What to do when patients and physicians disagree? Qualitative research among physicians with different working experiences

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Received: 4 October 2019 / Accepted: 16 March 2020 / Published online: 2 April 2020
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Key summary points

Aim To understand the challenges experienced by physicians when their opinion on medical decisions differ from those of patients or relatives.

Findings Physicians felt uncomfortable when there was disagreement between themselves and patients or relatives. Frustration was felt when relatives spoke on behalf of the patient, while there was no evidence the desired decision was ever expressed by the patient. A disagreement with a patient was described as being less frustrating, when the patient was able to explain the reasons for making a decision. Differences in background, especially religious, were often mentioned as complicating communication.

Message Efforts must be made to establish a bond of trust between patient, relatives and physician. The use of advance directives should be encouraged. In case of an impasse between a physician and patient or relative, advice can be sought from other professionals.

Abstract

Purpose Impasses between patients, relatives and physicians occur frequently. With the growing attention for shared decision making, it is valuable to know how impasses arise. To understand the challenges experienced by physicians when their opinion on medical decisions differ from those of patients or relatives.

Methods Fifteen physicians with different working experiences, from five medical specialties were interviewed using a narrative approach. Interviews were based on two patient stories provided by the physician. First of a patient (or relative) who did not want to adhere to a treatment the physician deemed necessary, and the second of a patient (or relative) who requested a treatment the physician felt was unnecessary. Data were analyzed using a bottom-up approach, with identification of five themes (autonomy of the patient, communication, emotions, circumstances and metaphors). Twenty subthemes were formed.

Results 693 references were made. Six major nodes were identified: frustration experienced by the physician, role of the relatives, agreement, cultural/religious aspects, comprehension by the patient of the situation and the existence of an established relationship between patient and physician.

Conclusions Physicians felt uncomfortable when there was disagreement between themselves and patients or relatives. Frustration was felt when relatives spoke on behalf of the patient, while there was no evidence the desired decision was ever expressed by the patient. A disagreement with a patient was described as being less frustrating, when the patient was able to explain the reasons for making a decision. Differences in background, especially religious, were often mentioned as complicating communication.

Keywords Distress · General medicine · Terminal care · Treatments

Introduction

In recent years, much effort was invested in implementing shared decision making (SDM). In this process, patients and health professionals together decide on clinical approaches, discussing all information, risks and benefits as well as patient values and preferences. The primary
reason to promote SDM is to respect the autonomy of the patient, being the subject of treatment. Another reason is that engaged, informed patients are more satisfied with the chosen treatment [1]. From the point of view of the physician, the goal is to decide on treatment options in a dialog with the patient, considering guidelines and with respect for his or her professional judgment.

As far back as the mid-1970s, attempts have been made to reduce medical overuse. Nowadays, this struggle continues. Expectations and demands of patients can result in clinicians feeling pressured to provide low-value care. Furthermore, physicians tend to opt more easily to start a treatment, as opposed to withholding treatment. Initiatives such as the Choosing Wisely campaigns seek to advance a dialog on avoiding unnecessary medical tests, treatments and procedures [2]. The goal of these initiatives is to improve quality of care. One of the most important findings in the International Health Policy (IHP) survey was that 57% of Dutch general practitioners (GPs) thought patients received too much health care [3]. A recent survey in the USA in over 2000 physicians revealed the belief among physicians was that 20.6% of overall medical care was unnecessary [4]. Reducing overuse also provides economic benefit. Previous studies reported that at least 20% of health-care spending in the USA was unnecessary [5].

This spiral of events has been described as a “Perfect Health Storm” [6]. Four physician-related factors driving overuse have been identified: physician culture, fee-for-service payment, marketing and the fear of being sued for medical malpractice [6]. Dutch investigators described 15 mechanisms that can lead to excessive and excessively prolonged treatment [7]. Besides suggested treatments, patients or relatives can also request other treatments, based on their own beliefs. Problems arise if in the professional judgment of the physician, the preferred treatment is unacceptable.

We interviewed physicians to gain insight into how they experienced an impasse between themselves and a patient concerning treatment options. The study goal was to understand the challenges experienced by physicians when their opinion on medical decisions differs from those of patients or their relatives.

**Methods**

We interviewed physicians using a narrative approach. This allows participants to share their subjective experiences and by doing so also reinterpret and give further meaning to these experiences [8, 9]. We used this to do an in-depth exploration of how physicians experienced a difference of opinion with a patient or relatives regarding medical decisions. This method particularly suited this study as so far little is known about physicians’ subjective experiences when being faced with patients with a different opinion about their clinical management.

**Recruitment of participants**

We purposively invited 18 physicians to participate. Invitees were working in the south-west of the Netherlands with different years of working experience. Fifteen physicians, from five different medical area’s (internal medicine, general practice (GP), intensive care, surgery, and oncology), agreed to participate. Their work experience ranged from 1 to 35 years. All interviews were conducted face to face. Before the interview, the participants were informed that the interview would deal with “disagreement between physician and patient”. Participants were asked to think of two different cases from their experience: one in which the patient did not want to receive treatment, whereas the physician judged it necessary; and one in which the patient wanted prolonged or more intense treatment, while the physician considered this unnecessary or harmful. No limitations were given regarding type of underlying disease, patients’ age, sex or cultural background. Participants were given written and verbal information and gave informed consent. Ethical approval was granted by the ethical committee of the Erasmus MC.

**Data collection and analysis**

Interviews were digitally recorded and transcribed verbatim with the interviewees’ permission. The interviewer guides were flexible, allowing prompt and open questions to encourage participants to talk in depth about their experiences and perceptions. Basic demographic information of the patients (age, sex, ethnicity and religion) was recorded (see Table 1). It was also noted whether or not there was an existing patient–physician relationship prior to the described situation) and whether or not an impasse between patient (or relatives) and the physician was described by the physician. Basic information of the physicians (sex, medical specialty, years of working experience) was also recorded (see Table 2). The study was facilitated by QSR NVivo 12 software (QRS International Pty Ltd, Melbourne, Victoria, Australia).

After conducting 15 interviews, data were analyzed to see if new themes were emerging. This was not the case; therefore data saturation was reached.

The transcripts were coded by two independent researchers (RvB and LD) to ensure rigorous analysis. The first five interviews were used to compare coding strategies. No significant differences were found. The first author developed further interpretation of the results with regular comments from the other authors.
We used a bottom-up approach to identify recurring themes. This was performed according to the method specified by Braun and Clarke [8].

1. Transcripts were read and re-read to familiarize the researchers with the data.
2. Systematic line by line coding to identify common features.
3. Codes were reviewed to determine potential themes.
4. Themes were reviewed for internal homogeneity and external heterogeneity to ensure coherence and distinction.
5. Themes were identified and named.

Five themes were identified: patient autonomy, communication, emotions, circumstances and metaphors. Twenty subthemes were formed based on the interviews (see Table 3). In total, 693 different references were identified as matching with at least one (sub)theme.

## Results

Fourteen physicians described two different patients. One physician only discussed a patient who wanted to continue treatment (interview no. 4).

In 14 out of the 15 patient stories where the physician wanted to continue treatment and the patient wanted to stop, there was direct communication between physician and the patient. Of the 14 cases where the physician wanted to stop treatment, there was direct communication between the patient in three cases. In the other 11 cases the relatives either played a prominent role, accompanying the patient (two cases) or the communication was between physician and relatives only (nine cases).

Six major nodes were identified: frustration experienced by the physician, the role of the relatives, agreement, cultural/religious aspects, comprehension by the patient of the situation and existing of an established relationship between patient and physician. Quotes are identified by participant number.

### Frustration experienced by the physician

In 16 out of 29 cases, physicians spontaneously reported frustration on their part while dealing with patients or relatives. These comments were made both in relation to cases in which patients wanted to continue treatment, as well as those in which the physicians wanted to continue (8 vs. 8 cases). Physicians talked about their frustration and feelings of helplessness when they were unable to convince the patient or relatives of their views.
I have no words to reach her, to relate to her inner world. (1.1)

In the cases where direct communication with the patient was possible, frustration occurred when the physician felt that the patient did not (or was not able to) understand the severity of the condition or the necessity of the proposed treatment. Physicians also expressed frustration when they felt relatives did not recognize them as being persons having emotions of their own.

And nobody pays any attention to the physician. Same as in the case of performing euthanasia, the one crying, it was me. (4.2)

In more than one interview, physicians described their challenges with relatives who demanded more intensive treatment than the physician felt comfortable with. In many of these cases, the patient had never consented to the requested invasive treatments, and this caused physicians’ discomfort. In other cases, the physician considered the requested treatments futile.

I gave her antibiotics, while that was already going against every fiber of my being, but the husband insisted, because that is what his wife would have wanted. And one of the sisters, the one that was most involved, she is a patient of mine, said “should we be giving all this treatments?” or something like that. (8.2)

Remarkably, in the cases where a patient wanted to stop, against advice of the physician, and no frustration was felt, the patient was described as mentally competent and well aware of the options.

### Role of the relatives

Almost all physicians mentioned the role of the relatives. In nine cases, the relatives requested treatments the physician thought would not benefit the patient. In all cases, this concerned a patient who was not able to speak for himself. Several physicians expressed their doubts whether the wish to prolong treatment was prompted by the concern of the relatives for the patient’s well-being or for their own sake.

We will never know what Mrs. X herself thought on the matter. It was entirely the wish of the relatives. (9.2)

The fear of getting blamed for the death of the patient was mentioned, as well as the influence of religion. In 12 interviews, physicians described that they tried to accommodate the requests made by the relatives, until they felt their actions were no longer professionally and humanly justifiable.

For me, that was the turning point, I was continuing (treatment), not for the sake of the patient, but for the sake of the relatives. And when I realized that, I thought: this must stop. This (treatment) should not be for the relatives, it should be about the patient. (5.2)

In some cases, the physicians wanted to continue treatment, but the relatives expressed a desire to stop, often referring to prior expressed wishes by the patient. They expressed a strong wish to honor the request made by the patient, even when both relatives and physician agreed continuation of treatment would be the better option. When a patient expressed a desire not to start a treatment, all physicians complied, be it reluctantly in some cases.

Physicians described relatives wanted to stop treatment because of complications, and they found it difficult to grant that request, as it was unsure what the point of view of the patient actually was. More importantly, the complications

| Table 3 | Themes and subthemes, number of references and files |
|---------|-----------------------------------------------------|
| Theme   | Subtheme | References | Files |
|---------|----------|------------|-------|
| Autonomy of the patient | Rights | 10 | 9 |
|         | Role of relatives | 72 | 22 |
|         | Responsibility | 33 | 19 |
| Communication | Providing information | 14 | 10 |
|         | Listening | 14 | 12 |
|         | Agreement | 74 | 28 |
|         | Talking about death | 22 | 9 |
| Emotions | Fear | 15 | 6 |
|         | Anger | 8 | 4 |
|         | Hope | 10 | 7 |
|         | Frustration by patient | 4 | 4 |
|         | Sadness | 4 | 4 |
|         | Frustration by physician | 37 | 16 |
| Circumstances | Culture/religion | 17 | 14 |
|         | Financial matters | 24 | 11 |
|         | Comprehension of situation | 9 | 4 |
|         | Environment/setting | 32 | 20 |
|         | Social network | 26 | 15 |
|         | Patients’ disease | 61 | 28 |
|         | Relationship with patient | 38 | 28 |
| Metaphors | | 43 | 16 |
were, according to their professional opinion, an unfortu-
nate but foreseeable result of a treatment the patient had 
consented to. As a result, sometimes the treatment was con-
tinued against the wishes of the relatives.

Agreement

In 28 of the 29 cases, physicians mentioned the topic of 
(dis)agreement. In most cases, they referred to reaching an 
agreement, or at least trying to do so, with patient or rela-
tives. This was sometimes a smooth process and in other 
cases unsuccessful.

And in the end, when he was extubated, we talked to 
him for quite some time, and as a result we agreed on 
what limitations he wanted regarding treatments. 
(14.1)

I had just become an oncologist, and had not yet real-
ized that people could also chose not to start a treat-
ment. So my assumption was that she would want the 
full chemo, and so I told her what the plans were. And 
that is when she turned completely furious. (3.1)

When faced with a difficult, sometimes emergency deci-
sion, agreement was sought with fellow specialists, and in 
most cases this helped the physician to decide what to do. 
Other described strategies were: revisiting the patient or 
relatives and investing time and effort to build a relationship.

Cultural/religious aspects

Frequently, the cultural or religious background of the 
patient and his relatives was mentioned. In the cases of the 
patient (or relatives) wanting to stop treatment, with the phy-
sician advising to continue, there was one comment regard-
ing a cultural background. The physician thought the child 
of a patient would decide on behalf of the patient, but was 
surprised the decision was left to the patient. In other cases, 
the relatives spoke on behalf of the patient, without consult-
ing the patient.

Several physicians described how a difference in back-
ground between the patient, relatives and physician caused 
difficulties in communication.

I think we still make a lot of mistakes, because we are 
not aware of the way bad news should be addressed 
in specific cultures. (…) So, in several years, I found 
out by trial and error that in some cultures it is not 
custumary for younger relatives members, or those 
lower in the hierarchy, to be allowed to deliver bad 
news. They are not allowed to say that someone will 
die (…), or there is an incurable disease. (…) So we 
only found out after a long time (that the patient was 
not told the diagnosis). Because the daughter, who had 
been in the Netherlands for a very long time, spoke 
fluent Dutch, and was highly educated, understood per-
factly well what was happening, but had not told her 
mother, because she just refused to tell her mother the 
bad news, the diagnosis that was made.(1.02)

In the description of the patients, physicians often sponta-
neously emphasized the role of the religious background 
of a patient or relatives. Of the 14 cases where the physician 
wanted to stop treatment, in ten cases a religious background 
was described by the physician. In five cases a Muslim back-
ground was described, in two cases a Christian background 
and in three cases the specific religion was not specified.

This young man stood by his decision, he did not want 
to receive any blood, and accepted the consequences 
that that would be the end of it. His parents struggled, 
but were also Jehovah’s witnesses, so in the end they 
supported his decision, but it was a real dilemma, he 
was so young. (3.2)

Comprehension of the situation

In multiple cases, physicians expressed concerns regarding 
the level of comprehension of the patient of their situation. 
Physicians described their efforts to inform their patient in 
these situations. Multiple visits were made to the patient, 
help was sought from colleagues and time was invested.

And then we try and talk to them (the relatives) a lot, 
explain it, show them the scans, as I do with all the 
patients who suffered a neurological trauma, to make 
it less abstract, and to explain why we are going to 
stop the treatment. But, if they have no clue of what is 
depicted on the scan, that does not help at all. So then 
we try to explain, what are the different functions of 
the brain, and what functions are now gone. (5.2)

Sometimes, patients expected the outcome to be worse 
than the physician anticipated. In those cases, similar efforts 
were made to convey the (in the words of the physician) 
“more objective, expected outcome” to the patient.

Often, people do not know, notwithstanding the severity 
of the situation, everything can turn out just fine. 
(4.1)

Established relationship between patient 
and physician

In the case of an established relationship between the patient 
and physician, impasses occurred sporadically. Physicians 
described agreement was reached in multiple conversations, 
with enough time for all parties to think about the options.
But my supervisor, who had known her for years, said: “she always talks that way when things are not going well”. (9.1)

However, in the case of no previously existing relationship, impasses were more often described.

But well, when you do not know the patient at all, that is a major pitfall (3.1)

Remarkably, an impasse arose in all the cases without an existing relationship in which the patient or relatives requested treatment, whereas the physician wanted to stop. Physicians told about their efforts to get to know the patient, who was in a number of cases unable to communicate. The acuteness of the situation was described as being a major contributor to arising impasses.

Discussion

In the present study, we found that for the physician the impossibility of direct communication with the patient was frustrating. The frustration seemed to stem from the fact that the right sparring partner, being the patient, was unable to take part in the discussion. If the patient was described as mentally competent and aware of treatment options, physicians still tried to convince them to “see the physicians’ way”, but accepted the divergent desired treatment strategy without frustration.

In several cases, physicians felt uncomfortable with starting the desired treatment, that they set clear boundaries. In almost all cases, here was a line the physician was not willing to cross, to preserve professional and moral standards.

Relatives frequently have to make medical decisions for an incompetent patient acting as surrogate decision maker. Advance directives, in which the patient describes treatment preferences could guide the relatives in this process. In this study, no advance directives were mentioned. This is congruent with studies that show the community prevalence of advance care directives remains low, with percentages ranging between 5 and 20% [11, 12]. So, in most cases, the relatives cannot rely on such a document. Possible problems in communicating with relatives were (among others) identified as the failure to reach a shared view of the patients’ medical condition (and prognosis), problems with applying the principle of substituted judgment, difficulties in addressing the full range of end-of-life-decisions and offering the relatives “the wrong choice” [13]. The wrong choice explained as a choice between care and cure, where no cure is possible, as opposed to the choice between prolonging life and quality of life [13].

Although it is to be expected relatives would be adequate in assessing the wishes of the patient, this is not shown by the existing literature. Several studies performed in different categories of patients such as cancer patients, patients with early dementia and dialysis patients, suggest that relatives are often unable to correctly assess the preferences of the patient [14–16]. An interview-based study including 750 patient–caregiver dyads showed that, in case of discordance, patients and caregivers often had an unrealistic optimistic view regarding extent of disease, treatment goals and prognosis [17].

It seems comprehensible that relatives primarily opt for life-prolonging treatment for the patient. In a study on patients on the intensive care, relatives struggled with the conflict between honoring the patients’ wishes and values, and their own emotional problems with “being the one letting a patient die” [18]. We found that agreement was more easily achieved if the patient and relatives were already known by the physician. Establishing a relationship with both patient and relatives in early stages of a disease is important. The triad patient, relatives and physician can prove to be very helpful in case of difficult decisions [19].

In our study, most impasses arose when patients were “new” to the physician as a result of a transfer into a practice or due to an acute presentation in a hospital. Therefore, there was no history between the patient, relatives and physician. Impasses arose especially in the cases in which there was an acute treatment decision to be made (for instance in the emergency department or in the intensive care unit). The acuteness of the situation and the inevitability of the decision are likely to be important factors.

In our study, physicians told the perceived level of understanding the patient (or relatives) seemed to have of the actual situation was an important factor in the arising of an impasse. In a study including interviews of dyads of physicians and patients, unclear expression of values by both patient and physician, Impasses arose especially in the cases in which there was an acute treatment decision to be made (for instance in the emergency department or in the intensive care unit). The acuteness of the situation and the inevitability of the decision are likely to be important factors.

In our study, physicians told the perceived level of understanding the patient (or relatives) seemed to have of the actual situation was an important factor in the arising of an impasse. In a study including interviews of dyads of physicians and patients, unclear expression of values by both patient and physician, as well as the feeling of being uninformed caused uncertainty in both parties [20].

It seems to be a challenging task to fully inform all patients. As described in the interviews, some patients seem to be unable to comprehend the information that is provided. This can be caused by cognitive impairment, fear or unwillingness to hear bad news. A study conducted in the Netherlands showed that patients with an incurable lung cancer “chose” to ignore bad news and showed a false optimism about recovery, not wanting to hear the bad news and only focus on treatment options [21]. A Belgian study advised to gradually deliver the message of a diagnosed incurable, terminal disease and prognosis [22]. Relatives of terminally ill patients told they were informed about the severity of the illness too late in the process (often within 1 month of the patient’s death). At the same time, relatives did not know to what extent they wanted to be informed, and expressed difficulty in comprehending and accepting the message that they were told [23]. In another study, among general practitioners
(GP’s) and patients receiving palliative care, apart from physicians’ availability, the hesitation of both patients and GPs to talk about a bad prognosis was a main barrier to good communication [24].

Cultural or religious differences in background between the patient (or relatives) and the physician played an important role. Physicians described they felt unable to reach patients or relatives from cultures they were not familiar with, and felt that often the relatives had a different perception of treatment or death. This was more often described in cases where the physician wanted to stop treatment, against the wishes of the patient or relatives. Different religions were described, with the majority being a Muslim background, or a specific Christian background (for instance Jehovah’s witnesses). The physicians’ lack of knowledge of the cultural and social background of their patients can complicate shared decision making. Those experiences will for a great part be the basis of the end-of-life preferences of that same patient [25]. Understanding the patients’ explanatory models about illness, treatments and death can help make sense of seemingly unreasonable actions and decisions. The emphasis on patient autonomy and informed consent can clash with family-oriented cultures, where decisions are made by relatives. There is no reason to question requests from the appointed legal representatives, if a patient expressed the wish to let that person make decisions for them. Complying to the legal representative is a direct expression of following patient preferences.

Language differences can cause problems. This did not seem to be an important factor in the interviews, but can most certainly be of importance when communicating with patients. Translation by a relative may influence the content of the conversation, either due to misinterpretation of medical information, or driven by cultural beliefs that patients should be shielded from bad news. Taboo topics may be left out in translation. The use of trained medical interpreters should perhaps be standard of care when dealing with patients who do not speak the same language the physician does.

In conclusion, we found that physicians felt uncomfortable when there was disagreement between themselves and patients or relatives. Frustration was especially felt when relatives spoke on behalf of the patient, while there was no evidence the desired decision was ever expressed by the patient. The physicians were in doubt whether or not the desire for treatment was prompted by the previous wishes of the patient, or stemmed from the personal wish of the family not to lose a relative. Although this was comprehensible seen from the point of view of the family, the patients’ best interest was still the most important factor in the physicians’ decision to not start or stop a treatment. Differences in background, especially religious ones, were often mentioned as complicating communication. Although all physicians were trained in the Netherlands and now working within the same country, with its multicultural diversity, results cannot automatically be extrapolated to other countries or cultures. However, it is likely many of the experienced situations and frustrations are recognizable to physicians and patients in countries and cultures all over the world.

Recommendations

Based on this qualitative research, impasses between physicians and patients are less likely to occur if the patient is well informed, capable of making treatment decisions and there is an existing relationship between patient and physician. Efforts must be made to establish a bond of trust between patient, relatives and physician. The use of advance directives should be encouraged. In case of an impasse between a physician and patient or relative, advice can be sought from other professionals. However, in the end it is still the physician who will have to decide whether or not a treatment is to be started, with the best interest of the patient at heart, even when this is not congruent with the wishes of the family.

Strengths

To the best of our knowledge, this is the first study investigating the reaction of physicians on an impasse between them and a patient or relatives of that patient.

A wide variety of physicians was interviewed, from hospitals as well as general practitioners.

Limitations

There are limitations to this study. First, participating physicians work in the same region in the Netherlands; therefore, the results cannot be extrapolated to other regions of the same country, other countries or different cultures. Second, in acquiring data from interviews, we express their narrative view of their experiences and perceptions.

Author contributions Lisanne Dommershuijzen and Prof. Suzanne van der Vathorst contributed to the study.

Compliance with ethical standards

Conflict of interest None of the authors have any disclosures.

Ethical approval Ethical approval was granted by the Medical Ethical Committee of the Erasmus MC.

Informed consent Participants were given written and verbal information and gave informed consent.
References

1. Stacey D, Bennett CL, Barry MJ et al (2011) Decision aids for people facing health treatment or screening decisions. Cochrane Database Syst Rev. https://doi.org/10.1002/14651858.CD001431.pub4
2. Cassel CK, Guest JA (2012) Choosing wisely: helping physicians and patients make smart decisions about their care. JAMA 307:1801–1802
3. Schoen C, Osborn R, Squires D et al (2012) A survey of primary care doctors in ten countries shows progress in use of health information technology, less in other areas. Health Aff 31:2805–2816
4. Lyu H, Xu T, Brotman D et al (2017) Overtreatment in the United States. PLoS ONE 12:e0181970
5. Berwick DM, Hackbart AD (2012) Eliminating waste in US health care. JAMA 307:1513–1516
6. Emanuel EJ, Fuchs VR (2008) The perfect storm of overutilization. JAMA 299:2789–2791
7. Wal van der G (2015) Just because we can, doesn’t mean we should (Niet alles wat kan hoeft). Steering Committee for Appropriate End-of-Life Care.
8. Braun V, Clarke V (2006) Using thematic analysis in psychology. Qual Res in Psychol 3(2):77–101
9. Gilbert KR (2002) Taking a narrative approach to grief research: finding meaning in stories. Death Stud 26:223–239
10. Liampittong P (2009) Qualitative data analysis: conceptual and practical considerations. Health Promot J Aust 20:133–139
11. White B, Willmott L, Tilse C et al (2019) Prevalence of advance care directives in the community: a telephone survey of three Australian States. Intern Med J. https://doi.org/10.1111/ijm.14261
12. van Wijmen MP, Rurup ML, Pasman HR, Kaspers PJ, Onwuteaka-Philipse BD (2010) Advance directives in the Netherlands: an empirical contribution to the exploration of a cross-cultural perspective on advance directives. Bioethics 24:118–126
13. Lang F, Quill T (2004) Making decisions with families at the end of life. Am Fam Physician 70:719–723
14. Abdul-Razzak A, Heyland DK, Simon J, Ghosh S, Day AG, You JJ (2017) Patient-family agreement on values and preferences for life-sustaining treatment: results of a multicentre observational study. BMJ Support Palliat Care 9(1):c20
15. Dening KH, King M, Jones L, Vickerstaff V, Sampson EL (2016) Advance care planning in dementia: do family carers know the treatment preferences of people with early dementia? PLoS ONE. https://doi.org/10.1371/journal.pone.0159056
16. Miura Y, Asai A, Matsushima M et al (2006) Families’ and physicians’ predictions of dialysis patients’ preferences regarding life-sustaining treatments in Japan. Am J Kidney Dis 47:122–130
17. Shin DW, Cho J, Kim SY et al (2018) Patients’ and family caregivers’ understanding of the cancer stage, treatment goal, and chance of cure: a study with patient-caregiver-physician triad. Psychonolgy 27:106–113
18. Schenker Y, Crowley-Matoka M, Dohan D, Tiver GA, Arnold RM, White DB (2012) I don’t want to be the one saying ‘we should just let him die’: interpersonal tensions experienced by surrogate decision makers in the ICU. J Gen Intern Med 27:1657–1665
19. Mitnick S, Leffler C, Hood VL, American College of Physicians Ethics, Professionalism, and Human Rights Committee (2010) Family caregivers, patients and physicians: ethical guidance to optimize relationships. J Gen Intern Med. 25(3):255–260
20. LeBlanc A, Kenny DA, O’Connor AM, Legare F (2009) Decisional conflict in patients and their physicians: a dyadic approach to shared decision making. Med Decis Mak 29:61–68
21. The AM, Hak T, Koeter G, van Der Wal G (2000) Collusion in doctor-patient communication about imminent death: an ethnographic study. BMJ 321:1376–1381
22. Deschepper R, Bernheim JL, Vander Stichele R et al (2008) Truth-telling at the end of life: a pilot study on the perspective of patients and professional caregivers. Patient Educ Couns 71:52–56
23. Cherlin E, Fried T, Prigerson HG, Schulman-Green D, Johnson-Hurzeler R, Bradley EH (2005) Communication between physicians and family caregivers about care at the end of life: when do discussions occur and what is said? J Palliat Med 8:1176–1185
24. Slort W, Schweitzer BP, Blankenstein AH et al (2011) Perceived barriers and facilitators for general practitioner-patient communication in palliative care: a systematic review. Palliat Med 25:613–629
25. Crawley LM, Marshall PA, Lo B, Koenig BA, End-of-Life Care Consensus Panel (2002) Strategies for culturally effective end-of-life care. Ann Intern Med 136:673–679

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