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Development of an adaptive, personalized, and scalable dementia care program: Early findings from the Care Ecosystem

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Summary points

- Our objective was to develop and test a scalable model of dementia specialty care that complements primary care with additional caregiver support and education, medication consultation, and support in planning for future medical, financial, and legal decisions consistent with patient values. Care is delivered via the phone and web by unlicensed Care Team Navigators (CTNs), who are trained and supervised by a dementia specialist nurse, social worker, and pharmacist.
- This “Care Ecosystem” is being tested via a pragmatic randomized controlled trial. The care model was iteratively improved during the trial based on input from caregivers, primary care providers, and clinical team members.
- Based on the inputs, the care model was revised to enhance caregiver support, to clarify triage protocols, to include more strategies for managing problematic patient behaviors, to personalize care protocols, to better inform caregivers and patients about the services available, to adjust recruitment efforts to target the underserved, and to address CTN stress and burnout.
- The trial continues with the revised care model and ongoing evaluation of patient, caregiver, and health care cost outcomes. New implementation projects are adapting the care model to fit the priorities and workflows of three health care delivery organizations.
The challenge

The prevailing crisis management approach to dementia care does not adequately prepare and support patients and caregivers to manage the evolving challenges they face. Providers are reimbursed for brief problem-focused, in-person visits during which they simultaneously address multiple comorbid conditions [1]. Recent Centers for Medicare and Medicaid Services (CMS) policy changes provide reimbursement for monthly chronic care management by clinical staff that could provide an added layer of support [2]. A comprehensive care plan must be established, implemented, monitored, and revised as needed. Our challenge was to develop, improve, and test a new model of dementia care that addresses unmet needs of patients and caregivers, and aligns with the CMS’s initiatives for better health, better care, and lower costs [3].

The Care Ecosystem model

The Care Ecosystem is a telephone-based supportive care intervention for patients with dementia and their caregivers. The “Care Team Navigator” (CTN) is an unlicensed, trained dementia care guide who serves as the patient and caregiver’s primary point of contact. CTNs screen for common problems and provide support and standardized education. Contact frequency is scaled based on each patient and caregiver’s (dyad’s) needs and preferences but is typically monthly. CTNs triage complex medical or social issues to a nurse, social worker, and pharmacist, who have expertise in dementia care, and coordinate care with other health providers (Fig 1). CTNs are selected for strong communication and listening skills. They undergo 40 hours of didactic training through video or in-person lectures, assigned reading, and clinical observation. Ongoing training occurs during weekly case reviews and supervision ("clinical debriefings") with the nurse, social worker, and pharmacist. Workflows and communication among the clinical team is supported by the Care Ecosystem Dashboard, a Salesforce software program developed with input from clinical staff.

The Care Ecosystem trial

The impact of the Care Ecosystem on patient, caregiver, and cost outcomes is being tested through a pragmatic, randomized controlled trial [8] funded by a Centers for Medicare & Medicaid Innovation (CMMI) Healthcare Innovations Award Round Two (September 1, 2014, to August 31, 2017; see Fig 2). Inclusion criteria are dementia diagnosis of any type by any medical provider; age ≥45; Medicare- or Medicaid-enrolled or pending; California, Nebraska, or Iowa residence; a caregiver who may or may not reside with the patient; and fluency in English, Spanish, or Cantonese. Exclusion criteria are persons with a life expectancy of less than 3 months or living in a nursing home at time of enrollment. Consent materials were approved by the University of California San Francisco (UCSF) and the University of Nebraska Medical Center (UNMC) Institutional Review Boards. Patients with capacity for written informed consent are given the opportunity to consent for themselves; otherwise, consent is obtained from a legally authorized representative. Primary caregivers provide written consent. Dyads are randomly assigned to the Care Ecosystem intervention or usual care at a 2:1 ratio. The intervention
continues for 1 year or until trial end. Target enrollment is 1,050 dyads. Baseline characteristics of dyads enrolled are presented in Table 1.

Prior to starting the trial, we worked with our respective Institutional Review Boards to ensure that appropriate human subject protections were in place. We review any incidents or adverse events with our Data Safety Monitoring Board and the UCSF Privacy Office. To date, we have had three reportable incidents, all involving a breach of privacy, in which a staff person accidentally sent confidential information to the wrong participant. We instituted a peer check protocol to prevent future incidents.

**Trial outcomes**

A 30-minute Outcome Survey is administered by phone prior to randomization and at 6 and 12 months post-randomization by a research coordinator blinded to randomization status ([S1 Table](#)). Medicare and Medicaid claims data are analyzed to evaluate cost of care. At the end of the 12-month outcomes survey, the coordinator is instantly unblinded so that they can administer a 10-minute Satisfaction Survey to caregivers in the intervention group before ending the call (see [S1 Text](#)). This Satisfaction Survey was also administered to the first 132 caregivers randomized to the intervention, 4 months post-randomization, to support agile development of the care model (see below).
Agile development of the care model

Without compromising standardization of core program elements, including the composition of the clinical team and our emphases on caregiver support and education, medications, and advance care planning, we applied “agile” software development principles [15,16] to refine the care model based on user input. These principles, as applied to our care model development, were a care model that is adapted efficiently based on user inputs, modular care protocols with well-defined integration points, collaborative development and revisions by a multidisciplinary team, incremental revisions with minor releases (approximately monthly) and major releases (Care Model 1.0, Care Model 2.0, and the three implementation releases), and users broadly defined to include caregivers, clinical team members, and primary care providers. Feedback was used to revise modular care protocols, CTN training, patient and caregiver educational materials, and our Care Ecosystem Dashboard. These revisions did not require consent form modifications. The user inputs included 4-month Caregiver Satisfaction Surveys, quarterly meetings with the UCSF Memory and Aging Center Family Advisory Council comprising 13 dementia family caregivers, three phases of interviews with all Care Ecosystem clinical staff and input gathered during weekly clinical debriefings, and nine primary care provider focus groups. The provider focus groups, conducted between February and June 2015, were led by a geriatrician, included a total of 43 physicians and 6 nurse practitioners, and asked about perspectives on care in dementia, preferred methods of care coordination, and their reaction to the Care Ecosystem model. Qualitative data analysis of user inputs guided program revisions, which were complete by September 2016, creating Care Model 2.0. See Fig 3.

Trial status

As of September 1, 2016, when Care Model 2.0 was released, 610 dyads were enrolled in the Care Ecosystem trial, with 400 randomized to treatment and 210 to usual care (Table 1).
Table 1. Characteristics of persons with dementia and caregivers enrolled as of September 1, 2016.

<table>
<thead>
<tr>
<th></th>
<th>A. Persons with Dementia</th>
<th>B. Caregivers</th>
<th></th>
<th></th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Care Ecosystem</td>
<td>Usual Care</td>
<td>Care Ecosystem</td>
<td>Usual Care</td>
</tr>
<tr>
<td>n</td>
<td>400</td>
<td>210</td>
<td>400</td>
<td>210</td>
</tr>
<tr>
<td>Age M(SD)</td>
<td>78.8 (8.7)</td>
<td>77.3 (10.1)</td>
<td>66.8 (11.7)</td>
<td>65.5 (11.2)</td>
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<tr>
<td>Ethnicity (%)</td>
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<tr>
<td>Caucasian</td>
<td>82.0</td>
<td>84.8</td>
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<td>81.5</td>
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<td>5.5</td>
<td>4.8</td>
<td>Asian</td>
<td>6.5</td>
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<tr>
<td>Black or African American</td>
<td>4.5</td>
<td>3.3</td>
<td>Black or African American</td>
<td>4.5</td>
</tr>
<tr>
<td>Other</td>
<td>8.0</td>
<td>7.1</td>
<td>Other</td>
<td>7.5</td>
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<tr>
<td>Education (%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;12 years</td>
<td>7.0</td>
<td>6.2</td>
<td>&lt;12 years</td>
<td>1.8</td>
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<tr>
<td>12 years</td>
<td>20.3</td>
<td>22.4</td>
<td>12 years</td>
<td>7.3</td>
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<tr>
<td>13–15 years</td>
<td>21.3</td>
<td>18.1</td>
<td>13–15 years</td>
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</tr>
<tr>
<td>≥16 years</td>
<td>51.5</td>
<td>53.3</td>
<td>≥16 years</td>
<td>65.4</td>
</tr>
<tr>
<td>Female (%)</td>
<td>58.3</td>
<td>51.0</td>
<td>Female (%)</td>
<td>67.2</td>
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<tr>
<td>State of residence (%)</td>
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<td></td>
<td>Caregiver relationship (%)</td>
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<td>California</td>
<td>56.8</td>
<td>53.8</td>
<td>Wife</td>
<td>34.5</td>
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<tr>
<td>Nebraska</td>
<td>38.3</td>
<td>38.1</td>
<td>Daughter</td>
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<tr>
<td>Iowa</td>
<td>5.0</td>
<td>8.1</td>
<td>Husband</td>
<td>24.8</td>
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<tr>
<td>Annual household income (%)</td>
<td></td>
<td></td>
<td>Son</td>
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<tr>
<td>&lt;$15,000</td>
<td>2.6</td>
<td>3.0</td>
<td>Sibling or Other</td>
<td>7.7</td>
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<td>23.1</td>
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<tr>
<td>$50,000–$99,999</td>
<td>31.4</td>
<td>32.0</td>
<td>Mild–severe depression (%)</td>
<td>39.5</td>
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<tr>
<td>$100,000–$149,999</td>
<td>16.7</td>
<td>19.5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>≥$150,000</td>
<td>14.4</td>
<td>12.4</td>
<td></td>
<td></td>
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<tr>
<td>Don’t know/refused</td>
<td>9.0</td>
<td>10.1</td>
<td></td>
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<tr>
<td>Stage</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mild</td>
<td>49.8</td>
<td>50.5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Moderate</td>
<td>30.8</td>
<td>31.4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Advanced</td>
<td>19.5</td>
<td>18.1</td>
<td></td>
<td></td>
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<tr>
<td>Comorbidities</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cardiovascular</td>
<td>54.5</td>
<td>45.7</td>
<td></td>
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<tr>
<td>Arthritis/rheumatism</td>
<td>43.8</td>
<td>41.9</td>
<td></td>
<td></td>
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<tr>
<td>Depression</td>
<td>42.8</td>
<td>39.5</td>
<td></td>
<td></td>
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<tr>
<td>Stroke</td>
<td>15.0</td>
<td>11.4</td>
<td></td>
<td></td>
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<tr>
<td>Diabetes</td>
<td>13.5</td>
<td>13.3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>QoL-AD Mean (SD)</td>
<td>32.9 (5.8)</td>
<td>33.4 (5.8)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Emergency care utilization, last 6 months</td>
<td></td>
<td></td>
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<tr>
<td>Hospitalization rate</td>
<td>0.31</td>
<td>0.27</td>
<td></td>
<td></td>
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<tr>
<td>ER visit rate</td>
<td>0.37</td>
<td>0.47</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Characteristics were collected during the baseline Outcomes Survey with the caregivers. Stage of dementia was determined using the Quick Dementia Rating Scale using published cut-offs that correspond to the Clinical Dementia Rating Scale [9]. The most common comorbidities are reported based on questions from the Charleston Comorbidity questionnaire [10,11]. Hospitalization and ER visit rates are the number of visits in the prior 6 months divided by the number of patients. On the QoL-AD, caregivers rate patient quality of life on 13 items as poor, fair, good, or excellent, and scores can range from 13–52 [12]. Ratings that are “fair” on average result in a score of 26 and “good” in a score of 39. Caregiver burden is based on the Zarit-12, with scores >17 indicating severe [13]. Caregiver depression is based on the PhQ-9, with scores >4 indicating mild depression [14].

ER, emergency room; M, mean; QoL-AD, quality of life in Alzheimer disease; SD, standard deviation.

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Enrollment continued through February 28, 2017. A UCSF clinical team provides services to 227 California residents, and an independent UNMC clinical team provides services to 153 Nebraska and 20 Iowa residents (see S1 Fig for locations of enrolled patients). The current maximum caseload for a CTN is 64, and the target maximum caseload is 80. Bilingual CTNs provide care in the recipient’s preferred language (English, Spanish, or Cantonese).

Caregiver satisfaction ratings
Satisfaction ratings were obtained during the 4-month Caregiver Satisfaction Survey. Ninety-two percent would recommend the Care Ecosystem to other caregivers. Seventy-seven percent said they needed help from their CTN at some point since enrollment, and 96% of these said the CTN was available when needed. Ninety-two percent said the amount of contact they had with their CTN was about right, 7% said that there was too little contact, and none said they had too much contact (1% did not respond). Fifty-nine percent reported that the CTN referred them to resources in their community, and 87% of these said the referrals were helpful. Forty-nine percent of caregivers indicated that the patient had problematic behavioral changes; of these, 65% said the CTN helped them manage those behaviors. Eleven percent of caregivers indicated that they thought the Care Ecosystem helped the patient avoid an unnecessary emergency room or hospital visit. Responses to open-ended questions on the Satisfaction Survey were analyzed using Atlas.Ti, a qualitative data coding software.
Care model 2.0: User input and program revisions

Increase emphasis on caregiver support

Caregivers, clinical staff, and primary care providers all highlighted the importance of caregiver support for reducing caregiver distress. This was emphasized in seven of the nine primary care focus groups and in all six meetings with the Family Advisory Council. On the Satisfaction Survey, 83 caregivers commented favorably on aspects of the Care Ecosystem intervention that focused on caregiver support, such as the provision of information and resources, referrals to respite programs, and general emotional support. Of these caregivers, 35 specifically referred to the benefits of the relationship with their particular CTN.

To enhance this support, the CTN training curriculum was updated to improve competency in balancing rapport building with direct problem-solving, and sample scripts were created to help guide conversations around difficult topics regularly raised by dementia caregivers. Open-ended questions are now employed in every call to identify needs. Additionally, CTNs were trained to identify and proactively find solutions to problems impacting caregiver well-being (e.g., inattention to personal health, need for respite, loneliness, and grief).

Clearer triage protocols

Our licensed clinical staff emphasized the need to clarify scope of each role on the team for best practices. On the Satisfaction Survey, 12 caregivers commented on deficits in CTN knowledge or abilities. Providers in six focus groups expressed that while they appreciated that CTNs could help identify health issues between visits and prepare families for provider appointments, all medical guidance must be deferred to licensed clinicians. This feedback highlighted the limits of using a nonlicensed staff member as the first point of contact in delivering specialty care. We clarified training protocols around when and how to triage to the Care Ecosystem nurse, social worker, and pharmacist, and to the patient’s primary care provider, as follows.

Advance care planning

The role of the CTN was limited to include identifying gaps in planning (e.g., identifying a health care agent), making legal referrals, and preparing the family for a billable advance care planning appointment with their primary care provider [17].

Monitoring health status

CTNs provide education and screen monthly for acute health changes that could signal delirium. Concerns identified by the CTNs are triaged to the nurse immediately.

Managing challenging behaviors

CTNs screen for challenging behaviors during the Intake Screen, listen for behavior problems mentioned by the caregiver at any time, and provide targeted educational handouts. If the situation does not improve or the caregiver is significantly distressed, the nurse consults directly with the caregiver.

Medication management

The CTNs reconcile what the patient is taking and check in monthly for possible side effects or adherence problems. The pharmacist provides medication reviews and recommendations to the prescribing providers and caregivers.
Personalization of protocol

On the Satisfaction Survey, 45 caregivers indicated that our medication and advance care planning protocols were duplicative with work they had done elsewhere, and CTNs noted that these structured protocols could interfere with building rapport. Now each protocol starts with a needs assessment and the CTNs address only identified needs.

Creation of service menu

Twelve caregivers expressed a lack of clarity around the extent of services the Care Ecosystem offered. Therefore, now we include a Care Ecosystem service list in our welcome packet, and the CTN refers to this list in communications with the dyad.

Adjustment of recruitment efforts

Initially, the most efficient recruitment pipelines were an academic medical center tertiary care clinic (the UCSF Memory and Aging Center Clinic) and self-referrals. This resulted in a disproportionately white, high-income, and English-speaking sample (see Table 1). Primary care providers in five of the focus groups highlighted care gaps for the underserved, and some of our high-income caregivers indicated that their needs were already well met. Now, we emphasize recruitment of low-income, ethnically diverse, monolingual Chinese and Spanish speakers, and rural-dwelling persons. Currently, we suspect that the benefit experienced by the underserved will be greater.

Addressing CTN stress and burnout

CTNs expressed that listening to patients and caregivers talk about grief, loss, and frustration carries an emotional toll. We now provide a monthly workshop facilitated by a mental health specialist or chaplain to provide CTNs with emotional support and coping strategies.

Next steps and future directions

We have initiated implementation projects with Allina Health primary care clinics in Minnesota, the San Francisco Health Network primary care clinics, and the UCSF Memory and Aging Center dementia specialty clinic. The Care Ecosystem will be integrated into these clinics with on-site CTNs who have direct access to electronic medical records, facilitating a level of care coordination that was not possible during the trial. Without the burdens of research measurement and randomization to a control group, we anticipate that the program will be more appealing to distressed caregivers. Also, at the San Francisco Health Network clinics, CTNs will help address a major barrier to care—late or missed diagnosis—by providing cognitive and functional screens when they are requested by the primary care provider. The three implementation projects are beginning with regular meetings to carefully understand each site’s workflows and priority improvement opportunities to guide adaptations of Care Model 2.0 that are specific for each health care delivery system [18]. Concurrently with these implementation projects, we hope to measure the long-term impacts of Care Ecosystem in the CMMI trial cohort.

The iterative design of our trial allowed us to improve the care model based on our learnings but will complicate the interpretation of treatment effects. Importantly, the core program elements remained standard to permit analyses in the full sample. Secondary analyses that include a time-varying covariate will help us understand and communicate the impact of the revisions on study outcomes.
Supporting information

S1 Table. Outcome Survey measures.  
(DOCX)

S1 Text. The Caregiver Satisfaction Survey.  
(DOCX)

S1 Fig. Zip codes in which Care Ecosystem participants with dementia are enrolled.  
(TIF)

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Formal analysis: KLP, AB.

Funding acquisition: KLP, BLM.

Investigation: JM, SJB, KL, SD, TB, JEF, AMC, JH, PO, KMC, ADB.

Methodology: KLP, JM, SJB, AB, WC, KL, LW, SMH, SD, MWS, AKS, JH, KMC, RG, AC, GN, ABB, DD, CR, BLM.

Project administration: KLP, JM, SJB, SL.

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Software: KLP, KL, SD, MWS, PO.

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Validation: KLP, AB.

Visualization: KLP, AB.

Writing – original draft: KLP, JM, SJB, AB.

Writing – review & editing: KLP, JM, SJB, AB, WC, KL, LW, SMH, SD, DD, CR, BLM.

References


