



# TRANEXAMIC ACID FOR INTRACEREBRAL HAEMORRHAGE: TICH-3 TRIAL

# Ireland Investigators

Professor Nikola Sprigg

On behalf TICH-3 Trial Team

22/04/2024



# **Funding:**



➤TICH-3 funded by National Institute of Health and Rare Research (Health Technology Assessment)

TICH-3 Trial Registration: ISRCTN9769535

TICH-3 EU CTIS: 2022-500587-35-01

TICH-3 CTA reference: 03057/0074/001-0001

TICH-3 IRAS Project ID: 297457

TICH-3 Trial Sponsor: University of Nottingham



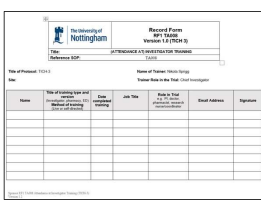




## **Overview**



- Welcome and introductions
- Presentation of TICH-3:
  - Inclusion criteria
  - Consent process
  - Randomisation
  - Safety outcomes
  - Pharmacy Drug storage and administration
  - Passwords, website and electronic case report forms (eCRFs)
- Updates on ethics/research/contracts
- Q&A PLEASE COMPLETE TRAINING LOG
- Future planning



Sel	f-referral checklist
	ase read through the following points carefully and icate which are applicable, then submit at the bottom.
0	I am a hospital researcher, administrator, pharmacist* or radiologis
0	I have completed the TICH-3 trial training.
0	I have my signed work CV.
0	I have a GCP certificate.
0	I do <u>not</u> already have a TICH-3 account. (If you do, please contact the co-ordinating office when another hospital needs to be added to your account).
0	I declare my intention to participate in the TICH-3 trial.
	Your country: [Select]
	Submit
	* - includes pharmacy technician



# **ACTION – Return Training Log**



 Please complete the investigator training log and return via email to the coordinating centre <u>Click here for direct download of training log</u> use the self referral form:

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

- Team members who could not attend live training can access training slides from TICH-3 website <a href="https://stroke.nottingham.ac.uk/tich-3/docs/#UK">https://stroke.nottingham.ac.uk/tich-3/docs/#UK</a> site training
- There are 3 versions of the training slides
  - Investigator training which gives a detailed description of the whole trial process, intended for the PI and research nurses/coordinators.
  - Enrolling investigator training this streamlined training is intended for team members who will only be taking enrolment consent i.e. consultants
  - Pharmacy training this streamlined training is intended for members of pharmacy team
- A short 3 ½ minute video is available to introduce team members to the TICH-3 trial <a href="http://tich-3.ac.uk/docs/#Videos">http://tich-3.ac.uk/docs/#Videos</a>





PLEASE SCAN ME

# BACKGROUND

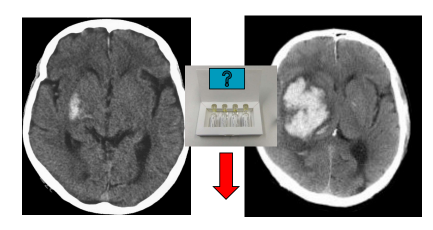


# Intracerebral Haemorrhage (ICH)



## Intracerebral haemorrhage can be devastating

- Haematoma expansion (HE) is common, occurs early and is main cause of death
- Predictors time, haematoma volume, anticoagulation and antiplatelets



 Drugs that stop bleeding (such as tranexamic acid), are effective in other bleeding conditions and could potentially reduce haematoma expansion

TICH-3: does giving tranexamic acid early after ICH prevent haematoma expansion and reduce death and disability



### Tranexamic acid in other trials



- TXA acts through antifibrinolytic mechanisms
- CRASH-2 In patients with traumatic haemorrhage (including from head injuries), TXA significantly reduces death due to bleeding and all-cause mortality, with no increase in vascular occlusive events.
- Analysis of the CRASH-2 trial showed that because death due to bleeding occurred early after trauma, hyperacute administration of TXA was necessary for patients to receive any benefit.
- A meta-analysis of TXA in traumatic intracranial haemorrhage showed that it was associated with a significant reduction in subsequent intracranial bleeding.
- CRASH-3, reduced head injury related deaths in patients with traumatic brain injury, with early treatment more effective than later treatment.
- In TICH-2 (in 2325 patients with ICH within 8 hours of onset) TXA was safe, reduced haematoma expansion and early death. It did not significantly change outcome at 3 months
- Tranexamic acid is inexpensive, easy to administer, seems to be safe, and is widely available, so even a modest treatment effect could have an important impact on the global scale.



# **Key changes from TICH-2**



## Target participants most likely to benefit

Change	TICH-2	TICH-3
Primary outcome	Function day 90	Death by day 7 Function day 180
Recruitment target	2000	5500
Recruitment time	8 hours since onset	4.5 hours since onset
Baseline ICH volume	No maximum	Exclude massive (usually Haematoma Volume> 60ml)
Consent	Written consent	Oral consent –followed by written consent
Randomisation	On-line	Simple – lowest pack number



# Other TXA in ICH trials

Completed trials	TXA timing after symptom onset (hours)	TXA Bolus	Total recruitment
Arumugam 2015	8	1g bolus 1g infusion	30
Liu TRAIGE 2021	8	1g bolus 1g infusion	171
Meretoja STOP – AUS 2020	4.5	1g bolus 1g infusion	100
Ni 2020	8	1g bolus 1g infusion	152
Seiffge TICH NOAC 2022	12	1g bolus 1g infusion	63 (stopped early)
Sprigg TICH 1 2014	24	1g bolus 1g infusion	24
Sprigg TICH 2 2018	8	1g bolus 1g infusion	2325
Zhao 2017 STOP-MSU	2	1g bolus 1g infusion	326

Ongoing trials	Registration	TXA timing after symptom onset (hrs)	TXA dose	Target recruitment
Ezati 2019	IRCT20191014045103N1	Not reported	1g bolus 1g infusion	Not reported
Jiang 2020 THE-ICH trial	ChiCTR1900027065	4.5	1g bolus 1g infusion	2400
Li Qi 2021 TARGET trial	ChiCTR2100045022	3	1g bolus 1g infusion	200
Pandian 2022 INTRINSIC trial	NCT05836831	4.5	2g bolus	3400
Pokhrel 2021	NCT04742205	24	1g bolus 1g infusion	142
Woo 2017 TRANSACT trial	NCT03044184	3	1g bolus 1g infusion	220

# PROTOCOL

EU Protocol v4.2 30.03.23 approved on 19.04.2023



# **TICH-3 Synopsis**



## ICH emergency condition - facilitate rapid enrolment

Design: Double blind randomised clinical trial, pragmatic streamlined design

**Participants:** Inclusion: < 4.5 hours of stroke onset

**Exclusion:** Massive ICH (Glasgow Coma Scale < 5 or Haematoma Volume > 60ml)

**Consent:** Rapid emergency process – oral consent followed by written consent

Intervention: Tranexamic 1g IV bolus added to 100ml sodium chloride over 10 mins then 1g

added to 250ml sodium chloride infusion over 8hrs or saline by identical regime Given alongside standard ICH care, including BP lowering as per clinical guidelines<sup>1</sup>

Randomisation: Simple - use the lowest available treatment pack number

**Primary Outcome:** Early death (day 7)

Secondary outcome: Function-Shift analysis modified Rankin Scale day at 6 months

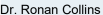
Sample size: 5500 (3900 UK and 1900 Internationally)

Cost/funder: UK NIHR plus others internationally

Duration: 7.25 years - Aim start UK recruitment early 2022

1: https://www.nice.org.uk/guidance/ng128/chapter/Recommendations







# **TICH-3 Sites**



### 6 sites in Ireland

Sites	PI
Tallaght University Hospital	Professor Ronan Collins
St. Vincent's University Hospital	Dr. Tim Cassidy
Mater Misericordiae University	Professor Peter Kelly
University Hospital Limerick	Dr. Margaret O Connor
University Hospital Waterford	Dr George Pope
Beaumont Hospital	Dr. Patricia Fearon

National Coordinator: Prof Ronan Collins

Legal representatives in Europe – Prof Peter Kelly, Katrina Tobin UCD



# Site updates

Site	Status	Contract	Notes
Tallaght University Hospital	Open	Signed	
Beaumont Hospital	None on delegation log	In Progress – DPIA being sorted	
Mater Misericordiae University Hospital	None on delegation log	In Progress	
St. Vincent's University Hospital	None on delegation log	Sent	Delays due to staffing
University Hospital Limerick	None on delegation log	Sent	PI mentioned possible capacity issues – will feedback to TICH-3 team
University Hospital Waterford	None on delegation log	UoN sponsor happy with modifications – awaiting response from Waterford team.	



# **TICH-3: Eligibility Criteria**



#### Inclusion criteria

Spontaneous ICH (confirmed on brain imaging) < 4.5 h of onset</li>

CT (or MRI) is conducted pre-recruitment in line with standard care, the haematoma volume measurement will help assess whether the participant is eligible.

Note - ICH secondary to ruptured aneurysm or vascular malformation or brain tumor or ischaemic stroke (haemorrhagic transformation of infarct, HTI) or thrombolysis or venous infarct is NOT spontaneous ICH

#### **Exclusion criteria**

- Known indication for TXA treatment (e.g. traumatic brain injury) in view of treating physician
- Known contra-indication for TXA treatment (e.g. active seizures) in view of treating physician
- Patient known to be taking therapeutic anticoagulation with warfarin or low molecular weight heparin at time of enrolment. Patients taking direct oral anticoagulants can be included and are not excluded.
- Massive ICH (usually when haematoma volume > 60ml)
- Severe coma, Glasgow Coma Scale <5</li>
- Decision for palliative (end of life) care

EU Protocol v4.2 approved on 19.04, 2023



# **TICH-3: Patients taking DOACs**



#### Which DOACs can the patients be on to be recruited for TICH 3?

- Direct thrombin inhibitor Dabigatran or Factor 10a inhibitor Apixaban, rivaroxaban, edoxaban
- If they have only recently started taking a DOAC, for as long as they have regularly taken it for the last 48 hours, we would enrol them as patients taking regular DOAC

#### Types of anticoagulation (blood thinners) that cannot be included:

 warfarin or therapeutic low molecular weight heparin. Note if LMWH is being used prophylactically at low dose to prevent DVT and PE these patients can be included in TICH-3

#### If patients on DOAC with ICH are enrolled to TICH-3 can they still have reversal agents?

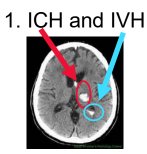
Yes, all patients should receive standard care, and be treated as per local DOAC ICH guidelines and given PCC or anticoagulation reversal agent (i.e Andexanet-alfa or Idarucizumab) in accordance with the treating physician. Please ensure you document which reversal agents were given in eCRF on day 7

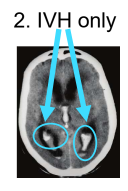


# Eligibility: Frequently asked questions



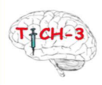
- If time of stroke onset is unknown?
   Patient can be enrolled if presenting within 4.5 hours of discovery if HV < 60mls on CT scan.</li>
- Can patients with intraventricular haemorrhage (IVH) be enrolled?
   Yes, so long as they have intracerebral haemorrhage, (fig 1) do not have other exclusion criteria. Isolated IVH (fig 2) should not be included.
- Can patient be enrolled if they are a candidate for neurosurgery?
   Yes, neurosurgery is not an exclusion.
- Can patient be enrolled if they have a DNAR/from care home/already dependent?
   Yes, so long as they are still for active care and consent is obtained
- Can patients with recurrent bleeds be enrolled? Yes, it is likely that most patients will have an arteriopathy due to hypertension or cerebral amyloid angiopathy.
- Can a nurse consultant assess eligibility? Confirming eligibility is defined as a medical decision, so must be undertake by a medically qualified doctor under the clinical trials regulations.







# **Investigator questions**



Q: If the patient has neurosurgery and is then given tranexamic acid is this a protocol violation?

 Only if it was known at the time of enrollment that the patient was to be given tranexamic acid (Indication for tranexamic acid is an exclusion criteria). However it should be noted that there is currently no evidence that TXA is effective in ICH and it is not routinely used.

Q: If the relatives want longer to make a decision (more than 10 minutes as suggested in the protocol) is that ok?

Yes – they should take as long as they need, but need to explain to them that in other bleeding conditions TXA is more effective if given as soon as possible after stroke onset, when the risk of bleeding (haematoma expansion) is greatest. If they cannot decide within the 4.5hours since stroke onset the patient should not be included.



## Haematoma volume measurement

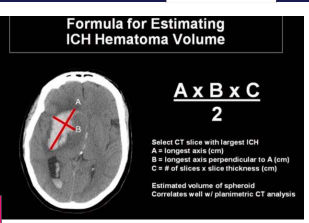


### **Exclude patients with massive haematoma (usually >60ml)**

- If CT scan uses automated software e.g. RAPID, BRAINOMIX patient can be enrolled if HV not greater than 60mls
- Calculate HV manually using TICH-3 HV=ABC/2 calculator on the website<sup>1</sup> or alternatives e.g. mdcalc app<sup>2</sup> Dimensions can be obtained from neuroradiology or measured directly.
- 3. If ABC/2 not possible: measure the maximum length of the haematoma. Exclude if max length A > 5cm
- Do not include IVH in measurements
- HV can be estimated by anyone trained to do so

The <u>ABC/2 calculator</u> can be used to calculate haematoma volumes during eligibility checks, without needing to be logged in.

Estimated volume of largest haematoma	1
View guide	•
Maximum haematoma length 'A' (up to 4 decimal places)	cm
Maximum haematoma width 'B' (up to 4 decimal places)	cm
Number of slices where haematoma visible	slices
Scan slice thickness (up to 3 decimal places)	mm



INSTRUCTIONS  Measure length and width or slices are typically measured		h the largest are	a of hemorrhage. NOTE: CT
When to Use 🗸	Pearls/Pi	itfalls 🗸	Why Use 🗸
Hemorrhage Shape		Round or Ellips	oid
		Irregular, Sepa	rated, or Multinodular
Hemorrhage Length			cn
Hemorrhage Width			cn
Number of CT Slices Slice with ≥75% Area of Hemorrh slice; Slice with 25-755¢ Area of H Counts as 0.5 slices; Slice with <2 Hemorrhage: Counts as 0 slices	emorrhage:		slices
CT Slice Thickness			mm

https://www.mdcalc.com/abc2-formula-intracerebral-hemorrhage-volume

# CONSENT



# **Emergency Consent Process**



# Rapid consent process, participants or relatives provide verbal consent

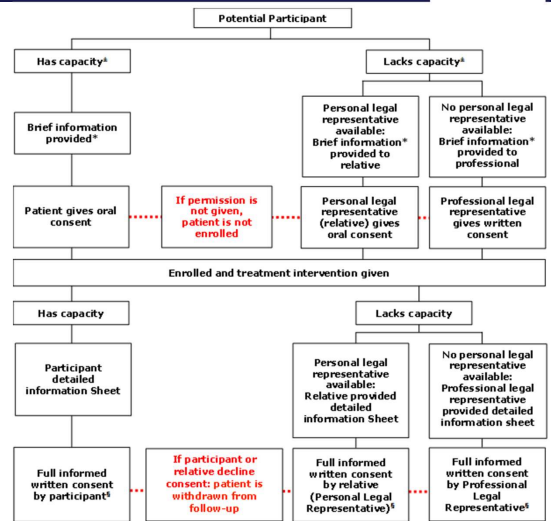
Full informed written consent to be obtained later after administration of IMP

#### Hierarchy approach

Patient has capacity – gives oral consent

- Patient does not have capacity relative or close friend likely to know patient wishes provides oral consent
- 2. Patient does not have capacity and no relatives available contact relatives via telephone to obtain consent.
- 3. Patient does not have capacity and no relatives available independent doctor provides written consent

The person taking consent must be appropriately trained and on the delegation log





## **Recent Approvals**

#### **HRCDC Approval**

- HRCDC Meeting 26th March 2024 | Decision Ref ID: 23-019-AF1/AMD1 [TICH-3]
   CONCLUDED ACCEPTED WITH CONDITIONS 5th of April 2024
- Scope of Amendment: Scope of amendment(s): The amendment covers the continued processing of personal data of participants who lack decision making capacity, in the rare situation where proxy assent from a relative/friend is not obtained and only initial permission for enrolment from the independent doctor is in place. As outlined in the information provided by the Applicant, the data to be processed in the event of this rare scenario is minimised (i.e., minimum safety data, alive and well status and modified rankin scale at 6 months)

### CTIS Amendment Approved 12 June 2024

- GP letter
- PILFs updated to remove UK references

#### **Documents Page link**

https://stroke.nottingham.ac.uk/tich-3/docs/?view=YMvDqkLgOSKgCU3lJgNVbr2gr4C6Kn2 SKPO

#### International countries / Republic of Ireland

#### **Republic of Ireland**

- » Please click here to view superseded documents
  - Family doctor letter international 20240213 v1.0.docx
  - International Ireland MASTER Investigator meeting 20230511 V1.0.pdf
  - · Participant Full Consent Form Ireland 20240116 v1.2.docx
- Participant Information Sheet Final Ireland 20240116 v1.2.docx
- Participant Short Information Sheet Final Ireland 20231221 v1.2.docx
- Professional Legal Rep Full Consent Form TICH3 Final Ireland 20240116 v1.2.docx
- Professional Legal Rep Information Sheet Final Ireland 20240116 v1.2.docx
- Professional Legal Rep Short Information Sheet and Consent Final Ireland 20240116 v1.2.docx
- Relative Legal Rep Full Consent Form Final Ireland 20240116 v1.2.docx
- Relative Legal Rep Information Sheet Final Ireland 20240116 v1.2.docx
- Relative Legal Rep Short Information TICH3 Final Ireland 20240116 v1.2.docx

Please add date and link to document page where all the approved Irish documents are Nikola Sprigg (staff), 2024-08-13T13:02:45.021 NS0



# Delegated roles for consent: J and Z

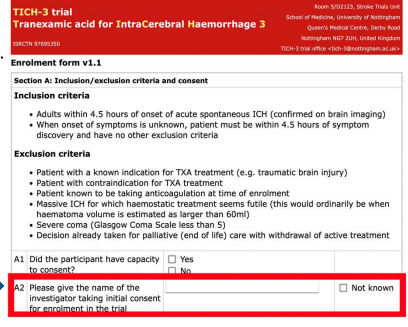
#### Person taking initial consent must be delegated role J

The PI must select whether code J should be applied as a delegated role.

J. Obtaining consent for enrolment (including oral consent, as appropriate to local policy and practice). I, PI, DPI

Site investigator (Not authorised)
Consent training
for enrolment (J)
No consent training

 Monitoring will check patient was consented by someone on delegation log



### Person taking written consent must be delegated role Z

Z assigned to those with relevant investigator roles (not pharmacists, radiology etc) and confirmed by PI.

Obtaining follow-on written consent (after initial consent) to continue in the study and for follow-up. I, PI, DPI



# Professional legal representative consent by an independent doctor



#### **Enrolment consent by independent doctor**

Short Information Sheet and Consent form should be used (pictured to the right). In this scenario this professional legal representative enrolment consent is handwritten and then a follow on written consent form is not required to be completed by the independent doctor. If the participant regains capacity or a relative becomes available they should complete the written follow on consent.

#### Follow on written consent by independent doctor

The follow on written consent form for professional legal representative should only be used if participant has capacity and consents for enrolment orally, then loses capacity and no relatives are contactable to provide the handwritten follow on consent. If the participant regains capacity or a relative becomes available they should complete the written follow on consent.

#### Informing relatives

The clinician at site has full responsibility for informing relatives of participant when professional legal representative consent has taken place. In event of a patient dying after being enrolled by a professional legal representative but before relatives can be contacted the clinical team should inform the relatives of the patient's involvement in the study and provide information about the study.

[Form to be printed on local headed paper]	
	[Form to be printed on local headed paper]
PROFESSIONAL LEGAL REPRESENTATIVE SHORT INFORMATION SHEET AND CONSENT	You have been asked to act as a professional legal representative to consider if you think the patient named above should take part in the TICH-3 study.  TICH-3 aims to assess whether the drug tranexamic acid reduces the risk of death and/
(Draft Version 0.2 / Final Version 1.0: 03/11/2021)	improves disability 6 months after stroke due to intracerebral haemorrhage (ICH).
e of Study: TICH-3	Because intracerebral haemorrhage is an emergency and the potential benefits of the stu
.S Project ID: 297457 CTA ref: 03057/0074/001-000	
me of Researcher:	given, every minute counts. We need to decide about giving the treatment as quickly possible. As the patient is not well enough to decide, and no relatives are immediately availal
ne of Participant:	you have been asked to decide on their behalf. You are able to make this decision in accordan
nfirm that I have been given a copy of the Short Professional Legal Representative Information Shee sion 1.0 dated 3/11/20(1) and I agree as professional Legal Representative on behalf of this stroke ent.  The patient will take part in the TICH-3 study and be given the study medication  For their medical records to be accessed  To be followed up at the moths  For their GP to be informed  For their contact details to be collected and used for the purpose of the study  For their contact details to the collected and used for the purpose of the study	with emergency consent procedures,  The patient has been identified because they have had a stroke caused by intracereb haemorrhage - and they fit the requirements for this research project. At present they are able to tall us whether to take part, so we are asking your opinion. If you do decide they wo take part you will be given this information sheet to keep and be asked to sign a consent for We are inviting approximately 5300 participants with intracerebral haemorrhage to take p from around the UK and worldwide.
destand that they are free to withdraw from the study at any point without giving a reason. participants who are enrolled following agreement by a professional legal representative as soon tives are available or when the patient regains capabily, a detailed information sheet will be provided, en consent sought for continuation in the trial.	
Professional nominee consent - to be completed if participant does not have capacity to consent	
	infusion through a venous cannula with a loading dose infusion over 10 minutes followed by an infusion over 8 hours.
capacity to consent  Name of Person giving Date Signature	infusion through a venous cannula with a loading dose infusion over 10 minutes followed by an infusion over 8 hours.  During the next 7 days members of the clinical and research team will monitor the potent participants condition and record relevant information from their medical notes.
Capacity to consent  Name of Person giving Date Signature nominine consent	infusion through a venous cannula with a loading dose infusion over 10 minutes followed by an infusion over 8 hours.  During the next 7 days members of the clinical and research team will monitor the potent participants condition and record relevant information from their medical notes.  For participants who are enrolled following agreement by a professional legal representative
Capacity to consent    Name of Person giving   Date   Signature	infusion through a venous cannula with a loading dose infusion over 10 minutes followed by an infusion over 6 hours.  During the next 7 days members of the clinical and research team will monitor the potent participants condition and record relevant information from their medical notes.  For participants who are enrolled following agreement by a professional legal representative soon as relatives are available or when the patient regains capacity, a detailed informatic sheet will be provided, and written consent sought for continuation in the trial.
Capacity to consent    Name of Person giving   Date   Signature	During the next 7 days members of the clinical and research team will monitor the potentit participants condition and record relevant information from their medical notes.  For participants who are enrolled following agreement by a professional legal representative: soon as relatives are available or when the patient regains capacity, a detailed informatic sheet will be provided, and written consent sought for continuation in the trial.  The participants' decision to withdraw would overrule the decision of either a professional or



### Written follow on consent

# The person taking written consent must be appropriately trained and delegated code Z by the PI to take consent on the delegation log

Full, written informed consent will be sought as soon as practicable, ideally within the next 24 hours. Written informed consent will be sought for access to medical notes and for participation in the trial follow up. The participant information sheet will be provided to the participant at this time if not already provided. Please localise the consent forms and participant information sheets prior to printing, see WPD preparing trial documentation to help you with this https://stroke.nottingham.ac.uk/sif/docs/?sid=TICH-3

University of Nottingham	FULL CONSENT FORM (Final version 1.1)		T+CH-3		
Title of Study: TICH	<del>1</del> -3		ém		
IRAS Project ID: 29	7457	CTA ref: 0305	7/0074/001-000		
Name of Researche	er:				
Name of Participan	t:		Please initial box		
	ve read and understand the in above study and have had the	formation sheet final version 1 opportunity to ask questions.	.0 dated		
without giving any understand that she	reason, and without my medic	d that I am free to withdraw at a cal care or legal rights being a nation collected so far cannot b roject analysis.	ffected. i		
be looked at by au group and regulato permission for the analyse and publish	I understand that relevant sections of my medical notes and data collected in the study may be looked at by authorised individuals from the University of Nottingham, the research group and regulatory authorities where it is relevant to my taking part in this study. I give permission for these individuals to have access to these records and to collect, store, analyse and publish information obtained from my participation in this study. I understand that my personal details will be kept confidential.				
box). I agree that the info Nottingham, for pos may be carried out including researche	rmation gathered about me car ssible use in future studies. I ur		YES/NO		
		ned by NHS Digital, (EDRIS in be used to help contact me or p	rovide		
	eing informed of my participation formation on my status for the	on in this study and who will be 180 Day follow up.			
asked to provide in		during the course of the study,			
7. If I lose the capacit		<b>3</b>	GE10000000000000		
If I lose the capacit     I'd be happy to con     raises an objection	to this.	5-507 89	YES/NO		
If I lose the capacit     I'd be happy to con     raises an objection     I agree to you send	to this. ing me a letter/email with a sun initial in box).	5-507 89	GE10000000000000		
7. If I lose the capacit I'd be happy to con raises an objection  8. I agree to you sendi (delete yes/no and)	to this. ing me a letter/email with a sun initial in box).	5-507 89	GE10000000000000		

University of Nottingham	RE	OR LEGAL REPRESENTATIVE LATIVE on 1.0: 03/11/2021	Tjan-3
Title of Study: TICH-3			
IRAS Project ID: 297457		CTA ref: 03057	//0074/001-0001
Name of Researcher:			
Name of Participant:		P	lease initial box
		e information sheet final version 1 the opportunity to ask questions.	.0 dated
at any time, without giving affected. I understand the	any reason, and wit at should they with	voluntary and that they are free to thout their medical care or legal rigi draw then the information collecte y still be used in the project analysi	nts being died so far
study may be looked at b research group and regula give permission for these i	y authorised individuatory authorities whe ndividuals to have a nation obtained from	ve's medical notes and data collect uals from the University of Notting rer it is relevant to taking part in this coess to these records and to colle participation in this study. I undersidential.	ham, the s study. I ct, store,
box). I agree that the information University of Nottingham, it these studies may be carrifirst study, including resea	n gathered about my for possible use in fu led out by researche rchers working for co	ch (Optional) (delete yes/no and inity relative can be stored by the ature studies. I understand that some irs other than the current team who ommercial companies. Any data us not be identified in anyway.	YES / NO
	I UK NHS bodies m	intained by NHS Digital, (EDRIS in ay be used to help contact my relat	ive or
		heir participation in this study and t status for the 180 Day follow up.	hat they
<ol> <li>I agree to you sending my (delete yes/no and initial ir</li> </ol>		il with a summary of the results	YES/NO
8. I agree to my relative takin	g part in the above s	study.	
Name of participant	Relationship	to participant	
Name of Relative	Date	Signature	<u>-12</u> ;

trainventity of Nottingham	Prof	R LEGAL REPRESENTAT essional n 1.0: 03/11/2021	TVE - TICH-3
Title of Study: TICH-3			
IRAS Project ID: 29745	7	CTA ref: 0:	3057/0074/001-00
Name of Researcher:			
Name of Participant:			Please initial bo
		e information sheet final versi he opportunity to ask question	
at any time, without give affected. I understand	ing any reason, and with that should they withd	roluntary and that they are fre hout their medical care or lega fraw then the information col still be used in the project an	I rights being lected so far
the study may be looked research group, and re- give permission for the	d at by authorised individual gulatory authorities whe se individuals to have ac primation obtained from	ipant's medical notes and dat duals from the University of No re it is relevant to taking part i ocess to these records and to participation in this study. I un infidential.	ttingham, the n this study. I collect, store,
box). I agree that the informs University of Nottinghar these studies may be c first study, including res	tion gathered about the m, for possible use in fut arried out by researcher	h (Optional) (delete yes/no an participant can be stored by t ture studies. I understand that is other than the current team immercial companies. Any dat httfied in anyway.	he YES
	ntral UK NHS bodies ma	ntained by NHS Digital, (EDRI ay be used to help contact the	
		of their participation in this student status for the 180 Day follows:	
7. I agree to you sending t (delete yes/no and initial		mail with a summary of the res	sults YES/No
8. I agree to the participar	nt taking part in the abov	ve study.	
Name of participant	Relationship	to participant	
Name of professional	Date	Signature	<u></u>
	consent Date	Signature	

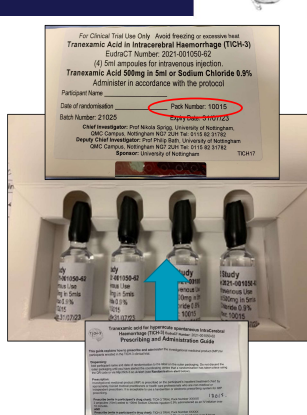
# RANDOMISATION



### Randomisation



- Blinded treatment packs will be randomly assigned to sites in blocks of 6 treatment packs
- TICH-3 will use simple randomisation
- After confirming eligibility and obtaining consent the investigator randomises the patient by selecting and opening the treatment pack with the lowest pack number.
- The prescribing and administration guide can be found inside each treatment pack.
- Due to emergency nature of trial randomisation is notified to the coordinating centre after the IMP has been administered by completing the randomisation alert (guidance for this is within the prescribing and administration guide).





## Prescribing and Administering the IMP



#### Prescribing the IMP

Investigational medicinal product (IMP) is prescribed on the participant's inpatient treatment chart by appropriately trained medical practitioners or health care professionals who are non-medical or independent prescribers. It is acceptable to use a handwritten or electronic prescribing system for IMP prescribing.

Do not need to be on delegation log to prescribe

Prescribe (write in participants drug chart):

TICH-3 - TRIAL Pack Number XXXXX

TRANEXAMIC ACID OR PLACEBO

2 ampoules (10ml) added to 100ml Sodium Chloride Injection 0.9% administered as an IV infusion over 10 minutes.

AND

TICH-3 TRIAL Pack Number XXXXX

TRANEXAMIC ACID OR PLACEBO

2 ampoules (10ml) added to 250ml Sodium Chloride Injection 0.9% as an IV infusion over 8 hours.

#### Administering the IMP

Administer in accordance with the prescription. The treatment can be administered by anyone qualified and appropriately trained to give intravenous injections. Do not need to be on delegation log to administer



#### Tranexamic acid for hyperacute spontaneous IntraCerebral Haemorrhage (TICH-3)

EudraCT Number: 2021-001050-82

#### Prescribing and Administration Guide

This guide explains how to prescribe and administer the investigational medicinal product (IMP) for participants enrolled in the TICH-3 clinical trial.

Add participant name and date of randomisation to the label on the outer packaging. Do not discard the outer packaging until you have alerted the coordinating centre that a randomisation has taken place using the QR code or via http://tich-3.ac.uk/alert (see Randomisation alert below).

#### Prescription

Investigational medicinal product (IMP) is prescribed on the participant's inpatient treatment chart by appropriately trained medical practitioners or health care professionals who are non-medical or independent prescribers. It is acceptable to use a handwritten or electronic prescribing system for IMP

Prescribe (write in participant's drug chart): TICH-3 TRIAL Pack Number XXXXX 2 ampoules (10ml) added to 100ml Sodium Chloride Injection 0.9% administered as an IV infusion over 10

Prescribe (write in participant's drug chart): TICH-3 TRIAL Pack Number XXXXX 2 ampoules (10ml) added to 250ml Sodium Chloride Injection 0.9% as an IV infusion over 8 hours.

Administer in accordance with the prescription. The treatment can be administered by anyone qualified and appropriately trained to give intravenous injections.



1. Scan the QR code or enter http://tich-3.ac.uk/alert in your web browser which will take you to the TICH-3 website. You do not need to have a TICH-3 investigator account 2. Enter the treatment pack ID number

Please notify the coordinating centre in Nottingham as soon as

3. Enter randomisation details as prompted by the system

If you cannot access the website, please call the trial office on 0115 8231782 with treatment pack ID, participant's initials, date of randomisation and name of person enrolling the participant (voicemail

4. Enter participant details on the IMP accountability log against the

Disposal of IMP packaging can take place only after the following tasks have been completed;

- Participant randomisation alert completed on the TICH-3 website Participant details added to the IMP accountability log at site.
- Prescription written on the participant's drug chart (using the guidance above)

Return any unused ampoules to clinical trials pharmacy AND record reason for non-use on the IMP accountability log.

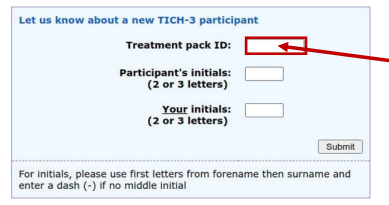


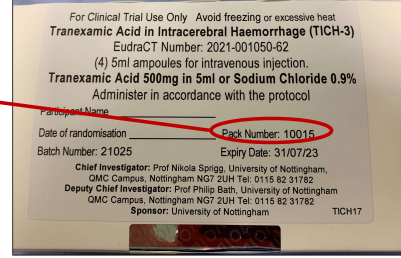
### **Randomisation Alert**

1. Investigator will enter the treatment pack ID (pack number), participant initials and their own initials to alert

the coordinating centre to a new randomisation.







2. Investigator will then confirm that the participant was randomised at the hospital shown in the alert box.





## **Broken vials:**



#### Broken prior to randomisation e.g. upon receipt in pharmacy

✓Inform the Nottingham coordinating centre and dispose of the pack(s) in accordance with WPD (Destruction of IMP).



#### Broken after randomisation, before treatment:

✓ Disregard this pack and use the lowest treatment pack ID that is available at your centre

#### Broken during treatment i.e. Bolus given but infusion vial breaks:

- ✓ Administer as much drug as possible
- ✓ Enter a protocol violation for 'participant does not receive all of the randomised treatment as per protocol'.
- x Do not open another treatment pack

Always record broken vials on the inventory or accountability log as appropriate

# SAFETY MONITORING

Safety outcomes
Serious adverse reaction (SAR)
Suspected Unexpected Serious Adverse Reaction (SUSAR)
Serious adverse event (SAE)



# **Good Clinical Practice (GCP)**



- TICH-3 is to be performed in line with all the principles of good clinical practice
- Investigators must adhere to the protocol at all times
- The safety and rights of the participant are paramount
- Training for investigators should be in proportion to their role within the trial and in accordance with their experience and skills
- The participant has the right to withdraw at any time without giving a reason, without it affecting their medical care
- Investigators eligible for NIHR GCP online training

https://portal.nihr.ac.uk/register





# **GCP** training



Free GCP training - In English, French and Spainish



https://globalhealthtrainingcentre.tghn.org/ich-good-clinical-practice/







## Safety Events, SARS and SUSARS



TXA has an established safety record – we only collect data on focused **safety outcomes** occurring within the **first 7 days or events suspected to be related to the IMP (SAR or SUSAR)**:

Safety outcomes: \*\*If a safety outcome (e.g. seizure) occurs during infusion, the infusion must be stopped immediately\*\*

- 1. Venous occlusive events: VTE (Pulmonary embolism, Deep vein thrombosis)
- 2.Ischaemic events (arterial thrombosis at any site, ischaemic stroke, transient ischaemic attack peripheral artery embolism, myocardial infarction, acute coronary syndrome)
- 3.Seizures
- 4. Fatal events up to discharge from hospital

#### Serious adverse reactions (SAR) or Suspected Unexpected Serious Adverse Reactions (SUSAR):

 All events suspected to be related to the IMP will be assessed for seriousness, expectedness and causality by local investigator. Section 4.8 of the SmPC, date of last revision 02 February 2021, will act as the Reference Safety Information: Tranexamic Acid https://Tranexamic Acid SmPC 20210202 REVISION.pdf

Serious Adverse Events (SAEs) that are not safety outcomes, SARS or SUSARS should not be reported E.g. Neurological deterioration, haematoma expansion, cerebral oedema that is NOT thought to be related to the IMP, and does not result in death does not need to be reported as an SAE



# SAE reporting cause of death



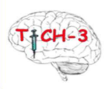
# IF PARTICIPANT DIES PRIOR TO DISCHARGE FROM HOSPITAL PLEASE REMEMBER TO COMPLETE SAE FORM

- Please complete SAE as soon as possible and also complete the discharge/death eCRF.
- Please provide details in question A4 and cause of death in question A5 e.g. Pneumonia, cerebral oedema
- Please only use death unattended in the rare situation when a patient has completely unexplained death e.g. in community

A4	Please describe the event, e.g. new limb weakness, crushing chest pain, bleeding gums, rash  Note: Death is an end result, not an independent event	
A5a	Event sub-categorisation  Please only enter a  code/description from the SAE  sub-category list	



# What to do in Case of Emergency



#### Safety events during the infusion:

If seizure, thrombosis or arterial occlusion occurs during infusion, the infusion must be stopped immediately. This will be recorded as part of the trial documentation and safety monitoring.

#### **Emergency Unblinding**

In general there should be no need to unblind the allocated treatment. If some contraindication to tranexamic acid develops after randomisation (e.g. clinical evidence of thrombosis), the trial treatment should simply be stopped. Unblinding should be done only in those rare cases when the doctor believes that clinical management depends importantly upon knowledge of whether the patient received TXA or placebo. In those few cases when urgent unblinding is considered necessary, the emergency telephone number should be telephoned, giving the name of the doctor authorising unblinding and the treatment pack number. The caller will then be told whether the patient received TXA or placebo.

#### Eligibility query or any other emergency query:

Call the emergency contact number listed on TICH-3 website

For urgent medical enquiries (including <u>unblinding</u>), and for randomisation problems, you can contact the following emergency mobile numbers. Please ensure that you have these written down.

+44 (0)7725 580 092 +44 (0)7736 843 592

+44 (0)7798 670 726 +44 (0)7810 540 604



### What to do in the event of a Protocol Violation



A protocol violation is a major variation in practice from the trial protocol, for example where a participant is enrolled in spite of not fulfilling all the inclusion and exclusion criteria (e.g. lack of consent, randomisation after 4.5 hours after ICH), or where deviations from the protocol could affect participant safety, the trial delivery or interpretation significantly.

\*\*Important to report any protocol violations to coordinating centre straight away\*\*

All protocol violations must be reported to the Chief Investigator, via the online electronic case report form and/or telephone call. The CI will notify the Sponsor if a violation has an impact on participant safety or integrity of the trial data. The Sponsor will advise on appropriate measures to address the occurrence, which may include reporting of a serious GCP breach, internal audit of the trial and seeking counsel of the trial committees.

## IMP AND PHARMACY



### Storage of IMP



Temperature monitoring is not required. The packs will be stored at room temperature and protected from excessive heat and freezing.

The IMP is stored in a secure, limited access storage area, this could be in the A&E, stroke ward or thrombolysis bag.

Each site will maintain an accountability log and be responsible for the storage and issue of trial treatment.

Ensure all members of the local team are aware of where the IMP and related documents (consent forms/PIS) are stored.







### **Drug dispatch: International**

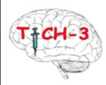


- Coordinating centre will order drug for dispatch once site is nearly ready to commence recruitment
- Blinded treatment packs will be randomly assigned to sites in blocks of 6 treatment packs
- Pharmacy informed of dispatch by email
- Delivery after noon next day of ordering
  - > No deliveries out of hours/weekends
- Pharmacy complete inventory log and part of accountability log and pass accountability log and treatment packs to research team for storage
- Investigator needs to 'mark available for randomisation' on TICH-3 website
- Coordinating centre will re-order/issue when stock running low or when drug due to expire

			ent packs fo Derby TEST	or hospital C002 hospital			
Block	Treatment pack IDs	Dates assigned/ dispatched to centre	Date at pharmacy	Date at stroke unit		mised/ naining	Comments
3	60157 60160 60174 <del>60188</del> 60191 60201	15 Sep 2021 -	15 Sep 2021	☐ Mark as available for randomisation	)	5	-
4	60215 60229 60232 <del>60246</del> 60263 60277	15 Sep 2021 -	15 Sep 2021	31 Jan 2022	1	5	ē
5	60280 60294 60304 60318 60321 60335	15 Sep 2021 -	15 Sep 2021	15 Sep 2021	0	6	
3 blocks	18 packs	18 assigned / 0 dispatched	18 received	11 available	2 use	d / 16 r	emaining



### **Monitoring of IMP**



The local sites pharmacy is responsible for the accountability and monitoring of the IMP.

The IMP will be shipped from Sharp directly to the site's pharmacy. The pharmacy will complete the inventory log and part of the accountability log and then distribute to the research team with the IMP to be placed in the agreed storage location (discussed and agreed when completing the assessment and monitoring of IMP storage form).

Once the IMP is in the storage location, pharmacy/research team will need to login to the TICH-3 web site and mark the treatment packs as available for randomisation.

The following forms are downloadable from the TICH-3 website and form part of the pharmacy's site file;

- 1. Assessment and Monitoring of IMP Storage to be completed prior to initiation
- 2. Inventory Log to be completed by pharmacy when IMP arrives at site
- 3. IMP Accountability Log to be completed by research team when IMP is used at site
- 4. IMP Check to be completed by research team to ensure IMP all present and accounted for



### IMP Paperwork (1): Set up, IMP receipt





#### Assessment and monitoring of remote IMP storage

Study Title:	Tranexamic acid for hyperacute spontaneous IntraCerebral Haemorrhage (TICH-3)
EudraCT No:	2021-001050-62
Chief Investigator:	Professor Nikola Sprigg
Site:	
Principal Investigator:	

Description of propo	osed area for IMP	Suitable for use (Yes/No)	Comments
Security measures in place (location, access controls etc)			
Size and description of proposed storage area (shelves, cupboards etc)			
If not for exclusive use, what controls are in place to segregate IMP from other medicines		9	This cabinet is for CLINICAL TRIAL USE only
Description of IMP ma	anagement. The follow	ing shou	and may contain
Dispensing procedure with documented training for research team	Select the next lowest numbered available treatment pack. Prescribing and administration guide to be followed		PLACEBO
Accountability procedure with documented training for research team	Prescribing and administration guide to be followed.		Anticophy annihilation of the Control of the Contro
A procedure for transfer of IMP between pharmacy and proposed storage facility		+3	The state of the s
Proposed methods of maintaining pharmacy oversight			TICH-3

#### 1. Assessment and monitoring of remote IMP storage

Pharmacy and trial team to complete form as part of site set up, Pharmacy and local trial team to complete, sign and then return to coordinating centre as part of site set up, before green light can be

issued.

#### 2. IMP Inventory Log

Pharmacy to complete inventory upon receipt of the IMP treatment packs (will be sent to sites in blocks of 6 treatment packs). Inventory log to be retained in the pharmacy site file.

EudraCT No: Principal Investigator:		2021-001050-62		Site:				
				Storage location:	Stroke unit / ED / other		/ other	
Date	Block number	Pack number	Do not use after	Received by	Date sent to str unit/ED from pharmacy	roke	Initials	Comments



### IMP Paperwork (2): Ongoing



EudraCT I	No:	2021-001050-62			Site:			
Principal Investigator:			Storage lo		Storage location:	Stroke unit / ED	/ other	
			Issu	Comments (reasons				
Pack number	Date sent to stroke unit/ED from pharmacy	Participant's Name	Participant's Hospital number/NHS number	Issued by	Checked by	Issue date and time	use & date returned to pharmacy)	

#### 3. IMP Accountability Log

Pharmacy to complete the first two left columns (pack number and date sent to stroke unit/ED). The accountability log will then be passed to the research team for them to complete when a treatment pack is randomised to a participant. Once the IMP is in the storage place please mark on the TICH-3 website that the treatment pack is ready for randomisation. Once this form is complete please return to local site pharmacy.

#### 4. IMP Check

The research team should complete checks of the IMP at least monthly. Any discrepancies are required to be investigated immediately and reported to the coordinating centre.

The inventory logs, completed accountability logs and IMP check should be stored in your site file. We do not require these to be uploaded to the TICH-3 website or sent to us unless requested.



#### IMP Check

4

#### \*\* CHECKS MUST BE COMPLETED AT LEAST MONTHLY \*\*

Study Title:	Tranexamic acid for hyperacute spontaneous IntraCerebral Haemorrhage (TICH-3)
EudraCT No:	2021-001050-62
Site:	
Principal Investigator:	

I confirm that I have checked that all treatment packs held in the locked cupboard matched the drug accountability log and the TICH-3 website, on the date shown below and that all are in date. NOTE: All expired IMP must be destroyed.

Any discrepancies must be recorded and investigated immediately to resolve the situation. The coordinating centre in Nottingham must also be informed immediately and kept up to date with any investigations.

DATE/TIME	SIGNATURE	COMMENTS	
	0	8	
	0	8	
		8	
	6	8	
	8	9	
	6)	8	
	8	8	
	2	8	
		<u> </u>	
	8	8	
	8	<u>×</u>	
	8		
	6		



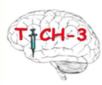
### **Pending from Mater**

- Pharmacy address to send IMP
- Online delegation log authorization

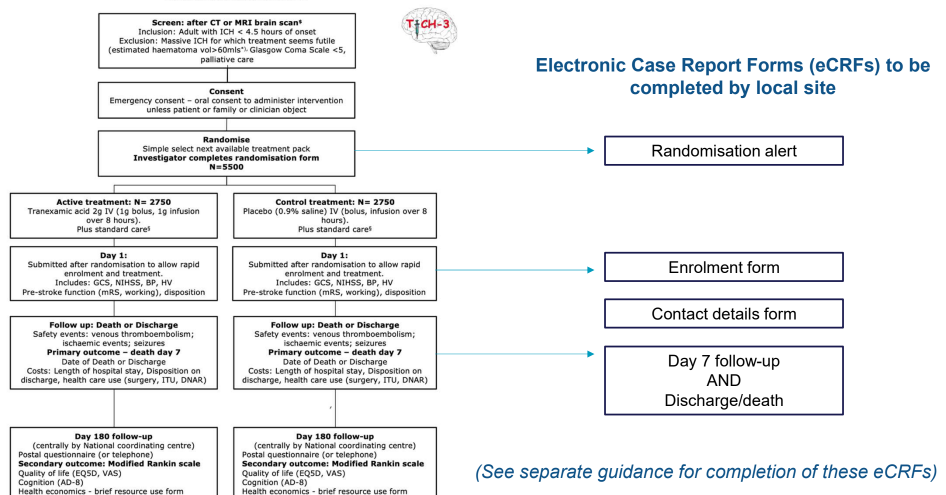
## DATA COLLECTION



### **Trial Flow Chart:**



TICH-3 Flow chart Protocol v1.3 4.Mar 21





### **Logging onto TICH-3 website**



TICH-3 trial Tranexamic acid for IntraCerebral Haemorrhage 3	Room S/02123, Stroke Trials Unit School of Medicine, University of Nottingham
	Queen's Medical Centre, Derby Road Nottingham NG7 2014, United Kingdom
ISRCTN 97695350	TIGH-3 total diffice such-3enothrophanacouts  Please log in if you need the TICH-3 emergency contact numbers. The recruitment total for the trial to date is: 0

Login using the investigator ID, password issued to you by the <u>TICH-3 trial office</u>. If you have forgotten your login details then please <u>click here</u>.

TICH-3 investigator ID:	
Password:	
	Login

Please ensure that your web browser has both cookies and JavaScript enabled.

NOTE: Serious Adverse Events (SAEs) — we have a legal responsibility to collect all safety events occurring within the first 7 days after randomisation (including SARs/SUSARs/fatal SAEs).

Safety events include: venous thromboembolism; ischaemic events (arterial thrombosis at any site, ischaemic stroke, transient ischaemic attack, peripheral artery embolism, myocardial infarction, acute coronary syndrome) and seizures.

Please remember that fatal SAEs need to be reported until discharge from hospital, even if this is after 7 days. Please assess if expected according to SmPC: <a href="https://medicines.org.uk/emc/product/1220/smpc">https://medicines.org.uk/emc/product/1220/smpc</a>

Investigators have a legal responsibility to report applicable SAEs to the chief investigator within 24 hours.

Documents

Switch to mobile site

#### TICH-3 trial

Tranexamic acid for IntraCerebral Haemorrhage 3

ISRCTN 97695350

Room S/D2123, Stroke Trials Unit School of Medicine, University of Nottingham Queen's Medical Centre, Derby Road Nottingham NG7 2UH, United Kingdom TICH-3 trial office < tich-3@nottingham.ac.uk>

Log out

Logged in as: Nikola Sprigg <nikola.sprigg@nottingham.ac.uk> (update email address)

For urgent medical enquiries (including unblinding), and for randomisation problems, you can contact the following emergency mobile numbers. Please ensure that you have these written down.

+44 (0)7725 580 092 +44 (0)7736 843 592 +44 (0)7798 670 726 +44 (0)7810 540 604



The <u>ABC/2 calculator</u> can be used to calculate haematoma volumes during eligibility checks, without needing to be logged in.

- Coordinating centre will set up an account for investigators we need the completed attendance at investigator training log completed (electronic signatures are accepted) and returned to us via email to know who needs logins for the TICH-3 website and subsequently added to the delegation log
- Additional investigators can be added later
- PI must activate before site can recruit
- Password reset on-line



### Adding a new participant to the database

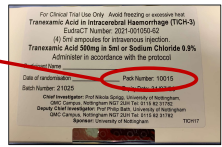


1. Complete randomisation alert





3. Need treatment pack ID number



4. Confirm randomisation site

#### 5. Complete enrollment form

Carti	This will be a record of a manual ra  The next available trial number wil		0.5.5		t pack 10 10015.
	usion criteria				
	Adults within 4.5 hours of onset of acute s When onset of symptoms is unknown, pat exclusion criteria				y and have no other
:	Patient with a known indication for TXA treat Patient with contraindication for TXA treat Patient known to be taking anticoagulation Massive ICH for which haemostatic treatm volume is estimated as larger than 60ml) Severe coma (Glasgow Coma Scale less th Decision already taken for palliative (end of	ment n at time lent seel nan 5) of life) c	e of enrolment ms futile (this would ordi are with withdrawal of a	narily be when hae	ematoma
A1 I	Did the participant have capacity to consent	17	Yes No		
	Please give the name of the investigator tak	ing	[Select]	*)	[Select] ‡
A2 i	initial consent for enrolment in the trial				[Select] *
į	initial consent for enrolment in the trial ion B: Participant details				[Selection]
Secti					(Ideleti)
į	ion B: Participant details Initials 3 letters from forenames then surname,	- Day -	\$ - Month - \$ - Year	•	[Josevi] *)
Secti B1	ion B: Participant details Initials 3 letters from forenames then surname, or 2 separated by a hyphen (-) Date of birth	- Day -		ī	[Joseph] Y



### **Contact Details Form**



	You can use the secure v	AUST NOT be entered into this pseudonymised database for TICH-3. rault (see below) to provide us with the participant's contact details and ition, which will be encrypted and stored separately.
0	Please indicate the docu	ure vault to upload required documents for the selected participant. ment type below, or use the 'Multiple document types' option (if you have ne file). If your file isn't a PDF or image, please 'print' it to a PDF first.
		Please choose a data type to access the secure vault
		Contact details  (*) applicable type only
	_	Switch to the secure vault site

It is really important to collect the participants identifiable data (as agreed on the consent form) so that the coordinating centre can contact the participant and the participant's GP for the Day 180 follow-up.

Identifiable data will be collected in the 'secure vault' which is accessed via the TICH-3 website. The secure vault site will encrypt the identifiable data so that it can only be accessed by the co-ordinating team, stored separately from the other data collected.



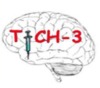


The information you enter into the form below will be encrypted and stored in the secure vault, held separately from the pseudonymised database used for TICH-3 CRFs.

A A A A D N O B A A S O O D O	This page will expire in 14 minutes and 50 seconds				
Please complete as much of this form as possible.  • Please make sure to include the participant's telephone number, which is required for follow-ups.					
TICH-3 participant ID:	C (female, 94 years old				
Surname:					
Forename(s):					
Middle initials:					
Permanent address:					
Post code:					
Country:	[Select]				
Follow-up telephone number:	[ecoca]				
Temporary residence:					
Alternate telephone number:					
Email address:					
Date of birth:					
NHS/CHI/H+C number:					
Hospital number:					
Name of hospital ward(s): (not hospital name)					
Place of birth:					
GP title/name:					
GP practice name:					
GP address:					
GP post code:					
GP telephone:					
Comments:					



### Enrolment, Day 7 followup and Discharge/death eCRF (2)



#### **Enrolment eCRF**

	anexamic acid for IntraCere	bral Haemorrhage 3	Queen's	ne, University of Nottinghi Medical Centre, Derby Ro n NG7 2UH, United Kingdi
ISRC	TN 97695350	TICH-3 trial office <tich-3@nottingham.ac.u< th=""></tich-3@nottingham.ac.u<>		
nr	olment form v1.1			
Sec	tion A: Inclusion/exclusion criteria	and consent		
Inc	clusion criteria			
	<ul> <li>Adults within 4.5 hours of onset</li> <li>When onset of symptoms is unk discovery and have no other exc</li> </ul>	nown, patient must be within		
Ex	clusion criteria			
	Patient with a known indication in Patient with contraindication for Patient known to be taking antic Massive ICH for which haemost haematoma volume is estimated. Severe coma (Glasgow Coma SC Decision already taken for pallia	TXA treatment coagulation at time of enrolmentic treatment seems futile (the das larger than 60ml) cale less than 5)	ent nis would ord	inarily be when
Α1	Did the participant have capacity to consent?	☐ Yes ☐ No		
A2	Please give the name of the investigator taking initial consent for enrolment in the trial			□ Not known
Sec	tion B: Participant details			
В1	Initials  3 letters from forenames then surname, or 2 separated by a hyphen (-)			
B2	Date of birth (dd-mmm-yyyy)	D/M/Y_	-	
ВЗ	Sex	☐ Male ☐ Female		
B4	Date/time of onset of index stroke (dd-mmm-yyyy hh:mm 24hr)	D/ M/ Y H: M		
В5	Date/time of randomisation (dd-mmm-yyyy hh:mm 24hr) If unknown, please use date/time of first dose	D / M / Y H : M		
В6	Allocated treatment pack ID			

#### Day 7 follow-up eCRF

Tranexamic acid for IntraCerebral Haemorrhage 3 No N			School of Medicine, University of Nottingham Queen's Medical Centre, Derby Road Nottingham NG7 2UH, United Kingdon	
Day	7 follow-up form v1.0		m	ICH-3 trial office <tich-3@nottingham.ac.uk< th=""></tich-3@nottingham.ac.uk<>
- 3	ion A: Day 7 follow-up			
A1	Participant status		☐ Alive and in hospital ☐ Discharged prior to day 7 ☐ Withdrawn from follow-up ☐ Died	
A2a	Was all randomised trea received?	itment	☐ Yes ☐ No	☐ Not known
A2b	Date/time of first dose (dd-mmm-yyyy hh:mm	24hr)	D / M / Y H : M	☐ Not done☐ Not known
A2c	Explanation if treatment received or data missing			☐ Not applicable
			Systolic / diastolic	
A3	Please enter BP recorde to 6 hours after stroke of			☐ Not done ☐ Not known
A4a	Blood pressure on day 7 reading 1	' -	/	☐ Not done ☐ Not known
A4b	Blood pressure on day 7 reading 2	' -	/	☐ Not done ☐ Not known
			_	
	ion B: Treatment during fi Was BP-lowering treatment given in the first 6 hours?	Yes No	•	☐ Not known
B1b	If yes, which antihypertensive drugs were given in the first 6 hours?	Glyceryl trinitrate (GTN) - patch Glyceryl trinitrate (GTN) - IV Sodium nitroprusside Other nitrate therapy (e.g. ISDN/ISMN) Urapidil Labetalol Other beta-blocker (e.g. atenolol, propranolol, bisoprolol) Calcium channel blocker (e.g. nifedipine, amlodipine) Diuretic (e.g. bendroflumethiazide, indapamide, hydrochlorothiazide)		

#### Discharge/death eCRF

TICH-3 trial Tranexamic acid for IntraCerebral Haemorrhage 3			Room S/D2123, Stroke Trials Unit School of Medicine, University of Nottingham Queen's Medical Centre, Derby Road Nottingham NG7 2UH, United Kingdom	
ISRCTI	N 97695350	TICH-	3 trial office	<tich-3@nottingham.ac.uk></tich-3@nottingham.ac.uk>
Disc	harge or death in hospital form v	1.0		
	r participants with a long stay in r as close as possible).	hospital, this form is to be	comple	eted by day 180
Sect	ion A: Discharge/death details			
A1	Date of discharge or death (dd-mmm-yyyy)	D/ M/ Y	_	
A2a	Discharge disposition	Home - independent, alon Home - independent, with partner/family/friend Warden-aided flat Residential home Home - needing care Carer's home Respite care Care home Nursing home Rehabilitation hospital In hospital Died Other		□ Not known
A2b	Did the participant return to their original place of residence?  If died, please select 'No'	☐ Yes ☐ No		□ Not known
А3	Please list any other trials into which the participant was co- enrolled			☐ Not applicable ☐ Not known
A4a	What was the final diagnosis of the randomising event?	☐ Intracerebral haemorrhage no known underlying cause ☐ Intracerebral haemorrhage underlying cause ☐ Ischaemic stroke with haemorrhagic transformatic ☐ Ischaemic stroke without ☐ Isc	e with	☐ Not known



# Enrolment, Day 7 follow-up and Discharge/death eCRF (1)



- The following eCRFs need to be completed in order on the TICH-3 website <a href="http://tich-3.ac.uk/live/">http://tich-3.ac.uk/live/</a>
  - 1. Enrolment form
  - 2. Day 7 follow-up
  - Discharge or death in hospital

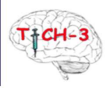


- Trial team members on the delegation log will have received an investigator ID and password to securely log in to the TICH-3 website
- You will need to enter the patients date of birth when entering data to confirm correct participant
- The forms can be completed early if the participant dies or is discharged before day 7
   e.g. If participant dies at day 2 you still need to complete day 7 form and discharge/death form
- If <u>repatriated before day 7 please complete forms early</u> and then check if patient still alive at day 7 and enter data

<sup>\*\*</sup>Only trial team members signed off on the delegation log can enter data\*\*



### **Uploading Consent forms**



The coordinating centre will complete ongoing monitoring of the eCRF data and consent forms.

#### Consent forms and contact details

Sites in Ireland will not be uploading consent and contact details to the TICH-3 database.

The National Coordinator of Ireland will oversee these documents.



### Day 180 Form

To be conducted by the National Coordinator team

To be confirmed if this is by post, phone or face to face

#### Includes

- Modified Rankin scale
- EQ-5D
- AD-8 cognition score

We will need to delegate follow up coordinator role to someone with appropriate access to complete forms on line

#### TICH-3 trial

#### Tranexamic acid for IntraCerebral Haemorrhage 3

RCTN 97695350

Room 5/D2123, Stroke Trials Ui School of Medicine, University of Nottingha Queen's Medical Centre, Derby Ro Nottingham NG7 2UH, United Kingdo TICH-3 trial office <mszlh@nottingham.ac.uk

Day 180 follow-up form - POSTAL VERSION v1.0

When you were in hospital approximately 180 days ago either you, or someone on your behalf, gave permission for you to participate in a clinical trial called TICH-3. This would have happened when you first attended hospital following your stroke.

It is important that we now collect information on how well you have recovered from your stroke. The following questionnaire asks important questions and we would be grateful if you would complete it to the best of your ability. For each question, please choose the answer that applies to you and put a tick in the box next to it. If you are unsure which answer to choose, please tick the box that seems closest. Even if you feel that the questions do not apply to you please would you answer them, as it will help us to answer important questions about strokes.

Some questions deal with personal matters. Your name and address will **not** be stored on a database with your answers, and we will keep the information that you give us in absolute confidence.

If you would prefer to speak to us and answer the questions by telephone, then please call the number provided on the covering letter. We can call you straight back to pay for the call. Alternatively, please post this questionnaire in the provided self-addressed envelope.

It is important that answers are given for all questions, to ensure completeness of data collection.

Section A: Basic information				
A1 Date of completion (dd-mmm-yyyy)	D	/M	/Y	

## LOCAL SITE FILE



### **Local Site File Contents**

- Please see the TICH-3 website <a href="http://tich-3.ac.uk/docs/">http://tich-3.ac.uk/docs/</a> where you can download a contents page for the local investigator site file.
- The coordinating centre will not be sending local (investigator) hardcopy sites files in the post for reasons due to sustainability and version control.
- All documents will be available on the TICH-3
  website <a href="http://tich-3.ac.uk/docs/">http://tich-3.ac.uk/docs/</a> if the local site want to
  print their own local site file that is their choice and their
  responsibility to keep the hardcopy site file up to date (this
  applies to electronic as well).
- The coordinating centre will send any amendment notifications electronically with guidance of if any documents need superseding, we will then put the updated documentation on the TICH-3 website.
- Sites need to ensure that there is an AUDIT trail for monitoring purposes – and all up to date documents are available.
- Safety file SAE forms in site file after sign off by PI



TICH-3 trial - Tranexamic acid for IntraCerebral Haemorrhage 3

#### **Trial documents**



**Emergency contacts** 

This page does not provide the emergency mobile numbers

Please log in to view them, or bookmark the main documents page instead of this one

Approved protocol

Protocol Final v1.0 03.11.2021 fully signed.pdf

Expression of interest

Online expression of interest form

**Trial documents** 

- Contact List 08.03.22.pdf
- File Note v1.0 01.05.21.docx
- Poster for ED v1.0 05.01.22.pdf
- Site File Index v1.0 20.10.21.pd

UK site training

- Enrolling Investigator Training Final v1.0 17.03.2022.pdf (updated 3 days ago)
- Investigator Training Final v1.7 17.03.2022.pdf (updated 3 days ago)
- Pharmacy Training Final v1.0 02.02.2022.pdf

Information sheets and consent forms

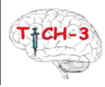
- Participant Full Consent Form -TICH-3 Final v1.0 03.11.2021.docx
- Participant Information Sheet TICH-3 Final v1.0 03.11.2021.docx
- Participant Short Information Sheet TICH-3 Final v1.0 03.11.2021.docx
- Professional (Legal Rep) Full Consent Form TICH-3 Final v1.0 03.11.2021.docx
- Professional (Legal Rep) Information Sheet TICH-3 Final v1.0 03.11.2021.doc
- Professional (Legal Rep) Short Information Sheet and Consent TICH-3 Final v1.0 03.11.2021.docx
- Relative (Legal Rep) Full Consent Form TICH-3 Final v1.0 03.11.2021.docx
- Relative (Legal Rep) Information Sheet TICH-3 Final v1.0 03.11.2021.docx
- Relative (Legal Rep) Short Information TICH-3 Final v1.0 03.11.2021.docx
- GP letter final v1.0 03.11.2021.docx

**Pharmacy documents** 

- Assessment and monitoring of remote IMP storage Final v1.0 20.12.2021.docx
- IMPD Final v2.0 09.03.2022.pdf
- IMP Accountability log Final v1.0 07.12.2021.docx
- IMP Check Final v1.0 20.12.2021.docx
- IMP Inventory Log Final v1.0 20.12.2021.docx
- Information for Pharmacy Final v1.0 20.12.2021.pdf
- Prescribing and administration guide Final v1.0 17.11.2021.pdf
- · Treatment packs specification.pdf



### **Electronic delegation Log**



Only people appropriately trained and delegated responsibility on the delegation log can take consent.

Anyone who is involved in the trial needs to be on the delegation log; nurses, admin entering data onto online platform, doctors, pharmacist handling the IMP. Can have as many people on the delegation log as required.

The training and roles delegated should be appropriate to the respective job role.

Completing training log will generate an email to the PI asking them to sign you on to the log

Requirements for the local team member to be able to go on the TICH-3 delegation log;

- Up to date investigator CV
- Evidence of GCP training
- Completion of TICH-3 training relevant to role in trial

It is the local PI's responsibility to check local teams investigator CV and GCP before they can be signed off on the delegation log

New members to the team need adding to the delegation log (meeting the requirements above) before they can start working on the TICH-3 trial.

If staff leave the team the PI is required to sign and date 'role finished' against their name.



For site initiation we require a minimum of the following team members signed off on the delegation log

- Principal Investigator
- Research Nurse/coordinator
- Pharmacist
- Please return the training log to us as soon as possible after training completed



### **Electronic Delegation Log**



#### TICH-3 delegation log for C001 Nottingham, Queen's Medical Centre

Chief investigator: Nikola Sprigg
Principal investigator: Kailash Krishnan

Thicipal investigator Randon Rhoman					
Log ID	Investigator name/ID	Certificate/ date trained	Roles and responsibilities*	Delegation log status	
1	Kailash Krishnan Consultant Physician (KKrishnan)	<u>G9L3P7</u> 2 Feb 2022	Principal investigator ABCDEFGHIJKLMNOPQRSTUVWXY	7 Mar 2022 08:23 <b>Authorised</b> <i>Kailash Krishnan</i>	
2	<b>Nikola Sprigg</b> <i>Professor of stroke medicine</i> (NSprigg)	<u>L9N9E7</u> 2 Feb 2022	Site investigator BFHIJKLNOPQRSYZ	7 Mar 2022 08:25 <b>Authorised</b> <i>Kailash Krishnan</i>	
3	Rachel Facilitator Researcher (RFacilitator)	<u>L3N3F7</u> 2 Feb 2022	<b>Site investigator</b> BFHIJKLNOPQRSTY	7 Mar 2022 08:25 <b>Authorised</b> <i>Kailash Krishnan</i>	
4	Clara Researcher Clinical Trials Researcher (CResearcher)	<u>K7H7C6</u> 4 Feb 2022	<b>Site investigator</b> BFHIJKLNOPQRSTY	7 Mar 2022 08:25 <b>Authorised</b> <i>Kailash Krishnan</i>	
5	Any Doctor  Researcher (ADoctor)	F3C9T7 2 Feb 2022	Site investigator BFHIJKLNOPQRSYZ	7 Mar 2022 08:25 <b>Authorised</b> <i>Kailash Krishnan</i>	
6	Zee Pharmacist Pharmacy DTO (ZPharmacist)	<u>Y7X6Y7</u> 2 Mar 2022	Pharmacist # FHLNPQSY	12 Mar 2022 08:49 <b>Authorised</b> <i>Kailash Krishnan</i>	

## SUMMARY



### **TICH-3 Key Points**

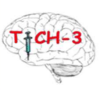


- Pragmatic design and methods
- Inclusion criteria ICH < 4.5 hours,</li>
   Exclusion massive ICH (low GCS < 5, HV < 60mls),</li>
   contraindication to tranexamic acid (e.g. seizures)
- Emergency consent initially oral followed by written consent
- Simple randomisation use the lowest available treatment pack number
- QR code randomisation alert inform trial office of enrolment
- Safety monitoring safety events for 7 days, SAR and SUSAR Venous and arterial occlusive events and seizures
- Central postal/telephone follow up at 6 months





### Site requirements before start up





Attendance at training for minimum PI, one research nurse/coordinator and one pharmacist

Paperwork - send documents to coordinating centre via email.

- > Signed and dated recent investigator CV and GCP certificate of the local Principal Investigator
- > Attendance investigator training log
- > Assessment and monitoring of remote IMP storage form
- > Fully executed non-commercial agreement and confirmation of local capacity and capability

#### Electronic delegation log

Local PI to authorise all local team members onto the online delegation log via the TICH-3 website.

#### **Drug dispatch**

Drug will be dispatched when the delegation log has been countersigned by the PI, drug needs to be marked as received and then marked as available for randomisation on the TICH-3 website.

#### Green light

We will advise once everything has been checked and confirm that the site can open to TICH-3 by sending the regulatory green light email from the University of Nottingham sponsor.

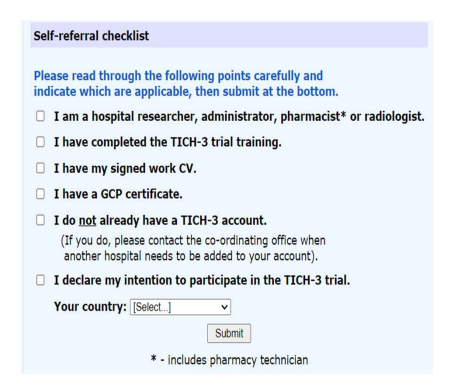
> All documents are required to be stored in your local investigator site file (electronic or hardcopy)

\*\*Investigators may only work on the trial once signed off on the delegation log and the site may only begin enrolling participants in the trial after green light has been received from the sponsor\*\* 59



### What happens next?

- CTIS approval Part I regulatory approval received 20/4/2023
- Part II Approved 26/04/2023
- Finalise contracts with each site
- Investigators will be added to the delegation log after completing the training log
- Please complete the investigator training log and return via email to the coordinating centre
- Or use the self referral form: <a href="http://tich-3.ac.uk/?ZSelfRef">http://tich-3.ac.uk/?ZSelfRef</a>
- IMP will be sent to designated sites
- DPIA to be completed for sites where required





### **ACTION – Return Training Log**



 Please complete the investigator training log and return via email to the coordinating centre <u>Click here for direct download of training log</u> use the self referral form:

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

- Team members who could not attend live training can access training slides from TICH-3 website <a href="https://stroke.nottingham.ac.uk/tich-3/docs/#UK">https://stroke.nottingham.ac.uk/tich-3/docs/#UK</a> site training
- There are 3 versions of the training slides
  - Investigator training which gives a detailed description of the whole trial process, intended for the PI and research nurses/coordinators.
  - Enrolling investigator training this streamlined training is intended for team members who will only be taking enrolment consent i.e. consultants
  - Pharmacy training this streamlined training is intended for members of pharmacy team
- A short 3 ½ minute video is available to introduce team members to the TICH-3 trial <a href="http://tich-3.ac.uk/docs/#Videos">http://tich-3.ac.uk/docs/#Videos</a>





PLEASE SCAN ME

## **CONTACT INFORMATION**



### **University of Nottingham Trial Team**



Name	Role	Contact Information
Chaamanti Menon	Clinical Research Fellow	E: chaamanti.menon@nottingham.ac.uk
<mark>TBC</mark>	Clinical Trials Manager (International Site Recruitment)	<mark>E:</mark>
Tiffany Hamilton	Senior Clinical Trials Manager	E: Tiffany.hamilton@nottingham.ac.uk
Kennedy Cadman	TICH 3 Administrator	E: kennedy.cadman@nottingham.ac.uk
Nikola Sprigg	Chief Investigator	E: nikola.sprigg@Nottingham.ac.uk

#### **Trial Coordinating Centre contact information:**



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