

Welcome to the TICH-3 trial!

- University Hospital of North Tees
- Darent Valley Hospital
- Ipswich Hospital
- Queen Elizabeth Hospital, Birmingham
- King's Mill Hospital, Sutton in Ashfield
- University Hospital Coventry
- Southend University Hospital
- Prince Philip Hospital, Wales
- Southmead Hospital, Bristol

Congratulations!

- Southend University Hospital recruiting within 1 week of greenlight
- King's Mill Hospital recruiting within 2 weeks of greenlight
- Queen Elizabeth, Birmingham recruiting 4 participants within 2 weeks of greenlight

Congratulations for recruiting within first month following greenlight!

- Darent Valley Hospital
- Ipswich Hospital
- University Hospital of North Tees



TICH-3 NEWSLETTER

ISRCTN: 97695350 March 2024, Issue 10

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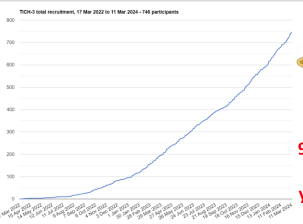
http://tich-3.ac.uk/

Useful documents: https://stroke.nottingham.ac.uk/tich-3/docs/



500 UK

263 International



TICH-3 has recruited a further 91 UK participants since the last newsletter, 20/12/2023 thank

you so much for your continued support!

STANDARD OF CARE FOR ICH

- Standard of care should be followed in addition to trial treatment, please administer any BP lowering medication or PCC at the same time but through separate cannulas.
- The standard of care in the UK for treatment of rivaroxaban associated ICH is 4F-PCC (octa/beriplex) and vitamin k.
- The use of tranexamic acid is not recommended as part of clinical guidelines for stroke—hence equipoise and the rational for TICH-3.
- If your clinical team plan to use tranexamic acid for patient, then they would not be appropriate to enrol the patients into TICH-3.
- Please discuss with us if you are planning to use TXA clinically.

ADDING PARTICIPANT TO TICH-3 WEBSITE

You can now click the 'add the randomisation record' link on the recruitment alert email to add the participant to the TICH-3 website which creates the official record for them by generating their study identifier, you can then proceed to upload contact details, consent form and begin the eCRF data collection.

Dear TICH-3 investigator,

A hospital investigator has notified us of a new TICH-3 participant.

Please make every effort to identify this participant, and locate their randomisation data and patient notes (if not already done), then add the randomisation record to the participant list.

ENROLLING INVESTIGATOR LIVE TRAINING WEBINARS

These Enrolling Investigator Webinars serves as a refresher to the existing team members on the delegation log but also can be attended by new team members to get onto your sites TICH-3 electronic delegation log e.g. new registrars, ED doctors. The more enrolling investigators there are on the delegation log maximises the chances of participant recruitment.

- Tuesday 16th April 2024 1 1.30pm
- Wednesday 15th May 2024 12.30 1pm

Please email us with your email address and date you would like to attend.



TRIAL DOCUMENTS UPDATES

These are not ethically approved documents and therefore there are not any approvals to coincide with this updates.

Training slides updated (no major changes to process) – audit list of changes available at the end of each training slide set

- Enrolling Investigator Training v2.3 08/03/2024
- Pharmacy Training v2.3 08/03/2024
- Investigator Training v3.4 08/03/2024
- Ward Training Final v1.1 08/03/2024
- CT scan upload training Final v1.2 08/03/2024

You are not required to re-read the training slides after any updates (unless we have specifically informed you of this) or complete a training log, however we recommend reading these very few months as a refresher to the trial.

New team members can read the appropriate slides for their role https://stroke.nottingham.ac.uk/tich-3/docs then complete self-referral to be added to the online delegation log https://tich-3.ac.uk/?ZSelfRef

eCRFs update The eCRFs data are intended to be inputted straight from the medical record electronically.

Enrolment v1.8

• Updated inclusion/exclusion criteria to add text relating to informed consent and anticoagulants.

Day 7 follow-up v1.7

- Changed A5a: Blood pressure on day 7 reading 1 to First recorded blood pressure on day 7.
- Changed A5b: Blood pressure on day 7 reading 2 to Second recorded blood pressure on day 7.

ED Poster Corrected ED poster footer version from 3.0 to 2.1.

Co enrolment log updated v3.1 to add in requirements for if needs localising at each site.

KEEPING IN TOUCH AND UPDATE MEETINGS

We would like to invite you all to join the TICH-3 trial team where we will discuss topics of interest for the trial:

Wednesday 17th April 12.30 – 1.30pm Click here to join the meeting

Invitations will also have been sent to your email address so look out for these!

If you have any points you would like to discuss, please email the team and we will add them to the meeting.



You can find us on Twitter: @tich3trial and @UoN_STU

Please feel free to tweet us or tag us in your Stroke related tweets

Conferences coming up

- European Stoke Organisation Conference (ESOC) 15-17 May 2024
- World Stroke Congress (WSC) 23 26 Oct 2024
- International Clinical Trials Methodology Conference (ICTMC) Edinburgh, Sept 2024

