Clustered domestic model of residential care is associated with better consumer rated quality of care

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Abstract

Objective: To compare consumer rated quality of care among individuals living long-term in homelike clustered domestic and standard models of residential care in Australia.

Design: Cross-sectional study.

Setting: Seventeen residential aged care facilities in four Australian states providing alternative models of care.

Study participants: A sample of individuals with high prevalence of cognitive impairment living in residential care for 12 months or longer, not immediately in palliative care and having a proxy available to provide consent and assist with data collection. Of 901 eligible participants, 541 consented and participated in the study.

Main outcome measure: Consumer rated quality of care was measured using the Consumer Choice Index–6 Dimension instrument (CCI-6D) providing a preference weighted summary score ranging from 0 to 1. The six dimensions of care time, shared-spaces, own-room, outside and gardens, meaningful activities and care flexibility were individually evaluated.

Results: Overall consumer rated quality of care (Mean Δ: 0.138, 95% CI 0.073–0.203 P < 0.001) was higher in clustered domestic models after adjusting for potential confounders. Individually, the dimensions of access to outside and gardens (P < 0.001) and flexibility of care (P < 0.001) were rated significantly better compared to those living in standard model of care.

Conclusions: Homelike, clustered domestic models of care are associated with better consumer rated quality of care, specifically the domains of access to outdoors and care flexibility, in a sample of individuals with cognitive impairment. Including consumer views on quality of care is feasible and should be standard in future evaluations of residential care.

Key words: dementia, nursing homes, quality of healthcare, Australia, self-report
‘Long term care is intimate care, and how it is given, when it is given, and by whom it is given shapes the biography of the long term care consumer...’[1]
—Rosaline Kane

Including consumer perspectives in the evaluation of quality of care is common in health but less common in residential aged care settings. Previous studies evaluating the quality of care of residents of aged care facilities have used measures such as pressure ulcers, restraints and falls, [2–5] that do not take into consideration the consumer perspective. Such an approach limits the definition of quality of care to medical and clinical components while ignoring the psychological and social aspects of care that are integral to overall health and wellbeing. Those indicators are also focused towards safety and quality compliance rather than promoting resident wellbeing and quality of life [6].

Evaluating the quality of care in residential care from a consumer perspective is critical, with the introduction of consumer directed care (and similar care philosophies) in long-term care worldwide [4]. Similarly, with care philosophies consistently moving from a medical model to a person-centred model, quality of care needs to be evaluated from the perspective of the care recipients.

Providing residential care for people with dementia has evolved in the recent few decades to include an increasing emphasis on domestic, small-scale environments and person-centred care [7]. In addition, the concept of personhood in dementia care has received increasing attention. Several innovative models of long-term residential care have been developed and use different terminologies and methods of practice. These models of care share several common themes or components including the design of the physical building, providing housing and services in smaller living units, continuity of care staff in the unit, additional training for staff, involvement of residents in administration of the organization, flexibility and personalization in care routines.

Efforts have been made to evaluate these models of care and improvements in outcomes for residents including quality of life, emotional and physical wellbeing, participation and interaction, food intake and behavioural and psychological symptoms of dementia have been demonstrated [3, 8]. A study of the Greenhouse model of care in the US found reduction in hospitalizations for residents, as well as reduction in measures of clinical care quality such as restraint use, pressure ulcers and catheter use [2].

Other studies evaluating this model of care have found mixed results with some finding benefits for physical function, fewer hospitalizations and clinical indicators of quality of care in comparison to standard care and other studies finding no evidence for benefits in measures such as ADL, pain, ulcers, mobility and other Minimum Data Set quality indicators [5, 8, 9]. However there is limited evidence examining the quality of this model from a consumer perspective [3, 10].

The Consumer Choice Index–6 Dimension (CCI-6D) is a recently developed instrument to measure quality of care in long-term facilities from a consumer perspective [11]. The instrument comprises a 6-item questionnaire that can be completed by participants and/or their family members, and better quality of care as measured by the instrument has been shown to be associated with better quality of life among residents. Its six dimensions include care time, shared-spaces, own-room, outside and gardens, meaningful activities and care flexibility. The CCI-6D has strong content and construct validity, and includes a scoring system weighted according to the preferences of people living in residential aged care, and their family member carers [11, 12]. The homelike models of care have never been evaluated with a standardized instrument of quality of care such as the CCI-6D.

The aim of this analysis is to evaluate quality of care from the consumer perspective in a population of older adults living long-term in residential care in Australia; specifically, to compare standard residential care facilities with a clustered, domestic model of care.

Methods

Study sample

The sample for this analysis emanates from the Investigating Services Provided in the Residential Care Environment for Dementia (INSPIRED) study which has been described elsewhere [3, 13]. Briefly, this cross-sectional study included individuals who had lived for at least 12 months in one of 17 care facilities across four states in Australia. Facilities providing different models of care were purposefully sampled to include those located in urban and rural locations and include residents living with cognitive impairment and, or dementia. Within facilities all individuals were assessed for eligibility, those in palliative care and for whom it was not possible to obtain consent and collect data either directly or through a proxy were excluded. A total of 1323 individuals were assessed for eligibility, 901 were eligible and 541 consented (majority (84%) with dementia or cognitive impairment) to participate in the study. Data were collected at a single time point in the period between January 2015 and February 2016. Individual level data on demographic and social characteristics, health and function, care provision, quality of life and quality of care were collected through questionnaires from participants or their proxies and from facility records. Facility level data were collected through a facility level questionnaire (adapted from a similar study [14]) from the administrators of each facility. Ethics approval for the study was obtained from the Flinders University Social and Behavioural Research Ethics Committee.

Variables

The main outcome for this analysis, i.e. consumer rated quality of care, was measured using the Consumer Choice Index–6 Dimension (CCI-6D) as introduced earlier. In this study responses to CCI-6D were collected from the participants or their proxies (where participants were assessed to have moderate to severe cognitive impairment (a PAS-Cog cognitive assessment score of ≥11)). A proxy needed to be in a close relationship with the resident, visiting them regularly and assisting them with decision making (i.e. usually this role was fulfilled by a close family member or friend). Several studies have indicated the difficulty using proxies to rate abstract concepts such as quality in people living with dementia, with poor agreement between the ratings given by proxies and the people with dementia [15]. However, in this population including proxy responses is the only opportunity to include individuals with severe cognitive impairment in research that involves them [16]. Family members are also preferred over clinicians as proxies [16]. A preference weighted scoring algorithm derived from resident ratings was applied to the CCI-6D responses to derive a summary quality of care score on a scale of 0–1 with higher scores indicating better quality of care [12]. Complete CCI-6D responses were not available for one participant and excluded from further analyses.

The main explanatory variable was the model of care [8, 17] provided at the facilities where these participants resided. The model of care was defined based on the following six criteria: small size living units, consistent allocation of staff to living units, accessibility to...
outdoor areas, meals cooked within the living unit, opportunity for self-service of meals or opportunity for residents to participate in meal preparations. These criteria were based on the literature [18–21] and consultations with industry (clinicians, researchers and care providers) and consumer representatives. Consultation involved initial discussions regarding concepts and evidence around the key features of these models of care, collation of the information considering the context of the evidence, drafting an initial list of criteria which was reviewed and then finalized by all parties. A binary variable to define the model of care was used since all facilities met either two or less and five or more of the six criteria. A facility was defined as providing a clustered domestic model of residential care if it met five of the six criteria. All other facilities were defined as those providing standard residential care as they met only one or two of the criteria.

Descriptive analysis compared means and proportions of participant characteristics between the two models of care using standard t-tests or chi-square tests as appropriate. Facility level characteristics were not compared statistically due to prohibitively low sample of 17 facilities. Analyses used multi-level random effect linear regression models adjusting for individual (age, gender, cognition (PAS-Cog) [22], ADL (Modified Barthel Index) [23], social interactions (frequency of visits from family and/or friends) and comorbidities (Cohen–Mansfield comorbidity index) [24]) and facility level (geographic location, size, staff training and direct care hours) characteristics. Geographical location was categorized as urban or rural based on standard Australian classifications, size of facilities was based on the total number of beds in each facility, staff training was defined based on the facility level expenditure on staff training in the preceding 2 years and direct care hours was defined as all care provided by nurses, care workers and allied health staff. Age of facility was the years since either initial construction, extension or major refurbishment. It was not possible to ascertain if the exact location the study participants’ lived in the facility was original build, new extension or that which was significantly refurbished. Hence age of facility was not included in the final adjusted model but only included in a sensitivity analysis to test its effect on the adjusted difference.

Unadjusted proportions of participants’ (or proxies’) responses and adjusted odds ratios of reporting best ratings in the 6 individual dimension of the CCI-6D among those living in the clustered domestic facilities were estimated using multi-level generalized linear regression models, with binary link functions. Confidence intervals were reported for all summary measures and statistical significance was determined at P < 0.05. All statistical analysis was performed using SAS software version 9.4. Copyright © 2018 SAS Institute Inc.

### Results

Four facilities (120 participants) of the 17 provided the clustered domestic model of care (Table 1). Overall the mean age of participants was 86 years, and 75% of the sample were female. Participants on average had 3.7 comorbid disease groups. Eighty-four percent had a medical diagnosis of dementia or were cognitively impaired (PAS-Cog score ≥5). Study participants living in facilities providing a clustered domestic model of care were significantly younger, had more neuropsychiatric symptoms, had fewer comorbid conditions, were more cognitively impaired, had fewer weekly social interactions, a higher proportion had dementia and the clear majority of the quality of care (CCI-6D) responses were by proxies. The clustered domestic model facilities had higher investment in staff training, higher hours of direct care staff allocated and were younger aged facilities.

Living in a facility providing clustered domestic model of care was significantly associated with better overall quality of care as

### Table 1 Baseline participant and facility characteristics

| Participant characteristics | Clusted domestic model of care (n = 120) | Standard model of care (n = 421) |
|-----------------------------|----------------------------------------|---------------------------------|
| Age, mean (SD)              | 83.3 (9.0)                             | 86.1 (8.5)*                     |
| Female, n (%)               | 90 (75.0)                              | 313 (74.4)                      |
| Married, n (%)              | 36 (30.0)                              | 101 (24.1)*                     |
| Modified Barthel Index, mean (SD) | 37.1 (31.1)                         | 41.3 (33.3)                     |
| Number of comorbid conditions, (Cohen–Mansfield Index), mean (SD) | 3.2 (1.4) | 3.8 (1.4)* |
| PAS-Cog, mean (SD)          | 17.4 (16.5)                            | 12.2 (11.4)*                    |
| Diagnosis of dementia in medical records, n (%) | 117 (97.5) | 231 (55.1%)* |
| PAS-Cog 0–4<10 (mild cognitive impairment), n (%) | 3 (2.5) | 90 (21.4) |
| PAS-Cog 4–<10 (mild cognitive impairment), n (%) | 12 (10.0) | 88 (20.9) |
| PAS-Cog 10–<16 (moderate cognitive impairment), n (%) | 18 (15.0) | 64 (15.2) |
| PAS-Cog 16–21 (severe cognitive impairment), n (%) | 87 (72.5) | 179 (42.5) |
| Dementia diagnosis or PAS-Cog ≥5, n (%) | 120 (100) | 333 (79)* |
| Weekly interaction with close social ties, n (%) | 66 (56.4) | 312 (75.2)* |
| CCI-6D proxy responses, n (%) | 114 (95.0) | 277 (65.8)* |

| Facility characteristics | Clusted domestic model of care (n = 4) | Standard model of care (n = 13) |
|--------------------------|----------------------------------------|---------------------------------|
| Metropolitan location, n (%) | 3 (75) | 10 (77) |
| Total facility size (No. beds), mean (SD) | 83 (6.6) | 83 (6.9) |
| High staff training costs (>$1000 per resident annually), n (%) | 4 (100) | 4 (33) |
| High direct care hours (>2.5 per resident per day), n (%) | 3 (75.0) | 6 (46) |
| Years since construction, extension or major refurbishment, mean (SD) | 10 (4.3) | 17 (25.1) |

SD, Standard Deviation; NPI-Q, Neuropsychiatric Inventory–Questionnaire; PAS-Cog, Psychogeriatric Assessment Scales–Cognitive Impairment Scale, CCI-6D, Consumer Choice Index–6 Dimensions, *P < 0.05.
measured by the CCI-6D (Mean Δ: 0.138, 95% CI 0.073–0.203, \( P < 0.001 \)) after adjusting for potential confounding factors (Table 2). Subgroup analyses of only those individuals who responded through a proxy (Supplementary Table 1) and those with severe cognitive impairment (Supplementary Table 2), despite reduced sample size showed similar significant differences. Inclusion of the age of facility in the adjusted model gave similar findings (Mean Δ: 0.128, 95% CI 0.033–0.223, \( P < 0.008 \) (not tabulated)).

Individually, a significantly higher proportion of those living in clustered domestic facilities reported better ratings for the dimensions of access to outside and gardens and flexibility of care. After adjusting for potential confounders, the probabilities of reporting better ratings for these two dimensions were significantly better among those living in the clustered domestic facilities compared to those living in standard model of care (Table 3 and Fig. 1).

**Discussion**

Living in a facility providing a clustered domestic model of care was associated with better consumer rated quality of care even though individuals living in this model had poorer health and function as indicated by several measures. This study used the Consumer Choice Index–6 Dimension (CCI-6D) instrument which not only provides a consumer perspective but was also developed with recipients of care [11]. Specifically, the individual dimensions of access to

### Table 2 Quality of care (CCI-6D total weighted score) by models of residential care

| Quality of care (CCI-6D) | Adjusted\(^a\) model predicted CCI-6D mean values (95% CI) | Mean difference (95% CI) | P-value (Adjusted\(^a\)) |
|-------------------------|------------------------------------------------------------|--------------------------|--------------------------|
| Clusters                | Standard (n = 420)                                         | (n = 120)                | Unadjusted | Adjusted\(^a\) |
| Consumer rated quality of care | 0.856 (0.792, 0.919) | 0.718 (0.674, 0.761) | 0.082* (0.009, 0.016) | 0.138 (0.073, 0.203) | <0.001 |

PAS-Cog, Psychogeriatric Assessment Scales–Cognitive Impairment Scale; CCI-6D, Consumer Choice Index–6 Dimensions.

\(^a\)Adjusted for age, gender, PAS-Cog, Barthel Index, Social ties, number of comorbidities, regional location, facility size, staff training and direct care hours. \(^*\)P < 0.05.

### Table 3 Unadjusted proportions and adjusted odds of individual dimensions of CCI-6D by models of care

| Dimension                                                                 | Frequency N(\%*) | Adjusted Odds ratio (95% CI) | P-value |
|---------------------------------------------------------------------------|------------------|-------------------------------|---------|
| Care Time: How much time are Caregiving staff able to spend with me (your family member)? |                  |                               |         |
| Always able to spend enough time attending to my individual needs          | 180 (43%)        | 57 (48%)                      | 1.48 (0.75, 2.96) | 0.261 |
| Sometimes able to spend enough time attending to my individual needs       | 197 (47%)        | 58 (48%)                      | 1.05 (0.52, 2.13) | 0.881 |
| Rarely able to spend enough time attending to my individual needs          | 43 (10%)         | 5 (4%)                        | 1.02 (0.49, 2.15) | 0.950 |
| Shared Spaces: Do the shared spaces of the Aged care home as a whole make you (or your family member) feel ‘at home’? |                  |                               |         |
| I feel very at home here                                                  | 251 (60%)        | 73 (61%)                      | 7.46 (3.35, 16.63) | <0.001 |
| I feel at home here sometimes                                             | 110 (26%)        | 30 (25%)                      | 1.47 (0.71, 3.04) | 0.297 |
| I feel at home here rarely                                                | 59 (14%)         | 17 (14%)                      | 7.81 (3.47, 17.60) | <0.001 |
| Own Room: Does your own room here make you (or your family member) feel ‘at home’? |                  |                               |         |
| I feel very at home in my room                                           | 318 (76%)        | 79 (66%)                      | 1.47 (0.71, 3.04) | 0.297 |
| I feel at home in my room sometimes                                       | 87 (21%)         | 33 (28%)                      | 7.46 (3.35, 16.63) | <0.001 |
| I feel at home in my room rarely                                          | 15 (4%)          | 8 (7%)                        | 1.47 (0.71, 3.04) | 0.297 |
| Outside and Gardens: Is there access to outside and gardens in this Aged care home? |                  |                               |         |
| I can get outside whenever I want                                        | 175 (42%)        | 93 (78%)                      | 1.47 (0.71, 3.04) | 0.297 |
| I can get outside sometimes                                               | 94 (23%)         | 12 (10%)                      | 7.81 (3.47, 17.60) | <0.001 |
| I cannot get outside easily                                               | 151 (36%)        | 15 (13%)                      | 7.81 (3.47, 17.60) | <0.001 |
| Meaningful Activities: How often does the facility offer me (or your family member) things to do that make me feel valued? |                  |                               |         |
| I can do things that make me feel valued often                            | 143 (34%)        | 45 (38%)                      | 1.47 (0.71, 3.04) | 0.297 |
| I can do things that make me feel valued sometimes                        | 149 (36%)        | 39 (33%)                      | 7.81 (3.47, 17.60) | <0.001 |
| I can only rarely or occasionally do things that make me feel valued       | 128 (31%)        | 36 (30%)                      | 7.81 (3.47, 17.60) | <0.001 |
| Care Flexibility: How flexible is the Aged care home with the care routines? |                  |                               |         |
| Care routines are very flexible                                           | 220 (52%)        | 102 (85%)                     | 1.47 (0.71, 3.04) | 0.297 |
| There is a little flexibility in the care routines                        | 141 (34%)        | 14 (12%)                      | 7.81 (3.47, 17.60) | <0.001 |
| There is not much flexibility in the care routines                        | 59 (14%)         | 4 (3%)                        | 7.81 (3.47, 17.60) | <0.001 |

*Some percentages summing up to more than 100 due to rounding up.
outdoors and flexibility in care were rated better among residents in the clustered domestic model of care.

Being a recently developed instrument, a minimally important difference (MID) for this quality of care measure has not been established. For the current analysis, the best opportunity to understand the importance of this association is by comparing it to a within sample distribution based MID [25] of 0.107 (0.5 SD of difference). Another option is to compare the observed difference to the MID of other preference-based measures in this population. The EQ-5D, a measure of health-related quality of life which is among the most widely used preference-based measures and is recommended for use in aged care populations has a published range of minimal importances were found not to be significantly different between the residents of the two models of care. All participating facilities in the study were not-for-profit organizations which are generally recognized to provide high quality of care [31]. It is also possible that of facilities that were approached for inclusion in this study, only those that provided a higher level of care were willing to participate. This may explain why there were minimal differences between the two models of care with respect to these domains.

Conversely, it is plausible that the distinguishing characteristics of the clustered domestic model of care have a larger impact on the two domains of access to outside and gardens and care flexibility. These two domains (along with ‘own room’) were also weighted higher for the most flexible choice in the resident-rating based algorithm used to calculate the overall quality of care score [12].

However, given the possibility that all the facilities in this study generally provide high quality of care, the observed differences between the two models of care are statistically (and potentially clinically) significant. This points to the strength of the evidence that the clustered domestic model of care is associated with better overall consumer rated quality of care.

The homelike models of care have also been shown to be associated with better quality of life outcomes in other studies, including within this study sample [3, 8, 21]. Hence, it is possible that providing a homelike model of care is associated with both an independent and complementary benefit in terms of quality of life and quality of

The Australian government’s productivity commission report [29] also recommends the need for increased choice and flexibility in residential care. This was echoed in the rapid review commissioned by the Australian Aged Care Quality Agency, which identified independence and autonomy as one of the main considerations of high quality residential care [30]. This study’s finding that the clustered homelike model of care is a better environment for flexibility and care from the consumer’s perspective, provides an opportunity to fulfil those identified recommendations.

In this analysis, the remaining four dimensions of the CCI-6D, namely, care time, shared spaces, own room and meaningful activities were found not to be significantly different between the residents of the two models of care. All participating facilities in the study were not-for-profit organizations which are generally recognized to provide high quality of care [31]. It is also possible that of facilities that were approached for inclusion in this study, only those that provided a higher level of care were willing to participate. This may explain why there were minimal differences between the two models of care with respect to these domains.

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Care. The findings of this study further strengthen the argument that investing in homelike models of care should be a priority for all stakeholders involved in the care of people with dementia. As the proportion of populations needing and living long-term in residential care increases worldwide, planning for increased supply of care services needs to prioritize incorporating these and other factors that characterize homelike models of care into development, management and delivery of care services. While the costs and consequences of this model of care have been evaluated [8], a full economic evaluation of these models of care is a future research priority.

This analysis has some strengths and limitations. This is the first study that examines quality of care from a consumer perspective in these homelike models of care. The sample of this study is largely representative of Australian residential aged care populations [13]. All facilities offering the homelike model of care in this study were operated by a single provider, hence it is not possible to delineate the effect the provider and its policies and resources have on the quality of care. However, one of the other facilities administered by this provider provided standard residential care. On the contrary one of the limitations of the larger evaluations of the Greenhouse model is the heterogeneity in the implementation of the model of care across organizations and facilities [21]. Our study reduces the effects of the heterogeneity in the implementation of the model of care.

The CCI-6D as described earlier has strong content and construct validity, and includes a scoring system weighted according to the preferences of people living in residential aged care, and their family member carers [11, 12]. However, it is still a relatively new instrument and is yet to undergo wider testing. In this analysis, the responses were predominantly reported by proxies in the clustered domestic model. Proxies are generally known to provide lower ratings than residents for subjective measures such as quality of life [15]. Hence, the higher proportion of proxy ratings in the clustered domestic model is unlikely to affect the significantly better rating, instead it has the potential to under estimate the ratings in that group. Irrespective of the resident or proxy reported quality of care this study shows that it is feasible to evaluate quality of care from a consumer perspective.

Another factor to consider in the interpretation of these findings is the potential bias due to self-selection of individuals into facilities. The Australian residential aged care system in theory provides some choice to entrants into the system. However, in practice for many people the decision to move into residential care is necessitated by a period of acute illness and/or loss of physical function that is sudden. Due to the shortage of available residential care places at a given point in time, the ability to self-select is limited [32]. Hence, the potential bias due to self-selection into one model of care or other is unknown but potentially limited.

This study is cross-sectional in design and hence no temporal or causative inferences can be drawn based on these associations. Confirmation of these findings in longitudinal studies is needed. Similarly, there were noted differences between facilities providing the two models of care and the individual characteristics of people living in the two groups. We have statistically controlled for observed differences, but it is possible that the quality of care may be affected by other factors which were not adjusted for or not measured as part of this study. One of those factors is the age of facility. The clustered domestic model facilities were newer than the standard care ones which has the potential to affect the perception of quality. However, the age of facility when adjusted for in the analysis remained statistically significant and only marginally reduced the difference in care quality between the two models of care.

Conclusion
This analysis suggests that clustered domestic models of care provide a higher quality of care from the consumer perspective, in a study which included a high proportion of residents with dementia. Specifically, the homelike, clustered domestic model of care is associated with better ratings for two of the six domains of consumer rated quality of care, namely access to outdoors and care flexibility. Changes to the way aged care is provided to better align with a homelike model of care has the potential to better meet consumer preferences. This analysis of the quality of this model of care provides preliminary evidence to warrant further study and investigation of its implementation. Including consumer views on the quality of care is feasible and should be considered essential in future evaluations of residential care.

Supplementary material
Supplementary material is available at International Journal for Quality in Health Care online.

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