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LETTER

brought to you







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Abstract

Objectives: To determine if a specific immunomodulatory intervention reduces progression of COVID-19-related disease to organ failure or death, compared to standard of care (SoC).

Trial design: Randomised, parallel 3-arm (1:1:1 ratio), open-label, Phase IV platform trial of immunomodulatory therapies in patients with late stage 1 or stage 2 COVID-19-related disease, with a diagnosis based either on a positive assay or high suspicion of COVID-19 infection by clinical and/or radiological assessment.

Participants: Patients aged 18 and over, with a clinical picture strongly suggestive of COVID-19-related disease (with/without a positive COVID-19 test) AND a Risk count (as defined below) >3 OR \geq 3 if risk count includes "Radiographic severity score >3". A risk count is calculated by the following features on admission (1 point for each): radiographic severity score >3, male gender, non-white ethnicity, diabetes, hypertension, neutrophils >8.0 x10⁹/L, age >40 years and CRP >40 mg/L.

Patients should be considered an appropriate subject for intervention with immunomodulatory therapies in the opinion of the investigator and be able to be maintained on venous thromboembolism prophylaxis during the inpatient dosing period, according to local guidelines. The complete inclusion and exclusion criteria as detailed in (Continued on next page)

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the additional file 1 should be fulfilled. Patients will be enrolled prior to the need for invasive mechanical ventilation, cardiac or renal support. Participants will be recruited across multiple centres including initially at Cambridge University Hospitals NHS Foundation Trust, King's College Hospital NHS Foundation Trust, Guy's and St Thomas' NHS Foundation Trust, University Hospital of Wales, Gloucestershire Royal Hospitals NHS Foundation Trust, and The Royal Wolverhampton NHS Trust.

Intervention and comparator: Each active comparator arm will be compared against standard of care (SoC). The immunomodulatory drugs were selected from a panel of licenced candidates by a drug evaluation committee, which considered potential efficacy, potential toxicity, scalability and novelty of each strategy. The initial active arms comprise baricitinib and ravulizumab.

Baricitinib will be given 4 mg orally (once daily (OD)) on days 1-14 or until day of discharge. The dose will be reduced to 2 mg OD for patients aged > 75 years and those with an estimated Cockcroft Gault creatinine clearance of 30-60 ml/min.

Ravulizumab will be administered intravenously once according to the licensed weight-based dosing regimen (see Additional file 1).

Each active arm will be compared with standard of care alone. No comparisons will be made between active arms in this platform trial.

Main outcomes: The primary outcome is the incidence (from baseline up to Day 14) of any one of the events (whichever comes first): death, invasive mechanical ventilation, extra corporeal membrane oxygenation, cardiovascular organ support (inotropes or balloon pump), or renal failure (estimated Cockcroft Gault creatinine clearance <15ml/min).

Randomisation: Eligible patients will be randomised using a central web-based randomisation service (Sealed Envelope) in a 1:1:1 ratio, stratified by site to one of the treatment arms or SoC.

Blinding (masking): This is an open-label trial. Data analysis will not be blinded.

Numbers to be randomised (sample size): There is no fixed sample size for this study. Serial interim analyses will be triggered by an Independent Data Monitoring Committee (IDMC), including analysis after 125 patients are recruited to each arm, 375 in total assuming 3 arms. Additional interim analyses are projected after 229 patients per arm, and potentially then after 469 per arm, but additional analyses may be triggered by the IDMC.

Trial Status: TACTIC-R Protocol version number 2.0 date May 20, 2020, recruitment began May 7, 2020 and the end trial will be the date 18 months after the last patient's last visit. The recruitment end date cannot yet be accurately predicted.

Trial registration: Registered on EU Clinical Trials Register EudraCT Number: 2020-001354-22 Registered: 6 May 2020 It was registered on ClinicalTrials.gov (NCT04390464) and on ISRCTN (ISRCTN11188345)

Full protocol: The full protocol is attached as an additional file, accessible from the Trials website (Additional file 1). In the interest in expediting dissemination of this material, the familiar formatting has been eliminated; this Letter serves as a summary of the key elements of the full protocol.

Keywords: COVID-19, Randomised controlled trial, Protocol, Baricitinib, Ravulizumab, Open-label, Adaptive trial, Repurposed drugs

Supplementary information

Supplementary information accompanies this paper at https://doi.org/10. 1186/s13063-020-04535-4.

Additional file 1. Full Study Protocol.

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Authors' contributions

SK and MF are sub-investigators and SK is the corresponding author. EHS is the trial coordinator. MK, EBH, AC, SN, SB contributed to writing of protocol. JC, AC, JG, FH, DJ and IW designed the study, contributed to protocol, comprise the Trial Management group (TMG) and are joint senior authors. FH is the Chief Investigator and DJ is the Co-Chief Investigator. The author(s) read and approved the final manuscript.

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Availability of data and materials

Not applicable. Ownership of the data arising from this trial resides with the trial team and the sponsor.

Ethics approval and consent to participate

The study received favourable ethical opinion by the East of England -Cambridge Central Research ethics committee. Ref: 20/EE/0135, Date of approval May 6, 2020. We certify that this trial has received ethical approval from the appropriate ethical committee as described above.

Full informed consent will be obtained from each patient prior to enrolment into the study. In line with other urgent COVID -19 trials, if the patient lacks capacity to give consent due to the severity of their medical condition (e.g. acute respiratory failure or delirium), then consent may be obtained from a relative acting as the patient's legally designated representative or from a professional legal representative.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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