The development of specialized palliative care in the community: A qualitative study of the evolution of 15 teams

Hsien Seow¹,²,³ and Daryl Bainbridge¹,³

Abstract

Background: Interprofessional specialized palliative care teams at home improve patient outcomes, reduce healthcare costs, and support many patients to die at home. However, practical details about how to develop home-based teams in different regions and health systems are scarce.

Aim: To examine how a variety of home-based specialized palliative care teams created and grew their team over time and to identify critical steps in their evolution.

Design: A qualitative study was designed based on a grounded theory approach, using semi-structured interviews and other documentation.

Setting/Participants: In all, 15 specialized palliative care teams from Ontario, Canada, representing rural and urban areas. Data were collected from core members of the teams, including nurses, physicians, personal support workers, spiritual counselors, and administrators.

Results: In all, 122 individuals where interviewed, ranging from 4 to 10 per team. The analysis revealed four stages in team evolution: Inception, Start-up (n = 4 teams), Growth (n = 5), and Mature (n = 6). In the Inception stage, a champion provider was required to leverage existing resources to form the team. Start-up teams were testing and adjusting care processes to solidify their presence in the community. Growth teams had core expertise, relationships with fellow providers, and 24/7 support. Mature teams were fully integrated in the community, but still engaged in continuous quality improvement.

Conclusion: Understanding the developmental stages of teams can help to inform the progress of other community-based teams. Appropriate outcome measures at each stage are also critical for team motivation and steady progress.

Keywords
Palliative care, terminal care, home care services, patient care team community health services, capacity building, qualitative research

What is already known about the topic?

- Community-based specialized palliative care has been shown to improve symptom management and quality of life, and reduce healthcare costs compared to usual homecare.
- Existing validated frameworks on community-based palliative care teams describe theoretical phases and antecedent factors but lack operational and practical details required for team development.

What this paper adds?

- Synthesizes experiences from 15 diverse specialized, community-based palliative care teams.
- Details the characterizing features, activities, milestones, and challenges unique to each of four stages of team evolution evident: Inception, Start-up, Growth, and Mature.
- Describes output and outcome measures that are appropriate to each stage.

¹Department of Oncology, McMaster University, Hamilton, ON, Canada
²Escarpment Cancer Research Institute, Hamilton, ON, Canada
³Juravinski Cancer Centre, Hamilton, ON, Canada

Corresponding author:
Hsien Seow, Department of Oncology, McMaster University, 699 Concession Street, Room 4-229, Hamilton, ON L8V 5C2, Canada.
Email: seowh@mcmaster.ca
Implications for practice, theory or policy

- Defining common stages in the evolution of community-based palliative care teams can inform the development of new teams.
- Using stage-appropriate milestones and measures can gauge progress and set realistic expectations for team growth.

Introduction

An estimated 70% of the dying population in Canada, and other high-income countries, will require palliative care. Developing these services in the community is important because many patients prefer to die at home, which can also be higher quality and less expensive than in hospital. Over a dozen randomized controlled trials and dozens more observational studies have shown that home-based palliative care can be effectively delivered using interprofessional specialized palliative care teams. These teams better managed symptoms, improved quality of life, and prevented late-life hospitalizations compared to usual care. Yet, community-based specialized palliative care teams are not commonplace. In part, this is because prior research has focused on demonstrating the health services outcomes of the teams, but has provided little description of how the teams developed and sustained themselves. As a result, we lack knowledge on effective strategies to build and replicate optimal models of teams in other regions.

A few evidence-based frameworks have been proposed that outline the development and growth of integrated palliative care programs in the community. Kelley’s Community Capacity-Building Model, based on research in rural areas, affirms that the initiation of a community-based palliative care program is founded upon four sequential phases or events: emergence of antecedent conditions, occurrence of a catalyst, creation of the team, and growing the program. Similarly, a grounded theory study by Ploeg et al. on the adoption and scaling of best practices in home care for older adults identified five process phases: committing to change, implementing on a small scale, adapting locally, spreading internally to multiple users and sites, and disseminating externally. Together, these models contribute to the understanding of the theoretical stages of local program development. Still there remains little evidence on practical details, such as key milestones and outcome measures at each stage, that would guide providers wanting to replicate an effective team model in different contexts.

To address this gap in knowledge, we capitalized on a natural experiment existing in Ontario, Canada, where 15 regions independently developed their own community-based, specialized palliative care teams, hereafter referred to as “Teams.” These teams consisted of palliative care “specialist” physicians (who had completed a 1-year fellowship in palliative care, whose billings were >50% for palliative care, and worked exclusively in the community) and nurses who completed additional training or a certification exam in palliative care. In a previous study, we found that exposure to any of the teams was associated with a 50% reduction in acute care use compared to usual care. This cohort represents a unique opportunity to investigate and compare several diverse and efficacious teams within one healthcare system. We undertook a qualitative examination of these teams to understand the process of their development over time.

Methods

Design

We used a grounded theory approach to understand the milestones and stages in the evolution of creating a home-based team. Data were collected and analyzed between February and August 2013. The study was approved by Hamilton Health Sciences/McMaster University research ethics review board, Ontario, Canada (11-403). Written informed consent was obtained from each participant.

Setting

Teams from Ontario, Canada, were examined. The majority of community-based palliative care is delivered by homecare providers (e.g., nurses and personal support workers). Ontario has a few dozen residential hospices (i.e., free-standing, home-like facilities in the community) but only about 4% of patients die in this setting. Some communities have a visiting hospice volunteer program and a small proportion have a hospital palliative care unit (19 units in ON).

To improve palliative care delivery and address the fragmentation between primary care and homecare, some communities developed interprofessional specialized palliative care teams. These teams provide services similar to the visiting hospice service under the United States’ Medicare Hospice Benefit or palliative homecare provided by MacMillan nurses in the United Kingdom. Patients referred to the Ontario teams are usually in the last months of life and are followed until death. The teams’ services are offered at no cost to patients, although the funding sources vary for the providers involved. Physicians are reimbursed through fee-for-service and/or salary arrangements, while homecare nurses are covered by homecare funding. Some of the teams also rely on fundraising to cover infrastructure costs.
Participants

In all, 15 teams and team members across Ontario were selected using a purposive sampling process, aimed at achieving a variety of profiles. The teams selected represent a diverse geography, including rural and urban communities. These teams met the study inclusion criteria of multidisciplinary team, having specialized palliative care expertise and training, providing care to patients in their homes, and having a focus on pain and symptom management.

The team models vary in the extent that the physicians consult or take over care. The “specialist” physicians sometimes worked in a consultative manner to family physicians, generalist homecare nurses and personal support workers, and other allied health providers to care for seriously ill patients in the home (consultation model). Other times, the “specialist” physicians would assume primary ownership of the patients (take over model). Eleven of the teams were previously studied as exemplars of specialized palliative care teams serving patients in their homes. The remaining teams were newly formed and in the initial stages of organizing a collaborative practice to serve palliative care patients in their area.

The researchers were known to the team leads through prior activities/events. Team leads introduced the study to their members, who were then approached by the researchers for participation using email and/or phone contacts. In teams with less than six members, all team members were interviewed. In larger teams, a purposive sample was taken to include interprofessional representation and those with the most experience with the team, until data saturation was reached. Core team members interviewed included palliative care specialist physicians, palliative care nurses, allied health professionals (e.g. generalist homecare nurses), homecare case managers, and team administrators. We did not interview family physicians who were supported by the teams. The lead from each team also provided formative information, team documents, and subsequent information requested for clarification by the research team.

Procedure

In-person semi-structured interviews were conducted with team members using an interview guide (Appendix 1). The guide was developed by the research team based on the relevant literature, existing palliative care provision frameworks, and expert opinion. The guide was designed to explore participants’ perspectives on the process of building a team to provide community-based palliative care. Based on responses, the trained interviewers followed up with additional or clarifying questions.

Interviews were conducted in the participants’ workplaces and took approximately one hour each. Each interview was lead jointly by a primary (H.S., D.Ba., D.Br., S.T.T. (see acknowledgements)) and a secondary interviewer who served as a note-taker. Interviews were audio taped and transcribed by the primary interviewer. Documents describing the history of the teams, relevant administrative data, and process data, that is, briefings, characteristics of caseload, tools incorporated into practice, were also compiled for the teams.

Data analysis

In accordance with Corbin et al.’s grounded theory approach, data analysis proceeded through stages of open, axial, and selective coding. Data for each participant were first coded by the primary interviewer, then reviewed by the secondary interviewer and discussed. Resulting codes and supplementary document data from the teams were then examined by the four primary interviewers together, to determine similarities, divergences, and associations. Codes, themes, and categories were generated first by individual respondent, then by each team, and then across all teams. A constant comparative method was used throughout the analytic process to compare new data to existing codes. To reduce analysis bias, we maintained an audit trail of decisions, used an external peer review process to substantiate the research process, and shared back preliminarily findings with the respondent teams for comment.

Results

A total of 122 palliative care providers and administrators involved in 15 home-based teams were interviewed (Table 1). Each team had a mean of seven providers interviewed (range of 4–10). The most common profession interviewed were nurses (37%). Almost a quarter (23%) of participants had 11 or more years of experience in their current role in the team. The 15 teams had characteristics that varied by geography, caseload size, number of physician and nurse providers on team, and year of inception (Table 2). For instance, the palliative care physician full time equivalent (FTE) varied from 0.5 to 11.5 between teams. Overall, 80% of the patients had cancer and were seen by the teams for a median of 1 to 2 months before death. The median patient age was 75 years. A specialist palliative care physician working with a team would typically have 50–100 patients under their care at one time, and see 125–175 new patients each year.

Stages of team evolution

Despite differences in characteristics, the analysis revealed that team development underwent a process of evolution that took time and determination:

It’s a gradual evolution, you can’t do it all at once, you have to be patient and learn from your mistakes. It’s very “two steps back and three forward,” hopefully in a way that
improves the program. As we’ve run into barriers we problem solve and try to bring it back to the team meetings and go forward one by one. (Interviewee 1–Team 11)

The thematic analysis of our data revealed four stages of team evolution: Inception, Start-Up (4 teams), Growth (5 teams), and Mature (6 teams). All teams in the study had evolved past an initial Inception stage. Characterizing features, milestones, challenges, and outcome measures of each stage are described below and summarized in Table 3.

**Stage 1: inception.** Most of the teams were initially spearheaded by a provider champion, usually a nurse or physician, rallying members of the local community to improve access to palliative care. The champion was able to leverage funding and/or obtain in-kind support from organizations and providers interested in fulfilling this “dream.”

The existence of at least one person to champion the cause and a community that perceives the value of palliative care, were seen as critical to the genesis of the team:

Without local champions and leaders, nothing would happen. (2–Team 12)

The teams built upon existing services when possible rather than duplicate those already in place. Many preliminary teams conducted some form of needs assessment to obtain a sense of what support is available for dying patients, what is needed, and who are the current “players.” Most teams first worked with local primary care physicians who were amenable to receiving support from the team for their dying patients. The teams gradually developed a reputation for expert care, which encouraged other primary care physicians to either work with or refer patients to the team:

**Table 1.** Participant Demographics (N = 122).

| Participant characteristic | Nurses | Executives<sup>a</sup> | Physicians | Other<sup>b</sup> | Overall (%) |
|---------------------------|--------|-------------------------|------------|-------------------|-------------|
| Sex (n)                   |        |                         |            |                   |             |
| Female                    | 42     | 34                      | 9          | 12                | 97 (82.9)   |
| Male                      | 3      | –                       | 13         | 4                 | 20 (17.1)   |
| No. years in current role (n) |        |                         |            |                   |             |
| 0–5                       | 19     | 13                      | 8          | 13                | 53 (43.4)   |
| 6–10                      | 12     | 8                       | 1          | 2                 | 23 (18.9)   |
| 11 +                      | 12     | 6                       | 7          | 3                 | 28 (23.0)   |
| Unknown                   | 2      | 7                       | 6          | 3                 | 18 (14.8)   |
| Total (%)                 | 45 (36.9) | 34 (27.9)            | 20 (16.4)  | 26 (21.3)         | 122 (100)   |

<sup>a</sup>Executives included directors, managers, and team leads, many had nursing or medical backgrounds.

<sup>b</sup>Other included homecare case managers, social workers, pharmacists, spiritual and bereavement staff, and data support staff.

**Table 2.** Characteristics of specialized teams.

| Team # | Approximate patient deaths per year (2012) | Median days in program until death | Palliative care physicians (FTE) | Nurses (FTE) | Other members<sup>a</sup> (FTE) | Date team established | Stage of team |
|--------|------------------------------------------|----------------------------------|---------------------------------|--------------|-------------------------------|----------------------|---------------|
| 1      | 90                                       | 32                               | 6                               | 2            | 4.7                           | 1979                 | Mature        |
| 2      | 340                                      | 45                               | 11.5                            | 1            | 5.9                           | 1986                 | Mature        |
| 3      | 390                                      | 38                               | 1.3                             | 3            | 1.7                           | 1996                 | Mature        |
| 4      | 135                                      | 23                               | 0.6                             | 1            | 2.5                           | 2004                 | Mature        |
| 5      | 250                                      | 49                               | 2                               | 2            | 1                             | 2007                 | Mature        |
| 6      | 110                                      | 53                               | 3                               | 3.5          | 5                             | 2009                 | Mature        |
| 7      | 45                                       | 63                               | 2                               | 2            | 1.2                           | 2009                 | Growth        |
| 8      | 55                                       | 36                               | 0.5                             | 1            | 0.2                           | 2009                 | Growth        |
| 9      | 65                                       | 40                               | 1                               | 2            | 1                             | 2009                 | Growth        |
| 10     | 70                                       | 38                               | 1                               | 1            | 0.6                           | 2009                 | Growth        |
| 11     | 415                                      | 40                               | 1                               | 8            | 2                             | 2009                 | Growth        |
| 12     | 55                                       | 45                               | 2<sup>b</sup>                    | 6<sup>b</sup> | 8<sup>b</sup>                | 2011                 | Start-up      |
| 13     | 70                                       | 45                               | 1<sup>b</sup>                    | 3<sup>b</sup> | 8<sup>b</sup>                | 2011                 | Start-up      |
| 14     | 71                                       | 32                               | 2<sup>b</sup>                    | 3<sup>b</sup> | 10<sup>b</sup>               | 2011                 | Start-up      |
| 15     | 84                                       | 31                               | 1<sup>b</sup>                    | 2<sup>b</sup> | 6<sup>b</sup>                | 2011                 | Start-up      |

<sup>a</sup>Other included homecare case managers, social workers, pharmacists, spiritual and bereavement staff, and data support staff.

<sup>b</sup>Total number of staff involved (not FTE); most of these staff were working on the team part-time.
Table 3. Evolution framework by stage.

| Summary statement | Inception | Start-Up | Growth | Mature |
|-------------------|-----------|----------|--------|--------|
| **What’s happening** | “The community has a dream to fulfill” | “Not fully functional, still proof-of-concept” | “Figuring out how to expand offering and build capacity” | “Integrated into the community” |
| | Local champion has a vision for improving palliative care. Begins building necessary infrastructure, resources and relationships through grassroots efforts. | Founding members begin to build team and ways of collaborating. They define and market services offered to patients and providers. | Stable core team with established relationships works to expand reach. | Team is trusted and valued. Seen as hub for knowledge-sharing and expertise. |
| **Key opportunities** | | | | | |
| | • Develop effective ways of sharing info among team and broader ecosystem that is not dependent on common electronic medical record | • Make all core and extended members feel part of the team | • Deliver seamless continuity of care between settings | • Prevent team member burnout |
| | • Build rapport with community providers | • Build palliative care capacity among health service providers | • Continue to build capacity of community providers | • Manage team turnover |
| | • Build an understanding in community of the team’s role and how they can support providers | • Gain providers’ trust by working side by side to support and educate them | • Advocate for growth of palliative care network, ecosystem, and resources in community | • Maintain role clarity and integration with other provider partners |
| | • Outreach to upstream partners in the community, to encourage timely referrals and collaboration | • Form partnerships with community institutions and leverage opportunities to share resources (i.e. residential hospice) | • Manage changes in health service provider contracts and policy changes | • Manage changes in health service provider contracts and policy changes |
| | • Build on existing palliative care relationships/network in community | • Find/hire compatible team members to work in dynamic environment | • Prevent team member burnout | • Maintain continual sources of funding and resources |
| | • Build local intelligence for a better understanding of needs, gaps, assets and what will succeed/fail | • Avoid turf wars with local homecare nurses and other health service providers | • Manage team turnover | |
| | • Create foundational relationships for a strong core team | • Manage patient caseload with limited teams members | • Maintain role clarity and integration with other provider partners | |
| | | • Lack of home care nurses and/or physicians with palliative care expertise | • Manage changes in health service provider contracts and policy changes | |
| | | • Work toward providing 24/7 care | • Manage travel and serving large regions with limited headcount | |
| | | • Get referrals from physicians and hospitals | • Get more referrals, earlier in patient trajectory | |
| | | • Secure enough funding to sustain team through start-up | • Find additional funding and resources to support growth | |

(Continued)
Table 3. (Continued)

| Milestones | Inception | Start-Up | Growth | Mature |
|------------|-----------|----------|--------|--------|
| • Complete a needs assessment: Identify existing palliative care infrastructure, untapped capacity, what unmet needs can be served, and how to more effectively use resources | • Assemble core team of essential roles, including nurses and physicians | • Team has established local role and dependable partnerships | • Successfully building community’s health service provider palliative care capacity |
| • Articulate a shared vision of team and how to connect to community | • Practice the culture of a patient-centered mission and vision during decision making and problem solving | • Able to offer 24/7 care | • Health service providers understand when/how to best utilize and involve team |
| • Start to build buy-in from key stakeholders and providers in the ecosystem | • Establish team processes such as communication tools and methods | • Work toward a full suite of compatible roles (nurse, spiritual care, palliative care specialist, personal support workers) | • Effective feedback loop with patients and families |
| • Determine the offering to complement what exists — not compete | • Create open ongoing dialogue of pain points and successes | • Team serves more patients directly and indirectly | • Team helps strengthen entire palliative network in community |

Measurement: main focus and potential outputs/outcomes

**Environmental Scan**

**Perceived gaps in care:** service inaccessibility, service bottlenecks and inefficiencies, communication gaps

**Taking inventory:** existing infrastructure, potential alliances, and resources including expertise and funding opportunities

**Team and Program Formation**

**Formative (administrative) data:** caseload, patients seen, ED visits, urgent calls, home visits, doctor visits, bereavement contacts, professional consults, triage statistics, referral sources and destinations, place of death

**Tracking process progress:** Identify most responsible physician, orphaned patients connected with a doctor, patients’ end-of-life preferences (including place of death) documented, advanced care planning, ongoing patient monitoring, physicians providing 24/7 care

**Tracking relationship formation:** perceptions of “team” collaboration, conflict resolution, power equity, role clarity, communication

**Spreading Quality to more Patients**

**Formal capture of patient experience:**

Start to collect and assess patient and family experiences: care need being met, preferences (including place of death) recorded and met, caregiver burden, provider continuity

**Adverse incidences:** urgent calls not responded to in a timely manner

**Health System Measures System outcomes:** Acute care use, acute care death, end-of-life emergency department use and re-admission rates, palliative care admissions to hospital deemed inappropriate (alternate level of care beds)

**Remaining gaps in care:** Palliative care at patient diagnosis, length of time in team’s care before death

ED: emergency department.
Complete a needs assessment. Begin with existing teams and experts. Develop a small team of interested and committed experts with a common goal and identify the critical roles and responsibilities of team members. Networking and profiling the program as it was developed was a key to our success. (4–Team 11)

When starting a palliative care team you have to look at your own community and see who’s there and where the interest lies. Look into which physicians in your community are doing palliative care, and then think how can you support that physician trying to do this work so they aren’t doing it alone. That is how our team evolved. (1–Team 6)

One of the most crucial pieces of advice offered by the teams was to begin small and build relationships; proceed even if the plan is not perfect:

Start somewhere, don’t wait until the pieces of the puzzle fit together—don’t be afraid. Just start and learn from your mistakes and then change and adapt. Start somewhere, because this person is dying now and they won’t be here tomorrow. If we wait until all the stars are aligned we will miss helping all those people who die while we wait around. Has it been perfect? No, probably some of my most difficult cases have been our greatest learning opportunities. (1–Team 6)

Stage 2: start-up. The teams in the Start-up stage were not fully functional and still in a proof-of-concept phase. They were caring for some patients in the community but still trying to solidify the team’s staff complement and test different care processes to improve the efficiency of their practice, the quality of care for dying patients, and access to the service. Approaches to care at this point were neither standardized nor consistent, but the teams had developed a clear mission of what they wanted to achieve. They remained adaptive since not everything they tried would work. Building a small team of capable core providers and further advancing working relationships with generalist and specialized providers in the community were essential to progressing through this stage:

Then you just start and you gain experience and confidence by doing the work and working through and solving the problems. Make a path with the help of team support and tapping into some external consultants you can call as you need, and by working very closely with community groups like hospice volunteers, church volunteers, and neighbours. You will quickly gain confidence that you can actually do this work very well. (2–Team 1)

Informal communication internally and externally to the team was integral for building rapport, improving care continuity, and establishing the team’s presence in the greater community. Many of the teams tested processes for the transfer and discussion of patient information between providers, to determine which methods worked best. Some of these strategies, for example, chart-in-the-home, worked well for some teams but not others. Core providers tended to communicate with one another by phone, and also offered their number to external providers to call the team for assistance or patient referral. Through the teams’ offering to collaborate, they strove to spread a culture of patient-focused practice. This including reaching out to “upstream” partners in the community, to encourage earlier referrals to palliative care:

Relationship building has happened informally [through] encouraging the nurses to take initiative to call and discuss if they think something’s missing and we work together. We are able to communicate and deal with things quickly. (1–Team 14)

Importantly, at this stage, most teams had not built enough capacity to offer around-the-clock-access to palliative care. However, they had begun proactively addressing potential crises among their patients to reduce the need for after-hours care. Part of this plan was being explicit with patients about the support they could depend upon from the team and from other community services. The development of role clarity and open communication, to facilitate collaboration of care, were key tenants of this stage.

Stage 3: growth. Teams in the Growth stage were figuring out how to both expand the offering and build capacity in supporting generalist providers. The teams had established dependable partnerships, including those with community institutions, to share information, resources, and responsibility for addressing palliative care needs. The teams, in conjunction with their partners, were able to provide 24/7 care in the home and help transfer patients to appropriate, more intensive, care settings if required. Growth teams made concerted efforts to promote their service among local health professionals and the larger community:

Knowing that there is this 24/7 service in place has enticed some [primary care] physicians. We went out to different family practice groups and provided presentations and offered our consultants to work collaboratively with them. There is still a lot of resistance but we’ve capitalized on the physicians that have shown interest in trying to make referrals to them. (1–Team 11)

At this point, most of the teams had implemented formal interprofessional platforms such as case rounds, as opportunities to plan and learn together. The teams had a desire to prove their worth and value to patients, funders, and their community; as a result, they continually strived to improve the patient and family experience:

The community nurses know we are here, we meet every two weeks as a broader team for rounds, we also meet with hospital discharge planners and have collaborated with the
ER [emergency room] to avoid admissions and facilitate continuity of care. (1–Team 8)

The teams incorporated more formal quality improvement into their practice, eliciting patient feedback and reflection on adverse incidents. Team members spread their expertise and gained trust by supporting, working with, and educating a growing number of generalist providers in the community to build their capacity for palliative care:

We have heard of nurses approaching the [primary care] physician and telling them that they are solely responsible for their patients. Usually this scares them [the physician] and if given an out to transfer care they will. We approach it differently and try to meet the physician where they are at and as a result have a much better response rate of them remaining involved. We don’t give them an out but work with them. (3–Team 14)

In the Growth stage, the core and extended members of the team constituted a full suite of compatible roles (nursing, personal support, therapy, primary care, emotional support, spiritual care), toward serving more patients collectively and initiating palliative care earlier in patient trajectory. Challenges such as turf wars with other organizations, prevailing bureaucracy, and ensuring the sustainability of the team were still being resolved. The teams stayed the course by reminding themselves and others of the ultimate goal of providing quality palliative care experiences for patients. A non-punitive environment which celebrates wins but also views failures as learning opportunities was regarded as key to growing the program while avoiding members becoming discouraged:

You need to name the problem, or at least someone needs to have the courage to name it. You need to be professional, gracious and clear. Transparency and clarity on an issue is important for trust. These “elephants in the room,” in my experience, are never one-offs so you need to develop a safe space to hash out the common ground and common vision. (1–Team 4)

The focus needs to stay on what we’re trying to do; provide the best care for someone at end of life. If there is something that is a consistent problem, that keeps on coming up, we have to be able to say it. (6–Team 6)

**Stage 4: mature.** The teams considered at the Mature stage were fully integrated into the community. They were recognized as a center of excellence for knowledge-sharing, expertise, and patient care. The contributions of these teams were both trusted and valued, which granted them authority in advocating for their palliative care patients and system:

The team is well known in the community and is very well respected. It [the team] is promoted by a number of organizations that help to fundraise. The turf wars may have happened in the early days but it is now well know that everyone plays an important role in the care of palliative patients and their families. All providers work together and I do not think there is any overlap or redundancy. (3–Team 1)

The sustainability of these teams was secure in tapping continual sources of funding but also in adaptability to top-down policy changes. The team’s role was clearly understood, and as such, external health service providers knew when/how to best utilize and involve the team. Established channels of formal and informal communication existed within the team, with external providers, and with relevant community organizations, to facilitate continuity and comprehensiveness of care in an efficient manner:

We talk to each other daily, sometimes even on weekends. We relay information to one another so nothing falls through the cracks. We have a great system. (2–Team 5)

Generally, teams evolved their model of care over time. At start-up, they worked more in a take-over model, as they were learning how to work together to provide comprehensive palliative care. However, over time, they realized the community’s need was too great for their team alone, and began to deliberately focus on building capacity of other primary care physicians and nurses. Thus, the majority of the mature teams did not assume full responsibility for most of the palliative care patients referred to them (take over model), but rather supported the primary care provider to remain significantly involved (consultation model). Teams had increased palliative care capacity through training other providers to the extent that often only the more complicated patients required team involvement and some of these other providers had become capable mentors themselves:

My mantra is that, first and foremost, we [the team] are here to serve primary care. I am here to put myself out of business. The more we are in a relationship with a family physician, the less they need us [the team]. We help them grow their skills. We remain engaged in care but may do very little—we hover but don’t do ongoing care. (1–Team 4)

**Stage appropriate measurement**

Outcome measurement was important to all the teams. The newer teams felt pressure to show patient and system improvements and were eager to do so quickly, resulting in frustration. In contrast, the more established teams had come to realize that expecting great impact too early can set the initiative up for failure. Mature teams emphasized that measurement needs to start simple, such as counting the number of patients seen (Table 3). Informal mechanisms of immediate feedback were often utilized:

We know we’re doing a good job because of the feedback that we get from not only families, but from each other working together as a team. It is very important to have that feedback
from one another. Sometimes when we’re having a bad day or a bad week or we’re about to lose one of our precious patients, we need feedback from each other to know that we’ve done a good job. (5–Team 5)

At formation, most teams had a sense of the local state of palliative care, that is, present gaps and assets. This knowledge of the environment is critical to build upon existing capacity and align the service to the needs of the community. Once the team had started serving patients, the primary focus of measurement was understanding how and why the team is contributing to the care experiences of dying patients. Outcomes at this Start-up stage were more like outputs: tracking patients, including the number and characteristics of patients seen; process counts, such as number of patients with a most responsible physician identified; and monitoring and reflecting on the team climate. Measurement becomes more sophisticated at the Growth stage to formally capture the patient and caregiver experience of care and examine adverse incidents, using these findings to improve care quality. Finally, at the Mature stage, health system outcomes were measured as evidence of effectiveness, as well as, patterns of care data to improve earlier and more inclusive patient access to the team:

Now we know the team is doing a good job based on some big measures such as reducing emergency visits or reducing time in the hospital and so on. (4–Team 2)

In the beginning of team development, measurement was more purposeful than perfect. They made sure to celebrate small successes at each stage, to help them to stay motivated, build momentum, and grow their program. As the teams matured, measurement became more formalized, structured, and integrated into the broader system. There is no formal system of palliative care measurement in Ontario, thus the teams had to measure their own outcomes using local data. By partnering with the authors in a related study, the Mature stage teams demonstrated significant patient and system impact, including increased home death rates and reduced emergency department and hospital use.13 Teams also sustained an informal feedback loop with patients and families to continually fine tune the care process:

We follow up with the families afterward and they tell us things went as well as could be expected. If the opposite is true, we ask how things could have been improved, and thank them for the opportunity to learn. (1–Team 8)

Discussion

Key findings

Our qualitative examination of 15 specialized palliative care teams at various degrees of development revealed four main stages of evolution: Inception, Start-up, Growth, and Mature. With each consecutive stage, the teams demonstrated more capacity to advise and educate providers external to the team; offer around-the-clock access to care; impact patient and system outcomes; work in a collaborative, interprofessional model; engage primary and tertiary care, incorporate quality improvement and measurement into practice; and influence the local palliative care system. Even though the teams represent different models and staff complements we observed the same patterns in evolution. Prior frameworks have proposed the essential elements to building quality palliative care provision28–30 but have not suggested how these features emerge over time. To our knowledge, this is one of the first studies to report on an in-depth analysis of the natural progression of palliative care teams, based on a large number of diverse examples.

Implications for policy and practice

The findings from this study are relevant for the many countries that utilize home-based palliative care, such as visiting hospice service teams in the United States or palliative care community provider teams in Australia or the United Kingdom.27–30 Although community-based palliative care services have expanded, consistency in access and quality remain variable, meaning teams have room to improve over time.31–33 Regardless of how teams are funded in different countries, the steps of how teams evolve and which challenges arise over time likely hold true. Initiating the team programs examined was dependent on a few individuals with the vision to work with others to provide better palliative care in the community. Many of the team founders reported that they did not wait for an external catalyst or the perfect opportunity, plan, or resources, but rather perceived a local need and started with small changes; leveraging available capacity—becoming the catalyst themselves. In the early stages, the teams placed less priority on choosing the best measures and collecting data, than serving the needs of patients who needed support at that moment. While growing, team attention was on implementing and adjusting their service. Simple forms of quality improvement measures were incorporated into practice. As teams matured, measurement became more formalized and directed at more ambitious outcomes. Most of the teams had taken a long time to mature; this evolution was directly related to the activities partaken and measures of success used, along the way.

What this study adds

The stages of evolution we identified among the teams share commonalities with the phases of palliative care development described in Kelley’s10,11 Capacity-Building model and those in Ploeg et al.’s12 model of scaling of best practices in home care programs. All three models describe
an iterative, non-linear process and emphasize the critical role of leaders to drive forward the vision for change. The models are different yet complementary in that the Capacity-Building Model focuses on teams in the earlier stages of evolution, while the scaling model conceptualizes the spread of standardized practices in general home care. In comparison, our model covers the progression from newly formed to well-established teams, across different geographies, and predicated on the formation of interprofessional collaborative partnerships.

Study limitations

A limitation of our study is that all the teams were from one province in Canada, possibly restricting the global applicability of the findings. However, our study sample includes a large and diverse number of teams, who face system challenges common to many countries (e.g. shortage of expertise, fragmentation of services, uncoordinated communication, lack of standardized outcome measurement, etc.).4 Another limitation is that we interviewed relatively few providers external to the teams, who may have a different perspective of team evolution and impact.

Conclusion

We found that specialized palliative care teams go through distinct stages of evolution, each with unique sets of opportunities and challenges. Despite each team being different in initial and present structures for palliative care delivery, four stages of development were evident across the team histories. Team establishment was an iterative process, largely based on relationships. Time was required to build the long-term trust, communication channels, and authority necessary to render the teams successful and sustainable. The teams learned and adapted as they grew, refining ways of working together and serving patients and their families. Understanding the steps that are associated with the developmental stages of these teams can help to inform the progress of other community-based teams. This information also assists planners to better anticipate and set expectations for how teams may advance toward strengthening the palliative care system.

Acknowledgements

The authors gratefully acknowledge funding received for this study from the Ontario Ministry of Health and Long Term Care (Contract No. 2011-0006) and the Canadian Institutes of Health Research (Grants No. 115112 and No. 148997). Dr. Seow is supported by the Canada Research Chair program. We also wish to thank Ms. Deanna Bryant (MPA) and Ms. Sue Tan Toyofuku (MSc) for their assistance in data collection and analysis. Hsien Seow contributed toward the study design, data collection, data analysis, writing of draft manuscript, and review of draft and final manuscript. Daryl Bainbridge contributed toward study design, data collection, data analysis, writing of draft manuscript, and review of draft and final manuscript.

Data management and sharing

The confidential nature of the qualitative data prohibits them from being released publicly.

Declaration of conflicting interests

The author(s) declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

Funding

The author(s) disclosed receipt of the following financial support for the research, authorship, and/or publication of this article: Funding was received for this study from the Ontario Ministry of Health and Long Term Care (Contract No. 2011-0006) and the Canadian Institutes of Health Research (Grants No. 115112 and No. 148997).

Research ethics and patient consent

The study was approved by Hamilton Health Sciences/McMaster University research ethics review board, Ontario, Canada. Written informed consent was obtained from each participant.

ORCID iD

Daryl Bainbridge https://orcid.org/0000-0001-9249-8229

References

1. Morin L, Aubry R, Frova L, et al. Estimating the need for palliative care at the population level: a cross-national study in 12 countries. Palliat Med 2017; 31(6): 526–536.
2. Bainbridge D, Giruparajah M, Zou H, et al. The care experiences of patients who die in residential hospice: a qualitative analysis of the last three months of life from the views of bereaved caregivers. Palliat Support Care 2017. DOI: 10.1017/S147895151700058X.
3. Gomes B, Calanzani N, Curiale V, et al. Effectiveness and cost-effectiveness of home palliative care services for adults with advanced illness and their caregivers. Cochrane Database Syst Rev 2013; 134(1): 93–94.
4. Gomes B, Calanzani N, Gysels M, et al. Heterogeneity and changes in preferences for dying at home: a systematic review. BMC Palliat Care 2013; 12: 7.
5. Higginson IJ and Evans CJ. What is the evidence that palliative care teams improve outcomes for cancer patients and their families? Cancer J 2010; 16(5): 423–435.
6. Bainbridge D, Seow H and Sussman J. Common components of efficacious in-home end-of-life care programs: a review of systematic reviews. J Am Geriatr Soc 2016; 64(3): 632–639.
7. Hearn J and Higginson IJ. Do specialist palliative care teams improve outcomes for cancer patients? A systematic literature review. Palliat Med 1998; 12(5): 317–332.
8. Higginson IJ, Finlay IG, Goodwin DM, et al. Is there evidence that palliative care teams alter end-of-life experiences...
of patients and their caregivers? J Pain Symptom Manage 2003; 25(2): 150–168.

9. Bhavsar NA, Bloom K, Nicolla J, et al. Delivery of community-based palliative care: findings from a time and motion study. J Palliat Med 2017; 20(10): 1120–1126.

10. Kelley ML, Williams A, DeMiglio L, et al. Developing rural palliative care: validating a conceptual model. Rural Remote Health 2011; 11(2): 1717.

11. Kelley ML. A framework to guide policy and program development for palliative care in First Nations communities. Thunder Bay, ON, Canada: Centre for Education and Research on Aging & Health, Lakehead University, 2015, http://olh.lakeheadu.ca/wp-content/uploads/2015/01/Framework_to_Guide_Policy_and_Program_Development_for_PC_in_FN_Communities_January_16_FINAL.pdf (accessed 4 April 2018).

12. Ploeg J, Markle-Reid M, Davies B, et al. Spreading and sustaining best practices for home care of older adults: a grounded theory study. Implement Sci 2014; 9: 162.

13. Seow H, Brazil K, Sussman J, et al. Impact of community based, specialist palliative care teams on hospitalisations and emergency department visits late in life and hospital deaths: a pooled analysis. BMJ 2014; 348: g3496.

14. Bainbridge D and Seow H. Palliative care experience in the last 3 months of life: a quantitative comparison of care provided in residential hospices, hospitals, and the home from the perspectives of bereaved caregivers. Am J Hosp Palliat Care 2017; 35(3): 456–463.

15. Ontario Palliative Care Network (OPCN). OPCN inaugural report 2016–17. Toronto, ON, Canada: OPCN, 2017, https://www.ontariopalliativecarenetwork.ca/sites/opcn/files/Inaugural_Report_ENGLISH.pdf (accessed 23 April 2018).

16. Pereira J, Seow H, Marshall D, et al. Consultation versus shared care roles in palliative care service delivery: deriving an empirically-based framework. In: Proceedings of the 13th world congress of the European Association for Palliative Care (EAPC), 2013, http://www.eapcnet.eu/LinkClick.aspx?fileticket=N_zLjCaQpVo= (accessed 4 April 2018).

17. Seow H, Bainbridge D, Brouwers M, et al. Common care practices among effective community-based specialist palliative care teams: a qualitative study. BMJ Support Palliat Care 2017. DOI: 10.1136/bmjspcare-2016-001221.

18. Quill TE and Abernethy AP. Generalist plus specialist palliative care — creating a more sustainable model. N Engl J Med 2013; 368(13): 1173–1175.

19. Bainbridge D, Brazil K, Krueger P, et al. A proposed systems approach to the evaluation of integrated palliative care. BMC Palliat Care 2010; 9: 8.

20. Borrill C, Carletta J, Carter A, et al. The effectiveness of health care teams in the National Health Service. Birmingham: University of Aston, 2001, http://ctraining.co.uk/documents/TheEffectivenessofHealthCareTeamsintheNHS_004.pdf (accessed 4 April 2018).

21. Sussman J, Barbara L, Bainbridge D, et al. Health system characteristics of quality care delivery: a comparative case study evaluation of palliative care for cancer patients in four regions in Ontario, Canada. Palliat Med 2012; 26(4): 322–335.

22. Corbin J, Strauss A and Strauss AL. Basics of qualitative research. Thousand Oaks, CA: SAGE, 2014.

23. Creswell J. Qualitative inquiry and research design: choosing among five traditions. 3rd ed. Thousand Oaks, CA: SAGE, 2013.

24. Ferrell B, Connor SR, Cordes A, et al. The national agenda for quality palliative care: the National Consensus Project and the National Quality Forum. J Pain Symptom Manage 2007; 33(6): 737–744.

25. Ferris FD, Gomez-Batiste X, Furst CJ, et al. Implementing quality palliative care. J Pain Symptom Manage 2007; 33(5): 533–541.

26. Luckett T, Phillips J, Agar M, et al. Elements of effective palliative care models: a rapid review. BMC Health Serv Res 2014; 14: 136.

27. Connor SR. Development of hospice and palliative care in the United States. OMEGA 2008; 56(1): 89–99.

28. Department of Health. End of life care strategy. Promoting high quality care for all adults at the end of life. London: Department of Health, 2008, http://www.cpa.org.uk/cpa/End_of_Life_Care_Strategy.pdf (accessed 4 April 2018).

29. Genet N, Boerma WG, Kringos DS, et al. Home care in Europe: a systematic literature review. BMC Health Serv Res 2011; 11(1): 207.

30. National EOL Framework Forum. Health system reform and care at the end of life: a guidance document. Canberra, ACT, Australia: Palliative Care Australia, 2010, http://palliativecare.org.au/wp-content/uploads/dlmUploads/2015/08/Health-system-reform-guidance-document-web-version.pdf (accessed 4 April 2018).

31. Hughes MT and Smith TJ. The growth of palliative care in the United States. Annu Rev Public Health 2014; 35: 459–475.

32. Luckett T, Davidson PM, Lam L, et al. Do community specialist palliative care services that provide home nursing increase rates of home death for people with life-limiting illnesses? A systematic review and meta-analysis of comparative studies. J Pain Symptom Manage 2013; 45(2): 279–297.

33. Sayma M, Saleh D, Kerwat DA, et al. A qualitative inquiry into the barriers and facilitators to achieving home death. BMJ Support Palliat Care 2017. DOI: 10.1136/bmjspcare-2016-001260.

34. Aldridge MD, Hasselaar J, Garralda E, et al. Education, implementation, and policy barriers to greater integration of palliative care: a literature review. Palliat Med 2016; 30(3): 224–239.

Appendix I

Indicative script for interviews

1. Tell us about yourself.

2. How do you get a specialized palliative care team started?

3. When you first joined the team can you tell us what it was like then? How are things different now?

4. Map out the milestones or changes the team encountered from inception to present (activity).

5. What were the highs and lows along the way? Can you tell us what made these points in time good/bad or challenging/successful?
6. How did the team evolve into what we see now?
7. What are some of the major lessons you or your team has learned that helped you operate and deliver better care?
8. Are there other key milestones or inflection points that you know off to help us understand the evolution of the team?
9. We often hear from teams that they don’t know where to begin without “boiling the ocean.” So how do you start without boiling the ocean?
10. Can you give examples of how you used existing resources differently, instead of relying on new money?
11. How do you sustain the program in the face of shifting priorities at the regional and provincial level?
12. Although the system may be broken, can you give examples about how your team makes it “work” despite system barriers?
13. How do you promote the team in the community?
14. How do you achieve role clarification and avoid turf wars among all organizations in hospice palliative care?
15. When there was an “elephant” in the room, how did you start talking about it and resolve it?
16. How do you foster relationships as well as the qualities or trust and mutual respect?
17. What have you learned about in terms of how to get buy-in from primary care physicians?
18. When trying to improve care delivery how do you move beyond naming, blaming, and shaming?
19. How do you know that your team is doing a good job?
20. What kind of things did your team do to improve care over time?
21. Where do you look to for inspiration for new ideas or ways of improving?
22. What piece of advice would you give to developing teams?
23. If the healthcare system wants to give more patients and families access to care teams like the one you work in, what advice do you have for expanding this model?
24. Is there anything else you would like to tell us about your team?