PROcalcitonin and NEWS2 evaluation for Timely identification of sepsis and Optimal use of antibiotics in the emergency department (PRONTO): protocol for a multicentre, open-label, randomised controlled trial

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ABSTRACT

Introduction Sepsis is a common, potentially life-threatening complication of infection. The optimal treatment for sepsis includes prompt antibiotics and intravenous fluids, facilitated by its early and accurate recognition. Currently, clinicians identify and assess severity of suspected sepsis using validated clinical scoring systems. In England, the National Early Warning Score 2 (NEWS2) has been mandated across all National Health Service (NHS) trusts and ambulance organisations. Like many clinical scoring systems, NEWS2 should not be used without clinical judgement to determine either the level of acuity or a diagnosis. Despite this, there is a tendency to overemphasise the score in isolation in patients with suspected infection, leading to the overprescription of antibiotics and potentially treatment-related complications and rising antimicrobial resistance. The biomarker procalcitonin (PCT) has been shown to be useful in specific circumstances to support appropriate antibiotics prescribing by identifying bacterial infection. PCT is not routinely used in the care of undifferentiated patients presenting to emergency departments (EDs), and the evidence base of its optimal usage is poor. The PROcalcitonin and NEWS2 evaluation for Timely identification of sepsis and Optimal (PRONTO) study is a randomised controlled trial (RCT) in adults with suspected sepsis presenting to the ED to compare standard clinical management based on NEWS2 scoring plus PCT-guided risk assessment with standard clinical management based on NEWS2 scoring alone and compare if this approach reduces prescriptions of antibiotics without increasing mortality.

Methods and analysis PRONTO is a parallel two-arm open-label individually RCT set in up to 20 NHS EDs in the UK with a target sample size of 7676 participants.

STRENGTHS AND LIMITATIONS OF THIS STUDY

⇒ Sepsis has a problem with both over and under diagnosis, and a major strength of PROcalcitonin and NEWS2 evaluation for Timely identification of sepsis and Optimal (PRONTO) is the use of coprimary outcomes to assess effectiveness as an antimicrobial stewardship intervention but also to ensure safety which is vital for widespread clinical adoption of this intervention.

⇒ PRONTO is designed to integrate into routine UK clinical pathways and includes assessment of acceptability and practicality in emergency department settings.

⇒ Limitations of the study design include the intervention being a change in risk assessment rather than a formal prescribe/do not prescribe rule for antibiotic use, which could lead to a higher rate of clinician preference in the study.

⇒ The use of deferred consent also has the potential to increase participant withdrawal from the trial, as not all patients would have agreed to prospective informed consent.

Participants will be randomised in a ratio of 1:1 to standard clinical management based on NEWS2 scoring or standard clinical management based on NEWS2 scoring plus PCT-guided risk assessment. We will compare whether the addition of PCT measurement to NEWS2 scoring can lead to a reduction in intravenous antibiotic initiation in ED patients managed as suspected sepsis, with at least no increase in 28-day mortality compared with NEWS2 scoring alone (in conjunction with local standard care pathways). PRONTO has two coprimary endpoints:
initiation of intravenous antibiotics at 3 hours (superiority comparison) and 28-day mortality (non-inferiority comparison). The study has an internal pilot phase and group-sequential stopping rules for effectiveness and futility/safety, as well as a qualitative substudy and a health economic evaluation.

Ethics and dissemination The trial protocol was approved by the Health Research Authority (HRA) and NHS Research Ethics Committee (Wales REC 2, reference 20/WA/0058). In England and Wales, the law allows the use of deferred consent in approved research situations (including ED studies) where the time dependent nature of intervention would not allow true informed consent to be obtained. PRONTO has approval for a deferred consent process to be used. Findings will be disseminated through peer-reviewed journals and presented at scientific conferences.

**Trial registration number ISRCTN54006056.**

**INTRODUCTION**

Sepsis is defined as life-threatening organ dysfunction caused by a dysregulated host response to infection and is a medical emergency requiring prompt antimicrobial therapy and physiological support. The identification, assessment and management of sepsis is challenging because of its many non-specific symptoms and signs, which can be caused by both infectious and non-infectious diseases. In line with international recommendations, the UK National Institute for Health and Care Excellence (NICE) sepsis guidelines suggest the administration of intravenous antibiotics within an hour in patients at risk of intensive care unit (ICU) admission and death. However, up to 50% of patients initially managed as sepsis in the emergency department (ED) do not have a final diagnosis of sepsis and often do not have an infection. The current approach leads to overdose of antibiotics with the associated risk of antimicrobial resistance, antibiotic-related adverse drug reactions (eg, *Clostridium difficile* infection) and extended hospital stays.

The challenge of delivering high-quality sepsis care in an ED setting has been well recognised. The third international consensus definition (Sepsis 3) recommended use of the quick Serial Organ Failure Assessment (qSOF) score, to identify patients at high risk of death and prolonged ICU stay. National Early Warning Score (NEWS) and NEWS2 are rapid physiology-based scoring systems which are used to detect and track the deteriorating patient. NEWS has been demonstrated to have better diagnostic accuracy to qSOF in detection of severe outcomes in sepsis. However, with its higher sensitivity comes reduced specificity which can result in significant increased numbers of patients being managed as high risk for suspected sepsis with a corresponding pressure on ED departments. NEWS2 replaced NEWS scoring system as the standard monitoring tool in the National Health Service (NHS) in 2015 and has been found to be comparable or superior to NEWS. In October 2021, Surviving Sepsis Campaign recommended that immediate antibiotics (within 1 hour) should be targeted to those with septic shock and others with suspected sepsis could wait for up to 3 hours for initial assessment to target antimicrobial choice or identify non-infectious mimics.

The emergence of COVID-19 has exacerbated this previously highlighted problem. COVID-19 is a viral infection which presents within the sepsis syndrome constellation. Secondary bacterial infections are uncommon at presentation to ED (3.5%), despite this up to 83% of patients with COVID-19 received antibiotics. NEWS2 scores are broadly predictive of COVID-19 outcome on presentation but does not appear to be predictive of bacterial coinfection. Initial investigations in the ED can be helpful in distinguishing between COVID-19 and bacterial pneumonia including typical radiographic change, and COVID-19 point-of-care diagnostics. These results would be available within 3 hours for assessment and could potentially reduce unnecessary antimicrobial usage in COVID-19 management.

**Aims and objectives**

**Primary objective**

To assess whether the addition of PCT measurement to NEWS2 scoring leads to a reduction in intravenous antibiotic initiation at 3 hours, with no increase in 28-day mortality compared with NEWS2 scoring alone in the management of patients presenting to hospital EDs in England and Wales with suspected sepsis.

**Secondary objective**

The assessment of (1) feasibility, (2) cost-effectiveness and (3) acceptability to healthcare practitioners, patients and their family

**METHODS AND ANALYSIS**

**Study design**

PROcalcitonin and NEWS2 evaluation for Timely identification of sepsis and Optimal (PRONTO) is a multicentre, parallel two-arm, open-label, individually randomised controlled trial with two coprimary endpoints, an internal pilot phase and group-sequential stopping rules for effectiveness and futility/safety. Participants will be randomised in a ratio of 1:1 to standard clinical management based on
NEWS2 scoring or standard clinical management based on NEWS2 scoring plus PCT-guided risk assessment.

**Internal pilot**

An internal pilot phase will be conducted over the first 9 months of the recruitment period with ten lead sites. Predefined progression criteria will be used to assess feasibility to progress to the full trial, such as site and patient absolute recruitment and consent rate, proportion of patients undergoing PCT assessments and the ability to collect coprimary outcome data.

**Eligibility**

**Inclusion criteria**

Up to 20 EDs from across England and Wales will recruit adults (≥16 years) who are being managed as suspected sepsis over a 24-month period. There is no minimum NEWS2 score for inclusion into the study.

**Exclusion criteria**

Patients already receiving intravenous antibiotics, currently receiving myeloablative chemotherapy, patients with solid-organ transplantation, allogeneic bone marrow or stem cell transplantation within 3 months prior to consent or patients known to require urgent surgical intervention at the time of randomisation.

Patients with an advance directive to withhold life-sustaining treatment or patients not wishing to receive cardiopulmonary resuscitation may qualify provided they receive all other resuscitative measures for example, respiratory support and fluid resuscitation.

**Study procedures and progress**

The trial schema is shown in figure 1.

The COVID-19 pandemic resulted in a delay to the original start date of June 2020. First participant was recruited on 20 November 2020. Current planned end date is 30 November 2022.

**Identification and screening**

Patients with suspected sepsis will be identified at ED triage. After initial NEWS2 scoring and assessment according to current standard of care the eligibility criteria will be assessed and if no exclusion criteria apply, patients will be enrolled into the trial and randomised. A screening log of all eligible and randomised patients will be kept at each site so that any biases from differential recruitment will be detected.

**Randomisation**

Participants will be individually randomised in a 1:1 ratio by delegated research staff within the ED to either standard clinical management based on NEWS2 scoring (control) or standard clinical management based on NEWS2 scoring plus PCT-guided risk assessment (intervention). We will use minimisation with NEWS2 score (≥10 or < 5) and site as balancing factors and add a random element to reduce the risk of subversion. This will be implemented in a secure 24-hour web-based randomisation programme controlled centrally by the Centre for Trials Research (CTR) in Cardiff. Full details are provided in the PRONTO randomisation strategy.

**Trial intervention**

The BRAHMS PCT-direct reader (ThermoFisher Diagnostics (Altrincham, Cheshire, UK) is a fully validated, CE-marked point-of-care test to determine levels of PCT in the blood. The test requires 20µL blood which will be obtained from either venous blood during standard care procedures at triage or via a finger-prick. This will be used in combination with NEWS2 assessment of adult patients with suspected sepsis in ED, using a guidance-only algorithm for clinicians (figure 1). The risk algorithm categorises individuals as low, medium or high risk, interpretation and management (table 1). Clinicians have oversight at all times as to whether to adhere to the algorithm. As currently mandated in UK, NICE clinical guidelines and quality standard QS161, urgent senior review within an hour will take place should any healthcare provider identify at least one risk factor indicating high risk of progression to severe illness or death regardless of underlying aetiology. This equates to a NEWS2 ≥9 or an individual having a single feature of the evidence-based ‘NICE high-risk criterion’.

**Informed consent**

Research carried out in emergency situations is challenging in terms of obtaining consent. Emergency research is when treatment needs to be given urgently, and it is necessary to take urgent action for the purposes of the study. In some emergency situations people may lack capacity to give consent themselves and obtaining consent from a legal representative or consulting others is not reasonably practicable. In England and Wales, the law allows adults who lack capacity to take part in emergency research without prior consent from a legal representative or consulting others, if certain conditions are met (Medicines for Human Use (Clinical Trials) Amendment (No 2) Regulations SI 2006 2984, Mental Capacity Act s32). Given the requirement for rapid clinical assessment and treatment in the management of suspected sepsis, for this trial we will use a deferred consent model. Patients and their relatives will be informed that a study is ongoing but a lengthy consent discussion will not be had so as not to delay treatment. Should the patient or consultee wish to take part at this point, then the decision will be respected and the patient will not be enrolled into the trial. Following randomisation an approach to obtain informed consent will be made as soon as is practicably feasible, ideally within 72 hours (figure 2). Where a participant lacks mental capacity, a maximum of three approaches will be made. After three approaches, or if the participant is not likely to regain mental capacity, a personal consultee will be approached. In extreme circumstances, where no personal consultee can be identified, a nominated consultee will be approached. Separate informed consent will be taken for participation in the
qualitative data collection. Patients who do not consent to continue in the study will be withdrawn completely from the study. A tiered consent model is used in this study and allows participants to consent to different aspects of the study (online supplemental appendix table 1). An example participant consent form is available in online supplemental appendix.

**Data collection during primary admission**

All data collection will be by electronic data capture using a bespoke database developed by the CTR and hosted by Cardiff University secure servers. It is encrypted and accessed by individual username and password. Paper copies of all case report forms will be available. Essential documents will be kept securely in a locked cupboard, and at the end of the trial, will be archived at an approved external storage facility for 10 years. A member of the research team in ED will undertake the data collection relating to the NEWS2 screening, trial intervention and whether clinical teams followed the intervention or standard of care risk assessment. Participants who consent to continue in the study will have daily information collected from the date of randomisation until they are discharged from hospital or until day 28, whichever is sooner. Trial data is collected from patients’ health records and no trial visits occur between consent and day 28. Key follow-up data are listed in online supplemental appendix table 2.

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**Figure 1** Trial schema. ED, emergency department; NEWS, National Early Warning Score; NICE, National Institute for Health and Care Excellence; PCT, procalcitonin.
FOLLOW-UP

Twenty-eight-day follow-up

Day 28 follow-ups will be conducted via telephone or in person if the participant remains an inpatient. These will comprise a European Quality of life five dimension, five level (EQ-5D/5L) validated questionnaire for participant or proxy completion, and a Health Economics questionnaire where patient outcomes (readmission, retreatment, hospital-acquired infection) and use of healthcare resource (hospital admissions, outpatient parenteral antimicrobial therapy, other prescribed medicines, privately purchased over-the-counter medicines, General Practitioner (GP) and hospital outpatient attendance) will be captured. In addition, direct non-medical costs borne by patients/carers as a result of attending hospital (travel costs, childcare costs, expenses incurred while in hospital, self-reported lost earnings and other direct non-medical expenses) will be collected.

Ninety-day follow-up

EQ-5D/5L questionnaires will be repeated and a shortened Health Economics questionnaire to capture any additional costs or hospital admissions since the day 28 questionnaires will be completed.

Withdrawal

Participants have the right to withdraw from the study at any time and can request that all data collected up to that point is not used.

Safety and pharmacovigilance

The trial population comprises unwell hospital inpatients. Events such as prolongation of existing hospitalisation, life threatening events and death are expected in this population and are recorded as part of routine data collection and therefore are not subject to expedited reporting. Serious adverse events will be reported if the event results in persistent or significant disability

Table 1  Clinical risk management interpretation

| Risk group | Interpretation |
|------------|----------------|
| High       | High risk of progression to sepsis. Likely benefit from immediate antibiotics (within 1 hour) |
| Medium     | Medium risk of progression to sepsis. Likely benefit from early antibiotics (within 3 hours) but consider non-bacterial sources and likely source. Allows clinical teams time to complete rapid assessment |
| Low        | Low risk of progression to sepsis. Consider non-bacterial sources, likely source and whether requires antibiotics |

NEWS2, National Early Warning Score 2; PCT, procalcitonin.

Figure 2  Consent procedures. CRF, case report form; ED, emergency department; HCP, Health Care Professional; NEWS, National Early Warning Score; PRONTO, PROcalcitonin and NEWS2 evaluation for Timely identification of sepsis and Optimal use of antibiotics in the emergency department; RN, Research Nurse.
or incapacity or consists of a congenital anomaly or birth defect. An assessment of causality between the event and the trial intervention will be carried out by the principal investigator or delegated clinician, and then independently by a clinical reviewer. If the clinical reviewer classifies the event as probably or definitely caused by the intervention, it will be classified as a serious adverse reaction. Non-serious Adverse Events (AEs) potentially attributable to the PCT test will be collected as part of routine follow-up at 28 days. Any other non-serious AEs will not be collected.

Data management
Details of data management procedures (such as checking for missing, illegible or unusual value (range checks) will be specified in the PRONTO Data Management Plan. Details of Monitoring procedures will be specified in the PRONTO Monitoring plan.

STATISTICAL ANALYSIS
Outcome measures
The coprimary outcomes of this study are the initiation of intravenous antibiotics at 3 hours (intervention arm to be shown superior to control) and 28-day mortality (intervention arm to be shown non-inferior to control). Coprimary and secondary outcomes are listed in box 1. Final decisions about the primary effectiveness of the intervention, using these coprimary outcomes will be made based on the decision matrix (table 2). All outcomes will be stratified by COVID-19 diagnosis (SARS-CoV2 PCR positive or high likelihood of clinical COVID-19 as determined by a senior clinician).

Sample size
The sample size calculation is based on two coprimary outcomes:29

1. Twenty-eight-day mortality, for which we want to show non-inferiority of the PCT guided assessment as compared with current standard practice, using an absolute 2.5% non-inferiority margin. Assuming a 28-day mortality of 15% in patients managed as suspected sepsis treated in the ED,3 30 this means that any increase in 28-day mortality from 15% to not more than 17.5% would be considered non-inferior. For 90% power and one-sided 5% significance level the sample size required is 532, which is substantially lower than what is needed for the non-inferiority endpoint. With 7002 patients we would be able to detect effects as small as a reduction from 90% to 87.6%, with 90% power.

Accounting for 5% drop-out, we would need a total sample size of 7372. The group-sequential design with O’Brien-Fleming stopping boundaries for both

| Table 2 | Decision matrix for coprimary outcomes |
|---------|----------------------------------------|
| **Reduced antibiotic initiation** | **Same or more antibiotic initiation** |
| **Decreased mortality** | Effective | Effective |
| **Equivalent mortality** | Effective | Not effective |
| **Increased mortality** | Not effective/harmful | Not effective/harmful |

Trust, unpublished data). A reduction to 80% would be considered a success. To detect such an effect with 90% power and two-sided 5% significance level the sample size required is 532, which is substantially lower than what is needed for the non-inferiority endpoint. With 7002 patients we would be able to detect effects as small as a reduction from 90% to 87.6%, with 90% power.

Accounting for 5% drop-out, we would need a total sample size of 7372. The group-sequential design with O’Brien-Fleming stopping boundaries for both
effectiveness and futility/safety will increase the total maximum sample size (if the study is not stopped after the interim analysis) by just over 4% to 7676 (inflated for 5% drop-out).

These sample sizes were calculated using SAS V.9.4 PROCOPOWER and PROC SEQDESIGN.

Interim analysis
A planned interim analysis of the coprimary outcomes will be conducted when 50% of patients have been recruited and followed up for 28 days. Stopping the study shall be recommended by the independent data monitoring committee (IDMC) based on group-sequential O’Brien-Fleming boundaries. They shall recommend stopping for effectiveness if:

- The PCT-guided assessment is superior in terms of 28-day mortality (ie, a significant reduction to less than 15%).
- The PCT-guided assessment is non-inferior in terms of 28-day mortality and superior in terms of initiation of antibiotics.

They shall recommend stopping for futility if the results of the interim analysis suggest futility for both endpoints. This strategy ensures overall type I error rate control.31 32

The exact stopping rules will be specified in an interim analysis plan.

Final analysis
The primary analysis will be intention to treat and will fit separate two-level logistic regression models (patients nested within sites) to both coprimary outcomes (antibiotic initiation and mortality), controlling for baseline NEWS2 score (minimisation factor). The intervention will be considered effective if there is both a significant reduction in antibiotic initiation (two-sided 5% level) and if the difference in mortality between the two groups is non-inferior (one-sided 5% level). In case the 28-day mortality rate in the control arm deviates from the assumed 15%, the absolute 2.5% non-inferiority margin will be replaced with an arcsine difference ‘non-inferior frontier’.33

The primary analysis will be adjusted to account for the group-sequential design. Imputation of missing data will be done as part of sensitivity analyses.

In a secondary analysis, complier adjusted causal effect models will be fitted to allow for non-adherence to the intervention. Two models will be fitted allowing for two different definitions of adherence:

1. Patients randomised to PCT-guided care in whom a PCT test is done and the clinician considers the results as part of their decision making.
2. Patients randomised to PCT-guided care in whom a PCT test is done and the clinician follows the algorithm exactly.

Analyses of secondary outcomes will also be performed as intention to treat and using appropriate two-level regression models depending on the type of outcome (eg, linear regression for continuous outcomes, Cox regression for time-to-event outcomes) to allow for patients nested within sites. This includes an HTA and economic evaluation as per CHEERS 2022 guidance. Analyses will be split by organ system of the infection (eg, lower urinary tract, lower respiratory, intra-abdominal, bacteraemia, skin and soft tissue). Stratified analyses will be undertaken at different levels of NEWS2 scoring ≤4, 5–6 and ≥7, and will also be undertaken by COVID-19 status. All further details will be specified in a statistical analysis plan which will be finalised prior to database lock for the planned interim analysis and subsequently published.

Missing primary outcome data are likely to be minimal, so complete-case analysis will be used. However, if this exceeds more than 20% of participants we will employ multiple imputation and report the impact on the treatment effect alongside the complete-case analysis.

QUALITATIVE STUDY
The qualitative work will have three components: interviews with clinicians, interviews with patients/carers, and observations of trial implementation (when appropriate during the ongoing current COVID-19 pandemic). Findings will be used to aid understanding of the quantitative data and provide areas for improvement in processes to enhance the efficiency of the trial.

Interviews with clinicians will take place at two time points. Interview 1 will take place during the pilot phase and will be a semistructured interview with 10–12 clinicians at ≤5 study sites (2–3 per site). This will explore the feasibility and acceptability of research processes and integration of the PCT algorithm into their ED setting. Interview 2 will be with clinicians towards the end of the trial when they have more experience of using the PCT algorithm and will identify barriers and facilitators to the use of the PCT test and algorithm in more detail, including reasons for deviating from the study algorithm.

We will conduct semistructured interviews with patients after the 90-day follow-up, in order to gain a detailed understanding of patients’ experiences of care to aid understanding of trial results. We will encourage patients to include a close family member in the interview also. This will allow us to capture an additional perspective on the patients’ care.

PATIENT AND PUBLIC INVOLVEMENT
The proposal has benefited from multiple interactions with patient and public involvement (PPI) groups to refine the research question and design. Author JC is a lay coapplicant/patient representative, who has coproduced and helped finalise the study design. As a coapplicant JC is a member of the trial management group (TMG) ensuring that all patient facing materials are presented in a suitable way. Her experience is invaluable throughout the project, including the promotion of the trial to potential participants and appropriate dissemination of findings to the lay public.

Euden J, et al. BMJ Open 2022;12:e063424. doi:10.1136/bmjopen-2022-063424
In addition, we have convened wider PPI advisory panels from both higher education institutions and NHS patient groups. We discussed the trial with the panel at the Royal Liverpool Hospital in August 2018, focusing on need, conception, design and trial management. The group fully supported the need for this trial recognising the potential for PCT measurement to improve outcomes for patients with suspected sepsis and supported the use of deferred consent. Specific feedback about these aspects has now been used to update the relevant parts of the proposal.

TRIAL MANAGEMENT

The trial is sponsored by the University of Liverpool and coordinated by Cardiff University CTR.

Trial management group

The TMG will meet monthly throughout the course of the trial and will include the cochief investigators, coapplicants, collaborators, trial manager, data manager and administrator. TMG members will be required to sign up to the remit and conditions as set out in the TMG charter.

Trial steering committee and IDMC

An independent trial steering committee (TSC) consisting of an independent chairperson, two independent members and a patient representative will provide oversight of the PRONTO trial. There will also be a separate IDMC to provide oversight of all matters relating to patient safety and data quality, and recommend continuing or stopping the trial depending on the results of the interim analysis. Members will be required to sign up to the remit and conditions as set out in the TSC and IDMC charters and will meet at least annually.

ETHICS AND DISSEMINATION

Ethics approvals and consent

The trial was approved by the NHS Research Ethics Committee (Wales REC 2, reference 20/WA/0058) on the 21 July 2020 and subsequent Health Research Authority (HRA) and Health and Care Research Wales approval was granted on 22 July 2020. In England and Wales, the law allows the use of deferred consent in approved research situations (including ED studies) where the time dependent nature of intervention would not allow true informed consent to be obtained. PRONTO has approval for a deferred consent process to be used, full details are in Informed Consent section above. The following substantial amendments were made to the trial and were communicated to all trial sites: Amendment 5 (23 October 2020); Amendment 7 (10 December 2020); Amendment 9 (25 February 2021); Amendment 12 (29 June 2021), Amendment 15 (15 October 2021), Amendment 17 (6 January 2022).

Dissemination plan

We will engage with patient groups and the wider public through relevant charities such as UK Sepsis Trust and Antibiotics Action, and seek to present trial updates at their annual conferences. We will use press releases and social media outlets to publicise the trial and disseminate findings. A 90 s animation outlining the PRONTO main aims was commissioned https://www.youtube.com/watch?v=H3x-rNVlwJF and accessed via posters and patient information leaflets via a scannable QR code. At the end of the trial, a final report will be prepared for the National Institute of Health Research Health Technology Assessment Journal series. The results will be disseminated locally, nationally and internationally among scientific, clinical and lay groups including participants and their families. All publications and presentations related to the trial will be authorised by the TMG in accordance with the PRONTO publication policy. Where appropriate, the results of this trial can be directly implemented in the revisions of the NICE guidelines.

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Contributors

NF and ST are co-chief investigators of this trial. NF and ST, along with ET-J, EC, LB-H, PH, MI-K, ML, FM, LWN, EN, PP, PS, DT-R, IW and KH led the development of the research question, study design, obtaining the funding and
implementation of the protocol. JE is the Trial Manager and ET-J is the senior trial manager who coordinate the operational delivery of the trial protocol and recruitment. LB-H is the lead qualitative researcher. PP is the trial statistician. SG is the data manager. All authors listed provided critical review and final approval of the manuscript.

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Disclaimer
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Competing interests
EC is co-Ci for the BATCH Trial (HTA 15/188/42) and the PEACH study (HTA Project: NIHR132254) on PCT use, and member of NICE Diagnostic advisory committee (2014–2020), and NICE Sepsis guideline development committee (2014-6). All other authors declare no competing interests.

Patient and public involvement
Patients and/or the public were involved in the design, or conduct, or reporting, or dissemination plans of this research. Refer to the Patient and public involvement section for further details.

Patient consent for publication
Not applicable.

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Supplemental material
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