TITLE: A qualitative exploration of seriously ill patients' experiences of goals of care discussions in Australian hospital settings

RUNNING TITLE: Patients and goals of care discussions

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ABSTRACT

Background: Goals of care (GOC) is a communication and decision-making process that occurs between a clinician and a patient (or surrogate decision-maker) during an episode of care to facilitate a plan of care that is consistent with the patient's preferences and values. Little is known about patients' experiences of these discussions.

Objective: This study explored patients' perspectives of the GOC discussion in the hospital setting.

Design: An explorative qualitative design was used within a social constructionist framework.

Participants: Adult patients were recruited from six Australian hospitals across two states. Eligible patients had had a GOC discussion and they were identified by the senior nurse or their doctor for informed consent and interview.

Approach: Semi-structured individual or dyadic interviews (with the carer/family member present) were conducted at the bedside or at the patient's home (for recently discharged patients). Interviews were audio-recorded and transcribed verbatim. Data were analysed for themes.

Key Results: Thirty-eight patient interviews were completed. The key themes identified were: 1. Values and expectations, and 2. Communication (sub-themes: i. facilitators of the conversation, ii. barriers to the conversation, and iii. influence of the environment). Most patients viewed the conversation as necessary and valued having their preferences heard. Effective communication strategies and a safe, private setting were facilitators of the GOC discussion. Deficits in any of these key elements functioned as a barrier to the process.

Conclusions: Effective communication, and patients' values and expectations set the stage for goals of care discussions, however environment plays a significant role. Communication skills training and education designed to equip clinicians to negotiate GOC interactions effectively, are essential. These interventions must also be accompanied by systemic changes including building a culture supportive of GOC, clear policies and guidelines, and champions who facilitate uptake of GOC discussions.

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Health care systems face a paradox (1) whereby advances in medical technology mean that life can be prolonged (2). Yet, seriously ill patients and their family members frequently prefer comfort care rather than aggressive interventions (1, 3), although there are cultural variations (4). Doctors tend to prioritise longevity (2) with the exception of palliative care providers whose focus is comfort and symptom control (5), and patients often continue to receive medical interventions as they approach end-of-life "EOL" (3, 6, 7). The outcomes for seriously ill patients who undergo interventions (e.g., cardiopulmonary resuscitation 'CPR') are frequently poor (8). Survival after discharge for patients who undergo in-hospital CPR ranges from 0 - 32%, with rates declining with increasing age (9, 10).

Goals of care (GOC) is a communication and decision-making process that occurs between a clinician and a patient (or surrogate decision-maker) to facilitate a medical care plan that is consistent with the patient's preferences and values in the event of clinical deterioration (11, 12). The GOC process focuses on guiding *current* health-care decisions during the episode of care (13), including whether to utilise life-sustaining interventions (14). This differs from the advance care planning (ACP) process which focuses on preparing for *future* health-care decisions when a person can no longer make these decisions themselves (13). The COVID-19 pandemic, where there is the potential for rapid clinical deterioration and respiratory complications, has highlighted GOC discussions as a priority for seriously ill patients (15). GOC discussions are associated with fewer aggressive interventions at EOL, and greater quality of life (7, 16-18).

Despite benefits, there are numerous barriers to effective GOC discussions (19-22). Clinicians report patient and family members' difficulties with accepting a poor prognosis, misunderstanding the limitations and complexity of life-sustaining interventions (19, 23), and

resistance to addressing EOL issues (19, 24, 25). Physician-related barriers include a lack of communication skills (22), discomfort in discussing death and dying (24), and fear of causing distress (26). System level barriers include time pressures (21, 26), lack of quality professional mentorship (24, 26), few guidelines on discussing EOL issues (24), and lack of training around communicating about EOL (21, 24, 26). These factors could explain why a recent Australian study found that only one quarter of patients referred to the intensive care unit (ICU) had a documented GOC discussion (27).

Previous research around GOC discussions has focused on the views of physicians and nurses (19-21, 23, 24, 26, 28) rather than patients and carers (14). Studies that examined the views of patients and families have frequently used questionnaires (2, 11, 29-31). A previous study used a validated questionnaire in interviews with older patients and family members and identified five (of 11) key elements that participants ranked most important for inclusion in a GOC discussion: preferences for care, prognosis, values, fears or worries, and questions (11). However, we still lack indepth knowledge about stakeholders' experiences of GOC discussions (32) including how well patients are prepared for GOC; the preferred timing of GOC, and environmental factors. In Australia, national healthcare standards and hospital accreditation schemes now require GOC discussions as part of comprehensive patient care (33, 34). The importance of aligning medical care with patients' values and preferences is also embedded in international health policies and standards (35, 36). The current study was needed to explore how well the GOC process is working, and to identify areas for improvement. Specifically, this research aimed to explore patients' experiences of GOC discussions in the hospital setting. The study findings will inform the development of an educational intervention for hospital-based clinicians tailored to patients' needs.

METHODS

Design: A descriptive, exploratory qualitative design was adopted using a social constructionist framework (37) which acknowledges the applied nature of the research and a focus on participants' views.

Setting: Participants were recruited from six hospitals in Western Australia and Victoria which had implemented the GOC process. The departments involved were: ICU, respiratory medicine, renal medicine, general medicine, geriatric medicine, internal medicine, and orthopedic surgery. Orthopedic surgical patients were included because these individuals were elderly patients with hip fractures who were at high risk of surgical complications and/or mortality (38).

GOC Discussion and Form: The usual practice is for the registrar or the consultant physician to conduct a GOC discussion with the patient and/or the carer, optimally within 48 hours of admission. Discussions are initiated for patients who present for an episode of care who are at risk of clinical deterioration. A completed GOC form is placed in the patient's notes (appendix 1). The form includes: patient information (Section 1); GOC and escalation plan [four options: all life sustaining treatment; life extending intensive treatment - with treatment ceiling; active ward based treatment – with symptom and comfort care, and optimal comfort treatment – including care of the dying person] (Section 2); summary of discussion (Section 3), and extended use (Section 4).

Recruitment and sampling: A convenience sample of adult patients who had completed a GOC discussion in their current or recent episode of care in hospital was utilised. Inclusion criteria were: Aged 18 years or over; a GOC form and discussion completed; able to

communicate in English. Interviews were conducted until saturation was reached. These findings are part of a parent program of research exploring the experiences of stakeholders in GOC discussions (patients, carers and health professionals).

Interviews: Semi-structured interviews were conducted by Authors 1, 2, 3, 6, and 8. All are experienced in sensitive interviewing and independent of the hospital setting. An interview schedule (appendix 2) was developed by the research team and informed by the literature (11, 26, 28, 39, 40).

Procedure: The researchers asked clinical contacts to identify eligible patients who had had a GOC discussion in their current or recent episode of care. The researcher approached the patient at the bedside or at home (for discharged patients), provided the study information sheet and consent form, and gave a verbal summary of the study. Written consent was obtained. Interviews were audio-recorded (range 8 to 38 minutes) and transcribed verbatim.

Data Analysis. Transcripts were thematically analysed using Braun and Clarke's six phases (table 1). Thematic analysis enables a rigorous data driven analysis (41). An inductive approach (42) was adopted. Authors 1 and 2 randomly selected four transcripts and coded these independently for initial concepts; they met to ensure consistency on the identified coding categories prior to Author 2 coding the remaining transcripts. Authors 1, 2, 6, 8, 9 and 15 regularly discussed emerging themes and examples; any disagreements were resolved by consensus discussion, as per usual practice (43, 44).

Quality: Authors 1 and 2 led the analysis of the transcripts; the other authors contributed to interpretation. Nine coding categories were identified and from these, two over-arching themes

and three sub-themes emerged (see Figure 1). The Consolidated criteria for Reporting Qualitative research (COREQ) checklist was used (45).

Ethics approval for this study was obtained from the relevant institutional Human Research Ethics Committees (EC00270) and (HRE2018-0404).

RESULTS

Participant Characteristics: Forty-nine interviews were initiated with patients, with 38 included in the final data set (table 2). The median time between the GOC discussion and the interview was three days (range 0 to 29 days). Participant characteristics are provided in Table 3; reasons for patients' hospital admission are summarised in Table 4.

Two main themes were identified: Values and expectations, and communication. Three sub-themes were identified within "communication": i) facilitators of the conversation, ii) barriers to the conversation, and iii) influence of the environment. "M" and "F" denote male and female gender, respectively.

1. Values and expectations

This theme centered on the importance of quality of life rather than longevity; having a sense of control; patients' comfort or discomfort discussing EOL issues, and family involvement in discussion of EOL preferences.

Most patients were aware of the seriousness of their condition, they accepted that death was possible, and were comfortable with the idea of life sustaining interventions being withheld if their condition deteriorated: *I do not want to be resuscitated. If I'm going, let me*

go. (F1) They wanted life sustaining interventions only if they could maintain a reasonable quality of life and some independence. Patients did not want their life prolonged if the likely outcome was poor function, impairment, or they perceived they would be a burden to others. Some patients said they were ready for death if it came, and that potentially futile medical and artificial interventions were unwelcome.

if it means you're going to be in hospital for years....with machines keeping you alive, then no, forget about it. (M2)

Most patients valued the opportunity to discuss their preferences and goals for care; for some this was the first time they had been asked about these issues. Having a say and being heard was important, with some expressing feelings of control by being able to articulate what they wanted. Participants used terms such as "empowerment": *It made me feel more empowered (F7)*, and being active rather than passive.

One patient suggested that stating preferences would reduce the burden of decisionmaking on others.

....other people need to know what you want. Otherwise, you're putting an unfair expectation on them.... (M2)

Several patients, however, stated they were not ready to die and would prefer to focus on living provided there was reasonable quality of life.

....if there's any possible way of being resuscitated and not a life-threatening hindrance to other people, I just want to keep going. (M3)

However, not all patients wanted the conversation. Some patients were not ready or comfortable with discussing EOL issues, they did not want to discuss death and dying, and they would have preferred not to have had the conversation. Not wanting to consider EOL was a major factor in this.

No, they didn't tell me what was going on, no I don't want to know, I'm too old now, and: they shouldnot say anything. (F9)

Death is unknown and scary, I don't want to think about it. (F10)

Many patients reported that they had not talked with their family or general physician about EOL issues, despite the majority of patients being elderly and seriously ill: *I don't* want to upset her (F4). Several patients assumed that their family knew their preferences. *I think she knows pretty well that I wouldn't want to be kept alive – have something breathing for me.* (F6)

Some patients who had tried to discuss EOL issues with family members said that the person was dismissive or had attempted to shut the conversation down suggesting family discomfort with or denial of the possibility of approaching death for their family member.

... they'll say, "Don't talk silly, Mum, you'll be here until you're a 100". (F5)

2. Communication

This theme centered on how the doctor's manner and communication style, and the setting influenced patients' perceptions of the discussion.

i) Facilitators of the conversation

Many patients reported that the doctor(s) had a very positive approach. Patients who experienced the GOC interaction positively emphasised patient-centered elements of the discussion including: normalising the topic; using non-verbal behaviours effectively; "avoiding medical jargon, and spending time with the patient. Feeling "heard" and sensing that the doctor was listening and demonstrating understanding of their situation and preferences, were very important to patients.

Introducing and normalising the discussion was experienced positively.

....he prefaced it by saying, "Oh this is a conversation we must have with seriously ill patients". (M11)

The discussion was also viewed positively by patients when the doctor used nonverbal behaviours to create rapport and trust and to show that they were fully engaged in the conversation:she was human. (F12). This reflected a need to feel "human".

[The doctor] came very close to me and spoke to me....She wasn't rough or tough. She was a gentle lady. (F13)

She sat by my side (F13).

....he brings his stature to your level. So there's none of this standing over you. (F15)

Patients appreciated a gentle, conversational style rather than being told what to do.

We just talked to each other, just person to person, that was very nice. (F14)

Time was considered important and most patients felt that the doctor spent enough time with them.

[I never] felt that he was rushed...You were his total focus until you had finished with him. (F15)

For most patients, the conversation was patient-driven and gave them a sense of a partnership and shared decision-making.

....together we did it....We got to the goal in the end. (F17)

It was also helpful when the doctor used simple, clear language: *I understood what he was getting at and what he was saying.* (M11)

He used common language. We don't want technical - I'm not in the medical profession. (F16)

ii) Barriers to the conversation

Several patients reported less positive experiences of GOC discussions because of the doctor's poor communication skills or inexperience; GOC interactions which were not patient-centered were associated with patient distress and dissatisfaction.

A failure by the clinician failure to build rapport during the conversation and failing to explore what was important to the patient was experienced negatively. This patient felt that the conversation was more about the doctor's agenda rather than focused on her needs and preferences.

....there is that kind of problem that she doesn't know me, she doesn't know my family, she doesn't know anything about me.... I don't think she was in tune enough....to have the conversation. It was quite hard...I felt like crying all the time.... (F19)

But it was what she wanted to say. Not what I wanted to say. (F19)

One patient did not feel her wishes were respected.

....no one took me seriously....the [specialist] said, "look, we've all just had lunch so I can assure you we won't be doing any extreme measures. We're too sleepy." (F18)

Several patients experienced an interaction where the doctor was sharp or dismissive.

It's like you're not there, you're an invisible person. (F20)

.... I don't want to see this person. He was very abrupt, really full on and, you know.

(F5)

One patient said her doctor could not communicate effectively and this impaired her capacity to participate in the conversation, and another felt that his doctor was inexperienced.

She doesn't know how to communicate effectively; Only from a medical point of view. (F19)

No. I suspect she hadn't had a lot of experience at this sort of thing. (M2)

Several patients said the doctor used medical jargon which made understanding challenging.

...but they need to remember that we don't talk doctor talk and sort of explain in layman's terms, so you can understand things. (F21)

One patient reported the doctors had a very task-oriented approach, focused on the process and technology not her, such as recording the patient's information onto the computer.

... he's talking to you, but he's directing his remarks to him [the other doctor] and his computer. (F22)

iii) Influence of the environment

Not having a private, quiet place where the doctor and patient could have the discussion without interruption or background noise was experienced negatively. Whilst recognising the constraints of the hospital environment, patients wanted some privacy around and time for the

conversation. One conversation was conducted in a corridor in the emergency department, which felt rushed with many distractions.

....there were nurses and everybody running all around but she sort of had me on a bed at the sideI don't think there was [a curtain]. (F12)

Interruptions to the discussions were also common, which again was distracting and resulted in feelings of being unimportant.

There were quite a few interruptions. (F6)

DISCUSSION

This study explored seriously ill patients' perspectives of a GOC discussion during or soon after an episode of care in hospital. Two overarching themes emerged from the patient transcripts: 1. Values and expectations, and 2. Communication.

Patients' willingness to engage in the conversation depended on how comfortable they were with death and dying, and whether they felt "ready to go". These views were embedded in a desire for quality of life including, minimum suffering, maintaining a level of independence, and not burdening others. These values are consistent with studies where community samples rated dignity, avoidance of pain and suffering, remaining independent (2) and not being a burden (29) as the most important treatment goals. Key elements of the GOC discussion include exploring patients' values and asking about preferences for care in the event of clinical deterioration (11). The clinician needs to work collaboratively with the patient (and/or family) to explore preferences and develop a treatment plan that is respectful

of, and responsive to, values (12). In relation to timing, previous research suggests it is better to initiate GOC early in a patient's episode of care (39). The doctor's clinical judgment alone is an unreliable trigger for GOC because clinicians frequently overestimate patients' prognosis (46). Clear guidelines about when and for whom the GOC process should be initiated need to be embedded within hospital policies and into communication skills training (CST) around GOC. Education around GOC for clinicians will need to cover more than just effective communication, but also when to initiate the conversation (39) and how to create an appropriate setting (47, 48).

Most patients viewed the conversation as *necessary* despite the challenges. It gave them the opportunity to voice their preferences and a sense of control. These findings accord with previous research examining attitudes toward ACP where 91% of older individuals reported wanting to discuss EOL care (49). Many patients report they are comfortable with the topic and are willing to discuss EOL preferences, including life sustaining treatments, if the doctor *just asks them* (30). Other research focussing on patient-centred care (50) and patients' participation in health care decision-making (51) has highlighted that providing patients with necessary information and involving them in decision-making increases feelings of empowerment (52, 53). Some patients, however, reported feeling distressed and several patients did not want a conversation about death and dying. The GOC conversation requires a skilled clinician (54) to prepare patients and to navigate this interaction (55) including responding to emotions expressed by patients and family members (56).

How the clinician communicated during the GOC discussion appeared to influence whether the interaction was a positive or negative experience for patients and their overall satisfaction with the discussion. However, the reverse might also be true, that the clinician's

communication is perceived more negatively if the patient is not ready or open to these discussions. Skilled communication and an interaction based upon trust where the clinician listens, builds rapport, speaks honestly and sensitively with patients about death and dying are important to patients and family members/carers (29, 54). Treating patients with respect and humanity, providing emotional support including compassion, hope and comfort, and being sensitive to cues from the patient (24, 54) are also critical elements of effective communication. These elements align with Epstein and Street's model of communication in cancer care (53) which describes the core functions of patient-centered interactions as: fostering a good patient-clinician relationship; information exchange; responding to emotions; managing uncertainty; decision-making, and enabling patient self-management. Their model also highlights the importance of the clinician's use of nonverbal behaviours to demonstrate an orientation toward the patient of genuine care and interest (53). This model could be a useful way of guiding conversations in this space together with the ReMAP (Reframe, Expect emotion, Map out patient values, Propose a plan) framework for GOC discussions (53, 56).

Physicians, junior doctors and nurses frequently report they feel ill-equipped to have GOC conversations because they lack formal training in communication skills regarding EOL care [23, 24, 48]. Providing physicians with access to CST early in their medical career (53), and in-house to facilitate access is key to effective GOC discussions (57). CST programs for health professionals are associated with improvements in communication skills, particularly interventions that adopt an experiential approach and utilise role-play to practice skills learned (58, 59). Several patients reported the doctor using medical jargon during the GOC conversation which may have compromised the patients' understanding. One study reported junior doctors frequently used jargon in a simulated patient-doctor encounter (60), and a

further study reported radiation therapists used various types of medical jargon in their patient education sessions (61). Effective GOC discussions need simple, jargon-free language (53); analogies to convey complex information; repetition (61), and patients reflecting in their own words (62).

There were challenges during the GOC discussion in relation to a lack of privacy, interruptions, and the doctor having limited time. Large patient volumes (47, 49), crowding in emergency departments (63), a focus on technology and efficiency, and spatial layouts in hospital departments do not support privacy (47), and are barriers to effective communication. These findings suggest that creating a sense of space, time and privacy should be addressed in GOC training and education (47). Organisations that are promoting GOC discussions need to ensure that appropriate time and suitable spaces for these interactions are made available.

Future Research

A research priority is the development and evaluation of CST targeted at equipping clinicians to conduct effective GOC discussions where there is immediate feedback given (58). Evaluations should include objective measures of the effectiveness and impact of CST programs such as videoing in situ GOC discussions between the treating team, patients, and family members, pre-and post-training. While it was beyond the scope of the current research, future studies could examine how different patient characteristics (e.g. age, gender, medical issues) and contextual factors influence patients' experiences of GOC discussions. The perspectives of family and non-family carers and health professionals, focussed on their experiences of the GOC process, also require further exploration.

Limitations

Patients who declined to participate may have been less willing to discuss their experiences of the GOC discussion or they may have differed from the participants in other ways, such as being more unwell or less comfortable with death and dying. Three interviews were brief but were retained because they contained valuable information. The patient sample was predominantly female and Anglo Celtic, so the findings may not reflect male views and those of culturally diverse patients. The study was also based upon retrospective reports of patients' experiences which may be influenced by recall bias. However, most interviews occurred within three days of the patient having the GOC conversation so this likely enhances the integrity of the findings.

CONCLUSIONS

Many patients were willing to engage in a conversation about treatment goals and EOL wishes. When the conversations were patient-centered and conducted in a clear, empathic way, patients were satisfied and the experience was perceived as positive. When the conversations were conducted in a rushed manner or the physician was dismissive, the experience was negative with people feeling fearful and invisible. Privacy was important to patients and they disliked interruptions. Physicians need to ensure some level of privacy and be "present" even when time poor. Patients' values and expectations set the stage for goals of care discussions, however the clinician's communication style and the environment play an important role.

At the individual level, CST for GOC discussions is needed using a range of strategies. Training should include ways of managing the environment to create privacy even in a busy ward. Organisational level changes should center on providing in-house training for

health professionals, sensitive to patients' needs. Systemic changes are also important and include building a culture of promoting and normalising discussions around GOC, clear policies and guidelines, and using champions to encourage uptake of GOC discussions.

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CONFLICTS OF INTEREST

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LEGEND FOR FIGURE

Figure 1. Visual representation of the nine coding categories (yellow), two themes and three sub-themes that emerged from the patient data

Table 1. Braun and Clarke's six phases of thematic analysis (41)

Phase	Title	Description
Phase 1	Familiarisation	Immersion and familiarisation with the data through repeated reading
Phase 2	Coding	Generation of initial codes by systematically identifying and labelling interesting features of the data
Phase 3	Searching for themes	Sorting the codes into themes and extracting representative data
Phase 4	Reviewing themes	Review and refinement of themes
Phase 5	Defining and naming themes	Defining the essence of each theme and naming them
Phase 6	Writing the report	Producing the final themes and writing a narrative

Table 2. Reasons for discontinuation of patient interviews

	n = 11
Patient could not recall the goals of care discussion	8
Patient became distressed and did not wish to continue	2
Patient could not continue due to cognitive impairment	1

Table 3. Patient sample characteristics (N = 38)

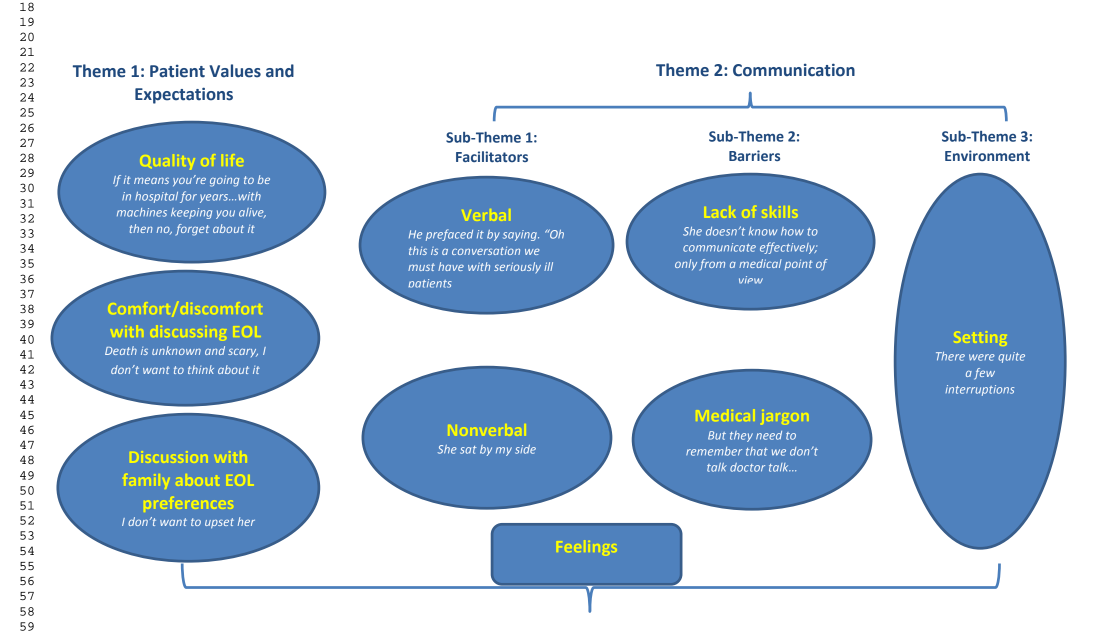
	Mean	SD
Age (in years)	76.2 ^a	10.9
	N	%
Gender		
Female	26	68
Male	12	32
Other	-	-
Country of birth		
Australia	20	53
Other	18	47
Language spoken at home		
English only	34	90
Other	4	10
Aboriginal or Torres Strait Islander		
No	37	97
Yes	1	3
Employment		
Currently employed	1	3
Not currently employed	3	8
Retired	31	81
Household management	1	3
Other	2	5
Relationship status		
Single	1	3
Married/defacto or in a relationship	21	55
Divorced or separated	4	10
Widowed	12	32
Residential location		
Metropolitan area	29	76
Regional area	5	13
Rural or remote area	3	8
Other (overseas)	1	3

Notes. ^aAge range 51-94 years.

Table 4. Summary of patients' self-reported medical conditions (N = 34)

Description	n	%
Heart and circulatory disease	11	29
Lung and breathing difficulties	10	26
Frailty and decline due to older age ^a	8	21
Cancer	7	18
Infection (e.g., urinary tract infection; cellulitis)	5	13
Kidney disease	2	5
Progressive neurological condition	1	3
Dementia	0	0
Other conditions (e.g., infection, diabetes, osteoporosis, anaemia)	10	26

Note. Data was not available for 4 patients. ^aIncludes fractures post-fall.



APPENDIX 1

	Hospital:		Family Name	UMRN	
	GOALS OF PA	TIENT CARE	First Name	DOB	Gender
000240	Ward: Dr / Consultant:		VIV. DOSEDNIKO POLICE		Postcode
+		ective (AHD) (ACP) Guardianship (EPG ve a registered orga e of the patient's do	eak for themselves, who nsible' Relationship: Yes No Yes No Yes No Phan donation decision?	one: No esignation: No	
DO NOT WRITE IN MARGIN	SECTION 2 GOAL Please tick one only and patient, person responsible patient care that will apply	OF CARE complete section 3 e and/or family/carer(: in the event of clinica	over the page to be vali s), please select the most	d. In discussion with t medically appropriate	e agreed goal of
ALATION PLAN	Life extending * Not for CPR	* For Rapid Res * For ventilatory * Specify maxim * For ICU/HDU a * Additional comme	support, including intub num level of support	oation Y	Yes No No
ESCAL	* Not for CPR * Not for ICU * Not for intubation	* For Rapid Res * For ventilatory * Specify maxim * Additional comme	- with symptom all ponse support (intent is symplem level of support	tom control)	Yes No Yes No
HCCZZFMR00H1	* Not for Rapid Response * Not for CPR * Not for intubation * Not for ICU	* For ongoing re * Ensure timely	including care of the eview to identify transition commencement of the e	on to the terminal ph Care Plan for the Dy	rase ving Person

All patients can have Rapid Response based on 'Worried Criteria' or to 'Summon Clinical Review'.

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		I			
GOALS OF PATIENT CARE	First Name	[ООВ	Geno	der
Ward:	Address			Posto	code
Dr / Consultant:	The state of the s				
SECTION 3 SUMMARY OF DISCUS: Goals of Patient Care has been discussed of Patient: Yes No Person Responsion: Name(s) of those present at this discussion:	with: Dat ible: ☐ Yes ☐ No	Family/ca	rer(s):	☐ Yes ☐	No
Is the patient able to fully participate in this d Comments:		□ No			
What is the patient's likely response to CPR	and critical intervention	?			
Patient preferences (needs, values and wish	es):				
☐ Medically-driven decision ☐	ent Care (please tick on Patient wishes	5319	Shared	decision-n	naking
Other information: Doctor's name (please print):	Patient wishes Des	signation:			
☐ Medically-driven decision ☐ Other information:	Patient wishes	signation:			

APPENDIX 2

Interview Guide (Patients)

Introduction

Do you remember having a conversation with your doctor recently about your illness and your preferences for your future care and treatment, especially if your condition gets worse? It probably included your doctor talking with you about what life sustaining interventions you would like if your condition deteriorates and your preferences and goals for your future care.

[You may need to give examples]. For example, your doctor might have talked about whether you want cardio pulmonary resuscitation, or a tube and a machine to help you breathe, or whether you want to go to ICU. Do you recall having that sort of conversation?

I'm going to ask you some questions about your experiences of these discussions. Do you have any questions before we start?

- 1. Tell me about the time your doctor talked to you about your illness and your preferences and goals for treatments and your future care?
- **2.** Who was present at the discussion? *Prompts: Which health professionals? Who else?*
- 3. Who led the discussion? How were you involved?
- 4. Who do you think is the <u>best person</u> to have this conversation with you? Why?
- 5. What things were discussed?
 - **a.** Around the purpose of your admission?

 Prompts: Stage of your illness? Stabilising your condition? Monitoring?
 - b. Around what happens if your condition worsens?

Prompts: Prognosis; options discussed e.g. CPR, insertion of a breathing tube or putting you on a machine to help you breathe; comfort care?

Were there any issues that you found difficult to discuss? Any topics or issues you wish had been discussed that were not raised?

6. How did you feel about the way the doctor communicated?

Prompts:

Was there anything you didn't understand?

Tell me about the way the doctor talked to you? What about the doctor's manner?

How did you feel about getting your preferences out in the open

- with the treating team?
- with your family?

Do you feel your doctor understood your situation and preferences?

How confident do you feel that your preferences and goals for care will be followed?

7. How did you feel <u>during</u> the discussion? <u>Looking back</u>, how do you feel about it now?

8. How satisfied were you with the discussion about your treatment and future care? *Prompts:*

Would you have liked a follow-up conversation with another health professional, for example, a nurse, a day or so later?

How was the timing of the discussion? Was it done at the right time? Would you have preferred the discussion to have occurred earlier or later? If so, when? Did you experience any problems during the discussion? E.g. location, privacy,

interruptions, time limitations?

- 9. What are the pros and cons of having a discussion with your doctor about your treatment and future care?
- 10. How could the discussion be improved?
- **11.** How does your family feel about your preferences for care?

 Prompt: Do you think your family's preferences for your care match your own?
- **12. How did the discussion with your doctor affect your care in hospital?**Prompt: After the discussion, how well does/did your care match your values and preferences?
- 13. Prior to your admission to hospital this time, have you ever had a discussion about your preferences for end of life care, and your values and goals for care?

 Prompts:
 - a. With your family?
 - b. With close friends?
 - c. With your GP?
 - d. Other?

Are there any other comments you would like to make? Do you have any questions for me?

Conclusion: Thank you very much for your time today.