A study to define the platelet count below which critically ill patients should receive a platelet transfusion before an invasive procedure

Submission date 01/09/2022	Recruitment status Recruiting	[X] Prospectively registered[X] Protocol
Registration date	Overall study status	Statistical analysis plan
30/09/2022	Ongoing	Results
Last Edited 10/06/2025	Condition category Other	☐ Individual participant data
		[X] Record updated in last year

Plain English summary of protocol

Background and study aims

Platelets are cells in the blood that help form clots and stop bleeding. People treated in a critical care unit often have a low number of platelets (platelet count) in their blood because they are very unwell. Platelet transfusions are made up of platelets collected from screened, healthy donors. Platelet transfusions are sometimes given before these procedures if the patient's platelet count is low. This is thought to reduce the possible risk of bleeding from the procedure. However, platelet transfusions also carry risks such as inflammation, infection, and allergic reactions, and may not work as effectively in unwell patients.

Currently, we do not know the platelet count below which giving a platelet transfusion might be beneficial. Surveys of doctors working in UK critical care units have shown uncertainty over the platelet count below which doctors should give a platelet transfusion. As a result, platelet transfusions are currently given to patients with a wide range of different platelet counts and there is no set threshold.

This study will test five different thresholds to find out the safest count below which platelet transfusions should be given before invasive procedures are carried out in intensive care.

Who can participate?

Patients aged 18 years and over who have accepted for admission or admitted to critical care, with a platelet count of less than $50 \times 10e9/L$ who are being considered for a platelet transfusion for a low bleeding risk invasive procedure

What does the study involve?

Patients will be allocated to one of five platelet count thresholds (less than 10, 20, 30, 40 or 50). If their platelet count is below their allocated threshold, then they will receive a platelet transfusion before a low bleeding risk invasive procedure. Patients will remain in their allocated 'group' (threshold) for the duration of their critical care unit stay.

Some information about the patients' hospital stay is collected from hospital medical records. Other important health information is collected from national health databases. Some patients

will also be sent a short health questionnaire around 90 days and 1 year after becoming involved in the study. At the end of the study, all this information will allow us to compare the different transfusion thresholds in the study to find out which is most beneficial.

What are the possible benefits and risks of participating?

The benefit of receiving a platelet transfusion is to possibly reduce the risk of bleeding during an invasive procedure. The possible risks of receiving a platelet transfusion include inflammation, infection and allergic reactions. The purpose of this study is to look at the best platelet count threshold at which the possible benefits of platelet transfusion outweigh the possible risks, as this is currently unclear.

Where is the study run from? University of Oxford (UK)

When is the study starting and how long is it expected to run for? January 2022 to December 2027

Who is funding the study? National Institute for Health Research (NIHR) – Health Technology Assessment Programme (UK)

Who is the main contact? Hayley Noble, T4P@icnarc.org

Contact information

Type(s)

Scientific

Contact name

Ms Hayley Noble

Contact details

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Type(s)

Scientific

Contact name

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Additional identifiers

Clinical Trials Information System (CTIS)

Nil known

Integrated Research Application System (IRAS)

312405

ClinicalTrials.gov (NCT)

Nil known

Protocol serial number

CPMS 53274, IRAS 312405

Study information

Scientific Title

The Threshold for Platelets (T4P) study: a prospective randomised trial to define the platelet count below which critically ill patients should receive a platelet transfusion prior to an invasive procedure

Acronym

T4P

Study objectives

That platelet transfusion in critically ill patients has net clinical and monetary benefit only below certain thresholds where any gain of preventing bleeding exceeds harm from exacerbating inflammatory and/or infective processes.

Ethics approval required

Ethics approval required

Ethics approval(s)

1. approved 08/07/2022, South Central – Oxford C Research Ethics Committee (Health Research Authority, Ground Floor, Temple Quay House, 2 The Square, Bristol, BS1 6PN, United Kingdom; +44 (0)207 104 8226; oxfordc.rec@hra.nhs.uk), ref: 22/SC/0186

2. approved 19/10/2023, Scotland A Research Ethics Committee (2nd Floor, Waverley Gate 2, 4 Waterloo Place, Edinburgh, EH13EG, United Kingdom; +44 (0)131 465 5680; Manx. Neill@nhslothian.scot.nhs.uk), ref: 23/SS/0082

3. approved 14/05/2024, Nepean Blue Mountains Local Health District HREC (Level 5, Block D (South Block), Nepean Hospital, Penrith, 2751, Australia; +61 (02) 4734 3441; NBMLHD-Ethics@health.nsw.gov.au), ref: 2024/ETH00464

Study design

Randomized interventional study

Primary study design

Interventional

Study type(s)

Treatment

Health condition(s) or problem(s) studied

Critical care

Interventions

Current interventions as of 10/06/2025:

T4P is a large-scale, multi-centre, data-enabled, registry-embedded, open-label, randomised, comparative effectiveness trial with an internal pilot across five equally spaced platelet count thresholds ($<10 - <50 \times 10e9/L$). There will be an integrated economic evaluation.

The trial plans to include 2550 critically ill patients recruited from 66 NHS adult critical care units over a period of 42 months.

The normal platelet count is 150-450 x 10e9/L. Patients whose platelet count is below 50 x 10e9 /L (at any time in their critical care unit stay) and requiring a low bleeding risk invasive procedure will be considered for the trial. Once a patient has been confirmed as eligible (i.e. they satisfy the inclusion and exclusion criteria), they will be randomised (see below) and the randomly allocated treatment commenced as soon as possible.

Prior to an invasive procedure, eligible patients will be randomised to one of five platelet thresholds below which they will receive a single adult equivalent dose (AED) of platelet transfusion for the index procedure and subsequent procedures during their critical care unit stay. The thresholds are:-

- 1. Platelet count <50 x 10e9/L
- 2. Platelet count<40 x 10e9/L
- 3. Platelet count <30 x 10e9/L
- 4. Platelet count <20 x 10e9/L
- 5. Platelet count <10 x 10e9/L

Patients will be given a platelet transfusion prior to the invasive procedure if their platelet count is below the threshold to which they have been allocated. Patients remain in their allocated 'group' (threshold) for the duration of their critical care unit stay.

In all groups, all other treatments and procedures will be carried out in accordance with standard NHS care and local practice.

CONSENT

As eligible patients will be critically ill at the point in which they become eligible for T4P – a model of research without prior consent (RWPC) (also known as 'deferred consent') is proposed. This model is believed to be the most appropriate as low bleeding risk interventional procedures are often initiated as a life-saving measure, during an emergency clinical situation. Patients will lack mental capacity due to their medical condition and by virtue of serious illness that required admission to a critical care unit (or continuing treatment in critical care) at the point that they become eligible for the trial. Any delay in commencing the trial treatment could be detrimental

to the patient, as well as to the scientific validity of the trial.

In brief, once a patient is screened as eligible for the trial (i.e. satisfies inclusion and exclusion criteria), they will be enrolled and randomised to receive the assigned treatment immediately. Patients in critical care units are monitored very closely and clinical/research staff working in this setting have extensive experience of assessing capacity in their patients. For patients recruited in England, Wales and Northern Ireland, once a patient has regained capacity, they will be approached by an authorised member of the site research team for informed deferred consent. This will be done as soon as practically possible (usually within 24 - 48 hours of the patient regaining capacity). In the interim period - once the patient's medical situation is deemed to no longer be an emergency, a Personal Consultee will be approached (in person or by telephone) to provide their opinion of the patient's wishes regarding participating in the trial. Telephone and postal mechanisms for consent is also in place for the situation where patients are discharged from hospital prior to confirming their consent decision.

This type of consent model is used in clinical trials comparing treatments in emergency clinical situations (such as this one) to find out which is best. The specific model proposed for T4P has been informed/approved by our Patient and Public Involvement (PPI) co-investigator.

For patients recruited in Scotland, consent must be in place prior to randomisation. This can be sought from the patient, or if they lack capacity, from a Personal Legal Representative. If consent sought from a Personal Legal Representative prior to randomisation, consent will then be sought after randomisation from the patient when they regain capacity. This consent model in Scotland has been reviewed and approved by Scotland A REC and is in accordance with the Adults with Incapacity (Scotland) Act 2000.

For patients recruited in the Republic of Ireland, once a patient has regained capacity, they will be approached for consent to continue. In the interim, a substitute decision maker (e.g., family, friend) will be approached for deferred assent.

For patients recruited in Australia, a waiver of consent for enrolment has been granted. Dependent on local jurisdictional requirements and legislation, consent for follow up at 90 days and 12 months will employ either an opt-out or a consent to continue approach. For participating sites using the opt-out approach, the patient, or if they lack capacity and in the interim, person responsible, will be provided with a brochure which will explain the trial and the procedure to decline or opt-out from follow-up. For participating sites using the consent to continue approach, the patient, or if they lack capacity and in the interim, person responsible, will be provided with an information sheet and the opportunity to provide consent.

At 90 days and 1 year, participants will be posted questionnaires about health-related quality of life and their use of health services since leaving hospital. These questionnaires have been used in previous critical care unit trials and will provide valuable information for the integrated economic evaluation. The questionnaires are designed to take no longer than 15 minutes to complete. A stamped addressed envelope and a pen will be included, so it will not cost the patient anything. A trained member of the T4P team at the ICNARC CTU will telephone participants who have not returned the questionnaire after three weeks, to check if they have received it and offer the option of resending the questionnaire (either by post or email) or going through the questionnaire over the telephone. Patient follow-up questionnaires will be administered by the participating site teams in the Republic of Ireland and Australia.

INTERNAL PILOT

The pilot phase will cover the first 12 months of recruitment, assessing recruitment, willingness to randomise, protocol adherence and data quality. Data will be analysed at the end of the internal pilot trial stage. The analysis will take place in month 20 of the trial to allow data to be

collected and entered to assess all progression criteria. The outcome of this analysis will be presented to the majority-independent Trial Steering Committee who will provide their recommendation as to whether the trial should continue to the Funder (National Institute for Health Research (NIHR), Health Technology Assessment (HTA) Programme). The final decision on progression from the pilot stage to the full trial will be made by the NIHR HTA programme after recommendation by the TSC.

INDEPENDENT COMMITTEES

Both a Trial Steering Committee and an independent Data Monitoring & Ethics Committee (DMEC) will be convened and will meet regularly during the trial. The DMEC will monitor recruitment and retention, protocol adherence (including adherence to treatment protocols) and patient safety (including serious adverse events), and will review the interim analysis.

TIMELINE

Months 1-6: Study set-up: all approvals & preparation for the start of the trial (site sign-up and local approvals, production of materials for participating sites, conduct site initiation meetings)

Months 7-48: Recruitment/follow-up period

Months 7-18: Internal pilot stage

Month 16: First annual REC report

Month 19: First follow-up questionnaires sent

Month 20: Second DMEC and TSC meetings to review internal pilot analysis Internal pilot report submitted to NIHR HTA

Month 48: Close to recruitment

Month 52: Final follow-up questionnaires

Months 48-60: Analysis and dissemination

Month 53: Database lock for primary analysis (clinical and economic evaluation) Commence primary analysis and write up

Month 55: Lock database for longer-term outcomes

Month 59: Submit primary outcome paper Collaborators' meeting Final DMEC and TSC meetings

Month 60: Submit longer-term outcomes paper and draft final report to NIHR

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Intervention Type

Other

Primary outcome(s)

All-cause mortality at 90 days measured through review of patient medical notes at 90 days post-randomisation and/or data linkage with nationally held death registrations.

Primary health economic outcome measure:

Incremental costs, quality-adjusted life year (QALYs) and net monetary benefit at 90 days, measured through combining Health-related Quality of Life (EuroQol EQ-5D-5L questionnaire) data, valued resource use data obtained via a health services questionnaire and data obtained through linkage with national hospital episode statistics, death registrations and the national clinical audit for adult critical care.

Key secondary outcome(s))

- 1. Mortality at discharge from critical care unit, hospital and at 1 year, measured through review of patient medical notes at the relevant timepoints and/or data linkage with nationally held death registrations and the national clinical audit for adult critical care (for mortality at discharge)
- 2. Survival to longest available follow-up, measured by review of patient medical notes and/or data linkage with nationally held death registrations
- 3. Rates of major and fatal bleeds classified according to the HEmorhage Measurement (HEME) bleeding score, measured through review of patient medical notes up until critical care unit discharge
- 4. Venous and arterial thromboses in hospital and to 1 year, measured through review of patient medical notes at hospital discharge, data obtained via a health services questionnaire and through data linkage with national hospital episodes statistics and the NICE-mandated hospital-acquired venous thromboembolism (VTE) audit
- 5. Duration of renal, advanced cardiovascular and advanced respiratory support according to UK Critical Care Minimum Data Set (CCMDS) criteria, measured through review of patient medical

notes during critical care admission and data obtained through linkage with the national clinical audit for adult critical care

- 6. Length of critical care unit and acute hospital stay, measured through review of patient medical notes and data obtained through linkage with the national clinical audit for adult critical care
- 7. Health-related quality of life measured through EQ-5D-5L questionnaire at 90 days and 1-year timepoints
- 8. Resource use and costs at 90 days and 1 year, measured by valuing resource use data obtained via a health services questionnaire administered to patients and through data linkage with national hospital episode statistics and the national clinical audit for adult critical care 9. Net monetary benefit (NMB) at 1 year, measured through combining health-related quality of life (EQ-5D-5L questionnaire) data, valued resource use data obtained via a health services questionnaire and data obtained through linkage with national hospital episode statistics, death

Completion date

31/12/2027

Eligibility

Key inclusion criteria

Current participant inclusion criteria as of 17/03/2023:

registrations and the national clinical audit for adult critical care

- 1. Adult (aged 18 years or older)
- 2. Accepted for admission or admitted to a participating critical care unit
- 3. Platelet count $<50 \times 10e9/l$
- 4. Planned to undergo a specified* low bleeding risk invasive procedure OR platelet transfusion being considered for an 'other' procedure
- *Specified low bleeding risk invasive procedures include the following:
- 1. Central venous vascular catheter insertion (including vascular access for renal replacement therapy)
- 2. Paracentesis/superficial abdominal fluid collection drainage
- 3. Pleural aspiration

'Other' procedures may be included if the clinician deems these to be a low bleeding risk invasive procedure and a platelet transfusion is being considered for the procedure. These include, but are not limited to, the following:

- 1. Arterial catheter insertion
- 2. Arterial or central venous catheter removal
- 3. Pleural drain
- 4. Interventional radiology (as defined by Society of Interventional Radiology guidelines)
- 5. Bronchoscopy with or without lavage
- 6. Wound dressing changes
- 7. Surgical procedures where the clinical team agree the risk of bleeding is low, e.g. re-look laparotomy, or wound closure

Previous participant inclusion criteria:

- 1. Adult (aged 18 years or older)
- 2. Accepted for admission or admitted to a participating critical care unit
- 3. Platelet count <50 x 10e9/l
- 4. Platelet transfusion being considered for a low bleeding risk invasive procedure*

- *Low bleeding risk invasive procedures include the following:
- 1. Vascular catheter insertion and removal (central venous including vascular access for renal replacement therapy)
- 2. Paracentesis/superficial abdominal fluid collection drainage
- 3. Pleural aspiration

'Other' procedures may be included if the clinician deems these to be a low bleeding risk invasive procedure. These include, but are not limited to, the following:

- 1. Arterial catheter line insertion
- 2. Pleural drain
- 3. Interventional radiology (as defined by Society of Interventional Radiology guidelines)
- 4. Bronchoscopy with or without lavage
- 5. Wound dressing changes
- 6. Surgical procedures where the clinical team agree risk of bleeding is low, e.g. re-look laparotomy, or wound closure

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Adult

Lower age limit

18 years

Sex

All

Key exclusion criteria

Current participant exclusion criteria as of 17/03/2023:

- 1. Ongoing major haemorrhage requiring blood products and/or surgical/radiological intervention*
- 2. Intercranial haemorrhage within prior 72 hours*
- 3. Contra-indication to platelet transfusion (such as thrombotic microangiopathies; heparin-induced thrombocytopaenia; immune thrombocytopaenia; congenital platelet function defects)
- 4. Acute promyelocytic leukaemia (APML)
- 5. Known advance decision refusing blood/blood component transfusions (e.g. Jehovah's Witnesses)
- 6. Death perceived as imminent or admission for palliation
- 7. Previously randomised into T4P
- 8. Fulfilled all the inclusion criteria and none of the other exclusion criteria >72 hours

*Exclusion criteria no. 1 and 2 are dynamic, and if resolved, the patient may be reconsidered for the trial

Previous participant exclusion criteria:

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- *Exclusion criteria no. 1 and 2 are dynamic, and if resolved, the patient may be reconsidered for the trial

Date of first enrolment 19/10/2022

Date of final enrolment 31/12/2026

Locations

Countries of recruitmentUnited Kingdom

England

Northern Ireland

Scotland

Wales

Australia

Ireland

Study participating centre
Barnet Hospital
Wellhouse Lane
Barnet
United Kingdom

EN5 3DJ

Study participating centre Victoria Hospital (blackpool) Whinney Heys Road

Blackpool
United Kingdom
FY3 8NR

Study participating centre Chelsea & Westminster Hospital

369 Fulham Road London United Kingdom SW10 9NH

Study participating centre Chesterfield Royal Hospital

Chesterfield Road Calow Chesterfield United Kingdom S44 5BL

Study participating centre Countess of Chester Hospital

Countess of Chester Health Park Liverpool Road Chester United Kingdom CH2 1UL

Study participating centre Croydon University Hospital

London Road Croydon United Kingdom CR7 7YE

Study participating centre

Doncaster and Bassetlaw Teaching Hospitals NHS Foundation Trust

Doncaster Royal Infirmary Armthorpe Road Doncaster United Kingdom DN2 5LT

Great Western Hospital

Marlborough Road Swindon United Kingdom SN3 6BB

Study participating centre Guy's and St Thomas' Hospitals

Trust Offices
Guy's Hospital
Great Maze Pond
London
United Kingdom
SE1 9RT

Study participating centre Heartlands Hospital

Bordesley Green East Bordesley Green Birmingham United Kingdom B9 5ST

Study participating centre Good Hope Hospital

Rectory Road Sutton Coldfield United Kingdom B75 7RR

Study participating centre Hull Royal Infirmary

Anlaby Road Hull United Kingdom HU3 2JZ

Study participating centre John Radcliffe Hospital

Headley Way Headington Oxford United Kingdom OX3 9DU

Study participating centre Kettering General Hospital

Rothwell Road Kettering United Kingdom NN16 8UZ

Study participating centre Kings College Hospital

Mapother House De Crespigny Park Denmark Hill London United Kingdom SE5 8AB

Study participating centre Kings Mill Hospital

Mansfield Road Sutton-in-ashfield United Kingdom NG17 4JL

Study participating centre Leicester Royal Infirmary

Infirmary Square Leicester United Kingdom LE1 5WW

Study participating centre Liverpool Heart & Chest Hospital

Broadgreen Hospital Thomas Drive Liverpool United Kingdom L14 3PE

Study participating centre Milton Keynes University Hospital

Standing Way Eaglestone Milton Keynes United Kingdom MK6 5LD

Study participating centre Northumbria Specialist Emergency Care Hospital

Northumbria Way Cramlington United Kingdom NE23 6NZ

Study participating centre Pilgrim Hospital

Sibsey Road Boston United Kingdom PE21 9QS

Study participating centre Poole Hospital

Longfleet Road Poole United Kingdom BH15 2JB

Study participating centre University Hospital Birmingham

Queen Elizabeth Hospital Edgbaston Birmingham United Kingdom B15 2TH

Queen Elizabeth Hospital

Woolwich Stadium Road Woolwich London United Kingdom SE18 4QH

Study participating centre Burton Hospital

Queens Hospital Belvedere Road Burton-on-trent United Kingdom DE13 0RB

Study participating centre Queens Medical Centre

Derby Road Nottingham United Kingdom NG7 2UH

Study participating centre Nottingham City Hospital

Hucknall Road Nottingham United Kingdom NG5 1PB

Study participating centre Royal Berkshire Hospital

Royal Berkshire Hospital London Road Reading United Kingdom RG1 5AN

Study participating centre Royal Hampshire County Hospital

Romsey Road Winchester United Kingdom SO22 5DG

Study participating centre Royal Liverpool University Hospital

Prescot Street Liverpool United Kingdom L7 8XP

Study participating centre Royal Papworth Hospital

Papworth Road
Cambridge Biomedical Campus
Cambridge
United Kingdom
CB2 0AY

Study participating centre New Cross Hospital Royal Wolverhampton

Wolverhampton Road Heath Town Wolverhampton United Kingdom WV10 0QP

Study participating centre Russells Hall Hospital

Pensnett Road Dudley United Kingdom DY1 2HQ

Study participating centre St Georges at Mayday University Hospital

530 London Road Thornton Heath United Kingdom CR7 7YE

Study participating centre St Richards Hospital

Spitalfield Lane Chichester United Kingdom PO19 6SE

Study participating centre Tameside General Hospital

Fountain Street Ashton-under-lyne United Kingdom OL6 9RW

Study participating centre Treliske Hospital

Treliske Truro United Kingdom TR1 3LJ

Study participating centre University Hospital Coventry

Clifford Bridge Road Coventry United Kingdom CV2 2DX

Study participating centre Warrington Hospital (site)

Warrington Hospital Lovely Lane Warrington United Kingdom WA5 1QG

Study participating centre West Middlesex University Hospital

Twickenham Road Isleworth United Kingdom TW7 6AF

Study participating centre Wexham Park Hospital

Wexham Street Wexham Slough United Kingdom SL2 4HL

Study participating centre The Whittington Hospital

Highgate Hill London United Kingdom N19 5NF

Study participating centre Aberdeen Royal Infirmary

Foresterhill Road Aberdeen United Kingdom AB25 2ZN

Study participating centre Western General Hospital

Crewe Road South Edinburgh Lothian United Kingdom EH4 2XU

Study participating centre University Hospital of North Durham

University Hospital of Durham Dryburn Hospital North Road Durham United Kingdom DH1 5TW

Study participating centre Darlington Memorial Hospital NHS Trust

Darlington Memorial Hospital Hollyhurst Road Darlington United Kingdom DL3 6HX

Study participating centre West Cumbria Health Care NHS Trust

West Cumberland Hospital Hensingham Whitehaven United Kingdom CA28 8JG

Study participating centre Salford Royal Hospital

Stott Lane Eccles Salford United Kingdom M6 8HD

Study participating centre Fairfield General Hospital

Fairfield General Hospital Rochdale Old Road Bury United Kingdom BL9 7TD

Study participating centre Medway NHS Foundation Trust

Medway Maritime Hospital Windmill Road Gillingham United Kingdom ME7 5NY

Study participating centre Newham University Hospital NHS Trust

Newham General Hospital Glen Road London United Kingdom E13 8SL

Study participating centre St Vincent's University Hospital

Elm Park Dublin 4 Dublin Ireland D04 T6F4

Study participating centre Royal Brisbane and Women's Hospital

Brisbane Australia 4029

Study participating centre Cairns Hospital

Cairns Australia 4870

Study participating centre Gold Coast University Hospital

Gold Coast Australia 4215

Mater Misericordiae Ltd

Newstead Australia 4006

Study participating centre Toowoomba Hospital

Toowoomba Australia 4350

Study participating centre Fiona Stanley Hospital

Perth Australia 6150

Study participating centre Royal Perth Hospital

Perth Australia 6000

Study participating centre Sir Charles Gairdner Hospital

Perth United Kingdom 6009

Study participating centre Bunbury Regional Hospital

Bunbury Australia 6230

The Canberra Hospital

Canberra Australia 2605

Study participating centre Royal North Shore Hospital

Sydney Australia 2065

Study participating centre Campbelltown Hospital

Sydney Australia 2560

Study participating centre Austin Hospital

Melbourne Australia 3084

Study participating centre Royal Melbourne Hospital

Melbourne Australia 3050

Study participating centre St Vincent's Hospital

Melbourne Australia 3065

Ballarat Hospital

Ballarat Australia 3350

Study participating centre Footscray Hospital

Footscray Australia 3011

Study participating centre Sunshine Hospital

St Albans Australia 3021

Study participating centre Monash Medical Centre

Melbourne Australia 3021

Study participating centre Victorian Heart Hospital

Clayton Australia 3168

Study participating centre The Alfred

Melbourne Australia 3004

St Vincent's Hospital

Sydney Australia 2010

Study participating centre Blacktown Hospital Blacktown

Australia 2148

Sponsor information

Organisation

University of Oxford

ROR

https://ror.org/052gg0110

Funder(s)

Funder type

Government

Funder Name

NIHR Evaluation, Trials and Studies Co-ordinating Centre (NETSCC); Grant Codes: NIHR131822

Results and Publications

Individual participant data (IPD) sharing plan

The datasets generated during and/or analysed during the current study are/will be available upon request from T4P@icnarc.org

IPD sharing plan summary

Data sharing statement to be made available at a later date

Details

Study outputs

Output type HRA research summary Date created Date added Peer reviewed? Patient-facing?

28/06/2023 No

No

Participant information sheet		11/11/2025	11/11/2025 No	Yes
Protocol file	version 2.0	25/11/2022	22/05/2023 No	No
Protocol file	version 3.1	16/01/2024	07/06/2024 No	No
<u>Protocol file</u>	Australian sites version 1.0	15/04/2024	10/06/2025 No	No
Study website	Study website	11/11/2025	11/11/2025 No	Yes