Medical oncologist perspectives on palliative care reveal physician-centered barriers to early integration

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Introduction

Early palliative care referral for patients with advanced cancer is associated with better quality of life, improved mood, and longer survival as compared to standard oncologic care (1-4). In addition, oncologists report that early and routine integration of palliative care for their patients with advanced cancer is acceptable, (5) and the American Society of Clinical Oncology (ASCO) definitively endorses early palliative care for all patients.
with advanced cancer (6). Despite strong clinical evidence and endorsement, early engagement with palliative care specialists occurs in fewer than 40% of patients with incurable cancer (7,8). It has been demonstrated that one reason for the low utilization rate of early palliative care is a perception from patients and caregivers that palliative care equates to end of life care and a terminal prognosis, and therefore there is a resistance to early palliative care integration from patients (9). As such, some have called for a “rebranding” of palliative care to highlight the benefits outside of end-of-life care domains that patients can expect from these interactions (10). Nonetheless, efforts to increase early palliative care referral for patients with advanced cancer are needed.

The objective of this study was to uncover important barriers to palliative care and end of life care planning from the perspective of medical oncologists. We hypothesized that medical oncologists would identify system-level barriers to palliative care referral [e.g., shortage of palliative care medicine specialists (11,12)] and would endorse interventions that increase collaboration among palliative care specialists, other oncologists, and other members of the interprofessional team to improve end-of-life care for patients with advanced cancer.

Methods

Participants

A sample of U.S. medical oncologists (N=31) was recruited through purposive, snowball sampling (13). Oncologists were recruited to participate in the study and were asked to refer other medical oncologists to the study investigators for recruitment. Physician study group participants were informed in the recruitment process that their responses were to be recorded, that their identities masked from the research team, and the transcripts were intended for research purposes. The study was reviewed and approved by the Northwestern University Institutional Review Board (STU00204433), and determined not to be human subjects research; hence informed consent was deemed unnecessary.

Focus groups

Four professionally-moderated focus groups were conducted during the ASCO annual meeting in Chicago, Illinois. Focus group moderators were non-physician qualitative researchers from the Northwestern Feinberg School of Medicine. Focus groups began with questions to elicit participants’ experiences with end-of-life care and palliative care in their own clinical practices. Next, moderators prompted participants to discuss the optimal timing of palliative care referral, barriers to palliative care referral, and the role of other clinical team members in end-of-life care. Portions pertaining in particular to radiation oncologist participation have been previously published (14).

Analysis

Each focus group was audio-recorded and transcribed verbatim, and identifying information was removed. We used NVIVO 10 software (QSR International, Melbourne, Australia) to organize and facilitate the analysis. Transcripts were analyzed using qualitative content analysis with an inductive approach. Four investigators from diverse healthcare backgrounds (SMK, medical oncology; TJK, radiation oncology; JMK, critical care medicine; MM, public health) independently reviewed each transcript, assigning a descriptive code to relevant sections of text. After the initial review of all focus group transcripts, the four study investigators met to review and collate preliminary codes into a coding taxonomy. The investigators then reviewed all four transcripts using the coding taxonomy and each coded section of text was assigned a consensus code during regular meetings of at least three investigators. The coding taxonomy was iteratively revised throughout the coding process. Higher-level analysis to evaluate relationships between codes and develop central themes was conducted through regular meetings of at least three investigators.

Results

Demographic characteristics of the 31 participating medical oncologists are shown in Table 1. The majority of participants practiced in academic settings (29/31, 94%) and in Midwestern states (22/31, 71%). Our analysis identified common, typical practice patterns for ACP among medical oncologists’ and two major types of barriers to regular adoption of early palliative care referral for patients with advanced cancer: patient-centered barriers and medical oncologist-centered barriers.

Typical practice patterns for end of life care

First, medical oncologists described meeting the patient and outlining the general intent of treatment (curative versus...
palliative). Medical oncologists universally endorsed the importance of early communication and clarification about the intent of chemotherapy or other cancer-directed therapy (palliative versus curative treatment) with patients and their families. For example, one oncologist said, “With the first visit I always set the expectation that either this is palliative or curative and I explain what those things mean”.

Next, oncologists typically recommend therapy (and subsequent therapy) with integrated discussions of end of life care. Most oncologists described integrating discussions about prognosis and end-of-life planning “within the first few visits” for incurable patients, but maintained this practice pattern as separate from palliative care referral. A representative oncologist stated “We start usually by the second or third visit—I start talking about discussing with your family what you would want... I don’t want it to be a surprise once we’ve finished (fourth line) chemotherapy and I say I don’t have anything else. That’s when the transition would happen, but my goal is that they’re not surprised.”

The final step is to integrate palliative care and “transition” to hospice once available treatments have failed—“... but then the really intense discussion about what the end of life entails and really getting the palliative care team involved and all that happens when a patient really gets symptomatic.”

System and patient-centered barriers to palliative care referral

Medical oncologists rarely described system level barriers to palliative care referral, such as specialist shortage. However, one participant did note, “I don’t send all my patients to palliative care and I think you can only send, you know, however many the palliative care physicians can see.” Patient-centered barriers to early palliative care referral were reported more often, related to logistical burdens placed on patients with advanced cancer who are referred to a palliative care specialist. For example, one medical oncologist noted the burden of parking when coming to see a palliative care physician, in addition to all the other medical visits:

- “I know there’s this study out there about early palliative care, and I don’t tend to refer a ton of my patients for palliative care, despite the fact I like the group I feel like they’re already coming having to park, coming to a lot of visits, and to me it’s almost a barrier of do you want to see another provider.”

Additional patient-centered factors were identified, including perceived anxiety caused by these referrals: “I have many of my patients referred to palliative care, but for some of them, it’s actually mentally quite difficult to see the palliative care, even the way, you know, the appointments are structured and how much sicker the patients in those clinics are, I think, has reverberations for them.”

Medical oncologist-centered barriers to early palliative care integration

Medical oncologists acknowledged studies that demonstrated the benefits of early palliative care referral for patients with incurable cancer. Nonetheless, many medical oncologists described personal practice patterns

| Table 1 Demographic characteristics of the 31 medical oncologist participants |
|---------------------------------|--------------------|-----|
| Characteristic                  | Number (N=31)     | %   |
| Gender                          |                    |     |
| Male                            | 13                 | 42  |
| Female                          | 18                 | 58  |
| Age                             |                    |     |
| 25–34                           | 10                 | 32  |
| 35–44                           | 20                 | 65  |
| 45–54                           | 1                  | 3   |
| 55+                             | 0                  |     |
| Practice setting                |                    |     |
| Academic/university system      | 29                 | 94  |
| Private practice/community-based system | 2           | 6   |
| State of practice               |                    |     |
| Illinois                        | 16                 | 52  |
| Virginia                        | 3                  | 10  |
| California                      | 3                  | 10  |
| Indiana, Ohio, Wisconsin        | 2 each             | 6 each |
| Tennessee                       | 2                  | 6   |
| New Jersey                      | 1                  | 3   |
| Race                            |                    |     |
| White                           | 17                 | 55  |
| Asian                           | 12                 | 39  |
| Hispanic                        | 2                  | 6   |

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Medical oncologists' position of authority

Medical oncologists voiced a unique position of authority related to palliative care referral and end-of-life planning for their patients (Table 2). Medical oncologists described themselves as “driving the bus,” “owning the patients,” and “holding the position of authority” when decisions about palliative care referral and end-of-life planning were under consideration. One participant explained:

“I think we know it’s been published a ton in literature and we know one of the barriers to advance care planning and goals of care conversations with patients is often times a medical oncologist thinking they can do it better than anyone else, or that you have this sort of possessiveness of your patients. I think that’s for a reason, again, for all the reasons everyone said. We know them, we know their course the best, things like that.”

Other physicians such as palliative care specialists and radiation oncologists were thought to lack this authority to independently consider end of life planning, such as discussions about hospice care, by the majority of the participating medical oncologists. Instead, study participants preferred that palliative care and other physicians acted as reporters who could make the medical oncologist aware of changes in the patient’s condition and prognosis, but not make independent decisions about palliative care referral or end-of-life care with patients (Figure 1). An illustrative quote regarding radiation oncologists stated “I think the times where I think it’s worked well with radiation oncologists is when they talk to the patient, where they see them every day and see them declining and then communicate that with us.”

Alternatively, other physicians could seek the permission of the medical oncologist to pursue end of life care discussions with patients (Table 2). For example, one participant said: “I think that the conversation needs to first be had with the oncologist. And if the oncologist says yes, absolutely you should have that conversation with my patient, then there shouldn’t be an issue. But if the oncologist says no, you’re crazy, that’s not what you should be talking about now, then that’s something different.”

Rationale for position of authority

Medical oncologists expressed multiple reasons to support their unique position of authority in end of life care (Table 3). First, medical oncologists viewed the typical long-term relationship and rapport between themselves and patients as conferring a responsibility or duty to...
guide discussion and decisions about end of life care. To illustrate, a participant said: “I think if you’re a medical oncologist this is what the expectation is and I think you have to be able to do it well, and I think you go along the journey with them.” Other physicians such as surgeons and radiation oncologists were described as having transient involvement in the patient’s cancer care and lacking the same level of rapport or trust from patients to introduce palliative care services: “I love all the radiation oncologists I’ve worked with, but… there’s an end to their treatment. So I assume for them too it can be sort of awkward to have these conversations with a patient that they may never see again.”

Second, medical oncologists noted their unique clinical expertise and knowledge as a source of authority related to end of life care and the timing of palliative care and hospice referrals (Table 3). Medical oncologists described having knowledge about experimental treatments, clinical trials, or second and third line treatments and expressed concerns about other physicians discussing end-of-life care without explicit permission (Table 4). For example, medical oncologists noted that they would stop referring to specific palliative care physicians that are viewed as encroaching on the medical oncologists’ role. Other consequences for physicians who were viewed as infringing on medical oncologists’ position of authority include

**Figure 1** A schematic of the decision-making framework (on the left) as described by medical oncologists, with the medical oncologist “driving the bus”, augmented by input from palliative care and radiation oncology physicians. The patient-centered model on the right reflects the patient as “the captain of the ship”, receiving counsel from their oncology providers working in a collaborative manner to enhance patient-centered decision making regarding end of life care planning.

Consequences of medical oncologists’ authority

Medical oncologists discussed several consequences for physicians from other specialties who discuss end-of-life care without explicit permission (Table 4). For example, medical oncologists noted that they would stop referring to specific palliative care physicians that are viewed as encroaching on the medical oncologists’ role. Other consequences for physicians who were viewed as infringing on medical oncologists’ position of authority include
Table 3 Rationale for medical oncologists’ authority in end of life care planning

Unique relationship with patient

“You can’t just be there for all the good times, you have to be there—I think if you’re a medical oncologist this is what the expectation is and I think you have to be able to do it well, and I think you go along the journey with them.”

“… he was just recently hospitalized and everyone was offering him antibiotics and blood transfusions… I was the only the person that he trusted enough to tell him, “You’re dying.” And I knew that his end of life preferences were to die at home and that this window of opportunity was very short.”

“So they’re viewing me as sort of the go-to person to say, you know, when it’s time to stop.”

Clinical knowledge

“I don’t pretend to tell patients whether they’d benefit more from SBRT versus IMRT, and at the same time I don’t expect radiation oncologists to know the intricacies of chemotherapy trials and what may or may not be an option.”

“If I’m ready to convey either a clinical trial or a second or a third line agent for a patient, and they are telling them that they need to enroll in hospice then this is sending mixed messages and it really hurts everyone.”

“Maybe there’s a new biologic coming out or something new that maybe we can think about offering, and then the idea of hospice comes up when it wasn’t necessarily what we were thinking long-term. And I think that derails things a little bit and is a setback, so I think that can get frustrating if it’s not what you’re planning.”

Clinical training

“I think medical oncologists, our job I think is to understand the whole treatment trajectory, the disease process…I think that’s the medical oncologist, that’s a core function of what we do.”

“I don’t know if palliative care is incorporated into radiation oncology training.”

System or culture

“For every practice there is a culture. Unfortunately, I haven’t been in any practice where the culture it’s team’s responsibility, it’s always the medical oncologist that they have to refer to.”

“I think there is a difference between who is supposed to and who ends up doing it. So most of the time the medical oncologist ends up being responsible.”

Table 4 Hierarchical communication dynamics related to medical oncologist authority

“We have one (palliative care physician) that will overstep and do hospice from palliative care, and I don’t refer to that person ever. I will never let them have a patient of mine, because of that. Because if you refer them for palliative care they have a hospice discussion behind your back.”

“I’d be cautious. I would probably then speak to (the radiation oncologist) before I sent them any patients. And I would then certainly inform the patients that they’re potentially trigger happy with the whole palliative care thing.”

“I know oncologists can be a little possessive [laughter] with their patients, so I’m not surprised at that feeling (fear) that they may have… Telling someone you need to enroll in hospice right now or you have two months to live, those types of conversations I think would be upsetting.”

“Often while our palliative care folks are really good, they know not to have those conversations without talking to us first, because it only takes a few times of having a doctor yell at you, “Why did you say this to my patient when we have five more months of treatment? We have two more (years) to go.” You know, so most of the time they’re pretty good.”

“I think we all have fabulous relationships with our radiation oncologists. We value them 100%. It’s like our surgeons—I work with a surgeon now who tells people whether or not they should get chemotherapy… I keep going back and I’m like shut up. Enough already, it’s not your job. This is why we’re trained all these years. And so I think they should be scared of the medical oncologist.”
interpersonal conflict. Said one medical oncologist “I think it’s a very common fear that general medicine docs have, hospitalists, everybody has is angering oncologists—I think that because oncologists tend to take such ownership of their patients so people don’t want to make decisions for the oncologists.”

Similarly, this position of authority led to the timing of palliative care referral and end-of-life care planning to be viewed as the domain of the medical oncologist alone; “I think there’s no role for radiation oncology in palliative care talk, at least for my patients.” An example quote regarding palliative care cited the concern that “They might not have the exact same vision of what’s going on. So I think I prefer to manage that.” The decision to continue chemotherapy or not was often framed as the “territory” of the medical oncologist, rather than a decision that a patient could make through consultation with varying providers (“So it’s the territorial idea of like you do your surgery yes or no and I’ll do my chemo yes or no”) while the integration of palliative care was described as a decision for the medical oncologist to make, not an option presented for patient-driven decision-making.

Divergent perspectives
Few medical oncologists who participated in this study emphasized the patient as the central figure in end of life care planning: “The patients will be the sort of captain of the ship (Figure 1).” Some medical oncologists described routine early utilization of palliative care: “I also introduce palliative care right out of the gate, so the idea is not waiting until we think they’re in their 11th hour and then try to bring in strangers.” Contrary to our hypothesis, an interprofessional, “all hands on deck” approach to early palliative care was rarely supported, but one medical oncologist stated “I think the more people who talk about advance care planning with patients and the more people who bring it up the better because then it’s something that becomes not as scary for people to talk about.”

Discussion
The care of patients with advanced cancer, especially near the end of life, is complex and is likely improved by an interprofessional team approach that serves to formulate optimal treatment plans that may include chemotherapy, radiation, and supportive oncology services such as palliative care and hospice. For patients with advanced disease, early palliative care is recognized as important yet challenging to implement due to multiple barriers. We hypothesized that medical oncologists would favor an interprofessional approach to palliative care integration and end-of-life care planning and would support early palliative care referral for patients with advanced cancer. Our results, however, did not support this hypothesis. Instead, we found that medical oncologists follow a practice pattern that reserves palliative care referral for late in a patient’s course of illness. In addition, we found that medical oncologists feel a strong sense of responsibility for and authority over decisions about palliative care referral and are skeptical of efforts by other clinical team members to initiate these referrals.

Previously identified physician barriers to earlier palliative care referral have included discomfort with end of life care (15), distress associated with the name “palliative” (9,16) as well as “clinician concern about taking away hope” and “unrealistic clinical expectations” about the efficacy of therapies (17). While physician barriers to high quality end-of-life care have been previously described, the most prevalent barrier (described by 97% of oncology survey respondents) was described as “unrealistic patient expectations” (17). However, our qualitative work finds that medical oncologists themselves often do not want to integrate palliative care services early. Uniquely, our results demonstrate that medical oncologists perceive themselves to have authority over the timing of palliative care referral and end of life care (their “territory”) thereby minimizing the potential input other providers can provide towards informed patient-centered decision making. Our findings are consistent with those of Gidwani and colleagues (18) who noted oncologists potentially viewing palliative care practitioners as a “team of outsiders”. Medical oncologists expressed concern that palliative care discussion of alternative options such as hospice enrollment could make the job of the medical oncologist more difficult, and result in inappropriately early termination of systemic therapy. However, a secondary analysis of the landmark Greer et al. trial demonstrated that the overall receipt of chemotherapy was not reduced by early integration of palliative care (19); similar findings have been demonstrated in patients undergoing phase I/II trials (20). These findings suggest an important misperception of medical oncologists that is a modifiable barrier to early palliative care referral.

This study highlights a critical opportunity for improved collaboration amongst oncologists (medical, surgical, radiation) and palliative care physicians, which may aid individual team members to feel autonomous in counseling patients regarding end-of-life care. Potential interventions to improve this collaboration could include a “tumor board” setting where multidisciplinary providers and supportive
oncology (i.e., nutrition, social work, psychologist) could discuss complex patients care plans (21) ensuring providers are “on the same page” for patient prognosis and care plan. Addressing these barriers to early palliative care integration may significantly decrease health care expenditures, as has been suggested by studies examining the timing of palliative care (22). In addition, leveraging national initiatives, such as the CMS Oncology Care Model (23), may provide health systems and providers the necessary momentum to address some of the shortcomings identified in this study. The Oncology Care Model (OCM) is a demonstration payment model program by CMS that aims to improve the quality of cancer care while reducing cost. The OCM mandates a 13-element care plan that includes improvement in end-of-life care by providing patients with information on prognosis, treatment goals, and advanced care planning. The OCM’s emphasis on improving end-of-life care highlights the high cost of this portion of the cancer continuum. Practicing practices are provided payments to improve elements of cancer care, and, based on our findings, we recommend a strong focus on increasing interprofessional collaboration to facilitate improved care for patients with advanced cancer.

Limitations of this study include our sample cohort, which mostly represented university-based practices and may therefore not be generalizable to other care settings. However, our study design did reveal important barriers to early palliative care integration, even though our sample would likely have better access to palliative care specialists compared to other care settings.

In conclusion, perceptions of medical oncologists towards palliative care, and their authority over this decision, appear to significantly limit early palliative care integration. Efforts to ensure cancer patients receive access to earlier palliative care services may be improved if end of life care is a patient-centered decision with input sought from the entire care team.

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Footnote

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Conflicts of Interest: All authors have completed the ICMJE uniform disclosure form (available at http://dx.doi.org/10.21037/apm-20-270). TJK reports personal fees from AstraZeneca, personal fees from OncLive, personal fees from Varian Medical Systems, and personal fees from Abbvie Inc., outside the submitted work. JMK reports grants from National Heart, Lung, and Blood Institute, during the conduct of the study. JPG, MM, KK, ES, and SMK have no conflicts of interest to declare.

Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. The project was reviewed by the Northwestern University IRB (STU00204433) and determined not to be human subjects research; hence informed consent was deemed unnecessary.

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References

5. Le BH, Mileskin L, Doan K, et al. Acceptability of early integration of palliative care in patients with incurable...


