Is it acceptable to video-record palliative care consultations for research and training purposes? A qualitative interview study exploring the views of hospice patients, carers and clinical staff

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Abstract

Background: Research using video recordings can advance understanding of healthcare communication and improve care, but making and using video recordings carries risks.

Aim: To explore views of hospice patients, carers and clinical staff about whether videoing patient–doctor consultations is acceptable for research and training purposes.

Design: We used semi-structured group and individual interviews to gather hospice patients, carers and clinical staff views. We used Braun and Clark’s thematic analysis.

Setting/participants: Interviews were conducted at one English hospice to inform the development of a larger video-based study. We invited patients with capacity to consent and whom the care team judged were neither acutely unwell nor severely distressed (11), carers of current or past patients (5), palliative medicine doctors (7), senior nurses (4) and communication skills educators (5).

Results: Participants viewed video-based research on communication as valuable because of its potential to improve communication, care and staff training. Video-based research raised concerns including its potential to affect the nature and content of the consultation and threats to confidentiality; however, these were not seen as sufficient grounds for rejecting video-based research. Video-based research was seen as acceptable and useful providing that measures are taken to reduce possible risks across the recruitment, recording and dissemination phases of the research process.

Conclusion: Video-based research is an acceptable and worthwhile way of investigating communication in palliative medicine. Situated judgements should be made about when it is appropriate to involve individual patients and carers in video-based research on the basis of their level of vulnerability and ability to freely consent.

Keywords
Communication, method acceptability, palliative medicine, stakeholder consultation, video recording

What is already known about the topic?

- Research and training using video recordings of actual practice can advance understandings of healthcare communication and improve care, but making and using video recordings raises concerns about threats to confidentiality and the risk of negatively affecting patient care.
- There is a concern that when patients are particularly vulnerable – including in palliative and end-of-life (EoL) care – dangers might be greater.
- A previous study in a hospice context found that patients support videoing of their consultation with a doctor in order to support that doctor in their own personal development.

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What this paper adds?

- Hospice patients, carers and clinical staff view video-based research on communication in palliative care as valuable because of its potential to improve communication, care and staff training.
- Hospice patients, carers and clinical staff express some concerns about video-based research including its potential to affect people's communication and threats to confidentiality; however, they do not see these as sufficient grounds for rejecting video-based research.
- They see video-based research as acceptable and useful providing that measures are taken to minimise its possible risks across the recruitment, recording and dissemination phases of the research process.

Implications for practice, theory or policy

- Video recording can be used in research to improve understanding and practice of palliative care.
- Involvement of people – particularly patients and carers – in video-based research is possible on condition that clear safeguards and operating procedures are adopted, including case-by-case judgements about whether individual patients and carers can be involved in research.

Background

Communication is a core part of good palliative and end-of-life (EoL) care.1–6 Previous studies explored healthcare professionals’ (HCPs), patients’ and carers’ views on communication retrospectively through interviews.7–17 Although these studies provided valuable insights on what can count as good EoL communication, they did not examine how such communication can be realised. Video-based research allows one to directly observe and examine how HCPs and patients communicate;18–22 findings from this kind of research can be used to improve healthcare communication.23,24 However, it raises several concerns about threats to confidentiality and the risk of negatively affecting HCP–patient communication, for instance by putting undue pressure on professionals or by inhibiting patients from voicing their concerns.25–35 These risks can be greater in settings where patients’ vulnerability is heightened, as in palliative and EoL care.36–38 Therefore, we need to ask whether it is acceptable to video-record interactions in palliative and EoL care settings.

There is one previous study that sheds some light on whether recording is acceptable. In that study, Hargreaves and Peppiatt39 surveyed 31 day hospice patients who had already had one of their conversations with a doctor video recorded. The majority (27/31) viewed the purpose of the recording (doctors wishing to improve their communication skills) as positive and regarded video recording as a good way of doing so; 30/31 said they would agree to another recording in future. Two of 31 of the patients reported that the camera had affected the consultations; 7/31 that they were unable to forget about the camera; but none felt that video recording had made the consultation less confidential, and none reported that it had made them feel nervous or less willing to talk. That study made some inroads into exploring views about the acceptability of video recording, but the purpose of the video recording was limited to improving doctors’ consultation skills – acceptability of video recording for research was not investigated. Additionally, only patients took part, leaving out other stakeholders. The questionnaire design meant relatively shallow exploration of perceived benefits and risks of video recording, whether anticipated benefits outweigh perceived costs and what safeguards might increase recording’s acceptability and safety. Our study addressed these questions in depth by examining stakeholders’ views on the acceptability of video recording patient and carer conversations with palliative medicine doctors for research and teaching purposes.

Methods

Design

We interviewed key stakeholders at one hospice. We used qualitative, semi-structured group and individual interviews40 to explore their views on the use of video-based research in the palliative care setting and to inform the design of a study where we recorded consultations at the same hospice for purposes of research and teaching about healthcare communication.41

NRES Committee Yorkshire & The Humber (UK) ethically approved the interview study on 29 April 2013 (reference: 13/YH/0127).

Setting

We interviewed stakeholders in 2013 at the same hospice where we planned to conduct the video-based study. The hospice provided outpatient, day therapy and inpatient services, as well as a substantial educational programme for health and social care staff in the surrounding region. Our decision to limit the enquiry to one hospice was dictated by the nature of the study as a stakeholder
consultation preliminary to a subsequent broader study in the same setting.

Recruitment and study sample

We interviewed hospice patients, carers with experience of accompanying patients to healthcare consultations, hospice doctors, senior hospice nurses and communication skills training educators. This was a sample of convenience, formed by stakeholders who were available at the hospice. Our sampling strategy was also purposeful; we involved representatives of all the groups that would later be affected by the video study (we would be recording consultations involving patients, carers and doctors; nurses would help with patient recruitment; educators would use the video recordings thereby generated in communication training). We sought to involve people with diverse characteristics (diagnoses for patients, types of relationship to patients for carers).

Members of the care team made a first face-to-face approach to patients who had capacity to consent and whom the team judged were neither acutely unwell nor severely distressed. Carers of current or past patients were also approached by the care team; several were members of the hospice’s carer and patient user group. The care team gave prospective participants an information sheet; R.P. approached them at least 24 h later to invite them to be interviewed. R.P. invited all educators at the hospice, and all doctors who had qualified as or were training to be palliative medicine specialists (i.e. those at UK consultant and specialty registrar levels). Totally, 11 patients, 5 carers, 4 senior nurses, 7 doctors and 5 educators took part (Table 1). The overall number of participants was dictated by time and financial constraints as well as participants’ availability. We excluded one patient interview from the analysis because the patient had severe dysarthria and their talk was not sufficiently intelligible for transcription. R.P. obtained written informed consent from the participants.

Data collection

We used small group interviews to promote discussion among participants, and so more comprehensively elicit their views. We did two group interviews for doctors, two for nurses, one for educators and one for carers. Individual interviews were more suitable for the patients’ health circumstances and were also conducted for participants unavailable on group interview dates (Table 1).

R.P. designed topic guides in consultation with a project advisory group (see Acknowledgements) and experienced clinicians and communication skills trainers at the hospice, some of whom also took part in the interviews. These multiple involvements were motivated by the collaborative nature of the stakeholder consultation, which was aimed at paving the way for a subsequent video-based study at the hospice. R.P. conducted the interviews. Table 2 contains information on how R.P. introduced herself and the research to the participants, and the topic guides. The breadth of views elicited through these guides was partly limited by the nature of the study, which also served as a stakeholder consultation to support the design of our subsequent video-based study. For instance, while we invited doctors to imagine being asked to take part in a video-based study, we did not do this with participating nurses because we were not planning to record patient–nurse interactions.

| Stakeholder group          | Number of participants and gender | Data collected                                      | Duration (h) | Additional information                                                                 |
|----------------------------|----------------------------------|----------------------------------------------------|--------------|---------------------------------------------------------------------------------------|
| Patients                   | 10 (7 F, 3 M)                    | 10 individual interviews                           | 2.57         | Diagnoses: Motor neurone disease (3), breast cancer + MS (1), cancer of the tongue (1), myotonic dystrophy (1), heart failure + ischaemic heart disease (1), multi system atrophy (1), gastric and ovarian cancers (1), COPD + bronchiectasis + heart failure + osteoporosis (1) |
| Carers                     | 5 (4 F, 1 M)                     | 1 group interview (2 participants)                 | 2.55         | Relationship to patient: wife (2), daughter (2), father (1)                           |
| Doctors                    | 7 (6 F, 1 M)                     | 2 group interviews (one of 5 participants, one of 2 participants) | 1.54         | Role: consultant (5), specialist registrar (2)                                       |
| Senior nurses              | 4 (4 F)                          | 2 group interviews (2 participants each)           | 1.41         | –                                                                                     |
| Communication skills educators | 5 (5 F)                         | 1 group interview (4 participants)                 | 2.24         | –                                                                                     |

M: male; F: female; MS: multiple sclerosis; COPD: chronic obstructive pulmonary disease.
R.P. conducted the interviews. At that time, she held a PhD in Social Sciences and was employed as a Senior Research Fellow at the University of Nottingham. At the start of the interviews, R.P. told the participants about her background in physiotherapy, her experience in conducting video research in healthcare settings and her interest in conducting video research in the future at the same hospice. The purpose of the research, as explained to participants, was to assess their views about the possibility of conducting a subsequent video-based study at the same hospice where the interviews were taking place. In that subsequent study we recorded conversations between hospice doctors, patients and carers (when present). R.P. further told them the purpose of the interviews was to gather views about whether or not collecting video data at the hospice would be acceptable, and why or why not; she also described measures, gleaned from the literature, that could be adopted to make video recording safer and asked the interviewees’ views on those and any other safeguards they deemed important.

### Interview guide

- Patients and companions were asked to imagine that they had been asked to consider allowing a video recording to be made of their next conversation with their hospice doctor, for research on how doctors communicate.
- Senior doctors, who were also potential participants in the subsequent study, were similarly asked to imagine they had been asked to consider allowing video recordings to be made of their next conversations with some of their patients.
- Senior nurses and educators were asked about the overall acceptability of the proposal to video-record doctor patient consultations at the hospice (because the subsequent study did not aim to record nurses and educators).

**Subsequent topics**

- As worded for patients and companions. The same topics, but with amended wording, were covered with other interviewees.

If they said video recording would be unacceptable for them, they were asked:

- To elaborate on their reasons.
- About whether they thought it would ever be acceptable, even though they would not participate personally.

If they said recording would be acceptable, they were asked about:

- What they (and their companions) would want to know about the study.
- How long they would need in order to decide whether to participate.
- The acceptability of prospective verbal assent with written consent sought after recording.
- Whether or not the camera operator should be present within the room.
- Their views on having a remote control available in the room allowing recording to be stopped at any time.
- How long after recording consent should be discussed.
- Whether participants should be given an opportunity to see the recording or receive a copy.
- Whether or not it would be acceptable for people beyond the small research team to view recordings – specifically: other experts viewing them with the aim of strengthening the analysis; other researchers attending presentations of findings; and closed groups of health and social care staff or trainees attending communication skills training. The fact that some participants in the recordings would have died by the time clips were used in such training was explicitly addressed unless the interviewer judged raising this would be distressing for the interviewee.
- Measures that could be adopted to make video recording safer.

Towards the end of interviews, participants were invited to add other thoughts and comments. Throughout, the interviewer attempted to facilitate the raising of relevant matters not specifically covered by the guide.

R.P. had a prior association with two interviewees, both doctors; R.P. had consulted them about conducting research at the hospice and had also observed their work as a formal visitor to the hospice. Four of the consultants she interviewed also took part in the subsequent video-based study. The interviews took place in quiet rooms at the hospice and were audio recorded. R.P. took brief field notes during the interviews and expanded these afterwards.

### Data analysis

M.P. transcribed the interviews verbatim and analysed the data using thematic analysis as described by Braun and Clarke; we broadly adopted thematic analysis as a ‘realist’ method, treating what people said as reflecting their views and perspectives. Nevertheless, we also endeavoured to take into account the way in which the interviewer’s questions shaped interviewee responses. We did not use specialist software; instead, we designed customised forms and tables with word-processing software. M.P. and R.P. met regularly to discuss and refine emerging analyses; differences of interpretation were discussed and resolved. We derived themes from the data inductively. We shared emerging findings with two carers, two doctors and two educators, all of whom were also study participants, seeking and building in their comments. We deemed that these stakeholders’ insider knowledge would help our understanding of the meanings conveyed in the interviews, which contained several references to processes and dynamics specific to the hospice setting. We did not
consult patients in this way because they all had such rapidly deteriorating conditions.

Results

We grouped our analyses into three themes: participants’ views about the value of video-based research, their concerns about its risks and their views on ways of mitigating those risks (Figure 1). The overarching theme is the central dilemma of protecting participants (particularly patients) from the risks associated with video-based research while giving them the opportunity to make their own decisions about involvement in research.

The value of video-based research

On being presented with the hypothetical scenario of their next medical consultation being recorded, 9 of the 10 patients said they would consent to this and to the recording being used in training. One of these nine said he would only agree to audio recording because of how the illness had changed his appearance. The one patient who said she would not consent said this was because of both her changed appearance and the fact that she found herself getting emotional and tearful during consultations and would not want this to be recorded: ‘I am so close to the edge all the while and I wouldn’t want anybody seeing that’ (Interviewee 3). She nevertheless supported the idea of doing the research and thought that others would have different reactions to the idea of being recorded.

Participants in all stakeholder groups commented that communication was a crucial aspect of good care, thought there was scope and need for improving knowledge and training and viewed video-based research as a good vehicle for doing so. Participants in different groups approached this theme from different angles. The educators reported the need for more resources for training. Several patients mentioned contributing to others as a reason for participating in video-based research. One patient said, ‘I want to know that by doing this [research, it] is gonna help someone at the end of the day’ (Interviewee 9). Another patient compared the authorisation to use her voice and image to organ donation: ‘This is the same when you ask if somebody died whether you donate your organs. You’re dead. So why not? If you’re going [it’s] going to help somebody else’ (Interviewee 6). Carers suggested that participating in research could in itself be beneficial to those who take part; for instance, a carer expressed that involvement in research can be one of the ways to make sense of a person’s dying: ‘something good has got to come out of something so bad’ (Interviewee 13).

What concerns does video-based research raise?

Interviewees expressed concerns about possible risks associated with the recruitment, recording and dissemination phases of video-based research.

Recruitment phase. Several participants referred to power imbalances between patients and staff, leading to a risk that patients and carers could feel obliged to take part. According to a carer, ‘they [patients] will probably say yes to a doctor. My mother-in-law would have said yes to any doctor’ (Interviewee 13). Some participants mentioned circumstances where patients’ and carers’ vulnerability would be heightened and informed consent could be at risk: ‘They will come in to that appointment, particularly if it’s a crisis, I could not retain anything’ (Interviewee 21 – carer); ‘You wouldn’t do that [seek written consent] immediately after the consultation […] because there may be a lot of distress’ (Interviewee 20 – senior nurse). These considerations led to discussions about whether and when participants should be approached and given a choice to take
part. There was consensus that some individuals would be so vulnerable that they should not even be approached, for example, people who were obviously very distressed:

If such a person had reached the door of [the hospice] it would perhaps not be appropriate to ask and that would just be another raft of questions and dilemmas to offer them [when] they were already dealing with a lot of stuff. (Interviewee 14 – carer)

At the same time, several participants emphasised that there should be equity of opportunities for patients to be involved and furthermore that they should be regarded as capable of making their own decisions. One senior nurse commented, ‘I don’t think you should discriminate amongst age or disfigurement […] you shouldn’t rule out people’ (Interviewee 20). At the other end of the spectrum, one educator said that sometimes it is not in a patient’s best interest to be invited to take part and that professionals should make those decisions on their behalf: ‘I think they say they want that but […] they don’t always know what will be helpful [for them]’ (Interviewee 18). Notably, none of the patients expressed concerns about feeling pressured in the recruitment stage; however, three mentioned that they would need time to consider whether to take part in the research.

Summarising the above, there was some divergence in the perspectives voiced within different stakeholder subgroups; some carers and HCPs voiced concerns relating to patients’ recruitment whereas patients did not express such concerns. This discrepancy connects to the overarching theme of this study (centre of Figure 1): the dilemma of protecting participants (particularly patients) from the risks associated with video-based research while giving them the opportunity to make their own decisions about involvement in research. While it is important to let patients make up their own mind about participation, HCPs and researchers have a duty to protect patients in circumstances where the initial approach to take part in research could be in itself a source of unjustifiable distress, or when there are reasons to believe that patients’ conditions could hamper their ability to provide informed consent.

Recording phase. Within each stakeholder group, there was disquiet that recording might affect communication within the consultation, particularly by inhibiting patients and carers from disclosing concerns and expressing distress. According to one carer, ‘I think my only concern would be that the person being asked to sit in on a video I might be anxious and I guess some patients themselves may be aware of it and […] that may therefore make them a bit more self-conscious or less able to open up or share sort of deepest feelings, some concerns. (Interviewee 16)

These views resonate with the responses of the patient, mentioned above, who had said she would not want to be recorded (Interviewee 3). She described conversations with her doctor as moments of heightened vulnerability which she would not want caught on camera (see above). Other patients expressed less concern about the impact of recording on within-consultation communication than did other participants. Indeed, 6 of the 10 patients said they anticipated that recording would have no effect on their communication with their doctor; another two expressed the same view while acknowledging that other patients might be affected.

Broadly, the level of concern about recording consultations varied, and no one opposed the overall idea of the research on these grounds. Rather, it was recommended that the risk of influencing in-consultation communication should be minimised through safeguards.

Dissemination phase. There were few direct comments about concerns relating to using participants’ images in the dissemination phase. One patient raised issues of anonymity and privacy: ‘I don’t want this to come back to me’ and ‘I don’t want it put on BBC’ (Interviewee 5). An educator alluded to the risk of objectification: ‘obviously you are using patients in this instance as means, not ends in themselves. So it will be of no benefit to the individual patient’ (Interviewee 18). A carer expressed some disquiet at the idea of seeing the recording after the patient’s death: ‘I’d think I’d find it bit weird down the line, but if I wasn’t to see it, and it was just medical students [that would be okay]’ (Interviewee 10). Some interviewees suggested the recordings might be distressing for some audiences if used in communication training. As previously noted, all nine patients who said they would agree to recording also said they would consent to use of the recording in training.

The vulnerability of professional participants was raised in doctors’ interviews: ‘I suppose partly I’m slightly worried with this project that actually what would come out [is that I am] not an expert on communication but that actually I’m doing it all wrong’ (Interviewee 16 – doctor). This linked to the idea of having one’s performance exposed and subjected to scrutiny: ‘It’s much more exposing to us than it is to the patient I think’ (Interviewee 17 – doctor). A carer, who had previously worked as a social care professional, said, ‘Perhaps if I was the professional person being asked to sit in on a video I might be anxious about how I would turn out’ (Interviewee 14).

What safeguards should be used?

For the recruitment phase, proposed safeguards included measures to ensure that patients and carers who were evidently in great distress would not be approached. For these
individuals, informed consent and recording procedures would impose an unjustifiable additional burden. Interviewees emphasised that people approached to participate would need to be fully informed about the purpose of the study and that any kind of pressure should be avoided. Explaining the research to potential participants and giving them time to reflect was seen as important:

I think it’s about being able to explain to patients and carers in fairly succinct terms what our aims are. And for them to be able to reflect on that. Not make a quick decision because actually it’s quite big really. (Interviewee 20 – senior nurse)

Some interviewees remarked that it should be clear to people that they were being invited but not obliged to participate:

A lot of people just tell people what to do, like ‘you’re going to do this’. [...] It has to be what they want to do and not what you want to do to me. So has to be clear that it is an invitation. (Interviewee 5 – patient)

Interviewees also observed that it would be important to make staff and hospice users aware that the study was taking place via briefing sessions and posters in public areas: ‘It’s about setting your stall out for everybody to understand what’s going on’ (Interviewee 20 – senior nurse). The opportunity to opt out of the research after the recording was also seen as important. Proposed safeguards for the recording phase included measures to make the recording unobtrusive, to stop the recording during the consultation and the offer of extra support to people who had been recorded. Safeguards for the dissemination phase included use of anonymisation techniques, restriction of data access to selected audiences and screening the data in safe environments where spectators (e.g. communication skills courses trainees) could be supported.

Discussion

We interviewed hospice patients, carers and clinical staff about the acceptability of video recording palliative medicine consultations for research and teaching purposes. Compared to previous research that has supported the acceptability of video recording for the more limited purpose of improving doctors’ communication skills, our study explored the views of a range of stakeholders, and – by virtue of its qualitative design – allowed a broader and deeper exploration of reasons underlying judgements around acceptability of video recording for communication research and training.

Participants expressed concerns about the risks associated with video-based research; although these concerns did not outweigh the overall judgement that video-based research was acceptable and worthwhile, they raised a more situated dilemma of whether some participants should be invited to take part in a video-based study. Our findings raise the question: who ultimately makes the decision of whether to invite patients and carers to take part in video-based research? Most interviewees acknowledged that patients and carers should be given the opportunity to decide for themselves about involvement in research; the patients we interviewed endorsed this view. However, our stakeholders also emphasised that a line had to be drawn somewhere; this meant that severely distressed patients and carers should not be approached. In our subsequent video-based study at the same hospice, we adopted this as an exclusion criterion for patient and carer involvement; the care team made judgements about whether or not patients and carers could be approached on a case-by-case basis.

Broadly, we believe that the dilemma of patient and carer recruitment to video-based research cannot be solved once and for all and that it involves local judgements taking into consideration situational factors. Elsewhere we have proposed a framework to support researchers’ situated decisions on these matters. This framework comprises several safeguards (e.g. the option for participants to re-evaluate their consent in time) and diversified modes of patient, carer and HCP involvement (e.g. the option to have a consultation audio recorded, rather than video recorded).

Our findings should be taken with some caution. The people we interviewed had obviously already agreed to take part in a study (in which they were being recorded!); therefore, their favourable take on video-based research does not necessarily reflect the full range of attitudes. Some may also have been prone to express positive views in deference to the researcher – knowing that she was hoping to do video-based research at the hospice in the future. Unlike Hargreaves and Peppiatt’s study where patients experienced having a consultation recorded, we presented our participants with a hypothetical scenario. We also acknowledge that the number of participants in each subgroup was small and that therefore we cannot claim to have captured the full breadth of possible views on the topics explored in this study. Finally, we did not explore nurses’ reactions to the idea of being recorded; this remains an area of exploration for future research. While these limitations invite some caution when interpreting our findings, our study suggests that video-based research is positively valued by stakeholders on condition that measures are taken to manage risks associated with it. We believe our study also demonstrates the value of investigating participant acceptability of video recording; here, we have addressed video recording for research and training. Future research should examine participant acceptability of video recording for other purposes, including surveillance and quality improvement.

This could generate significant insights on the ethical implications of video recording people’s activities – an increasingly pervasive presence in our daily lives.
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Data availability
The primary data for the study consist of a corpus of audio recorded interviews. The authors’ study protocol does not allow them to share the data beyond the research team in order to protect the participants’ confidentiality. However, the pseudonymised transcripts of the interviews examined in this paper can be obtained upon request by contacting the first author.

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