Supportive Care: Communication Strategies to Improve Cultural Competence in Shared Decision Making

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Abstract

Historic migration and the ever-increasing current migration into Western countries have greatly changed the ethnic and cultural patterns of patient populations. Because health care beliefs of minority groups may follow their religion and country of origin, inevitable conflict can arise with decision making at the end of life. The principles of truth telling and patient autonomy are embedded in the framework of Anglo–American medical ethics. In contrast, in many parts of the world, the cultural norm is protection of the patient from the truth, decision making by the family, and a tradition of familial piety, where it is dishonorable not to do as much as possible for parents. The challenge for health care professionals is to understand how culture has enormous potential to influence patients’ responses to medical issues, such as healing and suffering, as well as the physician-patient relationship. Our paper provides a framework of communication strategies that enhance crosscultural competency within nephrology teams. Shared decision making also enables clinicians to be culturally competent communicators by providing a model where clinicians and patients jointly consider best clinical evidence in light of a patient’s specific health characteristics and values when choosing health care. The development of decision aids to include cultural awareness could avoid conflict proactively, more productively address it when it occurs, and enable decision making within the framework of the patient and family cultural beliefs.


Introduction

This Moving Points in Nephrology describes the principles and challenges of providing person-centered care for patients with advanced kidney disease at the end of life. Communication between the practitioner and the patient (including their wider family/social support) is key to achieving these aims. As discussed by Davison and Jassal (1), this includes sharing prognosis, determining symptoms, and providing care aligned to the preferences and goals of the individual patient. These principles of truth telling and patient autonomy are embedded in the framework of Anglo–American medical ethics. In contrast, in many parts of the world, medical practice is on the basis of family decision making and medical beneficence. These differences can inevitably lead to conflict between patients, families, and clinicians and therefore, the need to develop strategies to reduce cross-cultural miscommunication (2). This paper aims to help clinicians become culturally aware and competent, thereby improving their communication with patients and families.

The American clinical practice guidelines on dialysis initiation and withdrawal from the Renal Physicians Association (RPA) (2) state that “they reflect the ethical principle of respect for autonomy because clinicians, family members, and others have an ethical duty to accept the decisions regarding medically indicated treatment made by competent patients and in the absence of competence, to formulate decisions that would respect patients’ wishes, or if wishes are unknown, advance the best interest of their patients” (2). Historic migration and the ever-increasing current migration into Western countries have, however, greatly changed the ethnic and cultural patterns of patient populations. This is particularly true when considering advanced kidney disease, which is much more common in many ethnic groups compared with white European populations (3). As an example, around 50% of patients on RRT in London are from ethnic minorities, predominantly South Asian and Afro-Caribbean (4).

Migration: Magnitude of the Issue

Throughout human history, individuals, families, and groups have emigrated from their native homes to other places globally for many reasons: the prospect of education, economic, or social advantage; the need to escape war, political torture, or other conflicts; or the desire to reunite with other family members. At one point in 2005, there were an estimated 191 million immigrants across the globe: approximately 64 million of these immigrants arrived in Europe, and 44 million arrived in North America, a tripling of the immigrant populations in these regions compared with 20 years earlier (5). Spain, Germany, and the United Kingdom were the European countries with the highest immigration, receiving more than one half of all immigrants in 2008 (6). Increasing diversity is a reality as witnessed by the daily news bulletins about dramatic increases in global economic and political
migration. This means that there is an enlarging proportion of people who do not live in their own native country or culture.

In many parts of the world, the cultural norm is protection of the patient from the truth, decision making by the family, and a tradition of familial piety, where it is dishonorable not to do as much as possible for parents (7); examples are given in Table 1. With evidence that the ethics of minority groups may follow their religion and country of origin (8), conflict with Anglo–American medical ethics structure may arise from both patients’ and physicians’ perspectives, because both have their own languages, explanatory illness models, religious beliefs, and ways of understanding the experience of suffering and dying (9,10).

The Effect of Diversity on ESRD and at the End of Life

Culture is but one of several typologies of difference that have been used to signify diversity among individuals and groups. Narrowly defined from an anthropologic perspective, culture can be thought of as that which refers to “…patterns, explicit and implicit, of and for behavior acquired and transmitted by symbols,” language, and rituals (7). Seen as a recipe for living in the world, this conceptual framework for culture explains the means of transmitting these recipes to the next generation (8). The challenge for health care professionals in an increasingly diverse society is to understand how culture has enormous potential to influence patients’ responses to medical issues, such as healing and suffering, as well as the physician-patient relationship. As a direct result, those from migrant communities may possess little knowledge of or have little exposure to palliative care. (Palliative care has been defined as an approach that improves the quality of life of patients [adults and children] and their families who are facing problems associated with a life-threatening illness. It prevents and relieves suffering through the early identification, correct assessment, and treatment of pain and other problems, whether physical, psychosocial, or spiritual.

Addressing suffering involves taking care of issues beyond physical symptoms. Palliative care uses a team approach to support patients and their caregivers. This includes addressing practical needs and providing bereavement counseling. It offers a support system to help patients live as actively as possible until death (11). For example, in the United Kingdom (12) and more recently, among people living with end stage kidney failure in Canada (13), those from black, Asian, or minority ethnic groups were identified as being statistically less likely, after taking all other factors in account, to understand the value of palliative care. Specific to kidney disease, the US Renal Data System data show that rates of dialysis withdrawal in minority ethnic groups are lower compared with the white population (14). It is also important to note that migrant communities may also have different cultural values regarding life and death compared with the Western approach to dying, which includes palliative care.

Identifying preferences for medical care in advance of untoward or terminal circumstances can be a difficult and emotional process. The decision-making model of advance care planning derived from bioethics practices assumes that choices made by the individual can be arrived at through rational processes that are unchanged by time, shifting social consequences, or disease and illness progression. Such a model may only appeal to certain subsets of groups, thus limiting the utility of instruments used for advance planning (living wills or durable powers of attorney for health care) (15,16). For some groups, speaking about the dying process or planning for death may represent a transgression of a strong cultural taboo and could create additional distress. Other patients, unfamiliar with or mistrustful of the legal system, may misconstrue the purpose or nature of formal advance care planning documents. In all cases, rather than abandon the goals of advance care planning, strategies should be sought that facilitate understanding. For example, a generic discussion to identify a health care proxy need not be cast as a discussion of death but rather, can be an opportunity to determine desired roles of various family members and support persons. Discussions about patient preferences for

<table>
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<tr>
<th>Country/Cultural Group</th>
<th>Attitudes toward Truth Telling</th>
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<tr>
<td>China</td>
<td>When fatal diagnosis or prognosis is given, the physician informs the family and hides it from the patient—it is up to the family to decide whether, when, and how to disclose the truth to the patient; families usually decide to conceal such information, and physicians are willing to follow such decisions and cooperate with families in deceiving patients (39)</td>
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<tr>
<td>Black</td>
<td>Only God has knowledge and power over life and death, and physicians cannot have access to this type of knowledge; the Christian religious view held by many in the black community holds that suffering is redemptive—it is to be endured rather than avoided; foisting life support to avoid pain and suffering, therefore, might be seen as failing a test of faith (9)</td>
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<tr>
<td>Italy</td>
<td>Trend of partial and nondisclosure persists; this arises within families independent of patient requests, although there is some evidence that physician preferences are moving toward full disclosure (40)</td>
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<td>Spain</td>
<td>Tradition of partial and nondisclosure; the majority of doctors state that they would inform the patient only in certain circumstances or if requested by the patient (40)</td>
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<tr>
<td>India (Hindu)</td>
<td>Tradition of nondisclosure and relatives protecting the individual from knowledge in case he/she gives up hope and dies prematurely; this is exacerbated by the belief that modern medicine often provides hope, however unrealistic, that a cure is possible (41)</td>
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end of life care should be culturally and linguistically appropriate and reflect sensitivity to patient values and beliefs.

Relevant knowledge and greater awareness of palliative care are, therefore, critical, particularly as growing evidence suggests that a significant number of people from ethnic communities, which disproportionately include those on low incomes, misses out on high-quality palliative care and end of life care. This situation exists even in the United Kingdom, despite palliative care being free at the point of delivery from the National Health Service and the independent charitable sector. Possible explanations for these disparities include (1) different referral patterns to specialist palliative care and lack of understanding among professionals about exactly which patients to refer and when; (2) gatekeeping by services; (3) complex linguistic and communication barriers; (4) different preferences, including for more aggressive or curative care at the end of life, or a cultural mistrust of end of life care; and (5) strong religious and familial support systems (17–19). Additionally, people from minority ethnic communities may also experience overt and inadvertent racial discrimination at individual and institutional levels (20). Identifying and eliminating vertical health inequality (inequality among households or individuals) and horizontal health inequality (inequality among culturally defined [or constructed] groups) in the delivery of high-quality palliative and related care, therefore, represent critical mandates. More sophisticated standards for monitoring and ensuring the cultural sensitivity and cultural competency within the palliative medicine workforce and more widely should be used along with strategies to increase community-based partnerships.

Avoiding and Coping with Conflict

Both the US RPA guidelines (3) and the General Medical Council (GMC) (21) in the United Kingdom give some guidance about avoiding conflict. Both are on the basis of the premise that individual patients should be aware of their prognosis to make decisions regarding their care. The GMC states clearly that physicians do not have to provide treatment that they consider nonbeneficial. They also address the issue of conflict between clinicians and patients/families regarding demands for dialysis that the clinician deems medically inappropriate. The cultural conflict of giving information (or not) to individual patients is not addressed. The suggested solutions to conflict are specifically a trial of dialysis (but this needs clear end points), second opinions, consultation with the hospital ethics committee or ethics consultants, and ultimately, potential transfer to another institution or physician to provide dialysis. If none are willing to accept the patient, the family/legal representative can be informed that dialysis will be withdrawn unless there is a court injunction to the contrary.

In reality, neither the United Kingdom GMC nor the US RPA guidelines avoid conflict. Nephrology teams, therefore, need to develop cultural awareness and crosscultural communication strategies, including use of decision aids, to enable shared decision making within the framework of the patient and family cultural beliefs. This approach should proactively reduce conflict (23) and productively address it when it occurs.

Crosscultural Communication Strategies

Culture defines the way that people make sense of the world and influences how individuals view the illness experience and approach decision making. Despite the importance of culture in health care, traditional medical training is deficient in crosscultural communication education. Strategies to improve skills and knowledge in cultural competence and better communication relevant to the care are required (24). For example, crosscultural communication includes strategies that acknowledge individual cultural traditions, avoid generalizing a patient’s beliefs or values on the basis of cultural norms, and take into account one’s own beliefs, values, and experiences (25). Clinician culture is multifaceted and largely shaped by the biomedical influences, which include the knowledge and experience that accompany becoming a physician as well as the influence of a given health care system in which one practices (9). A recent qualitative study of United States and United Kingdom academic medical centers examined the influence of institutional culture on do not resuscitate decision making at end of life (26). The way that the physicians in training approached decision making was directly influenced by whether the hospital policies prioritized patient autonomy versus best interest. For instance, physicians training in a hospital that prioritized autonomy would be more likely to neutrally offer resuscitation, regardless of whether they believed resuscitation to be clinically appropriate.

To address these challenges, crosscultural communication strategies must be reflective and individualized (23,27). The first step to crosscultural competency involves becoming aware of the inherent beliefs, values, and biases within ourselves as clinicians and the influence of the health care system in which one practices. When clinicians become conscious of their own beliefs and values, they may become more receptive and open to those of the patients, especially when differences exist. Figure 1 visualizes the complex cultural influences within the patient and clinician relationship.

The second step involves effective communication strategies that are evocative, nonjudgmental, and respectful. Kagawa-Singer and Kassim-Lakha (23) have proposed a strategy to elicit information about the patient’s Resources, Identity, Skills and Knowledge (RISK), known as the RISK reduction assessment. This is a helpful strategy to learn and support the particular cultural influence and beliefs of a given patient and family. The RISK reduction assessment includes resources for patients and families to navigate the health care system, individual circumstances and migration...
experience, skills available to the patient and family to navigate the health care system and cope with the disease itself, and knowledge about the ethnic groups health beliefs, values, practices, and cultural communication etiquette (23). Teal and Street (27) developed a culturally competent communication model from existing models that incorporate

**Table 2. Cultural competency communication skills and examples (21–23)**

<table>
<thead>
<tr>
<th>Communication Task and Communication Strategy</th>
<th>Example</th>
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<tbody>
<tr>
<td><strong>Understand the patient’s experience</strong></td>
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<tr>
<td>Rapport building/ask about the patient as a person</td>
<td>Can you tell me about your life? Where were you born and raised? How has your experience been coming to a new country?</td>
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<td>Invite curiosity</td>
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<tr>
<td>Assess how the patient interprets her condition</td>
<td>As your clinician, what would be helpful for me to know about you and your life?</td>
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<tr>
<td><strong>Giving information</strong></td>
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<tr>
<td>Assess for health knowledge needs</td>
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<tr>
<td>Ask what kinds of information desired</td>
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<tr>
<td>Give information concisely without medical jargon and check in for understanding</td>
<td>What is your understanding of your condition?</td>
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<tr>
<td><strong>Determine patient involvement in medical decision making</strong></td>
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<tr>
<td>Assess decision-making preferences</td>
<td>How would you like decisions to be made about your health care?</td>
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<tr>
<td>Determine the patient’s preferred decision maker</td>
<td>Who is the person that you trust to help make medical decisions if you were unable to do so?</td>
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<tr>
<td><strong>Understand the patients beliefs and values</strong></td>
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<tr>
<td>Ask about what is important to the patient and loved ones</td>
<td>As we talk about how best to care for you, what are you hoping for? What concerns you most?</td>
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<td>Address spiritual concerns</td>
<td></td>
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<tr>
<td><strong>Address trust concerns</strong></td>
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<tr>
<td>Be transparent; avoid judgement and defensiveness</td>
<td>Faith can be a source of strength. Can you tell me about your faith?</td>
</tr>
<tr>
<td>Explore experience</td>
<td></td>
</tr>
<tr>
<td><strong>Address resource/needs</strong></td>
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<tr>
<td>Actively inquire about ways to support the patient and family</td>
<td>Some people are uncomfortable discussing their health with a clinician from a different background. Please feel comfortable sharing with me your concerns when it comes to your health, have you ever felt that you have been treated unfairly?</td>
</tr>
<tr>
<td>Actively assess for concerns about the plan</td>
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**Figure 1. Cultural aspects that influence the clinician-patient interaction.** When clinicians become conscious of their own beliefs and values, they may become more receptive and open to those of the patients, especially when differences exist.
critical elements of cultural communication. This model highlights five key communication skill sets: nonverbal skill, verbal skill, recognition of potential cultural differences, incorporation of and adaptation of cultural knowledge, and negotiation/collaboration (28).

Ask-tell-ask is a helpful communication strategy to engage in culturally competent communication (29). This framework encourages a two-way conversation, in which the clinician first asks for the patient’s and/or family’s input rather than reflexively disclosing information. The usefulness of this strategy extends beyond giving information to include asking about cultural experience, decision-making preferences, and prior experiences with health care and exploring values and preferences. Table 2 includes examples of open-ended questions to better understand the cultural preferences and values of a given patient and family (23,28,29).

After the clinician has an understanding of the kinds of information desired and ways to communicate this information, how the information is told is equally important. Patients and families may have language barriers and low health literacy, further complicating their ability to process and act on critical medical information (30). Clinicians should use clear language, with only one to three pieces of information without medical jargon. The final ask allows the patient and family to teach back what they have heard to ensure that the clinician gave information in a way that was easily understood: “To ensure that I did a good job giving you information, can you tell me what you will take away from our discussion today?” This question is nonjudgmental and invites the opportunity for the patient and family to correct information or perceptions that the clinician may have shared. Additionally, this final ask invites the patient and family to share any concerns or lingering questions.

Shared Decision Making

Shared decision making is seen as useful in enabling practitioners to be culturally competent communicators (28) by providing a model where clinicians and patients jointly consider best clinical evidence in light of a patient’s specific health characteristics and values when choosing health care (31). It is important to recognize that some patients would rather have different levels of engagement in the decision-making process, with some preferring a recommendation from the clinician (28). In practice, it requires patients, professionals, and health care systems to think differently about the delivery of and engagement with evidence-based care. Both practitioner and patient are expected to collaborate proactively in this decision-making process by exchanging information about the illness, diagnosis, and treatment from their areas of expertise; making explicit values and preferences in the context of care pathways and/or lifestyle; reasoning together the best option for the patient; and agreeing and implementing the choice that aligns best with clinical evidence and patient preference (32).

Patients, professionals, and health care infrastructures can be enabled to engage in shared decision making (31,33). Interventions enabling people to collaborate more effectively within this complex system are informed by findings from the applied social sciences, medical communication, evidence-based practice, and health professional training. Different types of interventions are used to support people in different stages of making a decision and reasoning with others (33): for example, evidence-based prompts for professionals to make accurate choices in the context of care pathways and patients to make informed decisions in the context of their lives (1 and 2 in Figure 2); health professional training, patient decision coaching, and consultation prompts for
more effective communication between patients and professionals (3 in Figure 2); and training and decision aids for others involved in implementing care in people’s lives (e.g., other health professionals and/or family) (Figure 2). When developed using systematic methods to identify patient, professionals, and health care needs and preferences, these resources are culturally relevant (34), because they make explicit the options, attributes, values, and evidence of importance to all people involved in making and implementing health care choices (Figure 2).

Evaluation of shared decision-making interventions within predialysis education programs suggests that they are acceptable to staff and patients and can be implemented across different health care systems: for example, shared decision-making training and prompts for use by health professionals to structure predialysis education consultations (35) and patient decision aids supporting patients’ engagement with predialysis programs (36). Furthermore, there is a range of patient-centered approaches used by others in the delivery of self-care, advance care planning, and palliative services that provide techniques (e.g., goal setting and training) and prompts (e.g., patient-reported outcome measures) to help professionals deliver and negotiate evidence-based care in a culturally appropriate way (37,38).

Conclusions

There is increasing awareness in both lay and medical circles that, for many patients, modern medicine fails to achieve the quality and dignity of death that most people would want when asked. To achieve this, it is essential to enhance communication between health care teams, patients, and families. The framework for doing this has mostly been developed around the Anglo-American model of truth telling and patient autonomy as essential components of the decision-making process. Many of our patients and families make decisions using different frameworks, and this may be further exacerbated by an underlying distrust of health care teams delivering care in culturally and linguistically different ways. The resulting conflict often disadvantages the individual patients concerned with failure to share prognosis, wishes, and goals. Increasing crosscultural competency with resulting enhanced use of shared decision making should avoid some of this conflict and improve the quality of medical care for patients throughout the continuum of their illness.

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Disclosures

None.

References


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