Advance Care Directives: A Herzl Clinic Quality Improvement Project on Patients’ Perspectives

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ABSTRACT

Background: Advance Care Planning (ACP) is beneficial to patients by ensuring their values and wishes regarding end-of-life care are respected. Despite the primary care setting being optimal for ACP discussions with tools available to facilitate these discussions, the initiation and documentation of Advanced Care Directives (ACD) in patients’ medical files were low and resident physicians perceived that patients were unwilling or unprepared for ACP discussions. The goal of this project was to understand the challenges and barriers that patients and their caregivers face in initiating and discussing ACD with their primary care team.

Methods: An online survey was conducted among 78 patients who are part of the Home Care program at the Herzl clinic. Participants were asked about the value placed on ACP and their preferences on various aspects surrounding the initiation of ACD discussions.

Results: 25 of 78 possible responses were received. This included survey responses from 6 patients, 13 caregivers, 4 family members and 2 physicians. Our results show that patients and their caregivers value Advance Care Planning discussions (>80%). Additionally, they endorse multiple benefits of ACP for themselves, their care teams, and families. Patients and caregivers prefer that medical professionals initiate and facilitate the discussions (70-80%) and are open to receive educational material to prepare for these discussions (68%).

Conclusion: Patients in a frail population are willing and open to discuss advance care planning with their primary care team. Family Medicine teaching clinics can support patients’ desire to engage in ACP by providing access to education material and initiating these discussions.

KEYWORDS
Advance Care Planning, Advance Care Directives, Patients
Advance care planning (ACP) is a proactive approach designed to facilitate comprehensive discussions and decisions regarding patients' values and wishes regarding end-of-life care. This process involves an open discussion between patients, their families, and their physician aiming to determine the desired level of care, goals of treatment and a surrogate medical decisions maker in case of loss of capacity. The s-32.0001 law (La Loi concernant les soins de fin de vie) is a Quebec law which underlines the importance of patients' agency with regards to end-of-life care through the establishment of the system of Advanced Care Directives (1). The Quebec Ministry of Health considers ACP a priority, making ACP discussions and documentation an integral part of the best practice of medical care.

ACP has been demonstrated to improve patients' end-of-life care, by respecting their consent and ensuring their quality of life should there be a loss of capacity (2). For example, it has been shown that a significant number of patients die in a medical setting, such as an Intensive Care Unit, despite a vast majority of people reporting that they would ideally prefer their death take place in their home (3). With well documented ACP, patients are able to dictate their preferences regarding end-of-life care, including the setting, lowering the risk of death in an unwanted setting. Additionally, ACP also benefits patients' families and surrogate decision maker, lightening the weight of certain medical decisions and lessening the burden of the bereavement process which can be turbulent, particularly if the patient's wishes were not felt to be respected.

In multiple studies, participants have stated their preferences regarding the discussion of ACP as being in the outpatient setting, with either their primary care physician or another healthcare professional they already had an established therapeutic relationship (4). These studies underline the fact that primary care is an optimal setting for ACP discussions. Discussed and well documented ACP minimizes urgent and invasive life-sustaining treatments that would otherwise be the default standard of care (5).
the process of initiating ACP discussions (by whom, when, where, and how).

The project was carried out by Drs Adrienne Poitras and Zhou Fang, residents in the Department of Family Medicine, McGill and supervised by Dr Hersson-Edery and Dr Keith Todd. Support was provided by Alexandru Ilie.

3 RESULTS

3.1 Characteristics of survey respondents

Survey response rates were 32% (25/78), among which 6 respondents are Herzl patients themselves. Barriers to completion of the survey included hearing and vision impairment, or declining cognitive function. The largest group of respondents were patients’ caregivers (13/25). Family members and physicians answered for the patients in 6/25.

96% of responders agreed that it is important to discuss ACP with healthcare professionals, yet more than half (54%) had not heard about ACP prior to this survey (Fig. 1A), and less than 25% have discussed ACP at Herzl.

A majority of respondents (24/25) answered ‘agree’ or ‘strongly agree’ to the statement “It is important to discuss Advance Care Planning with a health professional” (Fig. 1A). Six of the remaining respondents agreed and one individual was neutral. The reasons for placing this value were multiple. Some reasons included a desire to understand the choices around end-of-life interventions, to prepare for end-of-life decisions, to pre-emptively make decisions to avoid burdening family members, to ensure that their wishes will be respected, and to ensure dignity at end of life. Most respondents (70%) indicated that ACD could help reduce disagreements between family members and health teams.
3.2 | Timing and How ACP discussions should happen

Respondents showed a clear preference towards retaining autonomy on when to initiating discussions. Another large proportion indicated they would want an invitation to discuss ACP when there is a change in their health status (16/25), whether there is a new diagnosis or deterioration in health status. Less than half of participants (10/25) indicated that they would want this discussion on a yearly basis, such as when a new resident physician took over care, or if they are healthy.

More than 50% of respondents indicated a preference for informal initiation of ACD during a previously scheduled medical visit, rather than a separately scheduled visit, an email or a phone invitation. Additionally, (15/25) 68% of respondents were open to receiving preparatory medical information on ACD or medical interventions such as CPR, intubation, or dialysis, either through paper or digital format, or for this information to be provided to their family or caregiver. A single participant indicated that they preferred to receive preparatory information via a website.

3.3 | Who Should Discuss ACP

Participants expressed a stronger preference for the resident, staff physician or family member to bring up the ACP discussion, rather than a nurse or a social worker or themselves (Fig. 1B). 16 out of 25 respondents agreed or strongly agreed that a resident or staff physician should initiate the discussion.

4 | DISCUSSION

This quality improvement initiative adds important insights into how people think about advance care directives and their planning. Our results were somewhat surprising in that, although one would expect some knowledge of ACP amongst a frail population such as the one surveyed here, fewer than half had heard of ACP prior to the survey. Unlike a larger survey in Canada (13), our results are not focused on the term per se, since a definition was provided to ensure understanding. The work by Teixeira et al. (13), however, illustrated that many Canadians are engaging in these informal discussions with family members, which did not seem to be the case with our patients.

Our study confirms findings in the medical literature that patients prefer having physicians initiate discussions regarding end-of-life care in addition to facilitating the discussion when the patient brings it up or when there is a change is health status (10,11,12). The preference for involvement of a medical doctor may reflect a familiarity with these members of the care team since in our context since they are the professionals making the home care visits. However, it is quite likely that for other patients who were more familiar with other members of the health care team, such as a nurse or social worker, that these individuals would be the preferred contact person for these discussions (12).

The surprisingly small proportion of people who were
aware of ACD and who had participated in discussions highlights the need for more patient education. This seems not to be unique to our population as survey administered across Canada found that only 16% of people were aware of the term and only 20% had a written advance care plan (13). Interestingly, the residents’ perception that patients, their caregivers, and their family members are reluctant to prepare and discuss end-of-life care in the outpatient setting (9) was not supported in our iterative follow-up quality improvement project. Our survey did reveal an openness by patients and families to receive educational material prior to discussions. This gives us clear opportunity and focus for changing patient awareness to facilitate more frequent ACP discussions. Patients and family members also indicated a preference for individual, in-person visits with their physician and family members, while there was little interest in group discussions on ACP. This also clarifies where we need to focus our implementation strategies.

Considering the limitations posed by the frailty and other possible barriers such as communication, auditory or cognitive difficulties, of our patient population we hope to extend this survey to the general older adult population at Herzl Clinic to see if their experience and perspectives on the initiation of Advance Care Planning and Directives differ.

4.1 Limitations of study

The low response rate of 32% was likely multifactorial. In addition to the fact that our Home Care patients form a frail population in which varying degrees of cognitive, language or hearing barriers are not uncommon, many do not have or use email, and some would have difficulty understanding how to access an online survey. Only 6 out of 25 respondents were the patients themselves and family members answered the survey in 4/25 surveys.

Due to these anticipated barriers, the authors followed up with phone calls and offered to administer the survey by phone, which increased our response rate considerably. Most of our respondents were caregivers (13/25), who despite knowing their care recipient well, do not necessarily have the same values and beliefs as their care recipient, and therefore may not always respond to the survey in the same way the patients would themselves. Indeed, caregivers did include their care recipient in answering the survey questions, when possible, but it is unclear how well the responses reflected the patients’ perceptions. We did not capture the prevalence of significant cognitive and sensory barriers to participating in this survey among our population.

In conclusion, this study advances our understanding of the challenges patients, caregivers, and family members in a frail Home Care population perceive in the initiation and discussion of Advance Care Directives with their physicians in the outpatient setting. Patients identified several areas of improvement that will be addressed in future iterations of this quality improvement project in order to improve the frequency and quality of Advance Care Planning discussions between Family Medicine residents or staff physicians, and their patients.
REFERENCES