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Triggered Palliative Care for Late-stage Dementia: a Pilot Randomized Trial

Laura C. Hanson, MD, MPH,

Cecil G. Sheps Center for Health Services Research, Division of Geriatric Medicine and Palliative Care Program, School of Medicine, University of North Carolina, Chapel Hill, NC USA

Christine E. Kistler, MD, MASc, MPH,

Cecil G. Sheps Center for Health Services Research, Department of Family Medicine and Palliative Care Program, School of Medicine, University of North Carolina, Chapel Hill, NC USA

Kyle Lavin, MD, MPH,

Department of Psychiatry and Palliative Care Program, School of Medicine, University of North Carolina, Chapel Hill, NC USA

Stacey L. Gabriel, MPH,

Cecil G. Sheps Center for Health Services Research, University of North Carolina, Chapel Hill, NC USA

Natalie C. Ernecoff, MPH,

Cecil G. Sheps Center for Health Services Research, Department of Health Policy and Management, Gillings School of Global Public Health, University of North Carolina, Chapel Hill, NC USA

Feng-Chang Lin, PhD,

Department of Biostatistics, School of Public Health, University of North Carolina at Chapel Hill, Chapel Hill, NC USA

Greg A. Sachs, MD, and

Division of General Internal Medicine & Geriatrics, Indiana University School of Medicine, Indiana University Center for Aging Research, Regenstrief Institute, Inc., Indianapolis, IN USA

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Corresponding author: Laura C. Hanson, MD, MPH, CB 7550, Old Clinic Building, Division of Geriatrics, University of North Carolina, Chapel Hill, NC 27599-7550, 919-843-4096, lhanson@med.unc.edu. Author Contributions:

Dr. Hanson and Dr. Lin had full access to all the data in the study and take responsibility for the integrity of the data and the accuracy of the data analysis.

Study concept and design: Hanson, Sachs, Mitchell

Acquisition of data: Hanson, Gabriel, Ernecoff

Analyses and interpretation of data: Hanson, Kistler, Lavin, Gabriel, Ernecoff, Lin, Sachs, Mitchell

Drafting of manuscript: Hanson

Critical revision of manuscript for important intellectual content: Hanson, Kistler, Lavin, Gabriel, Ernecoff, Lin, Sachs, Mitchell Statistical analyses: Hanson, Gabriel, Lin

Administrative, technical or material support: Hanson

Study supervision: Hanson

Susan L. Mitchell, MD, MPH

Hebrew SeniorLife Institute for Aging Research and Department of Medicine, Beth Israel Deaconess Medical Center, Boston, MA USA

Abstract

Context: Persons with late-stage dementia have limited access to palliative care.

Objective: To test dementia-specific specialty palliative care triggered by hospitalization.

Methods: This pilot randomized controlled trial enrolled 62 dyads of persons with late-stage dementia and family decision-makers on admission to hospital. Intervention dyads received dementia-specific specialty palliative care consultation plus post-acute transitional care. Control dyads received usual care and educational information. The primary outcome was 60-day hospital or emergency department visits. Secondary patient and family-centered outcomes were patient comfort, family distress, palliative care domains addressed in the treatment plan, and access to hospice or community-based palliative care. Secondary decision-making outcomes were discussion of prognosis, goals of care, completion of Medical Orders for Scope of Treatment (MOST), and treatment decisions.

Results: Of 137 eligible dyads, 62 (45%) enrolled. The intervention proved feasible, with protocol completion ranging from 77% (family 2-week call) to 93% (initial consultation). Hospital and emergency department visits did not differ (intervention vs control, 0.68 vs 0.53 transfers per 60 days, p=0.415). Intervention patients had more palliative care domains addressed, and were more likely to receive hospice (25% vs 3%, p<0.019). Intervention families were more likely to discuss prognosis (90% vs 3%, p<0.001) and goals of care (90% vs 25%, p<0.001), and to have a MOST at 60-day follow-up (79% vs 30%, p<0.001). More intervention families made decisions to avoid re-hospitalization (13% vs 0%, p=0.033).

Conclusion: Specialty palliative care consultation for hospitalized patients with for late-stage dementia is feasible and promising to improve decision-making and some treatment outcomes.

Clinicaltrials.gov: NCT02719938 (3/21/16)

Keywords

dementia; randomized trial

Introduction

Alzheimer's disease and related dementias are leading causes of functional dependency and death. Over 5 million Americans live with dementia; more than 1 million have late-stage dementia with profound cognitive deficits and dependence for activities of daily living. Family caregivers provide extraordinary practical and emotional support, and face average out-of-pocket expenses of \$66,000 in the last year of life. A Societal costs for dementia care exceeded \$250 billion in 2018, and the affected population is projected to double by 2030.

Dementia is the only leading cause of death in the United States without meaningful treatment to prevent, cure or slow disease progression. Palliation should be an important

focus of dementia care, yet the evidence shows unmet need. Among nursing home residents with dementia, half experience frequent and unrelieved pain. ^{7,8,9} Distressing symptoms -- shortness of breath, neuropsychiatric symptoms, feeding problems, and problems with personal cleanliness -- are common. ^{10,11,12,13,14} Decisions about life-sustaining treatment are delayed, and access to hospice limited by lack of a discrete terminal phase of illness. ^{15,16,17,18}

Hospitalization is common in late-stage dementia, creating an opportunity to improve access to palliative care. ¹⁹ While only 27% of US nursing homes report any type of specialized palliative or endof-life care, 85% of large hospitals have specialty palliative care. ^{20,21} Infections, hip fracture and nutritional decline are common reasons for hospital transfer, and signal worsened prognosis with six-month mortality risk of 20–50%. ^{15,22,23,24,25}

Specialty palliative care improves treatment decision-making and other patient-centered outcomes for persons with cancer and other serious illnesses, but has rarely been studied for persons with dementia. 26,27,28,29,30 Palliative care clinicians rarely have training for dementia-specific symptom management, decision-making and caregiver support, yet their fundamental skills and interdisciplinary team approach is well matched to needs in late-stage dementia. We therefore developed a model of specialty palliative care consultation specific to the needs of persons with late-stage dementia and their families. Specific aims were 1) to develop a best-practice model of specialty palliative care for late-stage dementia, and 2) to conduct a pilot randomized trial of specialty palliative care for late-stage dementia triggered by hospitalization for acute illness. We hypothesized this model would be feasible, and show potential to improve outcomes of subsequent hospital or emergency department visits, hospice and community-based palliative care use, enhanced palliative care content of care plans, increased completion of MOST (Medical Orders for Scope of Treatment) forms, and decisions to avoid future hospitalization and potentially burdensome treatments.

Methods

Trial design:

Investigators reviewed evidence on palliative care needs and interventions in late-stage dementia. They held 3 meetings with key stakeholders – 3 family caregivers for persons with late-stage dementia and 4 clinicians with extensive experience in dementia care. Using these sources, investigators designed a dementia-specific protocol for specialty palliative care consultation with post-acute telephone support.

Research staff enrolled dyads of hospitalized persons with late-stage dementia with their primary family decision-makers, and randomized them to receive specialty palliative care consultation during hospitalization with post-discharge telephone support by a palliative care nurse practitioner (intervention) vs usual hospital care with educational material on dementia caregiving (control). The University of North Carolina institutional review board approved the protocol, and a Data Safety Monitor reviewed study procedures and preliminary data every six months; the study is a registered clinical trial (NCT 02719938).

Participant enrollment and randomization:

From March 31, 2016 to August 31, 2017 research staff enrolled dyads of hospitalized patients with late-stage dementia and family decision-makers. Investigators developed and refined a dementia phenotype within the hospital electronic health record (EHR) based on patient age, hospital admission date and ever use of ICD-9 or ICD-10 codes for dementia diagnoses. ³¹ Potentially eligible patients were thus identified within 24 hours of admission, reviewed by research staff for eligibility. A palliative care physician then confirmed dementia diagnosis and stage with the patient's attending physician, and sought permission to approach the family decision-maker about participation.

Patients were eligible if they were aged 65 or older, hospitalized with an acute illness, had a diagnosis of dementia stage 5–7 on the Global Deterioration Scale (GDS) verified by their attending physician, and had an eligible family decision-maker. Persons with GDS 5 dementia and later cannot live independently and are frequently disoriented, while those who progress to GDS 7 have sparse speech, dependency for all activities, and cannot recognize family. Family decision-makers were eligible if they were legally authorized representatives for healthcare decisions and could complete interviews in English. They provided written consent for themselves and the person with late-stage dementia. After informed consent, the study biostatistician randomized each dyad in a 1:1 ratio to intervention vs. control arms prior to the baseline interview. Allocation was revealed to the baseline interviewer, family decision-maker and to the attending physician, but concealed from the investigators and research staff collecting interview outcome data. Family decision-makers were compensated for time to complete interviews.

Intervention and control conditions:

Patient-family dyads randomized to the intervention received protocolized specialty palliative care consultation while hospitalized, plus 2-week post-discharge transitional telephone support by a palliative care nurse practitioner. Clinicians delivering the intervention were board-certified physicians and nurse practitioners in palliative care. The consult protocol addressed a) stage, prognosis and trajectory of dementia, b) assessment and treatment of pain and other physical symptoms, c) assessment and management of neuropsychiatric symptoms, d) social support for caregiver stress, e) spiritual needs assessment, f) cultural concerns framing care, g) goals of care decision-making, and h) key clinical decisions such as feeding options, antibiotic use, and re-hospitalization. Consultants provided families with a copy of the informational booklet "Advanced Dementia: A Guide for Families."33 Based on their assessments, consultants provided individualized recommendations for palliative care domains, offered to assist with completion of a Medical Orders for Scope of Treatment (MOST) order set, the North Carolina version of POLST (Physician Orders for Life Sustaining Treatment), and recommended referrals to postdischarge services. 34,35 Transitional care included a) provision of consult recommendations and MOST form to the post-acute primary provider, and b) follow-up supportive calls by a Palliative Care nurse practitioner to the family decision-maker at 72 hours and 2 weeks postdischarge. Investigators provided a 1-hour training session to Palliative Care physicians and nurse practitioners to teach them the dementia protocol, and to provide access to an EHRtemplated consult note.

Family decision-makers randomized to control received information on caregiving for late-stage dementia from the Alzheimer's Association, and patients received usual hospital care. ³⁶ Specialty palliative care consultation was allowed, if requested by attending physicians. All other procedures were identical for intervention and control participants.

Data Collection:

Research staff masked to study arm collected data using 30- and 60-day post-discharge telephone interviews with family decision-makers. When an enrolled patient died, staff conducted family interviews with modifications for care during dying. A separate research staff member conducted chart reviews at 60 days post-discharge, but could not be masked due to the differential documentation between study arms.

Feasibility:

Investigators evaluated feasibility based on dyad enrollment and follow-up, and on fidelity to components of the intervention. Research staff monitored six components (initial consult, call to post-acute provider, palliative care consult records to post-acute provider, MOST discussion, 72-hour call to family and 2-week call to family), with a goal of 80% completion.

Outcomes:

The pre-specified primary outcome was hospital or emergency department visits in the 60 days post-discharge, defined as hospital re-admissions or emergency room visits reported in family interviews. Secondary patient and family-centered outcomes were patient comfort, family distress, palliative care domains addressed in the treatment plan, and access to hospice or community-based palliative care services. Secondary decision-making outcomes were discussion of prognosis, discussion of goals of care, completion of MOST (Medical Orders for Scope of Treatment) order set, and documented decisions against re-hospitalization or other potentially burdensome treatments.

Patient comfort was measured in follow-up interviews using the reliable and valid Comfort at the End of Life in Dementia (CAD-EOLD). Fourteen items are rated 1-3 and summed, for a range of 14–42 with higher scores indicating better symptom control over the past week. It has good internal consistency (Cronbach's alpha 0.83-0.90) and convergent validity (r=0.81 with the Decision Satisfaction Inventory). ^{37,38} Family distress was measured using the Family Distress in Advanced Dementia (FDAD) scale, with 21 items rated and averaged on a 1–5 scale to create a potential range of 1–5, with higher scores indicating greater distress. The FDAD has good internal consistency in 3 domain scores for emotional distress (alpha=0.82), dementia preparedness (0.75) and clinician relations (0.83).³⁹ Palliative care domains (range 0-10) were measured in chart review as count of the presence or absence of 10 domains of palliative care addressed in the medical treatment plan -- prognosis, overall goals of care, physical symptoms, psychiatric symptoms, spiritual needs, and 5 treatment presuscitation, artificial feeding, intravenous fluids, antibiotics, and hospitalization. This measure has been used in a large clinical trial, and has good inter-rater reliability (kappa =0.90)⁴⁰ Access to hospice or community-based palliative care, MOST completion and decision-making about specific treatments was measured using family report at 60 days,

with verification in medical record review. Finally, research assistants reviewed charts for evidence of adherence to process measures for dementia quality of care using the Physician Quality Reporting System (PQRS) 2015 measures for dementia staging (#280), cognitive assessment (#281), functional assessment (#282), neuropsychiatric symptom assessment (#283), safety counseling (#286), and caregiver education and support (#288).

Additional Variables:

Research staff recorded data on patient and family decision-maker demographics, prior hospital transfers, and advance directives from baseline family interviews. Patient functional status was measured in baseline family interviews using the valid and reliable Bedford Alzheimer Nursing Severity scale (BANS-S), framed to represent function in the weeks just prior to acute illness. ⁴² The BAN-S ranges 7–28, with higher scores indicating greater functional impairment. During baseline and follow-up interviews, research staff asked family respondents about their perception of prognosis for the person with dementia, and perception of their involvement in decision-making. During chart review, staff recorded data on major co-morbid diagnoses and the acute illness causing hospital admission, post-acute site of care and patient's vital status at 60 days follow-up.

Analysis:

Patient-family dyads were the primary unit of analysis. Baseline characteristics of the dyads were reported in either means and standard deviations or medians and ranges if the variables are continuous, and in percentages if the variables are categorical. Intervention and control dyads were then compared on the baseline characteristics to explore possible differences between arms after randomization. All of the comparisons were tested using either chisquare tests or two-sample t-tests except hospital length of stay, which was compared using Mann-Whitney test because of non-normality. The primary outcome of hospital and emergency department visits was reported as the number of events per 60 days of follow-up, with censoring of eligible follow-up days if the patient died or the dyad withdrew from the study. In a priori power calculation, the sample size of 60 patients with 30 patients per group was estimated to give 81% power to detect a relative risk of 2.7. Poisson regression, which accounts for the length of follow-up, was used to compare the rate of hospital and emergency department visits between the two arms. Comparisons of the secondary outcomes were implemented using either chi-square tests or two-sample t-tests, when appropriate. Comparisons were not further adjusted since there was no apparent imbalance between arms. All analyses used intention-to-treat assignment, with a p value <0.05 threshold for statistical significance. Software SAS 9.4 (Cary, NC) was used to implement statistical tests.

Results

Study Enrollment and Feasibility:

Automated screening generated 3296 admissions with a dementia diagnosis code during the 18-month enrollment period, of whom 426 had late-stage dementia. (Figure 1) One-third of patients were discharged prior to contact due to very short lengths of stay (n=161), and others were admitted during periods when study personnel were not available (n=50). Some were not eligible due to active dying or current hospice enrollment (n=32), attending

physician refusal (n=29), or lack of eligible decision-maker (n=17). Of 137 eligible dyads contacted, 62 (45%) enrolled and randomized (32 control, 30 intervention). One-third of family caregivers who refused participation stated this was due to feeling overwhelmed by caregiving and other roles. After enrollment, one family decision-maker withdrew and 4 were lost to follow-up. Fidelity to the intervention was high, with completion of each of six intervention protocol components ranging from 77% (family 2-week call) to 93% (palliative care consultation).

Characteristics of enrolled patients and families:

Hospitalized patients with late-stage dementia had an average age of 83.9 years, 56% were female, 71% white and 24% African American. (Table 1) Thirty-seven percent had GDS Stage 5, 42% Stage 6 and 21% Stage 7 dementia. The most common admitting diagnoses were infections and neuropsychiatric symptoms. Two-thirds of patients had advance directives. Baseline characteristics did not differ significantly between study arms.

Family decision-makers had an average age of 59.7 years, 79% were female, and more than half were daughters. At enrollment, 92% of caregivers felt they were very involved in treatment decisions for the person with dementia, and 60% expected that person to get worse or possibly die in the next 6 months.

Patient and family-centered outcomes:

One of three enrolled patients with late-stage dementia visited an emergency department or was hospitalized in the 60 days after discharge. The primary outcome of 60-day hospital or emergency department visits did not differ significantly between intervention and control arms (0.68 vs 0.53 visits, p=0.415). (Table 2) Family caregivers' ratings of comfort for the person with dementia and of distress for themselves did not differ between arms at 60 days. However, dementia patients in the intervention arm had more elements of clinical palliative care addressed in their hospital treatment plan, as measured on the 10-point Palliative Care Domain score (7.6 vs 2.7, p<0.001). Patients with the intervention were more likely to have assessment and treatment for physical symptoms of dyspnea, constipation and nausea, and for neuropsychiatric symptoms. Spiritual needs were addressed for 47% of patient-family dyads in the intervention arm, and for 0% of those in the control arm. In the 60 days after discharge, family reported that patients in the intervention arm were more likely to receive hospice (25% vs 3%, p<0.019), with a trend toward greater access to community-based palliative care. In the intervention arm, more patients with dementia discharged to home or assisted living settings, while patients in the control arm were more likely to enter nursing homes (p=0.046). As anticipated, few patients died during follow-up and mortality did not differ between arms.

Decision-making outcomes:

Specialty palliative care during hospitalization also resulted in increased communication and decision-making about treatments relevant to late-stage dementia. Family decision-makers in the intervention arm were more likely to participate in discussions of prognosis (90% vs 3%, p<0.001) and goals of care (90% vs 25%, p<0.001). These families were more likely to complete a MOST in hospital (70% vs 13%, p<0.001) and have an active MOST in use at 60

day follow-up (79% vs 30%, p<0.001). While Do-Not-Resuscitate orders were common for persons with dementia in both arms, more intervention families made decisions to avoid future burdensome treatments such as tube feeding, antibiotics and IV fluids. Decisions to avoid re-hospitalization were also increased (13% vs 0%, p=0.033).

Dementia quality of care:

Compared to controls, persons with dementia in the intervention arm were more likely to receive formal dementia staging (93% vs 9%, p<0.001) and structured assessment of cognition (73% vs 9%, p<0.001) and function (97% vs 25%, p<0.001). Neuropsychiatric symptoms were more commonly addressed, as were caregivers' safety concerns. Finally, this intervention resulted in more education of family caregivers about late-stage dementia (80% vs 25%, p<0.001).

Discussion

Specialty palliative care consultation specific to late-stage dementia, initiated during hospitalization for acute illness, is feasible and promising to improve decision-making and clinical outcomes for persons with dementia and their families. In this pilot randomized trial, investigators used an innovative, efficient method to identify hospitalized patients with late-stage dementia. Compared to usual hospital care, triggered palliative care consultation was effective to improve decision-making, dementia neuropsychiatric symptom management, and use of hospice. While the intervention did not change the primary outcome of 60-day hospital or emergency department visits, it did increase decisions to forego further hospitalization and thus may affect this outcome over longer follow-up. Family decision-makers were more likely to discuss prognosis and goals of care, and make decisions to avoid burdensome treatments for the person with dementia. Furthermore, they were much more likely to record these decisions in a MOST order set, and these orders remained in use 60 days after hospital discharge.

Persons with dementia and their families need intervention research to reduce or moderate the effects of frequent healthcare transitions – including hospitalizations and admissions to nursing facilities. We found that one in three persons with dementia returned to a hospital or emergency room in the 60 days after discharge. Stakeholders interviewed in preparation for this trial likened the experience of late-stage dementia healthcare to "being nomads" and "becoming homeless." They referenced frequent hospitalizations and transitions to nursing facility care, resulting in the loss of a trustworthy primary clinician to guide treatment. While facility placement was not a pre-specified outcome, an unexpected finding in this study was that fewer persons with dementia in the intervention arm entered nursing home care. It is possible that skilled discussions of prognosis and goals of care supported families to opt for home or home-like assisted living facilities with dementia care units.

This pilot trial is an important addition to very limited research on specialty palliative care for dementia, and its promising intermediate outcomes warrant testing in a larger trial. A small randomized trial of specialty palliative care consultations for hospitalized patients with dementia found increased use of palliative care plans (23% vs 4%, p=0.008) and reduction in intravenous therapies (66% vs 81%, p=0.025), but no effect on life-sustaining treatment

decisions.⁴³ In a small pre-post study (n=52), proactive case-finding of intensive care unit patients with advanced dementia for palliative care was associated with fewer hospital days (7.4 vs 12.1, p<0.007) and ICU days (3.5 vs 6.8, p<0.004), with no difference in overall treatment intensity.⁴⁴ Mitchell conducted a single site pre-post pilot study of an Advanced Dementia Consult Service, which provided post-acute recommendations to the primary physician, and a follow-up call to the family caregiver. Investigators enrolled 24 controls followed by 5 intervention participants, and found promising trends in increased hospice use (40% vs 25%), reduced 30-day re-admissions (0% vs 29%), and improved family understanding of advanced dementia (50% vs 0%).⁴⁵

Investigators have also tested interventions in which non-specialty clinicians deliver elements of palliative care, termed "primary palliative care." Our research team has extensive experience in primary palliative care interventions for dementia, delivered by interdisciplinary nursing home staff and geriatric care teams in nursing homes, assisted living, and in the community. In the nursing home setting, we have shown that dementia-specific decision aids can improve communication and decision-making for family caregivers, enhance palliative care treatment plans, improve goal-concordant care, and reduce hospitalizations. ^{46,47,48,49,50} Ongoing clinical trials include primary palliative care interventions to improve decision-making about treatment for infection in advanced dementia (TRAIN-AD AG032982) in nursing homes and dementia palliative care delivered community-dwelling patients and their families (IN PEACE AG057733). Given the large population of persons with Alzheimer's and related dementias, and the prolonged suffering associated with this condition, improvements in both specialty and primary palliative care are necessary to improve outcomes.

Meaningful interpretation of our results requires consideration of limitations. This pilot clinical trial provides promising evidence for feasibility and efficacy, yet a statistically powered efficacy trial is necessary to prove this intervention can reduce hospital transfers and improve other outcomes important to persons with dementia and caregivers such as comfort and caregiver distress. As expected in a pilot trial this study had limited statistical power. In addition, the duration of specialty palliative care may need to be extended beyond discharge to affect these important outcomes. The study site has robust clinical resources in palliative care and geriatric care, and multi-site testing is needed to ensure this model can be exported and replicated. Study participation was good, but many persons with dementia could not be enrolled due to short hospital stays and caregiver stresses. Enrollment and outcomes may be improved if the intervention includes extended post-discharge specialty palliative care services, or access to services by telemedicine. As persons with dementia experience transitions between hospital, emergency room, nursing home, home or assisted living settings it may be important to offer enhanced elements of transitional care. 51,52

Specialty palliative care tailored to late-stage dementia, triggered by hospitalization for acute illness, is feasible and demonstrates potential to improve decision-making and clinical outcomes for persons with dementia and their families. This promising new model for dementia palliative care was designed using stakeholder input and evidence-based standards for dementia care. Results from this research will be used to develop a multi-site randomized controlled trial of specialty palliative care for hospitalized patients with late-stage dementia.

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References

- 1. Reisberg B, Ferris SH, De Leon MJ et al. The global deterioration scale for assessment of primary degenerative dementia. Am J Psych 1982; 139:1136–1139
- 2. Hebert LE, Weuve J, Scherr PA, Evans DA. Alzheimer disease in the United States (2010–2050) estimated using the 2010 census. Neurol 2013; 80:1778–1783.
- Alzheimer's Association. Latest Facts and Figures Report, 2018 https://www.alz.org/ alzheimersdementia/facts-figures. Accessed July 12, 2018.
- 4. Kelley AS, McGarry K, Fahle S, Marshall SM, Du Q, Skinner JS. Out-of-pocket spending in the last five years of life. J Gen Intern Med 2012; 28:304–309. [PubMed: 22948931]
- 5. Prince M, Bryce R, Albanese E, Wimo A, Ribeiro W, Ferri CP. The global prevalence of dementia: a systematic review and meta-analysis. Alzheimers Dement 2013; 9(1):63–75. [PubMed: 23305823]
- Hurd MD, Martorell P, Delavande A, Mullen KJ, Langa KM. Monetary costs of dementia in the United States. N Engl J Med 2013; 368:1326–1334. [PubMed: 23550670]
- Ferrell BA, Ferrell BR, Rivera L. Pain in cognitively impaired nursing home patients. J Pain Symptom Manage 1995; 10:591–598. [PubMed: 8594119]
- 8. Reynolds KS, Hanson LC, DeVellis RF, Henderson M, Steinhauser KE. Disparities in pain management between cognitively intact and cognitively impaired nursing home residents. J Pain Sym Manage 2008;35: 388–396.
- 9. Schulz R, Mendelsohn AB, Haley WE, Mahoney D, Allen RS, Zhang S, Thompson L, Belle SH. Resources for Enhancing Alzheimer's Caregiver Health (REACH) Investigators. End-of-life care and the effects of bereavement on family caregivers of persons with dementia. N Engl J Med 2003; 349:1936–1942. [PubMed: 14614169]
- 10. Black BS, Finucane T, Baker A et al. Health problems and correlates with pain in nursing home residents with advanced dementia. Alzheimer Dis Assoc Disord 2006; 20:283–290. [PubMed: 17132974]
- 11. Shega JW, Hougham GW, Stocking CB, Cox-Hayley D, Sachs GA. Management of noncancer pain in community-dwelling persons with dementia. J Am Geriatr Soc 2006; 54:1892–1897. [PubMed: 17198495]
- 12. Kverno KS, Rabins PV, Blass DM, Hicks KL, Black BS. Prevalence and treatment of neuropsychiatric symptoms in advanced dementia. J Gerotological Nurs 2008; 34:8–15.
- Hanson LC, Eckert JK, Dobbs D, Williams CS, Caprio AJ, Sloane PD, Zimmerman S. Symptom experience of dying long-term care residents. J Am Geriatr Soc 2008; 56:91–98. [PubMed: 17727647]

14. Hendriks SA, Smalbrugge M, Hertogh CMPM, van der Steen JT. Dying with dementia: symptoms, treatment, and quality of life in the last week of life. J Pain Symptom Manage 2014; 47:710–720. [PubMed: 23916680]

- Mitchell SL, Teno JM, Kiely DK, Shaffer ML, Jones RN, Prigerson HG, Volicer L, Givens JL, Hamel MB. The clinical course of advanced dementia. N Engl J Med 2009; 361:1529–1538.
 [PubMed: 19828530]
- 16. Miller SC, Lima JC, Mitchell SL. Hospice care for persons with dementia: the growth of access in US nursing homes. Am J Alz Dis Other Dem 2010; 25:666–673.
- 17. Ahronheim JC, Morrison RS, Baskin SA, Morris J, Meier DE. Treatment of the dying in the acute care hospital: advanced dementia and metastatic cancer. Arch Intern Med 1996; 156:2094–2100. [PubMed: 8862102]
- 18. Mitchell SL, Kiely DK, Hamel MB. Dying with advanced dementia in the nursing home. Arch Intern Med 2004; 164:321–326. [PubMed: 14769629]
- 19. Callahan CM, Arling G, Tu W, Rosenman MB, Counsell SR, Stump TE, Hendrie HC. Transitions in care for older adults with and without dementia. J Am Geriatr Soc 2012; 60:813–820. [PubMed: 22587849]
- 20. Miller SC, Han B. End-of-life care in US nursing homes: nursing homes with special programs and trained staff for hospice or palliative / end-of-life care. J Palliat Med 2008; 11:866–877. [PubMed: 18715179]
- Morrison RS, Augustin R, Souvanna P, Meier DE. America's care of serious illness: a state-by-state report card on access to palliative care in our nation's hospitals. J Palliat Med 2011; 14:1094

 1096. [PubMed: 21923412]
- 22. Givens JL, Selby K, Goldfeld KS, Mitchell SL. Hospital transfers of nursing home residents with advanced dementia. J Am Geriatr Soc 2012; 60:905–909. [PubMed: 22428661]
- Teno JM, Gozalo P, Mitchell SL, Tyler D, Mor V. Survival after multiple hospitalizations for infections and dehydration in nursing home residents with advanced cognitive impairment. JAMA 2013; 310:319–320. [PubMed: 23860991]
- 24. Hanson LC, Ersek M, Lin FC, Carey TS. Outcomes of feeding problems in advanced dementia in a nursing home population. J Am Geriatr Soc 2013; 61:1692–1697. [PubMed: 24083403]
- Morrison RS, Siu AL. Survival in end-stage dementia following acute illness. JAMA 2000; 284:47–52. [PubMed: 10872012]
- Murphy E, Froggatt K, Connolly S, O'Sea E, Sampson EL, Casey D, Devane D. Palliative care interventions in advanced dementia. Cochrane Database of Systematic Reviews 2016, Issue 12 Art. No.: CD011513. DOI: 10.1002/14651858.CD011513.pub2.
- 27. Gade G, Venohr I, Conner D, McGrady K, Beane J, Richardson RH, Williams MP, Liberson M, Blum M, Della Penna R. Impact of an inpatient palliative care team: a randomized controlled trial. J Palliat Med 2008; 11:180–190. [PubMed: 18333732]
- 28. Casarett D, Pickard A, Bailey FA, Ritchie C, Furman C, Rosenfeld K, Shreve S, Chen Z, Shea JA. Do palliative consultations improve patient outcomes? J Am Geriatr Soc 2008; 56:a593–599.
- Morrison RS, Penrod JD, Cassel JB, Caust-Ellenbogen M, Litke A, Spragens L, Meier DE, for the Palliative Care Leadership Centers' Outcomes Group. Cost savings associated with US hospital palliative care consultation programs. Arch Intern Med 2008; 168:1783–1790. [PubMed: 18779466]
- 30. Hanson LC, Usher B, Spragens L, Bernard S. Clinical and economic impact of palliative care consultation. J Pain Symptom Manage 2008;35:340–346. [PubMed: 18261877]
- 31. Ernecoff NC, Wessell KL, Gabriel S, Carey TS, Hanson LC. A novel screening to identify latestage dementia patients for palliative care research and practice. J Pain Symptom Manage 2018; 55:1152–1158. [PubMed: 29288881]
- 32. Reisberg B, Ferris SH, de Leon MJ, Crook T. The Global Deterioration Scale for assessment of primary degenerative dementia. The American journal of psychiatry. 1982 Sep.
- 33. Mitchell SL, Catic AG, Givens JL, Knopp J, Moran JA. Advanced dementia: a guide for families. Institute for Aging Research, Jamaica Plain, MA, 2011.

34. Hickman SE, Keevern E, Hammes BJ. Use of the Physician Orders for Life-Sustaining Treatment program in the clinical settings: a systematic review of the literature. J Am Geriatr Soc 2015; 63:341–350. [PubMed: 25644280]

- 35. Mitchell SL, Catic AG, Givens JL, Knopp J, Moran JA. Advanced Dementia: A Guide for Families. Institute for Aging Research, Hebrew SeniorLife, Boston MA, 2011.
- Alzheimer's Association. https://www.alz.org/help-support/caregiving/stages-behaviors/late-stage. viewed September 28, 2018.
- 37. Volicer L, Hurley AC, Blasi ZV. Scales for evaluation of end of life care in dementia. Alzheimer Dis Assoc Disord 2001; 15:194–200. [PubMed: 11723370]
- 38. Kiely DK, Volicer L, Teno J, Jones RN, Prigerson HG, Mitchell SL. The validity and reliability of scales for the evaluation of end of life care in advanced dementia. Alzheimer Dis Assoc Disord 2006 Jul-Sep; 20:176–181. [PubMed: 16917188]
- 39. Givens JL, Jones RN, Mazor KM, Prigerson HG, Mitchell SL. Development and psychometric properties of the Family Distress in Advanced Dementia scale. JAMDA 2015; 16:775–780. [PubMed: 25940236]
- 40. Hanson LC, Zimmerman S, Song MK, Lin FC, Rosemond C, Carey TS, Mitchell SL. Effect of the Goals of Care intervention for advanced dementia: a randomized clinical trial. JAMA Internal Medicine 2017; 177:24–31. [PubMed: 27893884]
- 41. Centers for Medicare and Medicaid Services, 2015 Physician Quality Reporting System (PQRS) Measures Groups Specifications Manual, version 8.0, 11/10/2014.
- 42. Volicer L, Hurley AC, Lathi DC, Kowall NW. Measurement of severity in advanced Alzheimer's disease. J Gerontol 1994; 49:M223–M226. [PubMed: 8056941]
- 43. Ahronheim JC, Morrison RS, Morris J, Baskin S, Meier DE. Palliative care in advanced dementia: a randomized controlled trial and descriptive analysis. J Palliat Med 2000; 3:265–273. [PubMed: 15859668]
- 44. Campbell ML, Guzman JA. A proactive approach to improve end-of-life care in a medical intensive care unit for patients with terminal dementia. Crit Care Med 2004; 32:1839–1843. [PubMed: 15343010]
- 45. Catic AG, Berg AI, Moran JA, Knopp JR, Givens JL, Kiely DK, Quinlan N, Mitchell SL. Preliminary data from an advanced dementia consult service: integrating research, education, and clinical expertise. J Am Geriatr Soc 2013; 61:2008–2012. [PubMed: 24219202]
- 46. Hanson LC, Carey TS, Caprio AJ, Lee TJ, Ersek M, Garrett J, Jackman A, Gilliam R, Wessell K, Mitchell SL. Improving decision making for feeding options in dementia care: a randomized trial. J Am Geriatr Soc, 2011; 59:2009–2016. [PubMed: 22091750]
- 47. Hanson LC, Zimmerman S, Song MK, Rosemond C, Gilliam R, Carey T, Chisholm L, Lin FC, Mitchell S. Effect of the Goals of Care intervention for advanced dementia: a randomized clinical trial. JAMA Intern Med, 2017; 177:24–31. [PubMed: 27893884]
- 48. Mitchell SL, Shaffer ML, Cohen S, Hanson LC, Habtemariam D, Volandes AE. An advance care planning video decision support tool for nursing home residents with advanced dementia: a cluster randomized clinical trial. JAMA Internal Med, 2018;178:961–969. [PubMed: 29868778]
- 49. Ernecoff NC, Zimmerman S, Mitchell SL, Song MK, Lin FC, Wessell K, Hanson LC. Concordance between goals of care and treatment decisions for persons with dementia. J Palliat Med, e-pub ahead of print 6 2018.
- 50. Cohen S, Volandes AE, Shaffer ML, Hanson LC, Habtemariam D, Mitchell SL. Concordance between proxy level of care preference and advance directives among nursing home residents with advanced dementia. J Pain Symptom Manage, in press 9 2018.
- 51. Callahan CM, Arling G, Tu W, Rosenman MB, Counsell SR, Stump TE, Hendrie HC. Transitions in care for older adults with and without dementia. J Am Geriatr Soc 2012; 60:813–820. [PubMed: 22587849]
- 52. Phelan EA, Debnam KJ, Anderson LA, Owens SB. A systematic review of intervention studies to prevent hospitalizations of community-dwelling older adults with dementia. Med Care 2015; 53:207–213. [PubMed: 25588136]

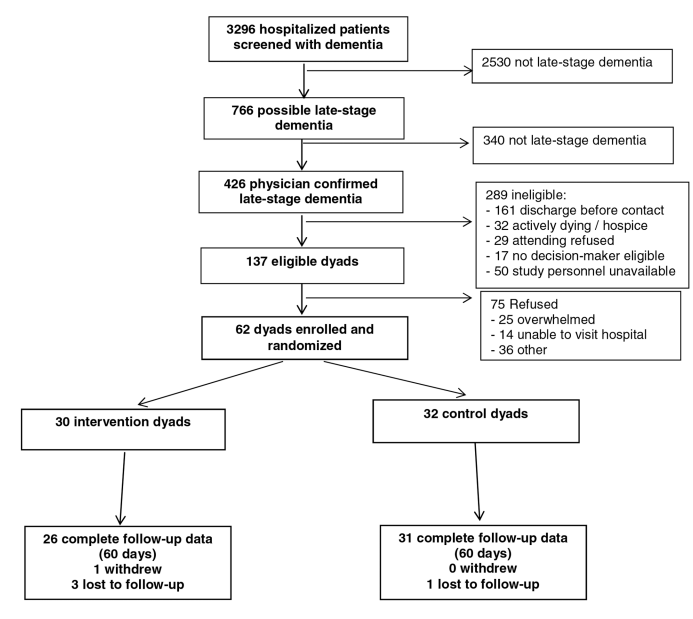


Figure 1: Enrollment and Participant Flow

 Table 1.

 Characteristics of Patients with Late-stage Dementia & Family Decision-makers.

	тоты	T44'	Control	
Patient Characteristics	TOTAL N=62	Intervention N=30	Control N=32	P value
Patient age, mean (SD)	83.9 (8.7)	83.0 (8.8)	84.7 (8.7)	0.434
Patient female, %	56%	67%	47%	0.116
Patient race, %				0.808
White	71%	70%	72%	
African American	24%	27%	22%	
Other	5%	3%	6%	
Patient Hispanic or Latino, %	5%	3%	6%	0.593
GDS *Stage, %				
5	37%	43%	31%	0.556
6	42%	40%	44%	
7	21%	17%	25%	
BANS-S **, mean (s.d.)	15.9 (3.6)	15.4 (3.5)	16.3 (3.6)	0.271
Admitting diagnosis in chart review				
Urinary tract infection	28%	37%	19%	0.114
Sepsis	16%	13%	19%	0.562
Neurospychiatric symptoms	15%	17%	13%	0.642
Hip fracture	11%	3%	19%	0.055
Pneumonia	8%	3%	13%	0.185
Dehydration/malnutrition	8%	7%	9%	0.700
Delirium	6%	7%	6%	0.947
Other	8%	13%	2%	0.298
Hospitalized in past 30 d.	10%	10%	9%	0.934
ER in past 30 d.	35%	40%	31%	0.472
Prior advance directive	63%	63%	63%	1.000
Pre-admission residence				
Home	51%	43%	60%	0.107
Assisted living	18%	30%	6%	
Nursing home	26%	23%	28%	
other	5%	4%	6%	
Hospital length of stay, median (range)	6 (2–36)	6 (2–36)	6 (2–32)	0.437
Family Decision-maker Characteristics				
Family decision-maker age, mean (SD)	59.7 (10.5)	60.0 (9.9)	59.5 (11.1)	0.844
Family decision-maker female, %	79%	73%	84%	0.286
Family decision-maker race, %				
White	71%	70%	72%	0.808
African American	24%	27%	22%	
Other	5%	3%	6%	
Hispanic or Latino, %	2%	0%	3%	0.329

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TOTAL N=62 Intervention Control **Patient Characteristics** N=32 P value N=30Relationship to person with dementia, % Daughter 47% 0.262 52% 56% Son 18% 27% 9% 19% 19% 20% Spouse Other 11% 16% 6% Family decision involvement, % 0.323 Not at all involved 0% 0% 0% Somewhat involved 10% 14% 6% 97% Very involved 90% 86% Family perceived 6-month prognosis, % Get better 0.926 17% 17% 17% Stay same 23% 23% 23% Get worse Likely to die 37% 40% 33% 23% 20% 27%

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^{*} GDS=Global Deterioration Scale, range 1–7, higher stage indicates more advanced dementia

^{**} BANS-S=Bedford Alzheimer's Nursing Severity-Subscale, range 7–28, higher scores indicate more functional disability

 Table 2.

 Outcomes of Triggered Palliative Care for Persons with Late-stage Dementia.

Primary Outcome N=30 do d N=32 do d P value Hospital / emergency visits per 60 days (no. of events/follow-up days) 0.68 (21/1843) 0.53 (20/2264) 0.415 Patient and Family Centered Secondary Outcomes Patient comfort (CAD-EOLD **, mean (SD) (range 14-42) 34.8 (4.2) 34.0 (4.1) 0.521 Family caregiver distress (FDAD ***, mean (SD) (range 14-5) 2.3 (0.5) 2.4 (0.5) 0.409 Palliative Care Domain Index (0-10) in hospital care plan 7.6 (2.5) 2.7 (1.7) <0.001 Phain 87% 72% 0.153 Dyspnea 77% 34% <0.001 Constipation 93% 2.5% <0.001 Poor appetite 37% 38% 0.946 Nausea 90% 2.5% <0.001 Neuropsychiatric symptom addressed 83% 2.5% <0.001 Depression 83% 2.5% <0.001 Anxiety 83% 1.6% <0.001 Decreased level of consciousness 47% 0.6 <0.001 Hyperactive delirium 80% 1.9%		Intervention	Control	
Hospital / emergency visits per 60 days (no. of events/follow-up days)	Outcomes			P value
Patient and Family Centered Secondary Outcomes	Primary Outcome	60 d	60 d	
Patient comfort (CAD-EOLD 5), mean (SD) (range 14-42) Family caregiver distress (FDAD **5), mean (SD) (range 1-5) Palliative Care Domain Index (0-10) in hospital care plan Palin (87), mean (SD) (range 1-5) Palliative Care Domain Index (0-10) in hospital care plan Physical symptom addressed Pain (87), mean (SD) (range 1-5) Palliative Care Domain Index (0-10) in hospital care plan Physical symptom addressed Pain (87), mean (SD) (range 1-5) Possible Care (1.5) Possible Care (1.5)		0.68 (21/1843)	0.53 (20/2264)	0.415
Family caregiver distress (FDAD ***), mean (SD) (range 1-5) Palliative Care Domain Index (0-10) in hospital care plan Physical symptom addressed Pain Pain 87% 72% 0.153 Dyspnea 77% 34% <0.001 Poor appetite 37% 38% 0.946 Nausea Pells 33% 16% 0.104 Neuropsychiatric symptom addressed Depression Anxiety Becreased level of consciousness 467% 22% 40.001 Spiritual needs addressed 47% 0% 40.001 Spiritual needs addressed Hospica at 60 days Community Palliative Care at 60 days Nursing home Death Other Death Other Patient vital status alive at 60 days, % Decision-making Secondary Outcomes Documented discussion of goals of care, % YES Documented discussion of goals of care, % YES DNR, present at admission 183% 22% 22% 0.0583 24.0.5) 2.7 (1.7) 2.0.001 2.7 (1.7) 2.0.001 2.7 (1.7) 2.0.001 2.7 (1.7) 2.0.001 2.7 (1.7) 2.0.001 2.7 (1.7) 2.0.001 2.7 (1.7) 2.0.001 2.7 (1.7) 2.0.001 2.0.001 2.5 (Patient and Family Centered Secondary Outcomes			
Palliative Care Domain Index (0–10) in hospital care plan Physical symptom addressed Pain Pain 87% 72% 0.153 Dyspnea 77% 34% <0.001 Constipation 93% 25% 0.946 Nausea 99% 25% 0.001 Falls Neuropsychiatric symptom addressed Depression Anxiety Becreased level of consciousness Hyperactive delirium Spiritual needs addressed Hospica at 60 days Community Palliative Care at 60 days Nursing home Death Other Patient vital status alive at 60 days, % Decision-making Secondary Outcomes Documented discussion of goals of care, % YES Documented discussion of goals of care, % YES DNR, present at admission 7.6 (2.5) 2.7 (1.7) 0.001 0.153 0.153 0.153 0.153 0.153 0.153 0.153 0.153 0.153 0.153 0.154 0.154 0.104 0.25% 0.104 0.10	Patient comfort (CAD-EOLD*), mean (SD) (range 14-42)	34.8 (4.2)	34.0 (4.1)	0.521
Physical symptom addressed 87% 72% 0.153 Dyspnea 77% 34% <0.001	Family caregiver distress (FDAD **, mean (SD) (range 1–5)	2.3 (0.5)	2.4 (0.5)	0.409
Pain 87% 72% 0.153 Dyspnea 77% 34% <0.001	Palliative Care Domain Index (0-10) in hospital care plan	7.6 (2.5)	2.7 (1.7)	< 0.001
Dyspnea	Physical symptom addressed			
Constipation 93% 25% <0.001 Poor appetite 37% 38% 0.946 Nausea 90% 25% <0.001 Falls 33% 16% 0.104 Neuropsychiatric symptom addressed Depression 83% 25% <0.001 Anxiety 83% 16% <0.001 Anxiety 83% 16% <0.001 Apperative delirium 80% 19% <0.001 Spiritual needs addressed 47% 0% <0.001 Hospica at 60 days 25% 30% 0.019 Community Palliative Care at 60 days 21% 7% 0.124 Hospital discharge location Home 33% 25% 0.046 Assisted living 24% 3% Nursing home 33% 63% Death 00% 0% Other 10% 9% Patient vital status alive at 60 days, YES Documented discussion of goals of care, % YES 90% 25% 0.001 MOST* completion at 60 days 79% 30% <0.001 MOST* completion at 60 days 79% 30% <0.001 Code status in hospital Full code 27% 22% 0.583 DNR, present at admission 53% 66%	Pain	87%	72%	0.153
Poor appetite 37% 38% 0.946 Nausea 90% 25% <0.001	Dyspnea	77%	34%	< 0.001
Nausea 90% 25% <0.001 Falls 33% 16% 0.104 Neuropsychiatric symptom addressed Depression 83% 25% <0.001 Anxiety 83% 16% <0.001 Decreased level of consciousness 67% 22% <0.001 Hyperactive delirium 80% 19% <0.001 Spiritual needs addressed 47% 0% <0.001 Hospice at 60 days 25% 3% 0.019 Community Palliative Care at 60 days 21% 7% 0.124 Hospital discharge location Home 33% 25% 0.046 Assisted living 24% 3% Nursing home 33% 63% Death 0% 0% 0% Other 10% 9% Patient vital status alive at 60 days, % 87% 94% 0.346 Decision-making Secondary Outcomes Documented discussion of prognosis, % YES 90% 3% <0.001 MOST completion at 60 days 79% 30% <0.001 MOST completion at 60 days 79% 30% <0.001 Full code 27% 22% 0.583 DNR, present at admission 53% 66%	Constipation	93%	25%	< 0.001
Falls 33% 16% 0.104 Neuropsychiatric symptom addressed 83% 25% <0.001	Poor appetite	37%	38%	0.946
Neuropsychiatric symptom addressed 83% 25% <0.001	Nausea	90%	25%	< 0.001
Depression 83% 25% <0.001	Falls	33%	16%	0.104
Anxiety Decreased level of consciousness 67% 22% <0.001 Hyperactive delirium 80% 19% <0.001 Spiritual needs addressed 47% 0% 0.001 Hospice at 60 days 25% 3% 0.019 Community Palliative Care at 60 days Hospital discharge location Home 33% 25% 0.046 Assisted living Nursing home Death 0% 0he Other 10% 9% Patient vital status alive at 60 days, % Pecision-making Secondary Outcomes Documented discussion of goals of care, % YES 90% 30% 0.001 A001	Neuropsychiatric symptom addressed			
Decreased level of consciousness 67% 22% <0.001	Depression	83%	25%	< 0.001
Hyperactive delirium Spiritual needs addressed 47% 0% <0.001 Spiritual needs addressed 47% 0% 0.001 Hospice at 60 days 25% 3% 0.019 Community Palliative Care at 60 days Hospital discharge location Home 33% 25% 0.046 Assisted living Nursing home Death Other Patient vital status alive at 60 days, % Pecision-making Secondary Outcomes Documented discussion of prognosis, % YES Documented discussion of goals of care, % YES 90% 30% 25% 0.046	Anxiety	83%	16%	< 0.001
Spiritual needs addressed 47% 0% <0.001	Decreased level of consciousness	67%	22%	< 0.001
Hospice at 60 days Community Palliative Care at 60 days Hospital discharge location Home Assisted living Nursing home Death Other Patient vital status alive at 60 days, % Decision-making Secondary Outcomes Documented discussion of prognosis, % YES Documented discussion of goals of care, % YES Pow MOST Completion at 60 days Code status in hospital Full code DNR, present at admission 25% 3% 0.019 0.019 0.019 0.0124 0.0246 0.046	Hyperactive delirium	80%	19%	< 0.001
Community Palliative Care at 60 days 21% 7% 0.124 Hospital discharge location 33% 25% 0.046 Home 33% 25% 0.046 Assisted living 24% 3% 3% Nursing home 33% 63% 63% Death 0% 0% 0% Other 10% 9% 24% 0.346 Decision-making Secondary Outcomes 87% 94% 0.346 Documented discussion of prognosis, % YES 90% 3% <0.001	Spiritual needs addressed	47%	0%	< 0.001
Hospital discharge location 33% 25% 0.046 Assisted living 24% 3% 3% Nursing home 33% 63% 63% Death 0% 0% 0% Other 10% 9% 9% Patient vital status alive at 60 days, % 87% 94% 0.346 Decision-making Secondary Outcomes 0cumented discussion of prognosis, % YES 90% 3% <0.001	Hospice at 60 days	25%	3%	0.019
Home 33% 25% 0.046 Assisted living 24% 3% Nursing home 33% 63% Death 0% 0% Other 10% 9% Patient vital status alive at 60 days, % 87% 94% 0.346 Decision-making Secondary Outcomes 0 3% <0.001	Community Palliative Care at 60 days	21%	7%	0.124
Assisted living Nursing home 33% 63% Death Other 10% 9% Patient vital status alive at 60 days, % 87% 94% 0.346 Decision-making Secondary Outcomes Documented discussion of prognosis, % YES 90% 3% <0.001 Documented discussion of goals of care, % YES 90% 30% <0.001 MOST + completion at 60 days Code status in hospital Full code DNR, present at admission 53% 66%	Hospital discharge location			
Nursing home 33% 63% Death 0% 0% Other 10% 9% Patient vital status alive at 60 days, % 87% 94% 0.346 Decision-making Secondary Outcomes 0 3% <0.001	Home	33%	25%	0.046
Death Other 0% 0% 0% Patient vital status alive at 60 days, % 87% 94% 0.346 Decision-making Secondary Outcomes Documented discussion of prognosis, % YES 90% 3% <0.001	Assisted living	24%	3%	
Other 10% 9% Patient vital status alive at 60 days, % 87% 94% 0.346 Decision-making Secondary Outcomes 0 3% <0.001	Nursing home	33%	63%	
Patient vital status alive at 60 days, % 87% 94% 0.346 Decision-making Secondary Outcomes 000 3% <0.001	Death	0%	0%	
Decision-making Secondary Outcomes 90% 3% <0.001	Other	10%	9%	
Documented discussion of prognosis, % YES 90% 3% <0.001	Patient vital status alive at 60 days, %	87%	94%	0.346
Documented discussion of goals of care, % YES 90% 25% <0.001	Decision-making Secondary Outcomes			
$MOST^{+}$ completion at 60 days79%30%<0.001Code status in hospital27%22%0.583DNR, present at admission53%66%	Documented discussion of prognosis, % YES	90%	3%	< 0.001
Code status in hospital	Documented discussion of goals of care, % YES	90%	25%	< 0.001
Code status in hospital 27% 22% 0.583 DNR, present at admission 53% 66%	MOST ⁺ completion at 60 days	79%	30%	< 0.001
Full code 27% 22% 0.583 DNR, present at admission 53% 66%				
DNR, present at admission 53% 66%	•	27%	22%	0.583
	DNR, present at admission			
	*			
Decision not to re-hospitalize 13% 0% 0.033			0%	0.033

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Intervention Control Outcomes P value N=30 N=32< 0.001 Decision made NOT to tube feed 53% 6% Decision made NOT to use antibiotics 7% 3% < 0.001 Decision made to LIMIT use of antibiotics 43% 0% Decision made NOT to use IV fluids < 0.001 13% 6% Decision made to LIMIT use of IV fluids 43% 3% Dementia Care Quality Metrics (chart review) 93% 9% < 0.001 Dementia staged Cognitive assessment 73% 9% < 0.001 Functional assessment 97% 25% < 0.001 Neuropsychiatric symptoms addressed 83% 25% < 0.001 0.001 Safety concerns addressed 83% 44% 80% < 0.001 Caregiver education 25%

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^{*}CAD-EOLD=Comfort Assessment in Dying, End of Life in Dementia, range overall 14–42, higher scores indicate more comfort

^{***}FDAD=Family Distress in Advanced Dementia, range overall and subscales 1–5, higher scores indicate more distress

^{*}MOST=Medical Orders for Scope of Treatment, the North Carolina version of POLST (Physician Orders for Life Sustaining Treatment) order set