Public Perceptions of Advance Care Planning, Palliative Care, and Hospice: A Scoping Review

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Abstract

Background: Although access to advance care planning (ACP), palliative care, and hospice has increased, public attitudes may still be barriers to their optimal use.

Purpose: To synthesize empirical research from disparate sources that describes public perceptions of ACP, palliative care, and hospice in ways that could inform public messaging.

Data Sources: Searches of PubMed and other databases were made from January 2011 to January 2020.

Study Selection: Studies reporting survey or interview data with the public that asked specifically about awareness and attitudes toward ACP, palliative care, or hospice were included.

Data Extraction and Synthesis: Two reviewers independently screened citations, read full texts, and performed data abstraction. Twelve studies met inclusion criteria and included >9800 participants. For ACP, 80% to 90% of participants reported awareness, and a similar proportion considered it important, but only 10% to 41% reported having named a proxy or completed a written document. For palliative care, 66% to 71% of participants reported no awareness of palliative care, and those who reported awareness often conflated it with end-of-life care. However, after being prompted with a tested definition, 95% rated palliative care favorably. For hospice, 86% of participants reported awareness and 70% to 91% rated it favorably, although 37% held significant misconceptions.

Limitations: A limited number of studies met inclusion criteria, and some were published in nonpeer reviewed sources. The studies reflect public perceptions pre-COVID-19.

Conclusion: Consumer perceptions of ACP, palliative care, and hospice each have a distinct profile of awareness, perceptions of importance, and reports of action taking, and these profiles represent three different challenges for public messaging.

Keywords: advance care planning; hospice; palliative care; perceptions; public

Introduction

Over the past 20 years, access and infrastructure for advance care planning (ACP), palliative care, and hospice have grown considerably.1–3 However, negative public attitudes toward these services are likely barriers to optimal use. In 2009, vice-presidential candidate Sarah Palin stated inaccurately that legislation for the Affordable Care Act would create “death panels,” and a similar proportion considered it important, but only 10% to 41% reported having named a proxy or completed a written document. For palliative care, 66% to 71% of participants reported no awareness of palliative care, and those who reported awareness often conflated it with end-of-life care. However, after being prompted with a tested definition, 95% rated palliative care favorably. For hospice, 86% of participants reported awareness and 70% to 91% rated it favorably, although 37% held significant misconceptions.

Limitations: A limited number of studies met inclusion criteria, and some were published in nonpeer reviewed sources. The studies reflect public perceptions pre-COVID-19.

Conclusion: Consumer perceptions of ACP, palliative care, and hospice each have a distinct profile of awareness, perceptions of importance, and reports of action taking, and these profiles represent three different challenges for public messaging.

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principles for health communication initiatives: (1) identify the targeted behavior, (2) develop an effective strategy for exposing people to the message, (3) take a comprehensive multichannel media approach, and (4) express the message in repeated consistent presentations. A single message may be expressed in a variety of formats, taglines, headlines, and sound bites. Successful models for population-based messaging approaches can be seen in campaigns to change public attitudes and health behaviors such as smoking cessation and HIV awareness. Those successful campaigns were grounded in empirical research, developed by messaging experts, tested with their specific public target audiences, and used in a coordinated and consistent way that sets the stage for more extensive conversations between clinicians and patients.

For serious illness care, public messaging that introduces ACP, palliative care, and hospice could increase demand for these services and lead to increased access and quality. But to create large scale culture change, messaging needs to work at the public level. The most effective messaging speaks of the needs of the audience who will need these services in the future—not to patients already familiar with these topics. To put it bluntly, improving public engagement with ACP, palliative care, and hospice needs to address existing perceptions held by the lay public (which in this review are referred to as the “public”) rather than the convictions and beliefs of patients who have already experienced the benefits of these services or dedicated clinicians who work with those patients. In addition, public messaging that introduces these services to the public should differ from the skilled communication that clinicians perform at the bedside of patients with a serious illness.

To inform future messaging to create more public engagement in ACP, palliative care, and hospice, we sought to analyze the empirical public research conducted on these services over the past decade. Our initial searches in medical databases, conducted in July 2019, suggested that the evidence base was too small to justify a systematic review. We felt that a more useful approach would be to describe the existing body of evidence in a way that could inform future research. Thus, we conducted a scoping review to synthesize data from disparate sources, including data not published in medical literature, to address a broad question, and address gaps in the literature. The question we chose for this scoping review was, “What evidence exists to describe how members of the public perceive ACP, palliative care, and hospice in ways that could inform public messaging that introduces the public to these services?” We completed this review in January 2020, before the full extent of the COVID-19 pandemic was widely appreciated.

Methods

Data sources and searches

We followed the Preferred Reporting Items for Systematic reviews and Meta-Analyses extension for Scoping Reviews (PRISMA-ScR) guidelines and the scoping review methods manual developed by the Joanna Briggs Institute. We searched the following databases: PubMed, Communication & Mass Media Complete, Rasmussen, and iPoll. The search strategy was designed to capture citations and reports from the search terms (hcpise OR palliative OR advance care) and (public awareness OR social marketing OR public opinion) from the time period January 2011 through January 2020. All searches and selection of search terms were performed by an experienced research librarian (N.S.D.). In addition, we sought unindexed reports known to experts in the field.

Study selection

We included studies that reported U.S. public perceptions about ACP, palliative care, or hospice. We excluded studies that asked patients with a serious illness to describe, report, or rate their communication with clinicians or indicate their attitudes toward ACP, palliative care, or hospice, because our objective was to synthesize evidence that could inform messaging that introduces the lay public to these services. Two investigators reviewed all search results, read all candidate reports, and agreed on the final selection of included studies (Fig. 1).

Data extraction

We designed a data charting form that two investigators (M.S.G. and A.L.B.) populated, iterated, and updated. Extracted variables included date of report, study methodology, population characteristics, and key outcomes. We did not confirm abstracted data with authors of the individual studies.

Data synthesis

We grouped study data by ACP, palliative care, or hospice because our early analyses indicated that the public has distinct views about each of these services.

Role of funding sources

The funding sources were not involved in the development of the research question, selection of search strategies, data extraction or analysis, or reporting of results.

Results

Our initial searches resulted in 34,208 citations of which 32 met our criteria to be screened and 12 met criteria for inclusion. The selection process is depicted in Figure 1, and the 12 included studies are summarized in Table 1.

Study methodologies and publication

The 12 studies spanned a variety of methodologies: 1 systematic review that addressed public perceptions, cross-sectional surveys, and surveys that also included focus groups. Of these 12 studies, 7 were published in peer-reviewed journals, and 5 were published as public reports by their funding organizations.

Study populations and settings

Most included studies involved a population of adults, mostly older with age >60 years, outside of health care settings. The total number of participants involved in studies of ACP, with the exclusion of the systematic review, was >3800. The number of participants in each ACP study ranged from 346 to 2500. The total number of participants involved in studies of palliative care was >5000, and ranged from 350 to 3504 per study. The total number of participants involved in studies of hospice was >1100, and ranged from 148 to 800 per study (Table 1).
Measurements and outcomes

The most common measurements involved awareness of a particular aspect of serious illness care, ratings of importance or favorability, participant misconceptions, and self-reports of direct engagement. The measurements included in these studies were heterogeneous, limiting comparisons that could be made across studies and topics. See Table 2 for key reported elements.

Perceptions of ACP

Four studies examining public awareness and perceptions about ACP show that a large majority (80–90%) of people are aware of ACP (described in two surveys as “end-of-life planning”)

and a similar proportion (80–90%) think it is important. However, taking action is a different matter. A large body of evidence shows that only a minority of the public report having named a health care proxy decision maker (10–41%) or completing a written directive (23–32%).

The gap between high awareness and low action taking for ACP is best explained in a 2019 survey that also involved 150 participants recruited into an online community. That study found five different subgroups of consumers based on their attitudes toward ACP. Two of these segments, representing a combined 34% of the participants, had taken action in ACP, and their reasons for doing so included past experiences—often negative instances of being asked to make decisions for a family member who had not left any indication of their wishes for end-of-life care. The remaining segments, representing a combined 66% of the participants, had not taken action and did not wish to, and their reasons for declining to act included mistrust of the health system that their wishes would be followed, confidence that those who mattered to them already knew what they wanted, and inability to act because of other life stresses.

Perceptions of palliative care

Six studies examining public awareness about palliative care were included (three of which were different analyses of the same dataset conducted by different investigators), and all demonstrate that the public are unfamiliar with palliative care. Over 70% of participants could not even rate palliative care for favorability. In addition, the small proportion of participants who felt familiar with palliative care held perceptions that were mostly inaccurate. For example, 59% of those who said they were familiar with palliative care thought it was the same as hospice or end-of-life care. However, in two studies, when participants received a definition of palliative care that emphasized the benefits and appropriateness at any stage of illness, virtually all (95%) rated it favorably.

Perceptions of hospice

Public awareness of hospice is high, at 83% to 85%, and public favorability is also high, at 71% to 90%. Hospice awareness and favorability are higher among participants with more education or personal experience. However, a significant minority of participants, 37%, had misconceptions about hospice, including beliefs that choosing hospice was tantamount to “giving up.”

Factors influencing public perceptions

These studies showed consistently that older white females with higher education and socioeconomic status had higher awareness and were more inclined to take action for ACP and hospice. Personal experience with a family member that was positive, trust in the health system, presence of a strong relationship with a primary care or other clinician also correlated with higher awareness.

For other ethnic and socioeconomic groups, lower awareness and action taking were correlated with barriers about discussing serious illness, distrust of documents, and mistrust in the health care system. Political party affiliation did not influence views of palliative care or hospice in the one study that included that information.

Gaps in existing research

There were no studies addressing awareness or perceptions of specific ethnic or cultural groups, and no studies with a
### Table 1. Overview of 12 Studies That Met Inclusion Criteria

<table>
<thead>
<tr>
<th>Title</th>
<th>Date published</th>
<th>Methods/study population</th>
<th>Key findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Perceptions About Hospice from a Community-Based Pilot Study</td>
<td>January 4, 2011</td>
<td>Survey of 148 adults from 8 churches in eastern Ohio</td>
<td>Respondents had favorable opinions about hospice but older tended to have more negative impressions than younger, with older more likely to see it as “giving up.”</td>
</tr>
<tr>
<td>Center to Advance Palliative Care 2011 PC Consumer and Physician Research</td>
<td>April 11, 2011</td>
<td>National survey 800 adults age &gt;25 years, focus groups with physicians</td>
<td>In total, 8% were knowledgeable about palliative care, up to 50% were unable to rate opinion of palliative care. After hearing definition, 94% would likely consider.</td>
</tr>
<tr>
<td>Knowledge About Hospice: Exploring Misconceptions, Attitudes, and Preferences for Care</td>
<td>August 18, 2014</td>
<td>Cross-sectional survey of 123 U.S. adults with minority oversampling</td>
<td>In total, 86% had heard about hospice, 54% had personal experience with it. In total, 62% did not know that hospice cannot provide concurrent cure-oriented care. Greater knowledge of hospice is associated with more favorable attitudes about hospice philosophy of care and greater preference for hospice.</td>
</tr>
<tr>
<td>Massachusetts Baseline 2016 Advance Care Planning Survey</td>
<td>March 8, 2016</td>
<td>Survey of 860 adults age &gt;18 years in Massachusetts</td>
<td>In total, 96% believe EOL planning is important to have conversations with loved ones and health care providers about wishes for care. Nearly 90% indicated that conversation with their loved ones was somewhat or very helpful.</td>
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<tr>
<td>Massachusetts 2017 ACP Survey</td>
<td>February 22, 2017</td>
<td>Survey of 346 adults age &gt;18 years in Massachusetts</td>
<td>In total, &gt;80% believed doctors should discuss EOL care issues with patients, but few participated in a discussion. Older participants and those with lower household incomes are more likely to have spoken to a health care provider about wishes. In total, &gt;33% did not want to have a conversation about EOL care wishes with a doctor or other health care providers. In total, &lt;50% had completed a health care proxy and named an agent.</td>
</tr>
<tr>
<td>Awareness and Misperceptions of Hospice and PC: A Population-Based Survey Study</td>
<td>July 20, 2017</td>
<td>Cross-sectional study of 800 New York State residents &gt;18 years who participated in the 2016 Empire State Poll</td>
<td>In total, 83% were able to provide a definition of hospice care and 27% that of palliative care. In total, 37% had misperception about hospice and 53% had misperception about palliative care. Of those who defined palliative care, 9% reported it was applicable in any course of an illness.</td>
</tr>
<tr>
<td>Awareness of PC Among a Nationally Representative Sample of U.S. Adults</td>
<td>March 28, 2019</td>
<td>In total, 3445 participants from the 2018 Health Information National Trends Survey</td>
<td>In total, 71% had never heard of palliative care. Older individuals, those with higher educational attainment, women, and whites (vs. nonwhites) had greater odds of palliative care awareness. Five consumer attitudinal segments: anxious action takers 10%, self-assured action takers 24%, disengaged worriers 34%, defiant independents 18%, and self-reliant skeptics 14%. Patients and carers preferred health professionals to initiate discussions and relationships with professionals particularly important. Mixed feelings about the best timing, with many preferring to defer discussions until they perceived them to be clinically relevant. ACP was felt to bring benefits including a greater sense of peace and less worry, but it could also be disruptive and distressing.</td>
</tr>
<tr>
<td>Massachusetts Serious Illness Coalition Attitudinal Research</td>
<td>May 16, 2019</td>
<td>Survey of national sample 2500 adults age &gt;18 years</td>
<td></td>
</tr>
<tr>
<td>How Should EOL/ACP Discussions Be Implemented According to Patients and Informal Carers</td>
<td>August 1, 2019</td>
<td>Systematic review of reviews: 55 literature reviews published between 2007 and 2018</td>
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target population of the uninsured. Although we excluded studies of patients with serious illness, the level of participant experience with prior serious illness care or caregiving was not reported.

Discussion

The purpose of this scoping review, conducted well before COVID-19 was declared a pandemic, was to synthesize evidence that addressed the question: “What evidence exists to introduce the public to these services?” The 12 studies that met our inclusion criteria included >9800 participants, mostly >60 years of age, outside of health care settings. Taken together these studies indicate that public perceptions of ACP, palliative care, and hospice each have a distinct profile of awareness, perceptions of importance, and reports of action taking. For ACP, 80% to 90% of participants report awareness of the need for end-of-life planning, and a similar proportion consider it important, but only 10% to 41% report having named a proxy or completed a written document. For palliative care, 66% to 71% of participants reported no awareness, and those who reported awareness of palliative care often conflated it with end-of-life care. For hospice, >80% of participants reported awareness and 70% to 91% rated it favorably, although a minority had serious reservations.

What does all this mean? Our pre-COVID-19 interpretation was that these distinct profiles indicate that ACP, palliative care, and hospice each face their own messaging challenge. The challenge facing messaging about ACP could be summarized as “high awareness/low action.” Most of the public is aware of ACP, and think it is important, but a minority have taken action. The challenge facing messaging for palliative care is “low awareness/common misconceptions.” Few among the public are aware of palliative care, and of those who are aware, most hold an incorrect view that palliative care is about dying. The challenge facing messaging about hospice is “high awareness/significant reservations.” Most of the public are aware of hospice, but a significant minority have had negative personal experiences with hospice or hold incorrect views about what hospice provides.

What ACP, palliative care, and hospice have in common in these pre-COVID-19 data, however, are that consumer misperceptions, inaccuracies, and conflation of one for the other are almost routine. The public confuses ACP with end-of-life care, palliative care with hospice, and hospice with hastening death. The evidence we reviewed did not characterize how the public feels about confronting death or dying, but a growing body of evidence collected from patients living with a serious illness indicates that a majority are not interested in talking about dying, at least early in their illness. If the public holds a similar reluctance toward talk about dying, the biggest messaging challenge of all might be for a field that has traditionally promised to provide “a good death” to find new ways to talk to the public. We hasten to add, though, that improving public messaging does not require that clinicians back away from cultivating prognostic awareness or talking

### Table 1. (Continued)

<table>
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<th>Key findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patterns of PC Beliefs Among U.S. Adults</td>
<td>August 1, 2019</td>
<td>In total, 3504 participants from the 2018 Health Information National Trends Survey</td>
<td>In total, 43% automatically thought of death when thinking about PC, 32% equate PC with hospice, 15% believed PC means giving up, 15% believed PC requires discontinuation of other treatments, and 40% were unable to rate palliative care for favorability. Patients and caregivers were more able to rate palliative care but rated it slightly lower than general public. After hearing tested definition of PC, 86% to 90% would likely consider.</td>
</tr>
<tr>
<td>CAPC 2019 PC Consumer and Physician Research</td>
<td>August 8, 2019</td>
<td>National survey 350 adults age &gt;25 years, online with 250 patients and carers, 317 physicians</td>
<td>In total, 34% of participants self-reported having at least some knowledge of PC, but of those, only 41% were able to answer all three basic PC questions correctly.</td>
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<tr>
<td>When Patients Say They Know About PC, How Much Do They Really Understand?</td>
<td>November 1, 2019</td>
<td>In total, 3504 participants from the 2018 Health Information National Trends Survey</td>
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</table>

ACP, advance care planning; EOL, end-of-life; PC, palliative care.

### Table 2. Comparison of Public Perception Findings by Type of Serious Illness Care

<table>
<thead>
<tr>
<th>Types of misimpressions</th>
<th>Advance care planning</th>
<th>Palliative care</th>
<th>Hospice</th>
</tr>
</thead>
<tbody>
<tr>
<td>EOL planning, withdrawing life support</td>
<td>80–90%</td>
<td>29–34%</td>
<td>86%</td>
</tr>
<tr>
<td>Giving up</td>
<td>80–90%</td>
<td>NA</td>
<td>70–91%</td>
</tr>
<tr>
<td>Types of misimpressions</td>
<td></td>
<td></td>
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<tr>
<td>Action taken</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Having a health care agent/proxy, or documents, or enrolling in programs</td>
<td>34%</td>
<td>NA</td>
<td>48%b</td>
</tr>
</tbody>
</table>
| These data were not from an included study but represent Medicare data and are included for comparison. NA, not applicable.
about dying in appropriate circumstances. Public messaging has a different purpose than patient–clinician communication.

This scoping review did not evaluate the effect of messaging strategies that could overcome the challenges already described. However, these challenges suggest that one approach may be for individual public campaigns to address a specific population-based goal for ACP or palliative care or hospice—and for public campaigns to treat each service separately rather than trying to show how they overlap or, at times, serve similar functions. For example, a goal for an ACP campaign could be a message that targets concerning consumer assumptions that reduce willingness to take action. Similarly, a goal for a palliative care public campaign could be a message that quality of life is better at any stage of illness if palliative care is accessed. Finally, a goal for a hospice public campaign could stress dependability, trust, and care in the setting best for that patient. (Note that consumer associations of hospice with end of life are so strong that raising death in an introductory public message is unnecessary, and possibly alienating to those reluctant to think about dying.)

We acknowledge that these suggestions about messaging raise questions. For example, a commonly used graphic stresses continuity between palliative care and hospice, and has been useful in defining palliative care.25 This graphic was designed for clinicians and policy makers, and was never tested for clarity with the public. We also acknowledge that ACP has been defined as supporting adults “at any age or stage of health.”26 The international consensus process that produced this definition was not designed for public messaging. Finally, variability in service availability and eligibility criteria across the United States may affect people’s perception of these services, which even effective messaging cannot overcome. These issues do not, however, preclude innovation in public messaging. Successful public campaigns point toward a single behavior, and they do not attempt to fill in all the details.

The evidence base that we were able to assemble for this scoping review was limited, so some caution should be given to this analysis. A little more than a third of this evidence was not published in the medical literature and was not peer reviewed. The cross-sectional surveys involved telephone interviewing, which can introduce biases. The evidence describing differences by ethnic, socioeconomic, or generational factors is limited. Finally, the effect of the COVID-19 pandemic may well change public perceptions in ways that cannot yet be measured.

Nonetheless, these findings provide important insights into challenges in the pre-COVID-19 era to effective messaging and public campaigns for ACP, palliative care, and hospice. Particularly notable are the two studies on public awareness about palliative care done eight years apart that demonstrate no improvement whatsoever in public awareness about palliative care. How the public experiences serious illness care in the age of COVID-19 could shape a whole new set of public perceptions.

In conclusion, this scoping review indicates that pre-COVID-19, ACP, palliative care, and hospice had distinct profiles of public awareness, ratings of importance, and action taking. The results underscore the importance of empirical research that measures public perceptions before the development of public messaging, and suggest that further research may be needed once the COVID-19 pandemic has stabilized.

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References


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