizens with dementia who campaign for social change’, *Disability and Society*, 29(8), pp. 1291–1304. doi: 10.1080/09687599.2014.924905.

Bartlett, R. and O’Connor, D. (2007) ‘From personhood to citizenship: Broadening the lens for dementia practice and research’, *Journal of Aging Studies*, 21(2), pp. 107–118. doi: 10.1016/j.jaging.2006.09.002.

Bartlett, R. and O’Connor, D. L. (2010) *Broadening the Dementia Debate Towards Social Citizenship*. Policy Press Bristol.

Belk, R. W. (1988) ‘Possessions and the extended self’.

Bernard, H. R. (1995) *Research Methods In Anthropology Qualitative and Quantitative Approaches*. 2nd Ed. London: AltaMira Press.

Brannelly, T. (2011) ‘Sustaining citizenship: People with dementia and the phenomenon of social death’, *Nursing Ethics*, 18(5), pp. 662–671. doi: 10.1177/0969733011408049.

Brownlie, J. and Spandler, H. (2018) ‘Materialities of mundane care and the art of holding one’s own’, *Sociology of Health and Illness*, 40(2), pp. 256–269. doi: 10.1111/1467-9566.12574.

Buse, C. and Twigg, J. (2014) ‘Looking “out of place”: Analysing the spatial and symbolic meanings of dementia care settings through dress’, *International Journal of Ageing and Later Life*, 9(1), pp. 69–95.

Ceci, C., Symonds Brown, H. and Purkis, M. E. (2019) ‘Seeing the collective: Family arrangements for care at home for older people with dementia’, *Ageing and Society*, 39(6), pp. 1200–1218. doi: 10.1017/S0144686X17001477.

Cleeve, H. (2020) ‘Markings: Boundaries and Borders in Dementia Care Units’, *Design and Culture*, 12(1), pp. 5–29. doi: 10.1080/17547075.2020.1688053.

Cleeve, H., Borell, L. and Rosenberg, L. (2020) ‘(In)visible materialities in the context of dementia care’, *Sociology of Health and Illness*, 42(1), pp. 126–142. doi: 10.1111/1467-9566.12988.

Cooney, A. (2012) ‘“Finding home”: A grounded theory on how older people “find home” in Long-term care settings’, *International Journal of Older People Nursing*, 7(3), pp. 188–199. doi: 10.1111/j.1748-3743.2011.00278.x.

Csikszentmihalyi, M. and Rochberg-Halton, E. (1981) *The meaning of things Domestic symbols and the self*. Cambridge University Press.

Driessen, A. E. (2019) *A good life with dementia*. Amsterdam Institute for Social Science Research. Available at: https://dare.uva.nl/search?identifier=dd0c2b9b-348d-4de8-9747-84363846fdd0.

England, K. and Dyck, I. (2011) ‘Managing the body work of home care’, *Sociology of Health and Illness*, 33(2), pp. 206–219. doi: 10.1111/j.1467-9566.2010.01331.x.

Fetherstonhaugh, D. *et al.* (2016) ‘“The Red Dress or the Blue?” How Do Staff Perceive That They Support Decision Making for People with Dementia Living in Residential Aged Care Facilities?’, *Journal of Applied Gerontology*, 35(2), pp. 209–226. doi: 10.1177/0733464814531089.

Innes, A., Kelly, F. and Dincarslan, O. (2011) ‘Care home design for people with dementia: What do people with dementia and their family carers value?’, *Aging and Mental Health*, 15(5), pp. 548–556. doi: 10.1080/13607863.2011.556601.

de Koning, A., Jaffe, R. and Koster, M. (2015) ‘Citizenship agendas in and beyond the nation-state: (en)countering framings of the good citizen’, *Citizenship Studies*, 19(2), pp. 121–127. doi: 10.1080/13621025.2015.1005940.

Kroger, J. and Adair, V. (2008) ‘Symbolic Meanings of Valued Personal Objects in Identity Transitions of Late Adulthood’, *Identity: An International Journal of Theory and Research*, 8(1984), pp. 5–24. doi: 10.1080/15283480701787251.

Latimer, J. (2018) ‘Afterword: materialities, care, “ordinary affects”, power and politics’, *Sociology of Health and Illness*, 40(2), pp. 379–391. doi: 10.1111/1467-9566.12678.

Lid, I. M. (2015) ‘Vulnerability and disability: a citizenship perspective’, *Disability and Society*, 30(10), pp. 1554–1567. doi: 10.1080/09687599.2015.1113162.

Lindley, S. and Wallace, J. (2015) ‘Placing in Age : Transitioning to a New Home in Later Life’, *ACM Transactions on Computer-Human Interaction*, 22(4), pp. 1–40.

Lovatt, M. (2018) ‘Becoming at home in residential care for older people: a material culture perspective’, *Sociology of Health and Illness*, 40(2), pp. 366–378. doi: 10.1111/1467-9566.12568.

Lovatt, M. (2020) ‘Relationships and material culture in a residential home for older people’, pp. 1–18. doi: 10.1017/S0144686X20000690.

Mccracken, G. (1987) ‘Culture and Consumption among the Elderly: Three Research Objectives in an Emerging Field’, *Ageing & Society*, 7(02), pp. 203–224. doi: doi:10.1017/S0144686X00012563.

Molony, S. L. (2010) ‘The Meaning of Home: A Qualitative Metasynthesis’, *Research in Gerontological Nursing*, 3(4), pp. 291–307.

Nedlund, A.-C., Bartlett, R. and Clarke, C. L. (2019) ‘Everyday Citizenship: A way to broaden our view of life with dementia’, in *Everyday Citizenship and People with Dementia*. Edinburgh/London: DUNEDIN.

Nettleton, S., Buse, C. and Martin, D. (2018) ‘“Essentially it’s just a lot of bedrooms”: architectural design, prescribed personalisation and the construction of care homes for later life’, *Sociology of Health and Illness*, 40(7), pp. 1156–1171. doi: 10.1111/1467-9566.12747.

Nord, C. (2013) ‘A day to be lived. Elderly peoples’ possessions for everyday life in assisted living’, *Journal of Aging Studies*, 27(2), pp. 135–142. doi: 10.1016/j.jaging.2012.12.002.

Österholm, J. H. and Hydén, L. C. (2014) ‘Citizenship as practice: Handling communication problems in encounters between persons with dementia and social workers’, *Dementia*, 15(6), pp. 1457–1473. doi: 10.1177/1471301214563959.

Österholm, J. H. and Hydén, L. C. (2016) ‘Citizenship as practice: Handling communication problems in encounters between persons with dementia and social workers’, *Dementia*, 15(6), pp. 1457–1473. doi: 10.1177/1471301214563959.

Pfister, T. (2012) ‘Citizenship and capability? Amartya Sen’s capabilities approach from a citizenship perspective’, *Citizenship Studies*, 16(2), pp. 241–254. doi: 10.1080/13621025.2012.667615.

Phinney, A. and Chelsa, C. . (2003) ‘The lived body in dementia’, *Journal of Aging Studies*, 3, pp. 283–299.

Ritchie, J. and Spencer, L. (1994) ‘Analysing qualitative data’, in Bryman, A. and Burgess, R. G. (eds) *Analysing qualitative data*. Routledge: London, pp. 173–194.

Rochberg-Halton, E. (1984) ‘Object Relations, Role, Models, and Cultivation of the Self’, *Environment and Behavior*, 16(3), pp. 335–368.

Rubenstein, R. L. (1987) ‘The significance of personal objects to older people’, *Journal of Aging Studies*, 7, pp. 225–238.

Shakespeare, T., Zeilig, H. and Mittler, P. (2019) ‘Rights in Mind: Thinking Differently About Dementia and Disability’, *Dementia*, 18(3), pp. 1075–1088. doi: 10.1177/1471301217701506.

Srivastava, A. and Thomson, S. B. (2009) ‘Framework Analysis: A Qualitative Methodology for’, *Applied Policy Research. JOAAG*, 4(2), pp. 72–79. doi: 10.7748/nr2011.01.18.2.52.c8284.

Ursin, G. and Lotherington, A. T. (2018) ‘Citizenship as Distributed Achievement: Shaping New Conditions for an Everyday Life with Dementia’, *Scandinavian Journal of Disability Research*, 20(1), pp. 62–71. doi: 10.16993/sjdr.35.

1. In line with HRA Social Care REC recommendations data from care home one was excluded from analysis and write up due to being a pilot site [↑](#footnote-ref-1)
2. Pseudonyms are used to disguise the identity of the care home and participants

   [↑](#footnote-ref-2)