Choices for life: how can we help?

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“May your choices reflect your hopes, not your fears”.

Nelson Mandela

HISTORICAL CONTEXT

Fifty years ago, Portugal lived under a dictatorship. A significant number of people were illiterate and there was no health care system to support the sick. Since then, a long road has been traveled in medical care: Serviço Nacional de Saúde (the National Health Service) has been created, medicine has evolved, and brand-new treatments are available for all kind of patients, including dialysis for elderly and frail ones. The spread of resources has not been followed by literacy in health, meaning doctors start to choose for their patients, who blindly believe them.

PRINCIPLES OF BIOMEDICAL ETHICS

In 1977, Thomas Beauchamp and James Childress developed the Principles of Biomedical Ethics1 that now guide medical clinical practice. Autonomy, beneficence, nonmaleficence and justice are the four pillars of modern medicine...in theory. But the concepts of beneficence and nonmaleficence may not be clearly black and white. No doctor offers dialysis if he does not believe it is the best option. Otherwise, not offering it would be denying treatment. Yet, for the same patient, it is possible that another colleague would recommend Conservative Care to respect the principle of nonmaleficence in an elderly frail patient. The same patient, in the same scenario, may have two completely different opinions based on the nephrologist’s experience, on personal beliefs, on the lack of accuracy in prognostic tools or even based on institutional policies and resources. Autonomy refers to self-determination for free choices and is the fundament of informed consent. Nevertheless, there are several ways autonomy can be limited. Pain, fear, economic dependence, and manipulation are all forms of restraint on freewill.

CHOOSING END STAGE KIDNEY DISEASE THERAPY

In nephrology, end stage kidney disease (ESKD) treatment options are a pinnacle of how bioethical principles are respected in Portugal. In 2011 a guideline from Direção Geral de Saúde2 established the obligation to present kidney replacement therapy (KRT) options to all stage 4 to 5 CKD patients. The purpose of this guideline is remarkable: to establish the need to enlighten all patients about their condition and present them with all options, including Conservative Care. The document is divided into two parts: the first is dedicated to nephologists, where the fundaments and framework for each KRT are presented and the second is provided to patients as information to help them choose an option – a patient decision aid (PtDA).

PtDA are tools or interventions that support shared decision-making. They come in several formats: written information, oral presentations, podcasts, videos or web resources. Analyses of these different options – their benefits and limitations – are barely found3. Further, they depend heavily on social and cultural issues. In Portugal, the guideline demands that a total of 17 pages should be supplied during an outpatient appointment where KRT is discussed. The model of this appointment varies from center to center but the brochure provided is standardized. But after all, are these 17 pages of full information really helpful for decision making? Most of our incident dialysis patients are elderly, with significant geriatric syndromes such as impaired vision or hearing, cognitive dysfunction, and social, familiar or economic constraints that hamper the ability to realize and benefit from these extensive tools. Informed consent cannot be simplified to dump all possible information (risks, advantages) that doctors are aware of. Most of our patients are not doctors, so they won’t be able to digest all this information. Informed consent must be elevated to a provision of adequate information that patients may understand, allowing them to choose which option will help them attain their goals.

The way this information is provided has been widely studied in CKD patients by Keren Ladin et al.4,5 She identified four distinct approaches:

- Paternalistic (the physician acts as a patient guardian and uses his expertise to choose for the patient);
- Informative (the physician is the holder of technical information that is provided to the patient who decides, with no influence of doctor’s opinion);
- Interpretive (the physician exploits the patient’s goals and based on them, guides him to treatment choice that may fit better with his values. It relies on a truly shared responsibility where doctors translate medical information into patient wishes);
- Institutionalist (information provided is influenced by institutional guidelines and norms).

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All these models have strengths and limitations: the paternalistic model may be adequate in an emergency situation where there is no time for opportune discussions. The informative model may be suitable for patients who have high health literacy and where influences should be excluded. For our elderly patients, a more person-centered care may be more appropriate. The problem of the interpretative model is that it requires skills that are not usually taught in residency, such as communication, and it is time-consuming.

Dr. Atul Gawande, an American surgeon dedicated to end-of-life care, who is also an adviser of the Conversation Project (a plan to help people share their wishes for care through the end of life) writes “We think our job is to ensure health and survival. But really, it is larger than that. It is to enable well-being. And well-being is about the reasons one wants to live.” In his most cited bestseller Being Mortal, he also discusses the above models of communication. He stresses that the paternalistic model is the most commonly used in “vulnerable patients – the frail, the poor, the elderly, and anyone else who tends to do what they’re told”. He also highlights that patients “want information and control, but [they] also want guidance”. And suggests how to do it: “doctor’s role is to help patients determine what they want. (...) What is most important to you? What are your worries? Then, when they know [patients’] answers, they tell [them] about the red pill and the blue pill and which one would most help achieve [patients’] priorities”.

So, in addition to the content of the information, its format matters. In other words, choosing ESKD therapy must be a person-centered care (PCC). PCC is defined by the Institute of Medicine as a care that responds to individual patient needs and values and guides treatment decisions. Eight principles are generally recognized (Figure 1). Respect for patient preferences, information and education are some of the most important which are regularly mentioned and supported by all guidelines and recommendations from nephrology societies. Even so, the fact is that it is not widely implemented. There are several barriers to that:

– Patient barriers (illiteracy in health, limited access or motivation to search for information, analphabetism, blindness, deafness, cognitive impairment, lack of familiar support...);

– Doctors’ barriers (time constraints, lack of confidence in discussing end-of-life conversations.);

– System barriers (limited incentives to education, lack of support, lack of human or logistic resources,...).

Some of these barriers are for us, nephrologists, to beat and settle the requirements to a true shared decision-making process.

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**BASIC CONTENTS OF A DECISION-MAKING APPROACH IN CLINICAL PRACTICE**

In clinical practice, guiding CKD patients in a Decision-Making Process starts with the selection of who should be approached and when. It
Table 1
Basic contents of a pre-dialysis education program

<table>
<thead>
<tr>
<th>Framework</th>
<th>Meaning</th>
<th>Example</th>
</tr>
</thead>
<tbody>
<tr>
<td>Set up the interview</td>
<td>Arrange a place where conversation would not be interrupted</td>
<td>Can I talk to you about your situation? When is it best for you and your family?</td>
</tr>
<tr>
<td>Perception assessment</td>
<td>Prepare yourself with all the information you can get on patient’s clinical situation.</td>
<td>What has already been told to you about your situation?</td>
</tr>
<tr>
<td>Perception assessment</td>
<td>Assess patient understanding about his situation</td>
<td>Is that okay if I talk to you more about your condition?</td>
</tr>
<tr>
<td>Perception assessment</td>
<td>Determine patient level of comprehension</td>
<td>I am afraid your kidneys are not working anymore. Do you understand what that means?</td>
</tr>
<tr>
<td>Invitation</td>
<td>Ask the patient what information he wants to know</td>
<td>Provide medical facts in an intelligible language: consider educational, socio-economic and cultural background</td>
</tr>
<tr>
<td>Knowledge and information provision</td>
<td>Check patient comprehension of what you said</td>
<td>Identify patient’s emotions and be empathic to them. Give the patient time to express his feelings, even in silence.</td>
</tr>
<tr>
<td>Address patients’ Emotions with Empathic responses</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Strategize and Summarize</td>
<td>Summarize information. Ask patient if he wants any other information.</td>
<td>I believe it is very difficult to hear this. How do you feel about it?</td>
</tr>
<tr>
<td></td>
<td>Schedule next meeting and offer your support.</td>
<td>It is a lot to deal today. For now we will (establish the plan). We will check in our next appointment how it’s going for you.</td>
</tr>
</tbody>
</table>

Table 2
SPIKES framework and example – How to break bad news

The ideal pathway to choose KRT should start with patient education. A formal and structured program to raise awareness of the disease, its consequences and options is fundamental (Table 1)\(^\text{16}\). To approach all these issues, the team should be multidisciplinary. In addition to a nephrologist, nurse and nutritionist, a social worker to make clear patients’ rights and social support is also required, as is a psychotherapist. The formats in which all these questions are addressed are as important as the content. It is not worth providing the most complete information if the patient cannot understand it. A peer session or support group, with other patients sharing their experiences, may be more useful than all other information provided. In times of pandemic such as we are living, social distancing may require other options, such as videotapes of patients contributing with their own experiences, but the importance of “testimony from people who are in the same boat should not be neglected. In a specific article that analyses pre-dialysis education programs, those with a single session were excluded, from the beginning because they didn’t meet the minimum requirements\(^\text{17}\). The fact that education is a time-consuming task should not be an obstacle, as long as nephrologists realize its importance. If we believe it is necessary to educate patients to reach objective parameters such as blood pressure which are easy to measure\(^\text{18}\) we should not forget that education programs may impact something bigger: patient-related outcomes, or in other words – well living!

Finally, pre-dialysis programs are also the appropriate moment to discuss advance care planning. In Portugal, living will has been available for about 10 years\(^\text{19}\), but by last year, fewer than 30,000 persons had made a living will. This reflects illiteracy in health, but also physicians’ reluctance to have end-of-life discussions.

**HARD CHOICES AND COMMUNICATION SKILLS**

That is why, in addition to patient education, patient-centered care requires physicians’ education. One of the most important barriers to provide it is lack of communication skills. In Anglo-Saxon countries, where patient-centered decisions are more implemented, doctors have training in communication so they can guide patients through difficult options. There are several available resources that are useful
to help clinicians in these activities. There are also websites which present them in an accessible way, such as tutorials where examples can be followed. I am familiar with some of them that I consider really valuable in clinical practice.

SPIKES (Table 2) is a useful tool to break bad news. It might be used when patients come to the KRT appointment to set up the patient comprehension of the situation and to further establish a plan. As stated in this article, the healthcare team should not only provide information but also build insight on the best modality that serves the patient’s wishes and goals. The ideal way to recognize these goals is also using structured tools. The REMAP framework (Table 3) provides a mnemonic that help clinicians to start these conversations, discussing prognosis, exploring patient values, attending emotions, and then using this information to develop the final plan. It helps doctors to enroll information in an interpretative model. Whenever a patient is undecided about the best KRT to follow, after judging benefits and limitations related to their goals, a “Best case/worst case” approach

Table 3
REMAP framework and example – How to manage a plan with the patient

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<tr>
<th>Framework</th>
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</thead>
<tbody>
<tr>
<td>REFRAME the big picture</td>
<td>Doctor should summarize the patient’s clinical situation and try to understand patient’s comprehension of it</td>
<td>You have been suffering from CKD for a long time, also from [list other serious comorbidities]. I believe it has been very hard to deal with all of this.</td>
</tr>
<tr>
<td>EXPECT emotion</td>
<td>Doctor should create an empathetic relation and respond to emotions</td>
<td>I can see how hard this is to hear. Is it okay for us to talk about what it means?</td>
</tr>
<tr>
<td>MAP out patient values</td>
<td>This is the moment to present different trajectories and elicit the patient’s concerns, worries and values</td>
<td>Given this situation, what’s most important for you? How would you like to live your life from now on?</td>
</tr>
<tr>
<td>ALIGN with values</td>
<td>In this step, patient’s values are reviewed to aligned with the next proposals</td>
<td>If I have correctly understood, it sounds the most important things to you are...</td>
</tr>
<tr>
<td>PROPOSE a plan</td>
<td>The last point is to offer possible directions that match patient’s goals while reassuring involvement and nonabandonment.</td>
<td>I think the best option to cope with your values is... because it will allow you to... What do you think of this plan?</td>
</tr>
</tbody>
</table>

Figure 2
Best scenario/worst scenario approach – adapted from

“Best Case/Worst Case” tool involves a drawing made by the physician. Each treatment option is represented by a vertical bar; the length of the bar represents the range of possible outcomes. The “best case” is represented by a star; the “worst case,” by a box; and the “most likely” outcome, by an oval. The physician describes each “case” using narrative derived from clinical experience and relevant evidence and writes key points on the diagram.
can be used\textsuperscript{23}; a personalized scheme is created (Figure 2) where the patient can visualize the best and worst scenario in each option.

\section*{CONCLUSION}

These tools are not designed to decrease the responsibility to choose but to help patients to understand what fits the best into their goals of care. If we do not expect the patient to choose between classes of antidiabetics, why do we expect them to choose a modality of KRT, a treatment on which their lives will be dependent, for the rest of their days? We help patients to understand why it is important to adhere to medication. We shall help them to opt for what’s best, when life is in danger. Our goal as doctors should be how to provide better care for our patients, so instead of drowning them in information that they cannot cope with, we should ask ourselves more frequently “How can we help?”

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\section*{References}


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