Recruiting to surgical trials in the emergency setting: using a mixed methods study to understand clinician and patient perspectives

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Abstract

Background: Undertaking randomized clinical trials (RCTs) in emergency surgical settings is associated with methodological and practical challenges. This study explored patients’ and clinicians’ perspectives associated with the conduct of an RCT comparing laparoscopic and open colorectal surgery in the acute setting.

Methods: All eligible patients screened and enrolled for the ‘Laparoscopic versus open colorectal surgery in the acute setting (LaCeS)’ multicentre, randomized clinical feasibility trial in five UK NHS Trusts were invited to respond to a survey. Patients and healthcare professionals were also invited to take part in semi-structured interviews. Survey and interviews explored the acceptability of the feasibility trial. Interviews were audio recorded, transcribed verbatim, and analysed using thematic analysis. Survey data were analysed descriptively to assess patient views of the trial and intervention.

Results: Out of 72 patients enrolled for the LaCeS RCT, survey data were collected from 28 patients (38.9 per cent), and interviews were conducted with 16 patients and 14 healthcare professionals. Thirteen out of 28 patients (46 per cent) had treatment preferences but these were not strong enough to deter participation. Twelve of the patients interviewed believed that their surgeon preferred laparoscopic surgery, but this did not deter them from participating in the trial. Half of the surgeons interviewed expressed the view that laparoscopic surgery was of benefit in this setting, but recognized that the need for research evidence outweighed their personal treatment preferences. Eight of the 14 recruiters reported that the emergency setting affected recruitment, especially in centres with fewer recruiting surgeons. Interviewees reported that recruitment was helped significantly by using surgical trainees to consent patients.

Conclusion: This study identified specific challenges for the LaCeS trial design to address and adds significant insights to our understanding of recruiting to emergency surgical trials more broadly.

Introduction

Recruitment to surgical randomized clinical trials (RCTs) continues to be challenging1-3, with 20 per cent discontinued early, most commonly due to recruitment difficulties4. Qualitative research provides an opportunity to identify recruitment challenges by exploring patients’ and healthcare professionals’ experiences5.

When recruiters are not in a state of personal clinical equipoise, they can be reluctant to approach patients who they feel are better suited to one treatment than the other. Recruitment efforts can also be undermined by several issues, including the difficulties communicating the trial aims, problems describing the concept of randomization, and the concern about the effect of recruitment on the doctor–patient relationship. There is also the added workload involved with a trial to be considered5,6.

Patient-level recruitment barriers are complex and often trial specific7, and include difficulties understanding the concept of randomization, and patients wanting their clinician to choose the ‘best’ treatment for them8,9. Clinical trials in acute medicine investigate seriously ill or injured patients, which makes it more difficult for them to reach an informed decision about trial involvement10.

Emergency general surgery is one of the most common reasons for admission to hospital. Of the 30 000 people a year undergoing major abdominal surgery, a third are for emergency colorectal pathology11.

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This is an Open Access article distributed under the terms of the Creative Commons Attribution License (https://creativecommons.org/licenses/by/4.0/), which permits unrestricted reuse, distribution, and reproduction in any medium, provided the original work is properly cited.
The aim of this study was to investigate patient and clinician perceptions of the intervention (laparoscopic surgery) and trial processes to improve trial recruitment.

### Method

The ‘Laparoscopic versus open colorectal surgery in the acute setting’ (LaCeS) multicentre randomized feasibility trial recruited from five UK NHS Trusts and randomized patients presenting with acute colorectal pathology requiring resection, to receive laparoscopic or open surgery. The trial was registered with registration number: ISRCTN15681041 (http://www.controlled-trials.com). The target sample size was 66 patients, and 64 patients were randomized to the trial.

Two previous emergency colorectal surgery trials closed early, in part due to poor recruitment. LaCeS was conducted between July 2016 and November 2017 and included qualitative work to understand patient and recruiter experiences to identify recruitment challenges to inform the development of a training package for a definitive trial. Interviews were undertaken by researchers who were independent of the trial recruitment team. Patient and staff interviews were audio recorded (notes were taken for one staff interview), then transcribed verbatim for analysis and anonymized. Supplementary material reports an overview of the thematic structure.

UK NHS Research Ethics Committee (REC) approval was granted for the study (REC reference: 15/HY/0542).

![Radar plots](image)

**Fig. 1 Radar plots**

a Patient views of the surgery options. b Patient views of the trial (survey responses).
Design
The data collection and analysis were underpinned by a pragmatic interpretative framework, with both quantitative (survey) and qualitative (interview) data collected to identify key barriers and levers to trial recruitment. The mixed-methods approach was used to gather the feedback from as many eligible patients as possible, accepting that many would not consent for interview. The survey provided an overview of patient views of the trial and informed the sampling for the interviews, including the identification of topics to explore at interview. The interviews provided the opportunity to explore issues identified by the survey in depth; both survey and interviews were given equal status. Interviews with staff explored the challenges to recruiting in the emergency setting and identified ways that we could support patient recruitment.

Patient survey and interviews
All patients approached for trial involvement were invited to complete a short (paper copy) survey about the recruitment and randomization process 7 days after surgery. Patients completed the questionnaires independently, and responses were anonymous. Questions asked patients their views about being approached for the trial and the intervention options, with participants asked to respond on a 1 to 5 Likert scale, with

Fig. 2 Balancing laparoscopy versus open surgery: patients’ perspectives

Fig. 3 Randomization and clinician equipoise: patients’ perspectives
strongly disagree, indicated by 1 and strongly agree indicated by 5 (Supplementary material). To encourage participation no demographic information was collected.

All patients approached for the trial were also eligible to be interviewed. Patients were invited to participate in a semi-structured face-to-face (own home) or telephone interview to discuss their experiences of trial recruitment and participation (Supplementary material). Interviews continued until no new concepts were found during analysis16. Interviews were conducted by one researcher between September 2016 and October 2017.

Staff interviews
E-mail invitations were sent to 20 staff from the five sites (three to five staff per site), to ensure that staff of all levels on the recruitment log were represented (consultants, surgical trainees, nurses, and research nurses), including surgeons on the recruitment log who had not approached patients for the trial. Two researchers conducted telephone or face-to-face (in hospital) interviews. Separate topic guides were developed for patients and staff using the existing literature2,17,18 (Supplementary material).

Outcomes of interest
The outcome of interest was to evaluate patient views of the surgery and experiences of being approached for and taking part in the trial; also, the staff experiences running the trial, and approaching and consenting patients to the trial were explored.

Data analysis
Survey data were analysed using descriptive statistics. The qualitative sample size was not pre-determined for this analysis, but purposively selected to ensure diversity in age, intervention received, and sex. Line-by-line coding used an inductive approach, with a focus on content and meaning. Codes were grouped and a matrix was used to structure the data using the principles of thematic analysis19 in NVivo20. Patient and staff interviews were analysed separately. In each case, the first four transcripts were independently coded using an inductive approach by two experienced, postgraduate qualitative researchers. Codes were compared and a coding framework was agreed. The coding framework was then applied to the remaining transcripts. Codes were discussed and recategorized and themes were generated by authors with regular checking of the data. The data quality from phone and face-to-face interviews was assessed by examining the length of time that topics were discussed for each format; phone interviews were marginally shorter, but all topics were covered. The survey data and interview findings were then cross-tabulated to identify where the qualitative data provided explanation for the survey results, and these are presented within the analysis. The analysis was further refined by discussion of the initial themes and subthemes in trial meetings that included clinical colleagues, and the patients’ representative involved in the study. Patients’ perspectives were analysed, and the following topics were identified: laparoscopy versus open surgery, randomization, and clinical equipoise. Clinicians’ perspectives were analysed for the commitment to the trial, patients’ exclusion, their response to patients’ preferences, and the impact of the on-call rota on recruitment.

Results
At the close of trial recruitment, 28 of 72 (38.9 per cent) patients approached for the trial had completed feedback questionnaires (27 consenters and one decliner)12.

Twenty-two patients (30.5 per cent) approached for the trial consented to an interview and 16 trial participants (22.2 per cent) were interviewed. In total, only eight people declined trial participation12, and no trial decliners consented to interview (Supplementary material). Interviewees reflected the balance between trial arms, with interviews conducted with 6 out of the 33 patients randomized to laparoscopic surgery, and with 7 out of 31 randomized to open surgery. Younger trial participants were more likely to consent to interview, with 5 out of the 16 people aged 18–49 years (31.2 per cent) who took part in the trial consenting to interview, compared with only 1 patient out of the 12 (8.3 per cent) who were aged more than 80 years. Patient interviews lasted 36 to 80 min, and no new information was gained after 13 interviews, but continued for a further three interviews to ensure no new ideas were missed. Fourteen of the 20 staff approached (70 per cent) consented to interview, all of whom were actively involved in trial recruitment.

Staff interviews lasted between 22 and 72 min and no new ideas were identified after 12 interviews so were stopped at 14 participants (Supplementary material).

Quotes are provided to support the analysis. Sex, age range, and procedure are provided for patients. The role of the clinician is provided alongside quotes to aid interpretation.

Patient perspectives
Interviewees were aged from 18 years to 80 years, with all age ranges represented (Supplementary material). All respondents were of white ethnicity, and all participated in the trial. Presenting problems included diverticular disease and inflammatory bowel disease (further details are not provided as this may identify individuals). Five patients opted for telephone interviews; the remainder were interviewed face to face in the patient’s home or on university premises. Thirteen out of 28 patients (46 per cent) patients had treatment preferences but these were not strong enough to deter participation.

The survey investigated whether the patients understood the aims of the trial and the difference between the treatment options. Most consenting patients believed that both surgical options were appropriate for them (on the Likert scale, the median for consenters was 4), although there was a slight treatment preference towards laparoscopic surgery (median for consenters of 3), compared with preference for open surgery (median of 2) (Fig. 1). Some interviewees reported that their previous experience of surgery (both open and laparoscopic) influenced their decision to take part in the trial, as the trial gave them a chance of receiving laparoscopic surgery. However, others focused on the importance of a good outcome, rather than the mode of access. Despite a general preference for laparoscopic surgery, more than half of interviewees believed that open surgery would facilitate better visualization of the surgical field and ensure that things would not be ‘missed’ but recognized the cost of open surgery was significant scarring and an increased risk of infection and hernia. A few agreed to be randomized even though they were convinced that they would need open surgery anyway. Most patients believed that laparoscopic surgery would shorten their hospital stay but valued the reassurance that their operation would be converted...
to open, if necessary. Fig. 2 provides quotes that demonstrate patient perspectives related to this topic.

The survey indicated that patients were generally not worried about randomization (on the Likert scale, median of 5), and most agreed (25 patients, 89 per cent) with the statement ‘I understood how my treatment option would be chosen’ (median of 4) (Supplementary material); however, the interviews revealed that a small minority misunderstood what randomization actually involved. Survey data showed that participants trusted their doctor’s explanation of the study (median of 5), and that they participated to help with medical research (median of 5), or so that other patients would benefit in the future (median of 4), a view that was echoed in the interviews. Patients were largely in personal equipoise, feeling that given the evidence, both options were appropriate for them. Patients expanded on their reasons in the interviews, with trust in their clinical team being a key driver to accepting the trial. Patients who were very unwell, or processing news of a serious diagnosis recalled little reassurance that the clinical team provided. For a small minority, a family member helped them make the decision to participate, and none regretted involvement.

The survey data suggested that on the whole patients did not feel that their surgeon had a particular preference for one treatment over the other, but some interviewees also noted that the surgeon was taking part in the trial and so they assumed that their surgeon felt the question needed answering, or that they were in favour of laparoscopic surgery. Patients did not think there was anything unusual about this and were not concerned. Figure 3 provides example quotes regarding patient views about equipoise.

Clinician perspectives

Half of the surgeons interviewed expressed the view that laparoscopic surgery was of benefit in this setting, and nine (64 per cent) believed that the need for research evidence outweighed their personal treatment preferences (Fig. 4).

Interviewees provided possible reasons for colleagues not participating in the trial, including: a lack of awareness of the trial, lack of personal clinical equipoise, and a belief that the trial will be difficult to recruit to. Surgeons occasionally expressed initial regret about offering the trial to a patient and were surprised by the positive outcome (Fig. 4). Despite their claim that they could set aside their own personal views, four out of eight consultants (50.0 per cent) expressed unwillingness to enter patients with ulcerative colitis into the trial as there was a strong belief that the benefits of reduced scarring from laparoscopic surgery outweighed any possibility of poorer outcomes, and so clinicians often did not raise the idea of the trial to these patients.

Recruiters recalled that few patients expressed a treatment preference, but when they did, recruiters were often reluctant to discuss these with the patient; some citing a belief that this was not ethical, others believing that if the patient provided a sensible justification, that this should be accepted (Fig. 4). Finally, at most sites, specialist trainees were involved in identifying and recruiting patients but the unpredictability of cases, together with the on-call rota meant that some eligible patients could not be approached because the surgeon on call could not operate laparoscopically on patients requiring lower gastrointestinal surgery. In other instances, identified patients could not be operated on because the on-call surgeon was not in personal equipoise and refused to enter the patient into the trial. When only one or two surgeons were involved in the trial, the on-call rota meant that recruitment patterns were sporadic, with significant intervals where no patients could be approached, making monthly recruitment rate targets inappropriate. Surgeons at these sites were frustrated about how many eligible patients were missed (Fig. 4).

Discussion

The aim of the LaCeS feasibility trial was to establish whether it is possible to recruit to a surgical trial in the emergency setting and
to ensure that trial processes and the intervention were acceptable to patients and healthcare professionals. The trial was acceptable to patients but some selection bias on the part of the recruiters, a lack of engagement by some clinicians, and a lack of personal clinical equipoise on the part of some clinicians were barriers to recruitment.

In this trial, potentially eligible patients were identified by specialist trainees and consultant surgeons. This approach worked well, but the complex on-call rota meant that some patients were not recruited because there was no participating surgeon on call. This is in sharp contrast to the elective setting where the flow of patients is steady and relatively predictable. In common with other trials, recruitment practices could be at odds with the trial protocol. Staff interviews revealed that when recruiting patients with ulcerative colitis, clinicians were reluctant to risk patients receiving open surgery. This meant that some patients were actively excluded from the trial by clinicians, a problem that has been reported elsewhere. The need for personal equipoise is debated, but staff interviewees suggested that some surgeons did not recruit due to an absence of personal equipoise, and clinician preference has been identified in a recent survey as a key reason that trials fail to recruit.

The survey showed that patients were not worried about randomization, but the interviews revealed some misconceptions; in particular around the nature of randomization. This phenomenon is well recognized. In the emergency setting time to discuss the trial is limited and patients may be anxious, and this may contribute to this difficulty, however, as recruitment interviews were not recorded we cannot determine where the issue lies.

Clinicians believed that patients would have a strong preference for laparoscopic over open surgery, but the survey and patient interview data, as well as the recruitment data, do not support this conclusion. The interviews provide insights into the reasons for this. Unlike studies with a placebo arm, which are often difficult to recruit to, all patients were to receive their planned surgery and only the mode of access to the abdomen varied. For most patients, the outcome of their surgery was viewed as more important than how surgical access was achieved. Although laparoscopic surgery was viewed as preferable to open surgery because of the reduced risk of infections and quicker healing, patients also believed a keyhole approach reduced the visual field, and some worried that things might be missed, and were reassured that their surgeon could revert to an open procedure if needed.

The findings reported are helpful for understanding some of the specific recruitment issues around recruiting to trials in the emergency setting. Only eight eligible patients declined trial participation, which speaks strongly to the nature of the interventions compared in this trial. In contrast to many surgical trials, both interventions were surgical, and so were viewed as comparable (and generally acceptable) by patients. Although patients participating in the trial were interviewed, no trial decliners consented to an interview, so the voices of this group are missing from the data. The response rate to the patient feedback survey was also low (38.9 per cent). Recruiters were invited to offer the survey to all eligible patients, but staff were reluctant to ask patients to complete the survey when they had declined the trial. The staff taking part in the trial were also interviewed, and their experiences was collected, but clinical staff who were on the recruitment log, but did not recruit any patients, declined participation, so this study offers only a partial understanding of the reasons why some staff did not engage well with the trial.

This paper adds to the literature on patient recruitment in the emergency setting. The recruitment was helped by the support of junior doctors identifying eligible patients and hindered by on-call rota, clinical pathways, and a lack of personal clinical equipoise. Monthly recruitment targets were unsuitable for the sporadic nature of emergency surgery and for trials in the emergency setting and so 3-monthly targets may be more helpful; unlike elective surgery, emergency surgery is responsive, so recruitment rates cannot be managed in the same way.

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### Disclosure

The authors declare no conflict of interest.

### Supplementary material

Supplementary material is available at BJS Open online.

### Data availability

Data are not available as permission was not requested from participants at the time of study.

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