

REQUEST FOR AUTHORISATION OF A CLINICAL TRIAL ON A MEDICINAL PRODUCT FOR HUMAN USE TO THE COMPETENT AUTHORITIES AND FOR OPINION OF THE ETHICS COMMITTEES IN THE COMMUNITY
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To be filled in by the applicant

The questions in this form for the request for authorisation from the Competent Authority are also relevant for the opinion from an Ethics Committee (it represents module 1 of the form for applying to an ethics committee) and can be used as part of that application. Please indicate the relevant purpose in a box below.

REQUEST FOR AUTHORISATION TO THE COMPETENT AUTHORITY: Yes ●
REQUEST FOR OPINION OF THE ETHICS COMMITTEE: No ●

A. TRIAL IDENTIFICATION

A.1	Member State in which the submission is being made:	UK - MHRA
A.2	EudraCT number:	2007-006749-42
A.3	Full title of the trial: English	Safety and efficacy of clopidogrel when added to aspirin and dipyridamole in high risk patients with recent ischaemic stroke: a randomised controlled trial
A.3.1	Title of the trial for lay people, in easily understood, i.e. non-technical, language: English	Is intensive blood thinning with three antiplatelet drugs better than the current guideline treatment in reducing recurrence after acute stroke?
A.3.2	Name or abbreviated title of the trial where available: English	TARDIS
A.4	Sponsor's protocol code number, version and date ¹ :	
A.4.1	Sponsor's protocol code number:	31350
A.4.2	Sponsor's protocol version:	1.0
A.4.3	Sponsor's protocol date:	2008-07-24
A.5	Additional international study identifiers (e.g. WHO, ISRCTN ² , US NCT Number ³) if available	
A.5.1	ISRCTN number:	ISRCTN47823388
A.5.2	US NCT number:	
A.5.3	WHO Universal Trial Reference Number (UTRN):	
A.5.4	Other Identifier:	
A.6	Is this a resubmission? If 'Yes', indicate the resubmission letter ⁴ :	No ● First Submission
A.7	Is the trial part of a Paediatric Investigation Plan?	No ●
A.8	EMA Decision number of Paediatric Investigation Plan:	

B. IDENTIFICATION OF THE SPONSOR RESPONSIBLE FOR THE REQUEST

B.1	SPONSOR	
B.1.1	Name of organisation:	University of Nottingham
B.1.2	Name of the person to contact:	
B.1.2.1	Given name	Paul
B.1.2.2	Middle name	
B.1.2.3	Family name	Cartledge
B.1.3	Address:	
B.1.3.1	Street address	Kings Meadow Campus, Lenton Lane
B.1.3.2	Town/city	Nottingham
B.1.3.3	Post code	NG7 2NR
B.1.3.4	Country	United Kingdom
B.1.4	Telephone number:	+44 0115 9515679
B.1.5	Fax number:	+44 0115 9513633
B.1.6	E-mail:	paul.cartledge@nottingham.ac.uk

B.2	LEGAL REPRESENTATIVE⁵ OF THE SPONSOR IN THE COMMUNITY FOR THE PURPOSE OF THIS TRIAL (if different from the sponsor)	
B.2.1	Name of organisation:	
B.2.2	Name of person to contact:	
B.2.2.1	Given name	
B.2.2.2	Middle name	
B.2.2.3	Family name	
B.2.3	Address:	
B.2.3.1	Street address	
B.2.3.2	Town/city	
B.2.3.3	Post code	
B.2.3.4	Country	
B.2.4	Telephone number:	
B.2.5	Fax number:	
B.2.6	E-mail:	

B.3	STATUS OF THE SPONSOR:	
B.3.1	Commercial:	No •
B.3.2	Non commercial:	Yes •

B.4	Source(s) of Monetary or Material Support for the clinical trial (repeat as necessary):	
B.4.1	Name of organisation:	British Heart Foundation
B.4.2	Country:	United Kingdom

B.5	Contact point⁶ designated by the sponsor for further information on the trial	
B.5.1	Name of organisation:	University of Nottingham
B.5.2	Functional name of contact point (e.g. "Clinical Trial Information Desk"):	TARDIS Trial Office
B.5.3	Address:	
B.5.3.1	Street address	Div Stroke, Cl.Sci.Bldg, Hucknall Rd
B.5.3.2	Town/city	Nottingham
B.5.3.3	Post code	NG5 1PB
B.5.3.4	Country	United Kingdom
B.5.4	Telephone number:	+44 0115 8230210
B.5.5	Fax number:	+44 0115 8230273
B.5.6	E-mail: (use a functional e-mail address rather than a personal one)	tardis@nottingham.ac.uk

C. APPLICANT IDENTIFICATION, (please tick the appropriate box)

C.1 REQUEST FOR THE COMPETENT AUTHORITY	
C.1.1	Sponsor
C.1.2	Legal representative of the sponsor
C.1.3	Person or organisation authorised by the sponsor to make the application Yes •
C.1.4	Complete the details of the applicant below even if they are provided elsewhere on the form:
C.1.4.1	Name of Organisation: University of Nottingham
C.1.4.2	Name of contact person:
C.1.4.2.1	Given name Philip
C.1.4.2.2	Middle name
C.1.4.2.3	Family name Bath
C.1.4.3	Address:
C.1.4.3.1	Street address Clinical Sciences Building, City Hospital, Hucknall Road
C.1.4.3.2	Town/city Nottingham
C.1.4.3.3	Post code NG5 1PB
C.1.4.3.4	Country United Kingdom
C.1.4.4	Telephone number: +44 0115 8231765
C.1.4.5	Fax number: +44 0115 8237167
C.1.4.6	E-mail: philip.bath@nottingham.ac.uk
C.1.5	Request to receive a copy of CTA data as XML:
C.1.5.1	Do you want a copy of the CTA form data saved on EudraCT as an XML file? No •
C.1.5.1.1	If Yes provide the e-mail address(es) to which it should be sent (up to 5 addresses):
C.1.5.1.2	Do you want to receive this via password protected link(s)? No •
If you answer No to question C.1.5.1.2 the .xml file will be transmitted by less secure e-mail link(s)	

D. INFORMATION ON EACH IMP

Information on each 'bulk product' before trial-specific operations (blinding, trial specific packaging and labelling) should be provided in this section for each investigational medicinal product (IMP) being tested including each comparator and each placebo, if applicable. **For placebo go directly to D8.** If the trial is performed with several products use extra pages and give each product a sequential number in D1.1. If the product is a combination product, information should be given for each active substance.

D.1 IMP IDENTIFICATION		
Indicate which of the following is described below, then repeat as necessary for each of the numbered IMPs to be used in the trial (assign numbers from 1-n):		
D.1.1	This refers to the IMP number:	PR1
D.1.2	IMP being tested	Yes •
D.1.3	IMP used as a comparator	No •
D.2 STATUS OF THE IMP		
D.2.1	Has the IMP to be used in the trial a marketing authorisation? Yes • If the IMP has a marketing authorisation in the Member State concerned by this application, but the trade name and marketing authorisation holder are not fixed in the protocol, go to section D.2.2.	
D.2.1.1	If 'Yes', specify the product to be used in the clinical trial:	
D.2.1.1.1	Trade name	
D.2.1.1.1.1	EV Product Code (where applicable)	
D.2.1.1.2	Name of the Marketing Authorisation Holder:	
D.2.1.1.3	Marketing Authorisation number (if Marketing Authorisation granted by a Member State):	
D.2.1.1.4	Is the IMP modified in relation to its Marketing Authorisation? Not Answered •	
D.2.1.1.4.1	If 'Yes', please specify:	
D.2.1.2	The country that granted the Marketing Authorisation	
D.2.1.2.1	Is this the Member State concerned with this application? Not Answered •	
D.2.2	Situations where an IMP to be used in the CT has a Marketing Authorisation in the Member State concerned, but the protocol allows that any brand of the IMP with a Marketing Authorisation in that Member State be administered to the trial subjects and it is not possible to clearly identify the IMP(s) in advance of the trial start	
D.2.2.1	In the protocol, is treatment defined only by active substance? Yes •	
D.2.2.1.1	If 'Yes', give active substance in D.3.8 or D.3.9	
D.2.2.2	In the protocol, do treatment regimens allow different combinations of marketed products used according to local clinical practice at some or all investigator sites in the MS? Yes •	
D.2.2.2.1	If 'Yes', give active substance in D.3.8 or D.3.9	
D.2.2.3	The products to be administered as IMPs are defined as belonging to an ATC group ⁹ No •	
D.2.2.3.1	If 'Yes', give the ATC group of the applicable authorised codes in the ATC code field (level 3 or the level that can be defined) in D.3.3	
D.2.2.4	Other: No •	
D.2.2.4.1	If 'Yes', please specify:	
D.2.3	IMPD submitted:	
D.2.3.1	Full IMPD: No •	
D.2.3.2	Simplified IMPD: No •	
D.2.3.3	Summary of product characteristics (SmPC) only: Yes •	
D.2.4	Has the use of the IMP been previously authorised in a Yes •	

D.2.4.1	clinical trial conducted by the sponsor in the Community? If 'Yes' specify which Member States: United Kingdom
D.2.5	Has the IMP been designated in this indication as an orphan drug in the Community? No •
D.2.5.1	If 'Yes', give the orphan drug designation number ¹⁰ :

D.2.6	Has the IMP been the subject of scientific advice related to this clinical trial? No •
D.2.6.1	If 'Yes' to D.2.6, please indicate source of advice and provide a copy in the CTA request:
D.2.6.1.1	CHMP ¹¹ ? Not Answered •
D.2.6.1.2	National Competent Authority? Not Answered •

D.3	DESCRIPTION OF THE IMP
D.3.1	Product name where applicable ¹² : Clopidogrel
D.3.2	Product code where applicable ¹³ :
D.3.3	ATC codes, if officially registered ¹⁴ : B01AC04
D.3.4	Pharmaceutical form (use standard terms): Tablet
D.3.4.1	Is this a specific paediatric formulation? No •
D.3.5	Maximum duration of treatment of a subject according to the protocol: 30
D.3.6	Dose allowed:
D.3.6.1	For first trial only: Specify per day or total Not Answered • Specify total dose (number and unit): Route of administration (relevant to the first dose):
D.3.6.2	For all trials: Specify per day or total Per day • Specify total dose (number and unit): 75 mg milligram(s) Route of administration (relevant to the maximum dose): Enteral use (Noncurrent)
D.3.7	Routes of administration (use standard terms): Oral use Enteral use (Noncurrent)

D.3.8	Name of each active substance (INN or proposed INN if available): CLOPIDOGREL
D.3.9	Other available name for each active substance (provide all available):
D.3.9.1	CAS ¹⁵ number 94188848
D.3.9.2	Current sponsor code
D.3.9.3	Other descriptive name
D.3.9.4	EV Substance code
D.3.9.5	Full Molecular formula
D.3.9.6	Chemical/biological description of the Active Substance
D.3.10	Strength (specify all strengths to be used):
D.3.10.1	Concentration unit: mg milligram(s)
D.3.10.2	Concentration type ("exact number", "range", "more than" or "up to"): range
D.3.10.3	Concentration (number). 75 - 300

D.3.11	Type of IMP
Does the IMP contain an active substance:	
D.3.11.1	Of chemical origin? Yes •
D.3.11.2	Of biological / biotechnological origin (other than Advanced Therapy IMP (ATIMP))? No •
Is this a:	

D.3.11.3	Advanced Therapy IMP (ATIMP)?	No ●
D.3.11.3.1	Somatic cell therapy medicinal product ¹⁶ ?	No ●
D.3.11.3.2	Gene therapy medicinal product ¹⁷ ?	No ●
D.3.11.3.3	Tissue Engineered Product ¹⁸ ?	No ●
D.3.11.3.4	Combination ATIMP (i.e. one involving a medical device ¹⁹)?	No ●
D.3.11.3.5	Has the Committee on Advanced Therapies issued a classification for this product?	No ●
D.3.11.3.5.1	If 'Yes' please provide that classification and its reference number:	
D.3.11.4	Combination product that includes a device, but does not involve an Advanced Therapy?	No ●
D.3.11.5	Radiopharmaceutical medicinal product?	No ●
D.3.11.6	Immunological medicinal product (such as vaccine, allergen, immune serum)?	No ●
D.3.11.7	Plasma derived medicinal product?	No ●
D.3.11.8	Extractive medicinal product?	No ●
D.3.11.9	Recombinant medicinal product?	No ●
D.3.11.10	Medicinal product containing genetically modified organisms?	No ●
D.3.11.10.1	Has the authorisation for contained use or release been granted?	Not Answered ●
D.3.11.10.2	Is it pending?	Not Answered ●
D.3.11.11	Herbal medicinal product?	No ●
D.3.11.12	Homeopathic medicinal product?	No ●
D.3.11.13	Another type of medicinal product?	No ●
D.3.11.13.1	If 'another type of medicinal product' specify the type of medicinal product:	
D.3.12	Mode of action (<i>free text</i> ²⁰) Platelets are tiny blood cells that join together to form a clot, at sites of injury, and thus prevent bleeding. However, clots can be dangerous if they form spontaneously in blood vessels, as they can lead to stroke or heart attacks. Clopidogrel is an antiplatelet drug that is able to stop a substance called ADP from sticking to platelets, which in turn inhibits the ability of platelets to join together and form a clot.	
D.3.13	Is it an IMP to be used in a first-in-human clinical trial?	No ●
D.3.13.1	If 'Yes', are there risk factors identified, according to the guidance FIH? ²¹	

D.4	SOMATIC CELL THERAPY INVESTIGATIONAL MEDICINAL PRODUCT (NO GENETIC MODIFICATION)	
D.4.1	Origin of cells	
D.4.1.1	Autologous	No ●
D.4.1.2	Allogeneic	No ●
D.4.1.3	Xenogeneic	No ●
D.4.1.3.1	If 'Yes', specify the species of origin:	
D.4.2	Type of cells	
D.4.2.1	Stem cells	No ●
D.4.2.2	Differentiated cells	No ●
D.4.2.2.1	If 'Yes', specify the type (e.g. keratinocytes, fibroblasts, chondrocytes...):	
D.4.2.3	Others:	No ●
D.4.2.3.1	If others, specify:	

D.5	GENE THERAPY INVESTIGATIONAL MEDICINAL PRODUCTS	
D.5.1	Gene(s) of interest:	
D.5.2	In vivo gene therapy:	No ●
D.5.3	Ex vivo gene therapy:	No ●

D.5.4	Type of gene transfer product	
D.5.4.1	Nucleic acid (e.g. plasmid): If 'Yes', specify if:	No ●
D.5.4.1.1	Naked:	No ●
D.5.4.1.2	Complexed	No ●
D.5.4.2	Viral vector:	No ●
D.5.4.2.1	If 'Yes', specify the type: adenovirus, retrovirus, AAV, ...:	
D.5.4.3	Others	No ●
D.5.4.3.1	If others, specify:	
D.5.5	Genetically modified somatic cells: If 'Yes', specify the origin of the cells:	No ●
D.5.5.1	Autologous:	No ●
D.5.5.2	Allogeneic:	No ●
D.5.5.3	Xenogeneic:	No ●
D.5.5.3.1	If 'Yes', specify the species of origin:	
D.5.5.4	Specify type of cells (hematopoietic stem cells...):	

D.6 TISSUE ENGINEERED PRODUCT		
The indication which determines that this is a Tissue Engineered Product as opposed to a Cell Therapy product is given in section E.1.1.		
D.6.1	Origin of cells	
D.6.1.1	Autologous	No ●
D.6.1.2	Allogeneic	No ●
D.6.1.3	Xenogeneic	No ●
D.6.1.3.1	If 'Yes', specify the species of origin:	
D.6.2	Type of cells	
D.6.2.1	Stem cells	No ●
D.6.2.2	Differentiated cells	No ●
D.6.2.2.1	If 'Yes', specify the type of cells(e.g. keratinocytes, fibroblasts, chondrocytes, ...):	
D.6.2.3	Others:	No ●
D.6.2.3.1	If others, specify:	

D.7 PRODUCTS CONTAINING DEVICES (i.e. MEDICAL DEVICES, SCAFFOLDS ETC.)		
D.7.1	Give a brief description of the device:	
D.7.2	What is the name of the device?	
D.7.3	Is the device implantable?	No ●
D.7.4	Does this product contain:	
D.7.4.1	A medical device?	No ●
D.7.4.1.1	Does this medical device have a CE mark?	No ●
D.7.4.1.1.1	The notified body is:	
D.7.4.2	Bio-materials?	No ●
D.7.4.3	Scaffolds?	No ●
D.7.4.4	Matrices?	No ●
D.7.4.5	Other?	No ●
D.7.4.5.1	If other, specify:	

D.8 INFORMATION ON PLACEBO (if relevant; repeat as necessary)		
D.8.1	Is there a placebo:	No ●

D.8.2	This refers to placebo number:
D.8.3	Pharmaceutical form:
D.8.4	Route of administration:
D.8.5	Which IMP is it a placebo for? Specify IMP Number(s) from D.1.1
D.8.5.1	Composition, apart from the active substance(s):
D.8.5.2	Is it otherwise identical to the IMP? Yes <input type="radio"/> No <input type="radio"/> Not Answered <input type="radio"/>
D.8.5.2.1	If not, specify major ingredients:

D.9 SITE(S) WHERE THE QUALIFIED PERSON CERTIFIES BATCH RELEASE²²

*This section is dedicated to **finished** IMPs, i.e. medicinal products randomised, packaged, labelled and certified for use in the clinical trial. If there is more than one site or more than one IMP is certified, use extra pages and give each IMP its number from section D.1.1 or D.8.2 In the case of multiple sites indicate the product certified by each site*

D.9.1 Do not fill in section D.9.2 for an IMP that:
*Has a MA in the EU **and***
*Is sourced from the EU market **and***
*Is used in the trial without modification(e.g. not overencapsulated) **and***
The packaging and labelling is carried out for local use only as per article 9.2. of the Directive 2005/28/EC (GCP Directive)
 If all these conditions are met tick and list the number(s) of each IMP including placebo from sections D.1.1 and D.8.2 to which this applies
PR1

D.9.2 Who is responsible in the Community for the certification of the finished IMPs?

This site is responsible for certification of (list the number(s) of each IMP including placebo from sections D.1.1 and D.8.2):
 please tick the appropriate box:

- D.9.2.1 Manufacturer
- D.9.2.2 Importer
- D.9.2.3 Name of the organisation:
- D.9.2.4 Address:
- D.9.2.4.1 Street Address
- D.9.2.4.2 Town/City
- D.9.2.4.3 Post Code
- D.9.2.4.4 Country
- D.9.2.5 Give the manufacturing authorisation number:
- D.9.2.5.1 If No authorisation, give the reasons:

Where the product does not have a MA in the EU, but is supplied in bulk and final packaging and labelling for local use is carried out in accordance with article 9.2 of Directive 2005/28/EC (GCP Directive) then enter the site where the product was finally certified for release by the Qualified Person for use in the clinical trial at D.9.2 above.

E. GENERAL INFORMATION ON THE TRIAL

This section should be used to provide information about the aims, scope and design of the trial. When the protocol includes a sub-study in the MS concerned section E.2.3 should be completed providing information about the sub-study. To identify it check the sub-study box in the 'Objective of the trial' question below.

E.1 MEDICAL CONDITION OR DISEASE UNDER INVESTIGATION					
E.1.1	Specify the medical condition(s) to be investigated ²³ (free text): English Ischaemic stroke and TIA (transient ischaemic attack)				
E.1.1.1	Medical condition in easily understood language English Stroke due to blood clot.				
E.1.1.2	Therapeutic area Diseases [C] - Cardiovascular Diseases [C14]				
E.1.2	MedDRA version, system organ class, level, term and classification code ²⁴ :				
	Version	System Organ Class	Classification Code	Term	Level
	14.1	10029205 - Nervous system disorders	10042244	Stroke	LLT
	14.1	10029205 - Nervous system disorders	10044390	Transient ischaemic attack	PT
E.1.3	Is any of the conditions being studied a rare disease ²⁵ ?				No •
E.2 OBJECTIVE OF THE TRIAL					
E.2.1	Main objective: English To assess the safety of short-term administration (1 month) of triple antiplatelet therapy (aspirin, dipyridamole and clopidogrel) versus standard dual therapy (aspirin and dipyridamole) in patients with very recent ischaemic stroke or TIA.				
E.2.2	Secondary objectives: English 1. <input type="checkbox"/> To further assess, in high risk patients with stroke/TIA, whether the addition of C to AD: i. <input type="checkbox"/> Is feasible to administer acutely and tolerable to take for 1 month, ii. <input type="checkbox"/> Is superior in respect of surrogate markers assessed as emboli (with transcranial doppler) and platelet function iii. <input type="checkbox"/> Improves functional outcome 2. <input type="checkbox"/> To assess whether ordinal outcomes are superior to binary events				
E.2.3	Is there a sub-study?				No •
E.2.3.1	If 'Yes', give the full title, date and version of each sub-study and their related objectives:				
E.3 PRINCIPAL INCLUSION CRITERIA (list the most important)					
	English	Adults at high risk of recurrent ischaemic stroke: 1. <input type="checkbox"/> Acute non-cardioembolic ischaemic stroke (<48 hours of onset). All strokes must have motor weakness or dysphasia at the time of randomisation. 2. <input type="checkbox"/> Acute TIA (<48 hours of onset) with an ABCD2 score >5. All TIAs must have motor weakness and/or dysphasia lasting at least 10 minutes. 3. <input type="checkbox"/> Meaningful consent, or consent from a relative, carer or legal representative if the patient is unable to give meaningful consent (e.g. in cases of dysphasia, confusion, or reduced conscious level).			

E.4	PRINCIPAL EXCLUSION CRITERIA (list the most important)	
	English	<ol style="list-style-type: none"> 1. <input type="checkbox"/> Age < 50; 2. <input type="checkbox"/> Motor weakness or dysphasia lasting < 10 minutes; 3. <input type="checkbox"/> Pure sensory, vertigo or dizziness, speech or visual disturbance symptoms without weakness or dysphasia; 4. <input type="checkbox"/> Patients with contraindications to, or intolerance of, aspirin, clopidogrel or dipyridamole; 5. <input type="checkbox"/> Patients with definite need for treatment with clopidogrel (e.g. recent MI) 6. <input type="checkbox"/> Pre-morbid dependency (mRS > 2); 7. <input type="checkbox"/> No enteral access; 8. <input type="checkbox"/> Parenchymal haemorrhagic transformation (PH I/II), subarachnoid haemorrhage or other non ischaemic cause for weakness; 9. <input type="checkbox"/> TIA not fulfilling inclusion criteria 10. <input type="checkbox"/> Definite need for full dose oral (e.g. warfarin) or parental (e.g. heparin or glycoprotein IIb IIIa inhibitors) anti-coagulation. NB Low dose heparin for DVT prophylaxis is allowed. 11. <input type="checkbox"/> Received thrombolysis within the last 30 hours; 12. <input type="checkbox"/> Presumed cardioembolic stroke (e.g. AF, recent MI, or other conditions need for anticoagulation); 13. <input type="checkbox"/> Severe high BP (BP > 185/110 mmHg); 14. <input type="checkbox"/> Known haemoglobin less than 10g/dL 15. <input type="checkbox"/> Known platelet count less than 100 x 10⁹ /L 16. <input type="checkbox"/> Known white cell count less than 3.5 x 10⁹ /L 17. <input type="checkbox"/> Bleeding within 1 year (e.g. peptic ulcer, intracerebral haemorrhage); 18. <input type="checkbox"/> Planned surgery during 3 month follow-up (e.g. carotid endarterectomy). 19. <input type="checkbox"/> Concomitant acute coronary syndrome; 20. <input type="checkbox"/> Stroke secondary to a procedure (e.g. carotid or coronary intervention); 21. <input type="checkbox"/> Coma (GCS < 8) 22. <input type="checkbox"/> Non-stroke life expectancy < 6 months; 23. <input type="checkbox"/> Dementia 24. <input type="checkbox"/> Participation in another drug trial concurrently or within 30 days. (Patients may be randomised into observational studies or non-drug trials) 25. <input type="checkbox"/> Not available for follow-up e.g. no fixed address, overseas visitor 26. <input type="checkbox"/> Females of childbearing potential, pregnancy or breastfeeding

E.5	END POINT(S):	
E.5.1	Primary End Point (repeat as necessary) ²⁶	
	English	This start-up phase will assess ordinal bleeding (fatal/major/minor/none²⁶) at 35 days (end of treatment) as adjudicated by an independent blinded panel.
E.5.1.1	Timepoint(s) of evaluation of this end point	
	English	8 years 3 months

E.5.2	Secondary End Point (repeat as necessary)	English	None
E.5.2.1	Timepoint(s) of evaluation of this end point	English	N/A

E.6 SCOPE OF THE TRIAL – Tick all boxes where applicable			
E.6.1	Diagnosis		No •
E.6.2	Prophylaxis		No •
E.6.3	Therapy		Yes •
E.6.4	Safety		Yes •
E.6.5	Efficacy		Yes •
E.6.6	Pharmacokinetic		No •
E.6.7	Pharmacodynamic		No •
E.6.8	Bioequivalence		No •
E.6.9	Dose Response		No •
E.6.10	Pharmacogenetic		No •
E.6.11	Pharmacogenomic		No •
E.6.12	Pharmacoeconomic		No •
E.6.13	Others		No •
E.6.13.1	If others, specify:		

E.7 TRIAL TYPE AND PHASE²⁷			
E.7.1	Human pharmacology (Phase I)		No •
Is it:			
E.7.1.1	First administration to humans		No •
E.7.1.2	Bioequivalence study		No •
E.7.1.3	Other:		No •
E.7.1.3.1	If other, please specify:		
E.7.2	Therapeutic exploratory (Phase II)		No •
E.7.3	Therapeutic confirmatory (Phase III)		Yes •
E.7.4	Therapeutic use(Phase IV)		No •

E.8 DESIGN OF THE TRIAL			
E.8.1	Controlled		Yes •
	If 'Yes', specify:		
E.8.1.1	Randomised:		Yes •
E.8.1.2	Open:		Yes •
E.8.1.3	Single blind:		No •
E.8.1.4	Double blind:		No •
E.8.1.5	Parallel group:		Yes •
E.8.1.6	Cross over:		No •
E.8.1.7	Other:		Yes •
E.8.1.7.1	If other specify:		
	English	Assessor blinded	
E.8.2	If controlled, specify the comparator:		
E.8.2.1	Other medicinal product(s)		Yes •
E.8.2.2	Placebo		No •
E.8.2.3	Other		No •
E.8.2.3.1	If 'Yes' to other, specify :		
E.8.2.4	Number of treatment arms in the trial	2	
E.8.3	Single site in the Member State concerned (see also section G):		No •
E.8.4	Multiple sites in the Member State concerned(see also section G):		Yes •
E.8.4.1	Number of sites anticipated in Member State concerned	100	
E.8.5	Multiple Member States:		No •
E.8.5.1	Number of sites anticipated in the EEA:		

E.8.6	Trial involving sites outside the EEA:	
E.8.6.1	Trial being conducted both within and outside the EEA:	No •
E.8.6.2	Trial being conducted completely outside of the EEA:	No •
E.8.6.3	If E.8.6.1 or E.8.6.2 are Yes, specify the regions in which trial sites are planned:	
E.8.6.4	If E.8.6.1 or E.8.6.2 are Yes, specify the number of sites anticipated outside of the EEA:	
E.8.7	Trial having an independent data monitoring committee:	Yes •
E.8.8	Definition of the end of trial: If it is the last visit of the last subject, please enter "LVLS". If it is not LVLS provide the definition: English As per protocol	
E.8.9	Initial estimate of the duration of the trial ²⁸ (years, months and days)	
E.8.9.1	In the Member State concerned	8 years 3 months 0 days
E.8.9.2	In all countries concerned by the trial	8 years 3 months 0 days
E.8.10	Proposed date of start of recruitment	
E.8.10.1	In the Member State concerned	2009-04-01
E.8.10.2	In any country	2009-04-01

F. POPULATION OF TRIAL SUBJECTS

F.1 AGE RANGE		
F.1.1	Are the trial subjects under 18? If 'Yes', specify the estimated number of subjects planned in each age range for the whole trial:	No •
	Approx. No. of patients ²⁹	
F.1.1.1	In utero	() No •
F.1.1.2	Preterm newborn infants (up to gestational age < 37 weeks)	() No •
F.1.1.3	Newborns (0-27 days)	() No •
F.1.1.4	Infants and toddlers (28 days - 23 months)	() No •
F.1.1.5	Children (2-11 years)	() No •
F.1.1.6	Adolescents (12-17 years)	() No •
F.1.2	Adults (18-64 years)	(350) Yes •
F.1.3	Elderly (>= 65 years)	(650) Yes •
F.2 GENDER		
F.2.1	Female	Yes •
F.2.2	Male	Yes •
F.3 GROUP OF TRIAL SUBJECTS		
F.3.1	Healthy volunteers	No •
F.3.2	Patients	Yes •
F.3.3	Specific vulnerable populations	Yes •
F.3.3.1	Women of child bearing potential not using contraception	No •
F.3.3.2	Women of child bearing potential using contraception	No •
F.3.3.3	Pregnant women	No •
F.3.3.4	Nursing women	No •
F.3.3.5	Emergency situation	No •
F.3.3.6	Subjects incapable of giving consent personally	Yes •
F.3.3.6.1	If 'Yes', specify: English Patients with a stroke are often incapacitated through confusion, dysphasia or coma.	
F.3.3.7	Others:	Yes •
F.3.3.7.1	If 'Yes', specify: English Elderly patients	
F.4 PLANNED NUMBER OF SUBJECTS TO BE INCLUDED:		
F.4.1	In the member state	750
F.4.2	For a multinational trial:	
F.4.2.1	In the EEA	
F.4.2.2	In the whole clinical trial	
F.5 PLANS FOR TREATMENT OR CARE AFTER THE SUBJECT HAS ENDED HIS/HER PARTICIPATION IN THE TRIAL. please specify (free text):		
	English As per protocol.	

G. CLINICAL TRIAL SITES/INVESTIGATORS IN THE MEMBER STATE CONCERNED BY THIS REQUEST

G.1	CO-ORDINATING INVESTIGATOR (for multicentre trial) and principal investigator (for single centre trial)	
G.1.1	Given name:	Philip
G.1.2	Middle name, if applicable:	
G.1.3	Family name:	Bath
G.1.4	Qualification (MD.....)	BSc MBBS MRCP MD FRCP FRCPATH
G.1.5	Professional address:	
G.1.5	Institution name	University of Nottingham
G.1.5	Institution department	Division of Stroke Medicine
G.1.5.1	Street address	Clinical Sciences Building, City Hospital Campus, Hucknall Rd
G.1.5.2	Town/city	Nottingham
G.1.5.3	Post code	NG5 1PB
G.1.5.4	Country	United Kingdom
G.1.6	Telephone number:	+44 0115 8231768
G.1.7	Fax number:	+44 0115 8231767
G.1.8	E-mail:	philip.bath@nottingham.ac.uk

G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
G.2.1	Given name:	Sunil
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Munshi
G.2.4	Qualification (MD.....)	MD(Medicine) Dip NBE FRCP Ed FRCP Lond
G.2.5	Professional address:	
G.2.5	Institution name	Nottingham University Hospitals NHS Trust
G.2.5	Institution department	Dept of Stroke & Health Care of the Elderly
G.2.5.1	Street address	City Hospital Campus
G.2.5.2	Town/city	Nottingham
G.2.5.3	Post code	NG5 1PB
G.2.5.4	Country	United Kingdom
G.2.6	Telephone number:	+44 0115 9691169 5565
G.2.7	Fax number:	+44 0115 9608409
G.2.8	E-mail:	sunil.munshi@nuh.nhs.uk

G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
G.2.1	Given name:	Amit
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Mistry
G.2.4	Qualification (MD.....)	MBBS, MRCP(UK)
G.2.5	Professional address:	
G.2.5	Institution name	University Hospitals of Leicester
G.2.5	Institution department	Dept of Cardiovascular Sciences
G.2.5.1	Street address	UHL NHS Trust, Gwendolen Rd
G.2.5.2	Town/city	Leicester
G.2.5.3	Post code	LE4 4PW
G.2.5.4	Country	United Kingdom
G.2.6	Telephone number:	+44 0116 25848731
G.2.7	Fax number:	+44 0116 2584187
G.2.8	E-mail:	akm17@le.ac.uk

G.2 PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)		
G.2.1	Given name:	Catrin
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Blank
G.2.4	Qualification (MD.....)	MBBS MRCP(UK)
G.2.5	Professional address:	
G.2.5	Institution name	Sheffield Teaching Hospitals
G.2.5	Institution department	Neurology Dept
G.2.5.1	Street address	Royal Hallamshire Hospital
G.2.5.2	Town/city	Sheffield
G.2.5.3	Post code	S10 2JF
G.2.5.4	Country	United Kingdom
G.2.6	Telephone number:	+44 0114 2268754
G.2.7	Fax number:	+44 0114 2268946
G.2.8	E-mail:	catrin.blank@sth.nhs.uk

G.2 PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)		
G.2.1	Given name:	Khulood
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Muhiddin
G.2.4	Qualification (MD.....)	MBChB, FRCP, PhD
G.2.5	Professional address:	
G.2.5	Institution name	Derby Hospitals NHS Foundation Trust
G.2.5	Institution department	Medicine for the Elderly
G.2.5.1	Street address	Uttoxeter Road
G.2.5.2	Town/city	Derby
G.2.5.3	Post code	DE22 3NE
G.2.5.4	Country	United Kingdom
G.2.6	Telephone number:	+44 01332 787724
G.2.7	Fax number:	+44 01332 787500
G.2.8	E-mail:	khulood.muhammad@derbyhospitals.nhs.uk

G.2 PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)		
G.2.1	Given name:	Martin
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Cooper
G.2.4	Qualification (MD.....)	MBB ChB, MRCP, CCST
G.2.5	Professional address:	
G.2.5	Institution name	Sherwood Forest Hospitals NHS Foundation Trust
G.2.5	Institution department	Medicine for the Elderly
G.2.5.1	Street address	King's Mill Hospital, Mansfield Rd
G.2.5.2	Town/city	Sutton-in -Ashfield, Nottinghamshire
G.2.5.3	Post code	NG17 4JL
G.2.5.4	Country	United Kingdom
G.2.6	Telephone number:	+44 01623 676160
G.2.7	Fax number:	+44 01623 672308
G.2.8	E-mail:	martin.cooper@sfh-tr.nhs.uk

G.2 PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)		
G.2.1	Given name:	Mohammad
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Albazzaz

G.2.4	Qualification (MD.....)	MBChB, FRCP(UK)
G.2.5	Professional address:	
G.2.5	Institution name	Barnsley Hospital NHS Foundation Trust
G.2.5	Institution department	General Medicine
G.2.5.1	Street address	Barnsley Hospital NHS Foundation Trust
G.2.5.2	Town/city	Gawber Road, Barnsley
G.2.5.3	Post code	S75 2EP
G.2.5.4	Country	United Kingdom
G.2.6	Telephone number:	+44 01226 432035
G.2.7	Fax number:	+44 01226 432035
G.2.8	E-mail:	m.albazzaz@nhs.net

G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
G.2.1	Given name:	Khalid
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Ayes
G.2.4	Qualification (MD.....)	MBBS, MRCP(UK), MSc
G.2.5	Professional address:	
G.2.5	Institution name	Kettering General Hospital NHS Foundation Trust
G.2.5	Institution department	Stroke Unit
G.2.5.1	Street address	Kettering General Hospital
G.2.5.2	Town/city	Rothwell Rd, Kettering
G.2.5.3	Post code	NN16 8UZ
G.2.5.4	Country	United Kingdom
G.2.6	Telephone number:	+44 01536 492254
G.2.7	Fax number:	+44 01536 492062
G.2.8	E-mail:	khalid.ayes@kgh.nhs.uk

G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
G.2.1	Given name:	Melanie
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Blake
G.2.4	Qualification (MD.....)	BMBCh, MRCP
G.2.5	Professional address:	
G.2.5	Institution name	Northampton General Hospital NHS Trust
G.2.5	Institution department	General Medicine
G.2.5.1	Street address	Cliftonville
G.2.5.2	Town/city	Northampton
G.2.5.3	Post code	NN1 5BD
G.2.5.4	Country	United Kingdom
G.2.6	Telephone number:	+44 01604 545963
G.2.7	Fax number:	+44 01604 545556
G.2.8	E-mail:	melanie.blake@ngh.nhs.uk

G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
G.2.1	Given name:	Dinesh
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Chadha
G.2.4	Qualification (MD.....)	MBBS, MD, FRCP
G.2.5	Professional address:	
G.2.5	Institution name	Doncaster and Bassetlaw Hospitals NHS Foundation Trust.
G.2.5	Institution department	Stroke/Health care of the elderly

G.2.5.1	Street address	Armthorpe Rd
G.2.5.2	Town/city	Doncaster
G.2.5.3	Post code	DN2 5LT
G.2.5.4	Country	United Kingdom
G.2.6	Telephone number:	+44 01302 366666
G.2.7	Fax number:	+44 01302 796139
G.2.8	E-mail:	dinesh.chadha@dbh.nhs.uk

G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
G.2.1	Given name:	Simon
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Leach
G.2.4	Qualification (MD.....)	MBChB, MRCP
G.2.5	Professional address:	
G.2.5	Institution name	United Lincolnshire Hospitals NHS Trust
G.2.5	Institution department	Lincoln County Hospital
G.2.5.1	Street address	Greetwell Road
G.2.5.2	Town/city	Lincoln
G.2.5.3	Post code	LN2 5QY
G.2.5.4	Country	United Kingdom
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G.2.7	Fax number:	+44 01522 543783
G.2.8	E-mail:	simon.leach@ulh.nhs.uk

G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
G.2.1	Given name:	David
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Mangion
G.2.4	Qualification (MD.....)	MRCS, LRCP, FRCP
G.2.5	Professional address:	
G.2.5	Institution name	United Lincolnshire Hospitals NHS Trust
G.2.5	Institution department	Pilgrim Hospital Boston
G.2.5.1	Street address	Sibsey Rd
G.2.5.2	Town/city	Boston
G.2.5.3	Post code	PE21 9QS
G.2.5.4	Country	United Kingdom
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G.2.7	Fax number:	+44 01205 353392
G.2.8	E-mail:	david.mangion@ulh.nhs.uk

G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
G.2.1	Given name:	James
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Okwera
G.2.4	Qualification (MD.....)	MBChB, FRCP
G.2.5	Professional address:	
G.2.5	Institution name	The Rotherham NHS Foundation Trust
G.2.5	Institution department	Rotherham General Hospital
G.2.5.1	Street address	Moorgate Road,
G.2.5.2	Town/city	Oakwood
G.2.5.3	Post code	S60 2UD
G.2.5.4	Country	United Kingdom
G.2.6	Telephone number:	+44 01709 304164

G.2.7	Fax number:	+44 01709 304283
G.2.8	E-mail:	james.okwera@rothgen.nhs.uk

G.2 PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)		
G.2.1	Given name:	Mahmud
G.2.2	Middle name, if applicable:	Ahmed
G.2.3	Family name:	Sajid
G.2.4	Qualification (MD.....)	MBBS FCPS MRCP UK
G.2.5	Professional address:	
G.2.5	Institution name	Chesterfield Royal Hospital NHS Foundation Trust
G.2.5	Institution department	Care of the Elderly/General Medicine
G.2.5.1	Street address	Calow, Chesterfield
G.2.5.2	Town/city	Derbyshire
G.2.5.3	Post code	S44 5BL
G.2.5.4	Country	United Kingdom
G.2.6	Telephone number:	+44 01246 512899
G.2.7	Fax number:	+44 01246 512670
G.2.8	E-mail:	mahmud.sajid@chesterfieldroyal.nhs.uk

G.2 PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)		
G.2.1	Given name:	Conrad
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Athulathmudali
G.2.4	Qualification (MD.....)	MBBS,FRCP
G.2.5	Professional address:	
G.2.5	Institution name	East Sussex Hospitals NHS Trust
G.2.5	Institution department	Stroke Unit
G.2.5.1	Street address	Eastbourne General Hospital, King's Drive
G.2.5.2	Town/city	Eastbourne
G.2.5.3	Post code	BN21 2UD
G.2.5.4	Country	United Kingdom
G.2.6	Telephone number:	+44 01323 435797
G.2.7	Fax number:	+44 01323 435797
G.2.8	E-mail:	Athulathmudali.conrad@esht.nhs.uk

G.2 PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)		
G.2.1	Given name:	Adrian
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Blight
G.2.4	Qualification (MD.....)	MBBS, MRCP
G.2.5	Professional address:	
G.2.5	Institution name	Royal Surrey County Hospital NHS Trust
G.2.5	Institution department	Stroke/elderly care
G.2.5.1	Street address	Guildford
G.2.5.2	Town/city	Surrey
G.2.5.3	Post code	GU2 7XX
G.2.5.4	Country	United Kingdom
G.2.6	Telephone number:	+44 01483 464105
G.2.7	Fax number:	+44 01483 302683
G.2.8	E-mail:	abligh@roysurrey.nhs.uk

G.2 PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)		
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forms)		
G.2.1	Given name:	Gunaratnam
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Gunathilagan
G.2.4	Qualification (MD.....)	FRCP
G.2.5	Professional address:	
G.2.5	Institution name	East Kent Hospitals University NHS Trust
G.2.5	Institution department	Stroke/Health care of the elderly(HCOOP)
G.2.5.1	Street address	QEQM Hospital, St Peter's Rd
G.2.5.2	Town/city	Margate
G.2.5.3	Post code	CT9 4NA
G.2.5.4	Country	United Kingdom
G.2.6	Telephone number:	+44 01843 225544 6309
G.2.7	Fax number:	+44 01843 234451
G.2.8	E-mail:	g.gunathilagan@ekht.nhs.uk

G.2 PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)		
G.2.1	Given name:	Barry
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Moniyhan
G.2.4	Qualification (MD.....)	MD
G.2.5	Professional address:	
G.2.5	Institution name	St George's Healthcare NHS Trust
G.2.5	Institution department	Neurology Dept
G.2.5.1	Street address	Blackshaw Rd
G.2.5.2	Town/city	Tooting
G.2.5.3	Post code	SW17 0QT
G.2.5.4	Country	United Kingdom
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G.2 PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)		
G.2.1	Given name:	Brian
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Clarke
G.2.4	Qualification (MD.....)	MBBCh, BAO, LRCPI, LRCSI
G.2.5	Professional address:	
G.2.5	Institution name	Frimley Park Hospital NHS Foundation Trust
G.2.5	Institution department	Stroke/HCE
G.2.5.1	Street address	Portsmouth Road
G.2.5.2	Town/city	Frimley, Surrey
G.2.5.3	Post code	GU16 7UJ
G.2.5.4	Country	United Kingdom
G.2.6	Telephone number:	+44 01276 604085
G.2.7	Fax number:	+44 01276 604862
G.2.8	E-mail:	Brian.clarke@fph-tr.nhs.uk

G.2 PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)		
G.2.1	Given name:	Mehool
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Patel

G.2.4	Qualification (MD.....)	MBBS
G.2.5	Professional address:	
G.2.5	Institution name	The Lewisham Hospital NHS Trust
G.2.5	Institution department	Stroke/HCE
G.2.5.1	Street address	Lewisham High Street
G.2.5.2	Town/city	London
G.2.5.3	Post code	SE13 6LH
G.2.5.4	Country	United Kingdom
G.2.6	Telephone number:	+44 0208 3333000 8185
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G.2.8	E-mail:	mehool.patel@nhs.net

G.2 PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)		
G.2.1	Given name:	Syed
G.2.2	Middle name, if applicable:	Muhammad Mazhar
G.2.3	Family name:	Zaidi
G.2.4	Qualification (MD.....)	MBBS, MRCP
G.2.5	Professional address:	
G.2.5	Institution name	James Paget University Hospitals NHS Foundation Trust
G.2.5	Institution department	Stroke
G.2.5.1	Street address	Lowestoft Road, Gorleston
G.2.5.2	Town/city	Great Yarmouth
G.2.5.3	Post code	NR31 6LA
G.2.5.4	Country	United Kingdom
G.2.6	Telephone number:	+44 01493 453776
G.2.7	Fax number:	+44 01493 452421
G.2.8	E-mail:	mazhar.zaidi@jpaget.nhs.uk

G.2 PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)		
G.2.1	Given name:	Frances
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Harrington
G.2.4	Qualification (MD.....)	MBChB, SRCP, CCST geriatric & general internal med
G.2.5	Professional address:	
G.2.5	Institution name	Royal Cornwall Hospitals NHS trust
G.2.5	Institution department	Stroke/Knowledge Spa Building
G.2.5.1	Street address	Royal Cornwall Hospital, Trelisk
G.2.5.2	Town/city	Truro
G.2.5.3	Post code	TR1 3LJ
G.2.5.4	Country	United Kingdom
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G.2.8	E-mail:	frances.harrington@rcht.cornwall.nhs.uk

G.2 PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)		
G.2.1	Given name:	Dulka
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Manawadu
G.2.4	Qualification (MD.....)	MSc MRCP MBChB
G.2.5	Professional address:	
G.2.5	Institution name	King's College Hospital NHS Foundation Trust
G.2.5	Institution department	Stroke Unit, Rushkin Wing, 5th floor
G.2.5.1	Street address	Denmark Hill

G.2.5.2	Town/city	London
G.2.5.3	Post code	SE5 9RS
G.2.5.4	Country	United Kingdom
G.2.6	Telephone number:	+44 07837 121299
G.2.7	Fax number:	+44 020 78485186
G.2.8	E-mail:	dulka.manawadu@nhs.net

G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
G.2.1	Given name:	Khalid
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Ali
G.2.4	Qualification (MD.....)	MBBS, MRCP MD
G.2.5	Professional address:	
G.2.5	Institution name	Brighton & Sussex University Hospitals NHS Trust
G.2.5	Institution department	Ac Dept of Geriatrics, Audrey Emerton Bldg
G.2.5.1	Street address	Princess Royal Hospital
G.2.5.2	Town/city	Haywards Heath, Brighton
G.2.5.3	Post code	BN2 5BE
G.2.5.4	Country	United Kingdom
G.2.6	Telephone number:	+44 01273 523360 3369
G.2.7	Fax number:	+44 01273 523366
G.2.8	E-mail:	khalid.ali@bsuh.nhs.uk

G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
G.2.1	Given name:	Raj
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Shekhar
G.2.4	Qualification (MD.....)	MBBS, MD, MRCP
G.2.5	Professional address:	
G.2.5	Institution name	Queen Elizabeth Hospital King's Lynn NHS Trust
G.2.5	Institution department	Stroke
G.2.5.1	Street address	Gayton Road
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G.2.5.4	Country	United Kingdom
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G.2.3	Family name:	Davies
G.2.4	Qualification (MD.....)	BSc(Hons) MD, FRCP
G.2.5	Professional address:	
G.2.5	Institution name	North Cumbria University Hospitals NHS Trust
G.2.5	Institution department	Stroke
G.2.5.1	Street address	Cumberland Infirmary (& west Cumberland Hospital
G.2.5.2	Town/city	Carlisle (& Whitehaven)
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G.2.2 Middle name, if applicable:
G.2.3 Family name: **Mohd Nor**
G.2.4 Qualification (MD.....): **MBBCh MD MRCPI MRCP(UK)**
G.2.5 Professional address:
G.2.5 Institution name: **Plymouth Hospitals NHS Trust**
G.2.5 Institution department: **Stroke**
G.2.5.1 Street address: **Derriford Hospital**
G.2.5.2 Town/city: **Plymouth**
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G.2 PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)

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G.2.2 Middle name, if applicable:
G.2.3 Family name: **Beavans**
G.2.4 Qualification (MD.....): **MD, MSc Geriatric Medicine, MRCP, MBChB**
G.2.5 Professional address:
G.2.5 Institution name: **Musgrove Park Foundation Trust**
G.2.5 Institution department: **Stroke**
G.2.5.1 Street address: **Musgrove Park**
G.2.5.2 Town/city: **Taunton**
G.2.5.3 Post code: **TA1 5DA**
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G.2.1 Given name: **Isam**
G.2.2 Middle name, if applicable: **Sayed Mahmoud**
G.2.3 Family name: **Salih**
G.2.4 Qualification (MD.....): **MBBS MRCP(UK)**
G.2.5 Professional address:
G.2.5 Institution name: **Torbay Hospital NHS Foundation Trust**
G.2.5 Institution department: **Stroke**
G.2.5.1 Street address: **Horizon Centre**
G.2.5.2 Town/city: **Torquay**
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G.2 PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)

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G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Rashed
G.2.4	Qualification (MD.....)	MBChB, MCRP, FRCP
G.2.5	Professional address:	
G.2.5	Institution name	Yeovil District Hospital NHS Foundation Trust
G.2.5	Institution department	Stroke
G.2.5.1	Street address	Higher Kingston
G.2.5.2	Town/city	Yeovil
G.2.5.3	Post code	BA21 4AT
G.2.5.4	Country	United Kingdom
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G.2.3	Family name:	Datta
G.2.4	Qualification (MD.....)	MBBS, MRCP
G.2.5	Professional address:	
G.2.5	Institution name	Mid Yorkshire Hosp.NHS Foundation Trust
G.2.5	Institution department	Dept of Stroke Medicine
G.2.5.1	Street address	Level 4, Bronte Towers, Halifax Rd
G.2.5.2	Town/city	Dewsbury
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G.2.3	Family name:	Chatterjee
G.2.4	Qualification (MD.....)	MD, MRCP, MBBS
G.2.5	Professional address:	
G.2.5	Institution name	Countess of Chester NHS Foundation Trust
G.2.5	Institution department	Stroke
G.2.5.1	Street address	Liverpool Road
G.2.5.2	Town/city	Chester
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G.2.5.4	Country	United Kingdom
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G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
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G.2.3	Family name:	Manoj
G.2.4	Qualification (MD.....)	MBBS, MRCP
G.2.5	Professional address:	

G.2.5	Institution name	Royal Liverpool & Broadgreen University Hospital NHS Trust
G.2.5	Institution department	Stroke/HCE
G.2.5.1	Street address	Prescott Street
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G.2.3	Family name:	Kumar
G.2.4	Qualification (MD.....)	MBBS(India)1994 MRCP(uk)1998 MD(UK)2007
G.2.5	Professional address:	
G.2.5	Institution name	Aintree Hospitals NHS foundation Trustintree
G.2.5	Institution department	Stroke Team for Audit & Research (STAR)
G.2.5.1	Street address	University Hospital Aintree, Lower Lane
G.2.5.2	Town/city	Liverpool
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G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Dizayee
G.2.4	Qualification (MD.....)	MBChB MSc FRCP
G.2.5	Professional address:	
G.2.5	Institution name	Stockport NHS Foundation Trust
G.2.5	Institution department	Stepping Hill Hospital
G.2.5.1	Street address	Poplara Grove, Hazel Grove
G.2.5.2	Town/city	Stockport
G.2.5.3	Post code	SK2 7JE
G.2.5.4	Country	United Kingdom
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G.2.3	Family name:	Hemsley
G.2.4	Qualification (MD.....)	BMedSci BM BS MRCP(UK) CCST MD SRCP
G.2.5	Professional address:	
G.2.5	Institution name	Royal Devon & Exeter Hospital Foundation Trust
G.2.5	Institution department	Stroke (in Child Health Building)
G.2.5.1	Street address	Barrack Road
G.2.5.2	Town/city	Exeter

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G.2.3	Family name:	Adiotomre
G.2.4	Qualification (MD.....)	FRCP FRCPI
G.2.5	Professional address:	
G.2.5	Institution name	North Lincolnshire & Goole Hospitals NHS Foundation Trust
G.2.5	Institution department	Stroke
G.2.5.1	Street address	Diana Princess of Wales Hospital, Scartho Road
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G.2.1	Given name:	Irfan
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Shakir
G.2.4	Qualification (MD.....)	BMedSci BMBS MRCP FRCP
G.2.5	Professional address:	
G.2.5	Institution name	Calderdale & Huddersfield NHS Foundation Trust
G.2.5	Institution department	Stroke
G.2.5.1	Street address	Calderdale Royal Hospital, Salterhebble
G.2.5.2	Town/city	Halifax
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G.2.5.4	Country	United Kingdom
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G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
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G.2.3	Family name:	Brotheridge
G.2.4	Qualification (MD.....)	MBBS, MRCP, FRCP, CCST gen & geriatric medicine
G.2.5	Professional address:	
G.2.5	Institution name	Harrogate & District NHS foundation Trust
G.2.5	Institution department	Medicine for the Elderly
G.2.5.1	Street address	Harrogate District Hospital, Lancaster Park Rd
G.2.5.2	Town/city	Harrogate
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G.2 PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)		
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G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Bamford
G.2.4	Qualification (MD.....)	MBChB(Hons), MRCP(UK), MD, FRCP
G.2.5	Professional address:	
G.2.5	Institution name	Leeds Teaching Hospitals NHS Trust
G.2.5	Institution department	Dept of Neurology, F Floor, Martin Wing
G.2.5.1	Street address	Leeds General Infirmary, George St
G.2.5.2	Town/city	Leeds
G.2.5.3	Post code	LS1 3EX
G.2.5.4	Country	United Kingdom
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G.2.7	Fax number:	+44 0113 3928070
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G.2 PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)		
G.2.1	Given name:	David
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Hargroves
G.2.4	Qualification (MD.....)	MB BSc
G.2.5	Professional address:	
G.2.5	Institution name	East Kent Hospitals University NHS Foundation Trust
G.2.5	Institution department	Stroke
G.2.5.1	Street address	William Harvey Hospital, Kennington Rd,
G.2.5.2	Town/city	Willesborough, Ashford Kent
G.2.5.3	Post code	TN24 0LZ
G.2.5.4	Country	United Kingdom
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G.2 PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)		
G.2.1	Given name:	Malcolm
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G.2.3	Family name:	Macleod
G.2.4	Qualification (MD.....)	BSc(Hons)(MBChB MRCP PhD
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G.2.5	Institution name	NHS Forth Valley
G.2.5	Institution department	Dept of Neurology
G.2.5.1	Street address	Forth Valley Royal Hospital, Stirling Rd
G.2.5.2	Town/city	Larbert
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G.2 PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)		
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G.2.1	Given name:	Mark
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Barber
G.2.4	Qualification (MD.....)	MD MRCP
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G.2.5	Institution name	NHS Lanarkshire
G.2.5	Institution department	Stroke/Dept of Medicine for the Elderly
G.2.5.1	Street address	Monklands Hospital, Monkscourt Avenue
G.2.5.2	Town/city	Airdrie
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G.2.5.4	Country	United Kingdom
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G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
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G.2.3	Family name:	Lees
G.2.4	Qualification (MD.....)	BSc(Hons) MBChB MRCP(UK) MD JCHMT FRCP
G.2.5	Professional address:	
G.2.5	Institution name	Western Infirmary
G.2.5	Institution department	Cardiovascular & Medical Sciences
G.2.5.1	Street address	Gardiner Institute 44 Church St
G.2.5.2	Town/city	Glasgow
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G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
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G.2.3	Family name:	MacWalter
G.2.4	Qualification (MD.....)	BMSc(Hons) MBChB(Hons) MD MRCP(UK) FRCP(Edin) FRCP
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G.2.5	Institution name	Ninewells Hospital & Medical School
G.2.5	Institution department	Acute Stroke Unit
G.2.5.1	Street address	off George Pine Way
G.2.5.2	Town/city	Dundee
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G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
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G.2.3	Family name:	Macleod
G.2.4	Qualification (MD.....)	MBChB, PhD, FRCP

G.2.5	Professional address:	
G.2.5	Institution name	Aberdeen Royal Infirmary
G.2.5	Institution department	Division of Applied Medicine/Stroke Unit
G.2.5.1	Street address	Foresterhill
G.2.5.2	Town/city	Aberdeen
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G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
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G.2.3	Family name:	Mandall
G.2.4	Qualification (MD.....)	Consultant in Elderly Care
G.2.5	Professional address:	
G.2.5	Institution name	Ashford & St Peters Hospital NHS trust
G.2.5	Institution department	Stroke Research/Duchess of Kent Wing level 4
G.2.5.1	Street address	St Peters Hospital, Guildford Rd
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G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
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G.2.3	Family name:	Ngeh
G.2.4	Qualification (MD.....)	MB(Hons) DME DGM MSc MD FRCP
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G.2.5	Institution department	Stroke
G.2.5.1	Street address	Heath Road
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G.2.5.4	Country	United Kingdom
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G.2.3	Family name:	McIlmoyle
G.2.4	Qualification (MD.....)	MBBCh MRCP FRCP
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G.2.5	Institution name	Blackpool Teaching Hospitals NHS Foundation Trust
G.2.5	Institution department	Stroke
G.2.5.1	Street address	Blackpool Victoria Hospital, Whinney Heys Road
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G.2.3	Family name:	Guyler
G.2.4	Qualification (MD.....)	BSc(Hons) MBBS MRCP
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G.2.5	Institution name	Southend University Hospital NHS Foundation Trust
G.2.5	Institution department	Stroke
G.2.5.1	Street address	Prittlewell Chase
G.2.5.2	Town/city	Westcliffe on Sea
G.2.5.3	Post code	SS0 0RY
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G.2.3	Family name:	Langhorne
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G.2.5	Institution name	Greater Glasgow NHS Board/Glasgow Royal Infirmary
G.2.5	Institution department	Academic Section of Geriatric Medicine
G.2.5.1	Street address	Level 3 University Block
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G.2.3	Family name:	Cvoro
G.2.4	Qualification (MD.....)	MBCHB
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G.2.5	Institution department	Stroke Unit, c/o Ward 11
G.2.5.1	Street address	Hayfield Road
G.2.5.2	Town/city	Kilcaldy
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G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
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G.2.3	Family name:	Sanmuganathan
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G.2.5	Institution department	Elderly Care, Level 1 Brown Zone
G.2.5.1	Street address	Windmill Road
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G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
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G.2.5	Institution name	Northumbria Healthcare NHS Found.Trust
G.2.5	Institution department	Wansbeck General Hospital(& N.Tyneside)
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G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
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G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Dixit
G.2.4	Qualification (MD.....)	MRCP MD DGM
G.2.5	Professional address:	
G.2.5	Institution name	Newcastle Upon Tyne Hospitals NHS trust
G.2.5	Institution department	Stroke/Clin.Research Facility, level 4
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G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
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G.2.3	Family name:	Dent
G.2.4	Qualification (MD.....)	MD MRCP MBChB
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G.2.5	Institution name	North Devon Healthcare NHS Trust/N.Devon District Hospital
G.2.5	Institution department	Stroke/Chichester House
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G.2.3	Family name:	Jenkinson
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G.2.5	Professional address:	
G.2.5	Institution name	Royal Bournemouth & Christchurch Hospitals NHS Foundation Trust
G.2.5	Institution department	Stroke
G.2.5.1	Street address	The Royal Bournemouth Hospital, Castle Lane East
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G.2.3	Family name:	Aghoram
G.2.4	Qualification (MD.....)	MBBS FRCP
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G.2.5	Institution name	Dartford & Gravesend NHS trust
G.2.5	Institution department	Stroke
G.2.5.1	Street address	Darenth Valley, Darenth Wood Rd
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G.2.3	Family name:	Williams

G.2.4	Qualification (MD.....)	FRCP MBBS BSc MD
G.2.5	Professional address:	
G.2.5	Institution name	Dorset County Hospital NHS Foundation Trust/Dorset County Hospital
G.2.5	Institution department	Stroke Unit
G.2.5.1	Street address	Williams Avenue
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G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
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G.2.3	Family name:	Paterson
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G.2.5	Professional address:	
G.2.5	Institution name	Scarborough & North East Yorkshire Healthcare NHS Trust
G.2.5	Institution department	Stroke Unit (1st Floor)
G.2.5.1	Street address	Scarborough General Hospital. Woodlands Drive
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G.2.3	Family name:	Scott
G.2.4	Qualification (MD.....)	BMedSci BM B MD FRCP
G.2.5	Professional address:	
G.2.5	Institution name	South Tyneside District Hospital
G.2.5	Institution department	Stroke
G.2.5.1	Street address	Harton Lane
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G.2.3	Family name:	Kerr
G.2.4	Qualification (MD.....)	MBCLB MRCP
G.2.5	Professional address:	
G.2.5	Institution name	NHS Borders - Borders General Hospital

G.2.5	Institution department	Stroke
G.2.5.1	Street address	off A6091
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G.2.4	Qualification (MD.....)	MBBS FRCP
G.2.5	Professional address:	
G.2.5	Institution name	Lancashire Teaching Hospitals NHS Foundation Trust / Royal Preston Hospital
G.2.5	Institution department	Stroke
G.2.5.1	Street address	Sharoe Green Lane, Fulwood
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G.2.3	Family name:	Church
G.2.4	Qualification (MD.....)	MD MBBS FRCP Ed Cert.Clin.Ed
G.2.5	Professional address:	
G.2.5	Institution name	Chester le Street Community Hospital
G.2.5	Institution department	Stroke
G.2.5.1	Street address	Front Street
G.2.5.2	Town/city	Chester le Street
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G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
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G.2.3	Family name:	Lawrence
G.2.4	Qualification (MD.....)	BSc MBBS MD FRCP
G.2.5	Professional address:	
G.2.5	Institution name	Croydon Healthcare NHS Trust / Croydon University Hospital
G.2.5	Institution department	Stroke
G.2.5.1	Street address	530 London Rd
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G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
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G.2.3	Family name:	Cassidy
G.2.4	Qualification (MD.....)	MB BCH BAO FRCP EDIN FRCP
G.2.5	Professional address:	
G.2.5	Institution name	Gateshead Health NHS Foundation Trust / Queen Elizabeth Hospital
G.2.5	Institution department	Stroke Research Team
G.2.5.1	Street address	Queen Elizabeth Avenue, Sheriff Hill
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G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
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G.2.3	Family name:	Roffe
G.2.4	Qualification (MD.....)	MD
G.2.5	Professional address:	
G.2.5	Institution name	University Hospital of North Staffordshire NHS TrustRoyal Infirmary
G.2.5	Institution department	Stroke Research office
G.2.5.1	Street address	North Staffordshire Royal Infirmary, Prince's Rd
G.2.5.2	Town/city	Hartshill, Stoke on Trent
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G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
G.2.1	Given name:	Anthony
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G.2.3	Family name:	Oke
G.2.4	Qualification (MD.....)	MBChB FRCP MPH
G.2.5	Professional address:	
G.2.5	Institution name	Cannock Chase Hospital
G.2.5	Institution department	Stroke
G.2.5.1	Street address	Brunswick Road
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G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
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G.2.3	Family name:	Farag
G.2.4	Qualification (MD.....)	MBBCh MRCP FRCP
G.2.5	Professional address:	
G.2.5	Institution name	West Hertfordshire Hospitals NHS Trust / Watford General Hospital
G.2.5	Institution department	Stroke Unit
G.2.5.1	Street address	Vicarage Road
G.2.5.2	Town/city	Watford
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G.2.5.4	Country	United Kingdom
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G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
G.2.1	Given name:	Elio
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G.2.3	Family name:	Giallombardo
G.2.4	Qualification (MD.....)	CSST FRCP
G.2.5	Professional address:	
G.2.5	Institution name	Basingstoke & North Hampshire Hospital NHS Foundation Trust
G.2.5	Institution department	Stroke
G.2.5.1	Street address	Aldermaston Road
G.2.5.2	Town/city	Basingstoke
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G.2.5.4	Country	United Kingdom
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G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
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G.2.3	Family name:	Choy
G.2.4	Qualification (MD.....)	BMedSci MBBS MRCP MSc
G.2.5	Professional address:	
G.2.5	Institution name	Epsom & St Helier University Hospitals NHS Trust / Kingston Hospital NHS Trust
G.2.5	Institution department	Stroke
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G.2.5.2	Town/city	Kingston upon Thames
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G.2 PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)		
G.2.1	Given name:	John
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G.2.3	Family name:	Coyle
G.2.4	Qualification (MD.....)	MBBCh FRCP
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G.2.5	Institution name	York Hospital
G.2.5	Institution department	Acute Stroke Unit
G.2.5.1	Street address	Wiggington Road
G.2.5.2	Town/city	York
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G.2 PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)		
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G.2.3	Family name:	Shaw
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G.2.5	Professional address:	
G.2.5	Institution name	Royal United Hospital NHS Trust
G.2.5	Institution department	Stroke
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G.2.3	Family name:	Findlay
G.2.4	Qualification (MD.....)	BSc (Hons) BM FRCP
G.2.5	Professional address:	
G.2.5	Institution name	NHS Highland / Raigmore Hospital
G.2.5	Institution department	Stroke
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G.2 PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)		
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G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Sein
G.2.4	Qualification (MD.....)	MBBS MRCP
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G.2.5	Institution name	Macclesfield District General Hospital
G.2.5	Institution department	Stroke
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G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
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G.2.4	Qualification (MD.....)	MBBS FRCP
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G.2.5	Institution name	Epsom & St Helier University Hospitals NHS Foundation Trust
G.2.5	Institution department	Stroke / HCE
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G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
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G.2.5	Institution name	Birmingham Heartlands Hospital
G.2.5	Institution department	Stroke
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G.2.5	Professional address:	

G.2.5	Institution name	Gloucestershire Royal Hospital
G.2.5	Institution department	Stroke Service
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G.2.4	Qualification (MD.....)	MBChB MSc
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G.2.5	Institution name	Selly Oak Hospital
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G.2.4	Qualification (MD.....)	BSc MBChB MMed FRCPI FRCP
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G.2.5	Institution department	Stroke
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G.2.5	Institution name	Mount Gould Hospital(Plymouth Teaching PCT)
G.2.5	Institution department	Stroke Rehab
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G.2.3	Family name:	Price
G.2.4	Qualification (MD.....)	MD FRCP
G.2.5	Professional address:	
G.2.5	Institution name	Northumbria Healthcare NHS Foundation Trust / North Tyneside General Hospital
G.2.5	Institution department	Stroke Dept
G.2.5.1	Street address	Rake Lane
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G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
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G.2.3	Family name:	Whitehead
G.2.4	Qualification (MD.....)	MRCP(UK)
G.2.5	Professional address:	
G.2.5	Institution name	NHS Lanarkshire, Wishaw General Hospital
G.2.5	Institution department	Stroke Dept
G.2.5.1	Street address	50 Netherton St
G.2.5.2	Town/city	Wishaw
G.2.5.3	Post code	ML2 0DP
G.2.5.4	Country	United Kingdom
G.2.6	Telephone number:	+44 01698 366164
G.2.7	Fax number:	+44 01698 366099
G.2.8	E-mail:	martin.whitehead@lanarkshire.scot.nhs.uk

G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
G.2.1	Given name:	Suzanne
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Ragab
G.2.4	Qualification (MD.....)	MPhil FRCP BM
G.2.5	Professional address:	
G.2.5	Institution name	Poole Hospital NHS Foundation Trust
G.2.5	Institution department	Geriatric & Stroke Medicine
G.2.5.1	Street address	Longfleet Road
G.2.5.2	Town/city	Poole
G.2.5.3	Post code	BH15 2NG
G.2.5.4	Country	United Kingdom
G.2.6	Telephone number:	+44 01202 448767
G.2.7	Fax number:	+44 01202 442993
G.2.8	E-mail:	suzanne.ragab@poole.nhs.uk

G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
G.2.1	Given name:	Sanjeevikumar
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Meenakshisundaram
G.2.4	Qualification (MD.....)	MBBS MRCP
G.2.5	Professional address:	
G.2.5	Institution name	St Helens & Knowsley Teaching Hospitals NHS Trust / Whiston Hospital
G.2.5	Institution department	Stroke / Medical Directorate
G.2.5.1	Street address	Warrington Road
G.2.5.2	Town/city	Prescot
G.2.5.3	Post code	L35 5DR
G.2.5.4	Country	United Kingdom
G.2.6	Telephone number:	+44 0151 4301224
G.2.7	Fax number:	+44 0151 4301224
G.2.8	E-mail:	sanjeevikumar.meenakshisundaram@sthk.nhs.uk

G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
G.2.1	Given name:	Ramachandran
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Sivakumar
G.2.4	Qualification (MD.....)	MBBS MRCP(UK) MD
G.2.5	Professional address:	
G.2.5	Institution name	Colchester Hospital University NHS Foundation Trust
G.2.5	Institution department	Stroke
G.2.5.1	Street address	Colchester General Hospital, Turner Rd
G.2.5.2	Town/city	Colchester
G.2.5.3	Post code	CO4 5JL
G.2.5.4	Country	United Kingdom
G.2.6	Telephone number:	+44 01206 742271
G.2.7	Fax number:	+44 01206 6742270
G.2.8	E-mail:	ramachandran.sivakumar@colchesterhospital.nhs.uk

G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
G.2.1	Given name:	Don
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Sims
G.2.4	Qualification (MD.....)	MBChB MSc
G.2.5	Professional address:	
G.2.5	Institution name	Moseley Hall Hosital (Sth Birmingham PCT)
G.2.5	Institution department	Stroke Rehab
G.2.5.1	Street address	181 Alcester Rd
G.2.5.2	Town/city	Birmingham
G.2.5.3	Post code	B13 8JL
G.2.5.4	Country	United Kingdom
G.2.6	Telephone number:	+44 0121 6278482
G.2.7	Fax number:	+44 0121 6278282
G.2.8	E-mail:	don.sims@uhb.nhs.uk

G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
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G.2.1	Given name:	Neil
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Baldwin
G.2.4	Qualification (MD.....)	BMedSci(Hons)BMBS MRCP(UK) MRCP(Lond)
G.2.5	Professional address:	
G.2.5	Institution name	North Bristol NHS Trust
G.2.5	Institution department	Frenchay Day Hospital
G.2.5.1	Street address	Frenchay Park Rd
G.2.5.2	Town/city	Bristol
G.2.5.3	Post code	BS16 1LE
G.2.5.4	Country	United Kingdom
G.2.6	Telephone number:	+44 0117 3406636
G.2.7	Fax number:	+44 0117 3406401
G.2.8	E-mail:	neil.baldwin@nbt.nhs.uk

G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
G.2.1	Given name:	Udayaraz
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Umasankar
G.2.4	Qualification (MD.....)	MBBS, MRCP, LAB ECFMG
G.2.5	Professional address:	
G.2.5	Institution name	Basildon & Thurrock University Hosp. NHS Trust
G.2.5	Institution department	Basildon University Hospital
G.2.5.1	Street address	Nethermayne
G.2.5.2	Town/city	Basildon
G.2.5.3	Post code	SS16 5NL
G.2.5.4	Country	United Kingdom
G.2.6	Telephone number:	+44 01268 524900
G.2.7	Fax number:	+44 01268 593085
G.2.8	E-mail:	Udayaraj.umasankar@btuh.nhs.uk

G.2	PRINCIPAL INVESTIGATORS (for multicentre trial ; where necessary, use additional forms)	
G.2.1	Given name:	Julie
G.2.2	Middle name, if applicable:	
G.2.3	Family name:	Dovey
G.2.4	Qualification (MD.....)	
G.2.5	Professional address:	
G.2.5	Institution name	University Hospitals Bristol Foundation Trust / Bristol
G.2.5	Institution department	Royal Infirmary
G.2.5.1	Street address	Dept Stroke Medicine
G.2.5.2	Town/city	Queen's Building, Upper Maudlin St
G.2.5.3	Post code	Bristol
G.2.5.4	Country	BS2 87HW
G.2.6	Telephone number:	United Kingdom
G.2.7	Fax number:	+44 01179 282315
G.2.8	E-mail:	+44 01179 284003
		julie.dovey@uhbristol.nhs.uk

G.3	CENTRAL TECHNICAL FACILITIES TO BE USED IN THE CONDUCT OF THE TRIAL	
	Laboratory or other technical facility, in which the measurement or assessment of the main evaluation criteria are centralised (repeat as needed for multiple organisations).	
G.3.1	Name of organisation:	University of Nottingham
G.3.2	Department	Cardiovascular Medicine

G.3.3	Name of contact person:	
G.3.3.1	Given name	Stanley
G.3.3.2	Middle name	
G.3.3.3	Family name	Heptinstall
G.3.4	Address:	
G.3.4.1	Street address	Cardiovascular Medicine, QMC
G.3.4.2	Town/city	Nottingham
G.3.4.3	Post code	NG7 1DR
G.3.4.4	Country	United Kingdom
G.3.5	Telephone number:	+44 0115 8231013
G.3.6	Fax number:	+44 0115 8231017
G.3.7	E-mail:	stan.heptinstall@nottingham.ac.uk
G.3.8	Enter the details of any duties subcontracted to this central technical facility in this trial	
G.3.8.1	Routine clinical pathology testing	No •
G.3.8.2	Clinical chemistry	No •
G.3.8.3	Clinical haematology	No •
G.3.8.4	Clinical microbiology	No •
G.3.8.5	Histopathology	No •
G.3.8.6	Serology/ endocrinology	No •
G.3.8.7	Analytical chemistry	No •
G.3.8.8	ECG analysis/ review	No •
G.3.8.9	Medical image analysis/ review - X-ray, MRI, ultrasound, etc.	No •
G.3.8.10	Primary/ surrogate endpoint test	Yes •
G.3.8.11	Other Duties subcontracted?	No •
G.3.8.11.1	If 'Yes', specify the other duties	

G.3	CENTRAL TECHNICAL FACILITIES TO BE USED IN THE CONDUCT OF THE TRIAL	
	Laboratory or other technical facility, in which the measurement or assessment of the main evaluation criteria are centralised (repeat as needed for multiple organisations).	
G.3.1	Name of organisation:	St George's University of London
G.3.2	Department	Clinical Neuroscience
G.3.3	Name of contact person:	
G.3.3.1	Given name	Hugh
G.3.3.2	Middle name	
G.3.3.3	Family name	Markus
G.3.4	Address:	
G.3.4.1	Street address	Cranmer Terrace
G.3.4.2	Town/city	London
G.3.4.3	Post code	SW17 ORE
G.3.4.4	Country	United Kingdom
G.3.5	Telephone number:	+44 0208 7252735
G.3.6	Fax number:	+44 0208 87252950
G.3.7	E-mail:	hmarkus@sgul.ac.uk
G.3.8	Enter the details of any duties subcontracted to this central technical facility in this trial	
G.3.8.1	Routine clinical pathology testing	No •
G.3.8.2	Clinical chemistry	No •
G.3.8.3	Clinical haematology	No •
G.3.8.4	Clinical microbiology	No •
G.3.8.5	Histopathology	No •
G.3.8.6	Serology/ endocrinology	No •
G.3.8.7	Analytical chemistry	No •
G.3.8.8	ECG analysis/ review	No •
G.3.8.9	Medical image analysis/ review - X-ray, MRI, ultrasound, etc.	No •
G.3.8.10	Primary/ surrogate endpoint test	Yes •
G.3.8.11	Other Duties subcontracted?	No •
G.3.8.11.1	If 'Yes', specify the other duties	

G.4 NETWORKS TO BE INVOLVED IN THE TRIAL (e.g. Paediatric Networks involved in the trial)	
G.4.1	Name of organisation:
G.4.2	Name of contact person:
G.4.2.1	Given name
G.4.2.2	Middle name
G.4.2.3	Family name
G.4.3	Address:
G.4.3.1	Street address
G.4.3.2	Town/city
G.4.3.3	Post code
G.4.3.4	Country
G.4.4	Telephone number:
G.4.5	Fax number:
G.4.6	E-mail:
G.4.7	Activities carried out by the network:

G.5 ORGANISATIONS TO WHOM THE SPONSOR HAS TRANSFERRED TRIAL RELATED DUTIES AND FUNCTIONS	
G.5.1	Has the sponsor transferred any major or all the sponsor's trial related duties and functions to another organisation or third party? Yes •
Repeat as necessary for multiple organisations:	
G.5.1.1	Organisation name: University of Nottingham
G.5.1.2	Organisation department: Stroke Medicine
G.5.1.3	Name of contact person :
G.5.1.3.1	Given name: Philip
G.5.1.3.2	Middle name
G.5.1.3.3	Family name: Bath
G.5.1.4	Address:
G.5.1.4.1	Street address: CSB, Nottingham City Hospital
G.5.1.4.2	Town/city: Nottingham
G.5.1.4.3	Post code: NG5 1PB
G.5.1.4.4	Country: United Kingdom
G.5.1.5	Telephone number: +44 0115 8231765
G.5.1.6	Fax number: +44 0115 8231767
G.5.1.7	E-mail: philip.bath@nottingham.ac.uk
G.5.1.8	All tasks of the sponsor: No •
G.5.1.9	Monitoring: Yes •
G.5.1.10	Regulatory (e.g. preparation of applications to CA and ethics committee): Yes •
G.5.1.11	Investigator recruitment: Yes •
G.5.1.12	IVRS ³⁰ – treatment randomisation: Yes •
G.5.1.13	Data management: Yes •
G.5.1.14	E-data capture: Yes •
G.5.1.15	SUSAR reporting: Yes •
G.5.1.16	Quality assurance auditing: Yes •
G.5.1.17	Statistical analysis: Yes •
G.5.1.18	Medical writing: Yes •
G.5.1.19	Other duties subcontracted?: No •
G.5.1.19.1	If 'Yes' to other, please specify:

H. COMPETENT AUTHORITY / ETHICS COMMITTEE IN THE MEMBER STATE CONCERNED BY THIS REQUEST

H.1 TYPE OF APPLICATION		
If this application is addressed to the Competent Authority, please tick the Ethics Committee box and give information on the Ethics committee concerned. If this application is addressed to the Ethics Committee, please tick the Competent Authority box and give the information on the Competent Authority concerned.		
H.1.1	Competent Authority	No ●
H.1.2	Ethics Committee	Yes ●
H.2 INFORMATION ON ETHICS COMMITTEE		
H.2.1	Name:	South East Research Ethics Committee
H.2.2	Address	
H.2.2.1	Street address	South East Coast Strategic Health Authority, Preston Hall, Aylesford,
H.2.2.2	Town/city	Kent
H.2.2.3	Post code	ME20 7NJ
H.2.2.4	Country	United Kingdom
H.2.3	Date of submission:	2009-05-22
H.3 OPINION		
H.3.1	To be requested	Yes ●
H.3.2	Pending	No ●
H.3.3	Given	No ●
	If 'Given', specify:	
H.3.3.1	Date of opinion:	
H.3.3.2	Opinion favourable	No ●
H.3.3.3	Opinion not favourable	No ●
	If not favourable, give:	
H.3.3.3.1	The reasons	
H.3.3.3.2	The eventual anticipated date of resubmission:	

I. SIGNATURE OF THE APPLICANT IN THE MEMBER STATE

I.1	I hereby confirm that /confirm on behalf of the sponsor (delete which is not applicable) that: <ul style="list-style-type: none">• the information provided is complete;• the attached documents contain an accurate account of the information available;• the clinical trial will be conducted in accordance with the protocol; and• the clinical trial will be conducted, and SUSARs and result-related information will be reported, in accordance with the applicable legislation.
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I.2	APPLICANT OF THE REQUEST FOR THE COMPETENT AUTHORITY (as stated in section C.1):
I.2.1	Date:
I.2.2	Signature ³¹ :
I.2.3	Print name:

I.3	APPLICANT OF THE REQUEST FOR THE ETHICS COMMITTEE (as stated in section C.2):
I.3.1	Date:
I.3.2	Signature ³² :
I.3.3	Print name:

ENDNOTES

- ¹ Any translation of the protocol should be assigned the same date and version as those in the original document.
- ² International Standard Randomised Controlled Trial Number. Sponsors may wish to use an International Standardised Random Controlled Trial Number (ISRCTN) to identify their trial in addition to the EudraCT number; for instance if their trial is part of a multinational trial with sites outside the Community. They can obtain the number and guidance from the Current Controlled Trials website <http://www.controlled-trials.com/isrctn> to which there is a link from the EudraCT database website <http://eudract.ema.europa.eu>. When available they should provide it in Section A.6 of the application form.
- ³ US National Clinical Trial (NCT) Numbers required on the FDA clinical trial application form.
- ⁴ For a resubmission following previous withdrawal of an application or unfavourable opinion of an ethics committee, or previous withdrawal of an application or refusal of a request by the competent authority, enter a letter in the sequence, A for first resubmission, B for second, C for third et seq.
- ⁵ In accordance with Article 19 of Directive 2001/20/EC.
- ⁶ The contact point should give functional information rather than details of one "person", in order to avoid the need for update and maintenance of these contact details.
- ⁷ This requires a EudraLink account. (See <https://eudract.ema.europa.eu/document.html> for details)
- ⁸ According to national legislation.
- ⁹ Available from the Summary of Product Characteristics (SmPC)
- ¹⁰ According to the Community register on orphan medicinal products (Regulation (EC) n° 141/2000): <http://ec.europa.eu/enterprise/pharmaceuticals/register/index.htm>
- ¹¹ Committee for Medicinal Products for Human Use of the European Medicines Agency
- ¹² To be provided only when there is No trade name. This is the name routinely used by a sponsor to identify the IMP in the CT documentation (protocol, IB...).
- ¹³ To be provided only when there is No trade name. This is a code designated by the sponsor which represents the name routinely used by the sponsor to identify the product in the CT documentation. For example, a code may be used for combinations of drugs or drugs and devices.
- ¹⁴ Available from the Summary of Product Characteristics (SmPC).
- ¹⁵ Chemical Abstracts Service.
- ¹⁶ Complete also section D.4 Cell therapy as defined in Annex 1 part IV of Directive 2001/83/EC as amended.
- ¹⁷ Complete also section D.5 Gene Therapy as defined in Annex 1 part IV of Directive 2001/83/EC as amended.
- ¹⁸ Complete also section D.6 - Tissue Engineered Product as defined in Article 2(1)(b) of Regulation 1394/2007/EC.
- ¹⁹ Complete also section D.7
- ²⁰ The mode of action should briefly describe the chemical, biochemical, immunological or biological means the IMP uses to effect its pharmaceutical action.
- ²¹ Guideline on strategies to identify and mitigate risks for first-in-human clinical trials with investigational medicinal products. EMEA/CHMP/SWP/28367/2007 19 July 2007
- ²² In accordance with paragraph 38 of Annex 13 of Volume 4 of the Rules Governing Medical Products in the European Union.
- ²³ In the case of healthy volunteer trials, the intended indication for the product under development should be provided.
- ²⁴ Applicants are encouraged to provide the MedDRA lower level term if applicable and classification code. These can be accessed from the EMEA EudraCT website (<http://eudract.ema.europa.eu/>).
- ²⁵ Points to consider on the calculation and reporting of the prevalence of a condition for Orphan drug designation: COM/436/01 (<http://www.ema.europa.eu/htms/human/orphans/intro.htm>).
- ²⁶ The protocol will usually identify a single primary end point but there may be a co-primary end point in some cases and/or a number of secondary end points.
- ²⁷ The descriptions of the trial types provided are those recommended in preference to Phases. See page 5 of Community guideline CPMP/ICH/291/95. The development of a new indication after initial approval of a medicine should be considered as a new development plan.
- ²⁸ From the first inclusion until the last visit of the last subject.
- ²⁹ These numbers will be initial estimates. Applicants will not be required to update this information nor do they constitute an authorisation or restriction on the inclusion of these numbers of patients in the trial. The numbers of subjects whose inclusion is authorised are those set out in the authorised version of the protocol, or subsequent authorised amendments.
- ³⁰ Interactive Voice Response System: commonly used for randomisation of treatment and controlling the shipment of stock of product.
- ³¹ On an application to the Competent Authority only, the applicant to the Competent Authority needs to sign.

³² On an application to the Ethics Committee only, the applicant to the Ethics Committee needs to sign.

TARDIS STUDY

SUBSTANTIAL PROTOCOL AMENDMENT

SUMMARY OF CHANGES VERSION 1.3

In the text below, protocol and information sheet changes, having implications for research design, conduct or participant safety, have been listed. Additional minor changes to text and formatting made to bring protocol, information sheets and consent forms up-to-date are not described below but can be viewed in the 'marked' version of the documents. The trial summary has been updated to reflect the changes in the protocol.

PROTOCOL VERSION 1.3: SUMMARY OF CHANGES

1. Title

Existing Protocol

Safety and efficacy of clopidogrel when added to aspirin and dipyridamole in high risk patients with recent ischaemic stroke: a randomised controlled trial

Revised Protocol

Safety and efficacy of intensive versus guideline antiplatelet therapy in high risk patients with recent ischaemic stroke and transient ischaemic attack: a randomised controlled trial

2. Trial Background Information and Rationale

The trial background information has been revised and updated to reflect the current evidence. These changes are highlighted in the document with tracked changes.

3. DETAILS OF INVESTIGATIONAL MEDICINAL PRODUCTS

Aspirin and dipyridamole will also be considered as investigational medicinal products in addition to clopidogrel.

Revised Protocol

Description

Aspirin (Asp)

International Non-Proprietary Name (INN): Aspirin

Chemical Abstracts Series (CAS) number: 50-78-2

Dose: Loading dose 300mg, then 75mg od.

Route: Enteral (including via nasogastric tube – dispersible or crushed tablets can be used) or rectal route.

For chemical and pharmacological properties, see summary of medical product characteristics (SmPC) at <http://www.medicines.org.uk/emc/>.

The IMP is defined by active substance only, so all authorised brands may be used.

Dipyridamole (Dip)

INN: Dipyridamole

CAS number: 58-32-2

Dose: 200mg modified release (MR), bd. Dysphagic patients with enteral access will take dipyridamole suspension 75mg tds. Patients with a headache from dipyridamole will have the dose weaned up from daily MR 200mg or standard release 75mg od to MR 200mg bd.

Fixed dose combinations of A and D can also be used, e.g. Asasantin Retard (Aspirin 25mg, Dipyridamole 200mg MR, bd)

Route: Enteral (including via nasogastric tube).

For chemical and pharmacological properties SmPC at <http://www.medicines.org.uk/emc/>.

The IMP is defined by active substance only, so all authorised brands in the UK can be used.

4. Packaging and labelling

This section has been revised and updated

Existing Protocol

Standard pharmacy supplies should be used as the IMP (clopidogrel) has marketing authorisation for use in stroke. Separate labelling and packaging details are not required but local sites can overlabel as they feel appropriate, in which case accountability logs for clopidogrel (batch numbers and expiry dates) should be recorded. Aspirin and dipyridamole are not IMPs as they are standard treatment for stroke and TIA; accountability logs for them are therefore unnecessary.

Revised Protocol

Standard pharmacy supplies should be used as all IMPs have marketing authorisation and are to be used in accordance with such authorisation. Hospitals/pharmacies should choose their own supplier for the IMPs and should be packaged according to local policy. All IMPs for the TARDIS trial should be labeled separately and pharmacies at the recruiting centre must have a written procedure in place for dispensing trial medications. The information on the label should include the trial name, EudraCT number, description of contents, batch number, expiry date, and quantity. There should be space for insertion of the trial number, name of the participant and the date of dispensing on the label (see appendix K). Under exceptional circumstances (e.g. out of hours) where labeled IMPs are not available, trial sites may choose to use ward stock without separate labeling if agreed locally and approved by the pharmacy.

5. Storage dispensing and return

This section has been revised and updated.

Existing Protocol

Standard pharmacy supplies will be prescribed and used.

Revised protocol

The IMPs must be stored in a secure location at room temperature (20°C to 25°C) with excursions permitted within 15°C to 30°C. Depending on local arrangement, this may be at the local pharmacy, the research department or the ward. Following recruitment and randomisation into the trial, the study treatment

should be prescribed on the drug chart and the IMPs dispensed by the principal investigator/qualified designee. An accountability log for all IMPs should be maintained by the pharmacy and/or the research team and should include the following information: hospital number, participant initials, trial number, date dispensed, brand manufacturer, batch number, expiry date, quantity dispensed, quantity returned and initials of personnel who dispense and check the log. This should be completed for every participant who is randomised into the study. Accountability logs must be available for inspection during trial monitoring and/or audit and open to regulatory authorities inspection at any time. A sample label is provided (Appendix K).

6. Known side effects

Side effects for aspirin and dipyridamole have now been added.

Revised protocol

Known Side Effects

Aspirin

Adverse reactions

- i. Bleeding: Aspirin prolongs bleeding time, and bleeding disorders, such as epistaxis, haematuria, purpura, ecchymoses, haemoptysis, gastrointestinal bleeding, haematoma and cerebral haemorrhage have been reported.
- ii. Blood and lymphatic system disorders - anaemia, haemolytic anaemia, hypoprothrombinaemia, thrombocytopenia, aplastic anaemia, pancytopenia, prolonged bleeding time, occult blood loss, elevated transaminase levels, agranulocytosis.
- iii. Gastrointestinal disorders - gastrointestinal bleeding, erosions, perforations or ulceration, which can occasionally be major (may develop bloody or black tarry stools, severe stomach pain and vomiting blood), gastrointestinal irritation (mild stomach pain, heartburn, vomiting and nausea). Fatalities have occurred.
- iv. Hepatic disorders - hepatitis (particularly in patients with SLE or connective tissue disease)
- v. Renal and urinary disorders – disturbances of renal function
- vi. Ear and labyrinth disorders - tinnitus.
- vii. Hypersensitivity reactions - rhinitis, urticaria, purpura, Stevens-Johnson syndrome, angio-oedema, asthma, worsening of asthma, bronchospasm.

Interaction with other medicinal products:

- i. Salicylates may enhance the effect of oral hypoglycaemic agents, phenytoin and sodium valproate.
- ii. They inhibit the uricosuric effect of probenecid and may increase the toxicity of sulphonamides.
- iii. Aspirin may potentiate the effect of heparin and increases the risk of bleeding with oral anticoagulants, antiplatelet agents and fibrinolytics.
- iv. The risk of gastrointestinal ulceration and bleeding may be increased when aspirin and corticosteroids are co-administered.
- v. Concurrent use of aspirin and other Non Steroidal Anti Inflammatory Drugs (NSAID) should be avoided. Use of two or more NSAID preparations increases the risk of serious gastrointestinal haemorrhage.
- vi. Concurrent administration of carbonic anhydrase inhibitors such as acetazolamide and salicylates may result in severe acidosis and increased central nervous system toxicity.
- vii. Alcohol may enhance the gastro-intestinal side effects of aspirin.

- viii. Patients using enteric-coated aspirin should be advised against ingesting antacids simultaneously to avoid premature drug release.
- ix. Selective Serotonin Reuptake Inhibitors (SSRI) may increase risk of gastrointestinal bleeding if coadministered.

Dipyridamole

Adverse reactions at therapeutic doses are usually mild.

- i. Bleeding: In very rare cases, increased bleeding during or after surgery has been observed.
- ii. Blood and lymphatic system disorders: Isolated cases of thrombocytopenia have been reported in conjunction with treatment with Dipyridamole.
- iii. Gastrointestinal disorders: Vomiting, diarrhoea and symptoms such as nausea, dyspepsia. These tend to occur early after initiating treatment and may disappear with continued treatment.
- iv. Cardiovascular: As a result of its vasodilating properties, dipyridamole may cause hypotension, hot flushes and tachycardia. Worsening of the symptoms of coronary heart disease such as angina and arrhythmias may occur.
- v. Central Nervous System Disorders: Dizziness, headache and myalgia may occur early after initiating treatment and may disappear with continued treatment.
- vi. Hypersensitivity reactions such as rash, urticaria, severe bronchospasm and angio-oedema have been reported.

Interaction with other medicinal products:

- i. Dipyridamole increases the plasma levels and cardiovascular effects of adenosine. Adjustment of adenosine dosage should therefore be considered if use with dipyridamole is unavoidable.
- ii. Dipyridamole may increase the hypotensive effect of blood pressure lowering drugs.
- iii. Dipyridamole may counteract the anticholinesterase effect of cholinesterase inhibitors thereby potentially aggravating myasthenia gravis.

TRIAL PURPOSE AND OBJECTIVES

7. PURPOSE

Existing Protocol

To perform a randomised trial assessing the efficacy, safety and tolerability of adding Clopidogrel to Aspirin and Dipyridamole in patients with recent ischaemic stroke or TIA and who are at high risk of recurrence. The study will comprise a start-up phase of 350 patients to then expand into a larger trial of 5000 patients assessing the efficacy, safety and health economics of this approach.

Revised Protocol

To perform a randomised trial assessing the efficacy, safety and tolerability of intensive antiplatelet therapy (Asp+Dip+Clop) versus guideline antiplatelet therapy (Asp+Dip or Clop) in patients with recent ischaemic stroke or TIA and who are at high risk of recurrence.

8. PRIMARY OBJECTIVE

Existing Protocol

To assess ordinal stroke severity at 90 days after short-term administration (1 month) of triple antiplatelet therapy (aspirin/clopidogrel/dipyridamole) versus standard dual therapy (aspirin/dipyridamole) in patients with very recent ischaemic stroke or TIA.

Revised Protocol

To assess ordinal stroke severity at 90 days after short-term administration (1 month) of intensive antiplatelet therapy versus guideline therapy in patients with very recent ischaemic stroke or TIA.

9. SECONDARY OBJECTIVE

Existing Protocol

1. To assess the safety of short-term administration (1 month) of triple antiplatelet therapy (aspirin/clopidogrel/dipyridamole) versus standard dual therapy (aspirin/dipyridamole) in patients with very recent ischaemic stroke or TIA.
2. To further assess, in high risk patients with stroke/TIA, whether the addition of clopidogrel to aspirin/dipyridamole:
 - i. Is feasible to administer acutely and tolerable to take for 1 month,
 - ii. Is superior in respect of surrogate markers such as emboli (with transcranial doppler) and platelet function.
 - iii. Improves functional outcome
3. To assess whether ordinal outcomes are superior to binary events

Revised Protocol

1. To assess the safety of short-term administration (1 month) of intensive antiplatelet therapy versus guideline therapy in patients with very recent ischaemic stroke or TIA.
2. To further assess, in high risk patients with stroke/TIA, whether:
 - iv. it is feasible to administer intensive therapy acutely and is tolerable to take for 1 month,
 - v. intensive therapy is superior in respect of surrogate markers such as platelet function.
 - vi. intensive therapy improves functional outcome

TRIAL DESIGN- TRIAL CONFIGURATION-SETTING

10. Design

Existing Protocol

Multicentre parallel group prospective randomised open-label blinded-endpoint controlled trial.

Revised Protocol

International, collaborative, multicentre, parallel group, prospective, randomised open-label, blinded-endpoint, controlled, Phase III trial.

11. Setting:

Existing Protocol

The trial comes from members of the SRN Prevention Clinical Study Group (PB, SH, HM, GV). Initially, 350 patients will be recruited from the UK Stroke Research Network. Each of the participating sites runs a stroke service with sufficient stroke/TIA patients to allow the planned recruitment rate (20+ centres x 0.6 patient/month [typical rate for academic stroke trials] x 12 months x 2.5 years = 360 patients). Expansion overseas and within the UK will occur for the main phase.

Revised Protocol

Initially, ~1000 patients will be recruited from the UK National Institute of Health Research (NIHR) Stroke Research Network (SRN) to the start-up phase. In the main phase, a further 3,100 participants from UK and overseas hospital-based stroke/TIA services will be recruited; UK participants (~2000) will be recruited from SRN sites (the trial is already adopted) including 55 sites who have been started-up and are recruiting in England and Scotland. These sites have dedicated SRN nurses to facilitate recruitment and follow-up. Philip Bath will run the trial from the University of Nottingham Stroke Trials Unit.

12. Trial Duration

Existing Protocol

The start-up phase will run for 3 years; months 0-3: development of trial systems (based on the internet site/database used in the ongoing ENOS trial) and training of LRN nurses at recruiting centres; months 4-31: patient recruitment; months 32-34: follow-up of the last recruited patients and data cleaning; months 35-36: analysis and report writing. There will then be a seamless transition from start-up to the main phase of the trial of the same design (as done with funding from BUPA Foundation to MRC for ENOS) so that recruitment does not stop. The main phase will last for an additional 5 years. Separate permission for funding from the appropriate bodies (e.g.HTA) will be sought for the second phase.

Revised Protocol

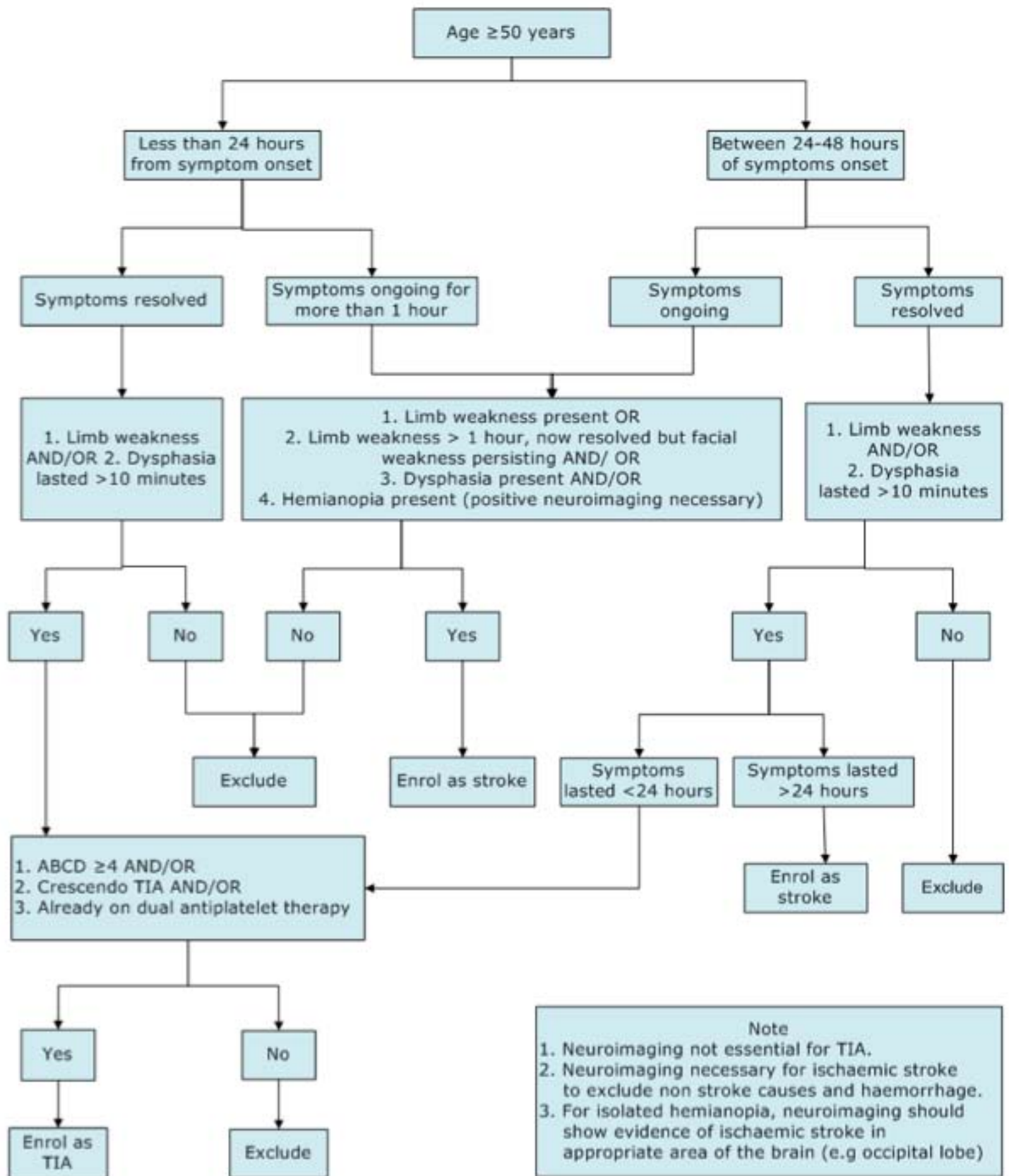
The start-up phase will run for ~4 years. If the start-up phase shows acceptable safety, there will then be a seamless transition to the main phase of the trial of the same design so that recruitment does not stop (**tables 2a, 2b**).

The main phase will recruit in the order of ~3,100 patients (depending on the rate and distribution of ordinal events) and will last an additional 5 years. Separate permission for funding from the appropriate bodies (e.g. HTA) is being sought for the main phase.

If the trial shows that intensive antiplatelet therapy is superior to guideline therapy (taking account of the balance between reduced stroke/vascular events and potentially increased bleeding), intensive antiplatelet therapy could be introduced rapidly for stroke prevention with immediate benefit to high risk

patients; each component is available now and licensed for secondary prevention. As the patent for clopidogrel has ended, NHS implementation of positive results will be based on generic costs, which will improve uptake and health economics.

A decision to proceed onto the main phase will be dependent on regular safety analyses during the start-up phase (by the Data Monitoring Committee), a successful funding application for the main phase, and the results of ongoing trials of dual antiplatelet therapy e.g. SPS-3 (Asp+Clop vs Asp), and ARCH (Asp+Clop vs. warfarin).





Safety and efficacy of intensive versus guideline antiplatelet therapy in high risk patients with recent ischaemic stroke or transient ischaemic attack: a randomised controlled trial

Short title: Triple Antiplatelets for Reducing Dependency after Ischaemic Stroke.

Acronym: TARDIS

EudraCT number: 2007-006749-42

ISRCTN: ISRCTN47823388

REC reference: 08/H1102/112

Trial Sponsor: University of Nottingham

Sponsor reference: 31350 and 08093

Funding Source: British Heart Foundation (start-up phase)

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ABBREVIATIONS

Asp	Aspirin
ADR	Adverse Drug Reaction
AE	Adverse Event
BI	Barthel Index
Clopidogrel	Clopidogrel
CAS	Chemical Abstract Series
CI	Chief Investigator
CRF	Case Report Form
Dip	Dipyridamole
GCP	Good Clinical Practice
HTA	Health Technology Assessment
IDMC	Independent Data Monitoring Committee
ICF	Informed Consent Form
IMP	Investigational Medicinal Product
INN	International Non-Proprietary Name
LMWH	Low Molecular Weight Heparin
LRN	Local Research Network
MHRA	Medicines and Healthcare products Regulatory Agency
MI	Myocardial Infarction
MR	Modified Release
mRS	Modified Rankin Scale
NIHR	National Institute of Health Research
NIHSS	National Institute of Stroke Health Scale
NHS	National Health Service
NICE	National Institute for Health and Clinical Excellence
NSAID	Non Steroidal Anti Inflammatory Drugs
NSTEMI	Non ST Elevation Myocardial Infarction
PI	Principal Investigator at a local centre
PIS	Participant Information Sheet
REC	Research Ethics Committee
R&D	Research and Development
SmPC	Summary of Medical Product Characteristics
SAE	Serious Adverse Event
SAP	Statistical Analysis Plan
SAR	Serious Adverse Reaction
SPC	Summary of Product Characteristics
STEMI	ST Elevation Myocardial Infarction
SUSAR	Suspected Unexpected Serious Adverse Reaction
SRN	Stroke Research Network
TIA	Transient Ischaemic Attack
TMC	Trial Management Committee
TSC	Trial Steering Committee

SYNOPSIS

Title	Safety and efficacy of intensive versus guideline antiplatelet therapy in high risk patients with recent ischaemic stroke or transient ischaemic attack (TIA): a randomised controlled trial
Acronym	TARDIS
Short title	<u>T</u> riple <u>A</u> ntiplatelets for <u>R</u> educing <u>D</u> ependency after <u>I</u> schaemic <u>S</u> troke
Trial Summary	Recurrence is greatest immediately after stroke or TIA; existing prevention strategies (antithrombotic, lipid/blood pressure lowering, carotid endarterectomy) reduce, not abolish, further events. Dual antiplatelet therapy – aspirin & clopidogrel for coronary disease, aspirin & dipyridamole for stroke - is superior to aspirin monotherapy. Triple antiplatelet therapy reduces MI and death in patients with coronary disease. We have shown that it is feasible to give triple antiplatelet therapy (aspirin, clopidogrel, dipyridamole) to patients with ischemic stroke/TIA. We will assess the efficacy, safety, tolerability and feasibility of intensive (combined aspirin, dipyridamole and clopidogrel) versus guideline antiplatelet therapy (combined aspirin and dipyridamole or clopidogrel) given for 1 month in ~1000 patients (over ~4 years) with acute stroke/TIA (i.e. at high risk of recurrence) in the start-up phase of a large randomised controlled trial. This will seamlessly run into the main phase of the trial (total 4100 patients) over the next 5 years providing safety information from the start up phase allows. The primary outcome is ordinal stroke severity at 90 days. Secondary outcomes include safety, serious adverse events, vascular events, death and platelet function.
Chief Investigator	Professor Philip Bath
Primary Objective	The trial will assess ordinal stroke severity: 5-level ordinal stroke and TIA scale with stroke ordered by its severity using the modified Rankin Scale (mRS): fatal stroke / severe non-fatal stroke (mRS 2-5) / mild stroke (mRS 0,1) / TIA / no stroke-TIA, measured at 90 days.
Trial Design	International, collaborative, multicentre, parallel group prospective, randomised, open-label blinded-endpoint, controlled Phase III trial.
Setting	In the start-up phase, patients will be recruited from the UK Stroke Research Network Centres. Further expansion within the UK and overseas will occur in the main phase.
Sample size estimate	The start-up phase is sized to assess safety and will inform the sample size calculation for the main trial phase, which will assess the efficacy of intensive versus guideline therapy. Assuming the distribution in 5 level recurrent stroke/TIA outcome (stroke with mRS 6 =0.1%/ mRS 2-5 = 0.7%/ mRS 0-1 = 1.53%/ TIA = 3.57%), odds ratio of 0.68, alpha 5%, power 90%, losses to follow-up 2%, treatment crossovers 5% the total sample size for the whole

	study is 4100.
Number of participants	Start up: ~1000; main phase: 3100
Eligibility criteria	Adults at high risk of recurrent ischaemic stroke: <ol style="list-style-type: none"> 1. Acute high risk TIAs \leq48 hours of onset All TIAs must have limb weakness and/or dysphasia lasting at least 10 minutes. 2. Ischaemic, non cardioembolic stroke with limb weakness, dysphasia or hemianopia \leq48 hours of onset with neuroimaging to rule out alternative causes. 3. Meaningful consent, or consent from a relative, carer or legal representative if the patient is unable to give consent (e.g. in cases of dysphasia, confusion, or reduced conscious level).
Description of interventions	Intensive versus guideline antiplatelet therapy will be given for 28 to 30 days along with standard 'best care' (including lifestyle advice, BP and lipid lowering). Randomised patients will receive clopidogrel (loading dose 300 mg, then 75 mg daily), aspirin (loading dose 300 mg, then 75 mg daily), and dipyridamole (modified release 200 mg twice daily), or guideline antiplatelet therapy (aspirin and dipyridamole or clopidogrel, doses as above), .
Duration of study	8 years
Randomisation and blinding	Patients will be randomised through the trial website with stratification and minimisation. Outcome assessments are blinded.
Outcome measures	Primary: Ordinal stroke severity at 90 days. Secondary: Binary and ordinal outcomes of stroke, TIA, MI, acute coronary syndrome, composite vascular outcome, death. Also safety (ordinal bleeding events), tolerability and feasibility. Additional measures include laboratory measures (FBC and P-Selectin), clinical efficacy (NIHSS), function (mRS, BI), cognition (TICS), quality of life (EuroQoL, EQ-5D), mood (Zung), disposition, days at home and economic activity.
Statistical methods	Ordinal logistic regression on ordered categorical outcomes, binary logistic regression on binary outcomes, analysis of covariance (ANCOVA) on continuous data and Kaplan-Meier and Cox proportional hazards regression on time to event data. Analyses will be adjusted for randomisation/minimisation factors. Subgroup analyses will only be performed in the main trial phase

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1 TRIAL BACKGROUND INFORMATION AND RATIONALE

Stroke is devastating to patients, carers and society through high mortality (~1-in-3 patients by 1 year), morbidity (dependency in ~1-in-3 patients often needing long term care) and cost (6% of NHS spend). Both stroke incidence and prevalence will increase as the UK population ages. Following stroke or transient ischaemic attack (TIA), the risk of recurrence is high, especially immediately after the event (~10% over weeks) after which it falls (~40% by 5 years). Importantly, recurrent strokes are usually more severe than earlier events. The Government has emphasised stroke as a clinical 'marker' condition and has supported its research importance through funding the UK Stroke Research Network (PB is prevention Director, TR and HM are Local Research Network Directors for Trent and South-east respectively).

TIA ('mini stroke') is a sudden, focal neurologic deficit that lasts for less than 24 hours (typically 10 minutes to 1 hour), is presumed to be of vascular origin, and is confined to an area of the brain or eye perfused by a specific artery. (A tissue-based definition¹ is not used since MR scanning is not universally available around the world in the participating sites.) TIAs are important because they are a key risk factor for subsequent stroke. Patients presenting with specific TIA features are at particularly high risk of a subsequent stroke, as assessed by the ABCD² score:² age ≥ 60 years (1 point); blood pressure $\geq 140/90$ mmHg (1); clinical symptoms of unilateral weakness,(2) or speech impairment without weakness (1); duration ≥ 60 minutes,(2) or ≥ 10 minutes (1); and diabetes (1).² The score ranges from 0 to 7 and the risk of stroke over the next 90 days increases exponentially: score 0 - risk <1%, 1 - 2%, 2 - 4%, 3 - 4.5%, 4 - 8%, 5 - 12%, 6 - 17%, and 7 - 22%.² Other groups have now validated the score. An important caveat is that data for the training databases used to derive and validate the ABCD² score were collected up to 1998 and 2005 respectively so the absolute risk rates of stroke now are likely to be lower as enhanced secondary prophylaxis with antithrombotics, BP and lipid lowering are now standard practice.

The risk of recurrence can be reduced, but not abolished with life style changes, carotid endarterectomy (large artery stroke) and drug interventions: antihypertensives and statin therapy. While oral anticoagulants are established for cardioembolic stroke,³ other patients with ischaemia (the majority) need antiplatelets.⁴⁻⁵ These interventions are cost-effective. The archetypal antiplatelet, aspirin (inhibitor of cyclooxygenase), reduces recurrence (relative risk reduction, RRR) by 17% in patients with prior stroke or TIA.⁶ Clopidogrel (adenosine diphosphate [ADP] receptor antagonist) was slight more efficacious than aspirin in CAPRIE.⁷ Importantly, the absolute difference in efficacy between A and C was highest in patients with prior stroke or MI.⁸ Dipyridamole (inhibits red cell uptake of adenosine) reduced recurrence by 16% in comparison with placebo in ESPS II.⁹ Evidence now suggests that stroke prevention is dependent on the number of antiplatelets, e.g. aspirin and dipyridamole reduces events by 23% in comparison to aspirin (or dipyridamole) alone without increasing the risk of bleeding, as seen in ESPS II and ESPRIT.⁹⁻¹⁰ As with clopidogrel alone, the difference in efficacy between aspirin and dipyridamole versus aspirin alone was largest in patients with highest baseline risk.¹¹ Similarly, aspirin and dipyridamole was superior to aspirin in cardiac patients (CURE, CREDO)¹²⁻¹³ but not in CHARISMA,¹⁴ probably because the apparent benefit in those with prior stroke or MI (high risk of recurrence) was diluted by lack of efficacy in those with no previous vascular events (low risk). The risk of bleeding with aspirin and dipyridamole vs. aspirin was 30-40% higher in these 3 trials. The MATCH trial (aspirin and clopidogrel vs. clopidogrel) found that dual aspirin and clopidogrel also increased bleeding.¹⁵⁻¹⁶

On the basis of this information and taking account of the prices of branded clopidogrel and dipyridamole-ER (£37 and £10 per month respectively), NICE recommended in 2005 that

patients should take combined AD after ischaemic stroke or TIA (TA90). In late 2010, NICE updated its recommendation to aspirin and dipyridamole for TIA, and clopidogrel for ischaemic stroke (TA210), this taking account of the massive drop in price of clopidogrel (£3.40, as a generic, British National Formulary [BNF] 61) but lack of significant randomised data and license for clopidogrel in patients with TIA. Former and current guidelines have not recommended dual aspirin and clopidogrel because of increased bleeding.¹⁷⁻¹⁸ The preference for dual aspirin and dipyridamole or clopidogrel alone over aspirin alone is also recommended by the European Stroke Organisation in its 2008 guidelines (Bath was Co-Chair of the Prevention section).¹⁹ In contrast, the 2011 American Stroke Association secondary prevention guidelines still give equal recommendations for aspirin (50-325 mg daily) alone, dual aspirin and dipyridamole, and clopidogrel (75 mg daily) alone,²⁰ thereby ignoring the results of recent trials.^{7, 9-10, 21}

The above data for stroke reflect long-term prophylaxis, a very different situation from the situation immediately after an event when the risk of recurrence is much higher. Conventional acute antiplatelet therapy is based on aspirin alone for ischaemic stroke reflecting the results of the IST-1 and CAST megatrials²²⁻²³ but the effect size is small (absolute risk reduction ~1.1%); until recently the acute treatment of TIA had not been investigated. Since risk of recurrence falls quickly after stroke or TIA, intensive antiplatelet specific treatment is only likely to be needed for a short period so that the exposure-time to hazard (mainly bleeding) is limited. While clopidogrel based dual therapy has not proved effective/safe in long-term stroke prophylaxis, early and short-term dual therapy may be useful, at least after TIA/minor stroke, as suggested by trials (FASTER, EARLY, PRoFESS early²⁴⁻²⁶) and observational studies (EXPRESS, SOS²⁷⁻²⁸). In FASTER (n=392), 90 days of aspirin and clopidogrel (vs aspirin) showed a trend to reduced stroke by absolute 3.7% (not significant (NS)) and increased symptomatic intracerebral haemorrhage (sICH) by absolute 1% (NS) leading to a net absolute benefit of 2.7%²⁴. Similarly, EARLY (n=543, acute ischaemic stroke/TIA) found a trend to reduced vascular events at day 90 with aspirin and dipyridamole (vs aspirin, NS) but no effect on functional outcome,²⁵ a pattern of observations also seen with aspirin and dipyridamole (vs clopidogrel) in the PRoFESS early subgroup (n=1,360, mild acute ischaemic stroke).²⁶

In a meta-analysis of all trials comparing dual with mono antiplatelet therapy in acute patients with stroke or TIA (including CARESS, CHARISMA, CLAIR, FASTER, EARLY, ESPRIT, ESPS-2, MATCH and PRoFESS early^{9-10, 14-15, 24-26, 29-30}), acute dual therapy versus monotherapy within 3 days of ictus significantly reduced subsequent vascular events,²⁴ stroke (ischaemic and haemorrhagic, **figure 1**), and composite vascular events (trend) (Geeganage & Bath; submitted *Stroke*). No significant differences were seen for MI, sICH, major bleeding or death (but there were few events, **table 1**). No heterogeneity existed in any analysis suggesting that the composition of dual and mono therapy was not of primary importance. None of the trials were large enough (each <1,400) to show individual significant differences in stroke or vascular events. Importantly, the magnitude of effect appeared to decline with time from ictus so trials recruiting early have greater reductions in their point estimates (albeit non-significant because of small sample size) than those recruiting later: range of odds ratio for stroke, early, OR 0.51 to 0.71 (EARLY, FASTER, PRoFESS early); later, OR 0.83 to 2.44 (CHARISMA, MATCH).

	Stroke, MI, Vascular death	Stroke, TIA, ACS, all death	Stroke recurrence	MI	sICH	Major bleed	Death
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Odds Ratio	0.75	0.71	0.67	0.71	1.39	2.09	1.34
95% Confidence intervals	0.56-0.99	0.56-0.91	0.49-0.93	0.25-2.03	0.22-8.75	0.86-5.06	0.76-2.34

Table 1. Meta-analysis of 12 trials of dual vs mono antiplatelets in patients with acute ischaemic stroke/TIA. Data were obtained from trialists for patients recruited within 72 hours of ictus (Geeganage & Bath; submitted *Stroke*).

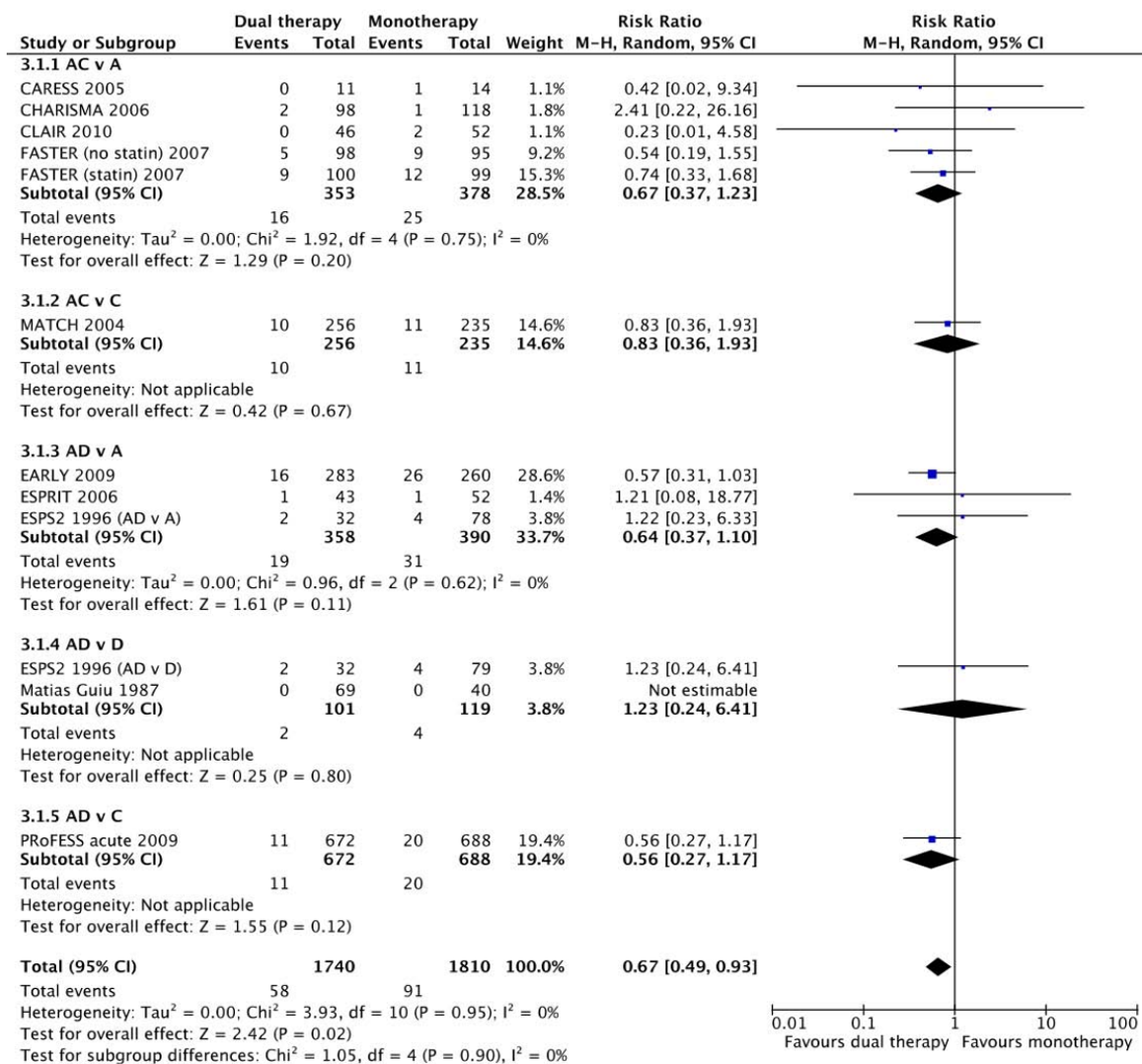


Figure 1. Meta-analysis of effect of dual versus mono antiplatelet therapy on stroke using data from 10 trials in patients with acute stroke or TIA. As compared with monotherapy, dual therapy reduced stroke: OR 0.67 (0.49-0.93). No heterogeneity was present suggesting that the composition of dual and mono therapy was less important than the number of antiplatelet agents (Geeganage and Bath; submitted *Stroke*).

Current stroke prevention is far from perfect: stroke is heterogeneous in type (ischaemic vs. haemorrhage; lacunar vs. cardioembolic vs. large artery), severity and outcome; treatments reduce, not abolish, events ('treatment failure'); and patients may be (relatively) insensitive to treatment ('treatment resistance', as identified for aspirin and clopidogrel³¹).

If aspirin and dipyridamole is superior to aspirin for long-term secondary prevention,^{9-10, 32} and aspirin and clopidogrel is probably superior to aspirin in acute minor stroke/TIA,^{24, 27} then triple antiplatelet therapy (aspirin+dipyridamole+clopidogrel) may be better still providing the risk of recurrence is high and bleeding does not become excessive. In this respect, the risk of bleeding when adding clopidogrel to aspirin and dipyridamole is likely to be similar to that when adding clopidogrel to aspirin since dual aspirin and dipyridamole does not increase bleeding over aspirin.⁹⁻¹⁰ We have performed a series of 'proof-of-concept' laboratory and clinical studies investigating this approach.³³⁻³⁷ In vitro studies found that triple therapy was most effective in inhibiting aggregation, platelet-leucocyte conjugation, and leucocyte activation.³³⁻³⁵ In multiway crossover phase I and II trials comparing short-term administration of mono dual, and triple antiplatelet platelet therapies, the combination of aspirin and clopidogrel, with or without dipyridamole, was most potent in inhibiting platelet function ex vivo in both normal volunteers (n=11) and patients with previous stroke/TIA (n=11).³⁶⁻³⁷

In the only parallel group trial of triple therapy in patients with stroke, triple therapy was feasible to administer (vs. aspirin, phase II trial, n=17) for up to 24 months.³⁸ [The comparator of aspirin was chosen since this was the UK standard of care at trial start. The trial was stopped early on publication of ESPRIT¹⁰ confirming the superiority of dual aspirin and dipyridamole over aspirin, i.e. it was unethical to continue patients on aspirin alone.] Predictably, there was a non-significant trend to increased bleeding with triple therapy vs aspirin. Although unintended, the patients were at low risk of recurrence (young/recruited months after the event/many lacunar strokes), a problem also seen in MATCH and CHARISMA.¹⁴⁻¹⁵ Future trials of triple antiplatelet therapy need to target patients at high risk of recurrence so that benefit is likely to outweigh hazard. We have also used chronic triple antiplatelet therapy in clinical practice in patients at high risk of recurrence, defined as recurrence on dual antiplatelet therapy.³⁹

Short-term randomised controlled trials of triple antiplatelet therapy have been reported in patients with acute coronary syndromes or to cover stent insertion (25 studies, 17,383 patients) . In our published meta-analysis and in comparison with dual antiplatelet therapy, GP IIb/IIIa based triple therapy reduced Myocardial Infarction (MI) in Non ST Elevation MI (NSTEMI) patients (OR 0.70, 95% CI 0.56-0.88) and ST Elevation MI (STEMI) (OR 0.26, 95% CI 0.17-0.38) patients , and vascular events in NSTEMI (OR 0.69, 95% CI 0.55-0.86) and STEMI (OR 0.39, 95% CI 0.30-0.51) patients⁴⁰. Death was also reduced after STEMI; major bleeding and transfusions were non-significantly increased and were few in number such that benefit outweighed hazard in absolute numbers of patients. The number of stroke events were too few to assess any trends, and insufficient or zero data were available for other antiplatelets (cilostazol, clopidogrel, dipyridamole)⁴⁰.

The proposed trial comes from members of the UK Stroke Research Network (SRN) Prevention Clinical Study Group (PB, SH, HM, GV) and is predicated on: (i) dual aspirin and dipyridamole is superior to aspirin after stroke; (ii) dual aspirin and dipyridamole or clopidogrel is the standard of care in the UK (NICE); (iii) dual aspirin and clopidogrel is superior to aspirin in patients with ischaemic heart disease (iv) some patients still 'fail' on aspirin and dipyridamole; and (v) Adding clopidogrel to aspirin may be useful in high risk patients, i.e. immediately after TIA/minor stroke. The results of our experimental medicine research (laboratory, phase I/II trials) and routine clinical use support this approach.³³⁻³⁷ Hence, triple therapy may be better still in high risk patients providing benefit exceeds bleeding.⁴¹

1.1 DETAILS OF INVESTIGATIONAL MEDICINAL PRODUCTS (IMP)

1.1.1 Description

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1.1.1.1 Aspirin (Asp)

International Non-Proprietary Name (INN): Aspirin

Chemical Abstracts Series (CAS) number: 50-78-2

Dose: Loading dose 300mg, then 75mg od.

Route: Enteral (including via nasogastric tube – dispersible or crushed tablets can be used) or rectal route.

For chemical and pharmacological properties, see summary of medical product characteristics (SmPC) at <http://www.medicines.org.uk/emc/>.

The IMP is defined by active substance only, so all authorised brands may be used.

1.1.1.2 Dipyridamole (Dip)

INN: Dipyridamole

CAS number: 58-32-2

Dose: 200mg modified release (MR), bd. Dysphagic patients with enteral access will take dipyridamole suspension 75mg tds. Patients with a headache from dipyridamole will have the dose weaned up from daily MR 200mg or standard release 75mg od to MR 200mg bd. Fixed dose combinations of A and D can also be used, e.g. Asasantin Retard (Aspirin 25mg, Dipyridamole 200mg MR, bd)

Route: Enteral (including via nasogastric tube).

For chemical and pharmacological properties SmPC at <http://www.medicines.org.uk/emc/>.

The IMP is defined by active substance only, so all authorised brands in the UK can be used.

1.1.1.3 Clopidogel (Clap)

INN: Clopidogrel

CAS number: 113665-84-2

Dose: Loading dose 300mg, then 75mg od.

Route: Enteral (including via nasogastric tube – crushed tablets can be used) or rectal route.

For chemical and pharmacological properties see SmPC at <http://www.medicines.org.uk/emc/>.

The IMP is defined by active substance only, so all authorised brands in the UK can be used.

1.1.2 Packaging and labelling

Standard pharmacy supplies should be used as all IMPs have marketing authorisation and are to be used in accordance with such authorisation. Hospitals/pharmacies should choose their own supplier for the IMPs and should be packaged according to local policy. All IMPs for the TARDIS trial should be labeled separately and pharmacies at the recruiting centre must have a written procedure in place for dispensing trial medications. The information on the label should include the trial name, EudraCT number, description of contents, batch number, expiry date, and quantity. There should be space for insertion of the trial number, name of the participant and the date of dispensing on the label (see appendix K). Under exceptional circumstances (e.g. out of hours) where labeled IMPs are not available, trial sites may choose to use ward stock without separate labeling if agreed locally and approved by the pharmacy.

1.1.3 Storage, dispensing and return

The IMPs must be stored in a secure location at room temperature (20°C to 25°C) with excursions permitted within 15°C to 30°C. Depending on local arrangement, this may be at the local pharmacy, the research department or the ward. Following recruitment and randomisation into the trial, the study treatment should be prescribed on the drug chart and the IMPs dispensed by the principal investigator/qualified designee. An accountability log for all IMPs should be maintained by the pharmacy and/or the research team and should include the following information: hospital number, participant initials, trial number, date dispensed, brand manufacturer, batch number, expiry date, quantity dispensed, quantity returned and initials of personnel who dispense and check the log. This should be completed for every participant who is randomised into the study. Accountability logs must be

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available for inspection during trial monitoring and/or audit and open to regulatory authorities inspection at any time. A sample label is provided (Appendix K).

1.1.4 Known Side Effects

1.1.4.1 Aspirin

Adverse reactions

- i. Bleeding: Aspirin prolongs bleeding time, and bleeding disorders, such as epistaxis, haematuria, purpura, ecchymoses, haemoptysis, gastrointestinal bleeding, haematoma and cerebral haemorrhage have been reported.
- ii. Blood and lymphatic system disorders - anaemia, haemolytic anaemia, hypoprothrombinaemia, thrombocytopenia, aplastic anaemia, pancytopenia, prolonged bleeding time, occult blood loss, elevated transaminase levels, agranulocytosis.
- iii. Gastrointestinal disorders - gastrointestinal bleeding, erosions, perforations or ulceration, which can occasionally be major (may develop bloody or black tarry stools, severe stomach pain and vomiting blood), gastrointestinal irritation (mild stomach pain, heartburn, vomiting and nausea). Fatalities have occurred.
- iv. Hepatic disorders - hepatitis (particularly in patients with SLE or connective tissue disease)
- v. Renal and urinary disorders – disturbances of renal function
- vi. Ear and labyrinth disorders - tinnitus.
- vii. Hypersensitivity reactions - rhinitis, urticaria, purpura, Stevens-Johnson syndrome, angio-oedema, asthma, worsening of asthma, bronchospasm.

Interaction with other medicinal products:

- i. Salicylates may enhance the effect of oral hypoglycaemic agents, phenytoin and sodium valproate.
- ii. They inhibit the uricosuric effect of probenecid and may increase the toxicity of sulphonamides.
- iii. Aspirin may potentiate the effect of heparin and increases the risk of bleeding with oral anticoagulants, antiplatelet agents and fibrinolytics.
- iv. The risk of gastrointestinal ulceration and bleeding may be increased when aspirin and corticosteroids are co-administered.
- v. Concurrent use of aspirin and other Non Steroidal Anti Inflammatory Drugs (NSAID) should be avoided. Use of two or more NSAID preparations increases the risk of serious gastrointestinal haemorrhage.
- vi. Concurrent administration of carbonic anhydrase inhibitors such as acetazolamide and salicylates may result in severe acidosis and increased central nervous system toxicity.
- vii. Alcohol may enhance the gastro-intestinal side effects of aspirin.
- viii. Patients using enteric-coated aspirin should be advised against ingesting antacids simultaneously to avoid premature drug release.
- ix. Selective Serotonin Reuptake Inhibitors (SSRI) may increase risk of gastrointestinal bleeding if coadministered.

1.1.4.2 Dipyridamole

Adverse reactions at therapeutic doses are usually mild.

- i. Bleeding: In very rare cases, increased bleeding during or after surgery has been observed.
- ii. Blood and lymphatic system disorders: Isolated cases of thrombocytopenia have been reported in conjunction with treatment with Dipyridamole.

- iii. Gastrointestinal disorders: Vomiting, diarrhoea and symptoms such as nausea, dyspepsia. These tend to occur early after initiating treatment and may disappear with continued treatment.
- iv. Cardiovascular: As a result of its vasodilating properties, dipyridamole may cause hypotension, hot flushes and tachycardia. Worsening of the symptoms of coronary heart disease such as angina and arrhythmias may occur.
- v. Central Nervous System Disorders: Dizziness, headache and myalgia may occur early after initiating treatment and may disappear with continued treatment.
- vi. Hypersensitivity reactions such as rash, urticaria, severe bronchospasm and angio-oedema have been reported.

Interaction with other medicinal products:

- i. Dipyridamole increases the plasma levels and cardiovascular effects of adenosine. Adjustment of adenosine dosage should therefore be considered if use with dipyridamole is unavoidable.
- ii. Dipyridamole may increase the hypotensive effect of blood pressure lowering drugs.
- iii. Dipyridamole may counteract the anticholinesterase effect of cholinesterase inhibitors thereby potentially aggravating myasthenia gravis.

1.1.4.3 Clopidogrel

Adverse Reactions

- i. Bleeding is the most common reaction reported and is mostly reported during the first month of treatment. Bleeding: some cases were reported with fatal outcome (especially intracranial, gastrointestinal and retroperitoneal haemorrhage); serious cases of skin bleeding (purpura), musculo-skeletal bleeding (haemarthrosis, haematoma), eye bleeding (conjunctival, ocular, retinal), epistaxis, respiratory tract bleeding (haemoptysis, pulmonary haemorrhage), haematuria and haemorrhage of operative wound have been reported; cases of serious haemorrhage have been reported in patients taking clopidogrel concomitantly with acetylsalicylic acid or clopidogrel with acetylsalicylic acid and heparin.

In addition to clinical studies experience, the following adverse reactions have been spontaneously reported. Within each system organ class (MedDRA classification), they are ranked under heading of frequency. "Very rare" corresponds to <1/10,000. Within each frequency grouping, undesirable effects are presented in order of decreasing seriousness.

- ii. Blood and lymphatic system disorders: very rare; Thrombotic Thrombocytopenic Purpura (TTP) (1/200,000 exposed patients), severe thrombocytopenia (platelet count $30 \times 10^9/l$), agranulocytosis, granulocytopenia, aplastic anaemia/pancytopenia, anaemia.
- iii. Immune system disorders: very rare; anaphylactoid reactions, serum sickness
- iv. Psychiatric disorders: very rare: confusion, hallucinations
- v. Nervous system disorders: very rare; taste disturbances
- vi. Vascular disorders: very rare; vasculitis, hypotension
- vii. Respiratory, thoracic and mediastinal disorders: very rare; bronchospasm, interstitial pneumonitis
- viii. Gastrointestinal disorders: very rare; pancreatitis, colitis (including ulcerative or lymphocytic colitis), stomatitis
- ix. Hepato-biliary disorders: very rare; acute liver failure, hepatitis
- x. Skin and subcutaneous tissue disorders: very rare; angioedema, bullous dermatitis (erythema multiforme, Stevens Johnson Syndrome, toxic epidermal necrolysis), rash erythematous, urticaria, eczema and lichen planus
- xi. Musculoskeletal, connective tissue and bone disorders: very rare; arthralgia, arthritis, myalgia.
- xii. Renal and urinary disorders: very rare; glomerulonephritis.

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Investigations: very rare; abnormal liver function test, blood creatinine increase

Interaction with other medicinal products:

- i. Clopidogrel should not be co-administered with warfarin due to increased bleeding risk. Caution should also be taken with corticosteroids, NSAIDs, heparin and thrombolytics.
- ii. Patients entered into clinical trials with clopidogrel have received a variety of concomitant medications including diuretics, beta blockers, ACEI, calcium antagonists, cholesterol lowering agents, coronary vasodilators, antidiabetic agents (including insulin), antiepileptic agents, hormone replacement therapy and GPIIb/IIIa antagonists without evidence of clinically significant adverse interactions.

2 TRIAL PURPOSE AND OBJECTIVES

2.1 Purpose

To perform a randomised trial assessing the efficacy, safety and tolerability of intensive antiplatelet therapy (Asp+Dip+Clop) versus guideline antiplatelet therapy (Asp+Dip or Clop) in patients with recent ischaemic stroke or TIA and who are at high risk of recurrence.

2.2 Primary Objective

To assess ordinal stroke severity at 90 days after short-term administration (1 month) of intensive antiplatelet therapy versus guideline therapy in patients with very recent ischaemic stroke or TIA.

2.3 Secondary Objectives

1. To assess the safety of short-term administration (1 month) of intensive antiplatelet therapy versus guideline therapy in patients with very recent ischaemic stroke or TIA.
2. To further assess, in high risk patients with stroke/TIA, whether:
 - ii. it is feasible to administer intensive therapy acutely and is tolerable to take for 1 month,
 - iii. intensive therapy is superior in respect of surrogate markers such as platelet function.
 - iv. intensive therapy improves functional outcome

3 TRIAL DESIGN

3.1 TRIAL /STUDY CONFIGURATION

3.1.1 Design

International, collaborative, multicentre, parallel group, prospective, randomised open-label, blinded-endpoint, controlled, Phase III trial.

3.1.2 Setting:

Initially, ~1000 patients will be recruited from the UK National Institute of Health Research (NIHR) Stroke Research Network (SRN) to the start-up phase. In the main phase, a further 3,100 participants from UK and overseas hospital-based stroke/TIA services will be recruited; UK participants (~2000) will be recruited from SRN sites (the trial is already adopted) including 55 sites who have been started-up and are recruiting in England and

Scotland. These sites have dedicated SRN nurses to facilitate recruitment and follow-up. Philip Bath will run the trial from the University of Nottingham Stroke Trials Unit.

3.1.3 Trial Duration:

The start-up phase will run for ~4 years. If the start-up phase shows acceptable safety, there will then be a seamless transition to the main phase of the trial of the same design so that recruitment does not stop (**tables 2a, 2b**).

The main phase will recruit in the order of ~3,100 patients (depending on the rate and distribution of ordinal events) and will last an additional 5 years. Separate permission for funding from the appropriate bodies (e.g. HTA) is being sought for the main phase.

If the trial shows that intensive antiplatelet therapy is superior to guideline therapy (taking account of the balance between reduced stroke/vascular events and potentially increased bleeding), intensive antiplatelet therapy could be introduced rapidly for stroke prevention with immediate benefit to high risk patients; each component is available now and licensed for secondary prevention. As the patent for clopidogrel has ended, NHS implementation of positive results will be based on generic costs, which will improve uptake and health economics.

A decision to proceed onto the main phase will be dependent on regular safety analyses during the start-up phase (by the Data Monitoring Committee), a successful funding application for the main phase, and the results of ongoing trials of dual antiplatelet therapy e.g. SPS-3 (Asp+Clop vs Asp), and ARCH (Asp+Clop vs. warfarin).

Table 2a: Trial timeline: Start-up phase

Time (months)	-6-0	0	0.25	0.5	1	1.5	2	2.5	3	3.25	3.5
Protocol	+										
Approvals	+										
Trial materials	+										
Site identification/training		+	+	+	+	+	+	+			
Recruit participants		+	+	+	+	+	+	+	+		
Day 90 follow-up			+	+	+	+	+	+	+	+	
DMC reviews				+	+	+	+	+	+		+
TSC meeting	+	+		+	+	+	+	+	+		+
Investigator meeting	+	+			+				+		
Feasibility reviews					<	+	+	+	>		
Database clean		+							+	+	
Finalise SAP										+	
Analysis											+

Table 2b: Trial timeline: Main phase

Year	0+	0.5	1	1.5	2	2.5	3	3.5	4	4.5	5	5.5
Further site identification	+	+	+	+	+	+	+	+	+			
Participant recruitment	+	+	+	+	+	+	+	+	+	+	+	
Day 90 follow-up	+	+	+	+	+	+	+	+	+	+	+	
DMC review		+		+		+		+		+		
TSC meeting	+		+		+		+		+		+	
Investigator meeting	+		+		+		+		+		+	
Publish protocol			+									
Database clean	+				+		+		+			
Data base close											+	
Analysis											+	+
Report writing											+	+

3.1.4 Primary outcome

5-level ordinal stroke and TIA scale with stroke ordered by its severity using the modified Rankin Scale (mRS): fatal stroke / severe non-fatal stroke (mRS 2-5) / mild stroke (mRS 0,1) / TIA / no stroke-TIA, measured at 90 days.; this approach allows for smaller sample sizes compared to binary outcomes such as stroke/no stroke.⁴²

3.1.5 Secondary outcomes

Days 35 and 90

Binary stroke; binary myocardial infarction; ordinal myocardial infarction (fatal MI/non-fatal MI/no MI);⁴² binary acute coronary syndrome; ordinal acute coronary syndrome (ACS - fatal/STEMI/NSTEMI/unstable angina/none); binary composite vascular outcome (non fatal MI & stroke, vascular death); ordinal composite vascular outcome;⁴² composite stroke, TIA, acute coronary syndromes and all cause death, incidence and type of infection.

Day 90, all participants:

Function (mRS, Barthel Index); Cognition (telephone mini mental state, TICS and animal naming); quality of life (EuroQoL/EQ-5D and EuroQOL VAS⁴³); Mood (Zung⁴⁴); disposition (home, carer, residential, nursing home); discharge from hospital (timing) days at home; economic activity.

Tolerability: Proportion of patients completing 30 days of randomised treatment.

Feasibility: Recruitment rate per week.

3.1.6 Safety Outcomes

Days 7 and 35

Full blood count by local investigator

Days 7, 35 and 90:

Ordinal bleeding (fatal/major/moderate/minor/none⁴²) as adjudicated by an independent blinded panel; death; binary major bleeding (fatal, symptomatic, causing fall in haemoglobin of $\geq 2\text{g/l}$, or leading to transfusion of ≥ 2 units of blood/red cells);⁴⁵ binary minor bleeding (e.g. bruising)
binary bleeding; all bleeding, symptomatic intracerebral haemorrhage, major extracranial bleeding, binary serious adverse events,
ordinal adverse events (fatal/serious/other/none⁴²);
thrombotic thrombocytopenic purpura; granulocytopenia.

3.2 Randomisation and Blinding

Patients will be randomised centrally using a secure internet site in real-time with stratification on index event (stroke/TIA) and country and minimisation on key prognostic/logistical baseline factors (age, gender, systolic blood pressure, cortical/lacunar syndrome, previous mono/dual antiplatelet, gastro-protection, use of low dose heparin, and time to randomisation, number of crescendo TIAs and ABCD2 score for TIAs and NIHSS and treatment with alteplase for strokes.) thereby maintaining concealment of allocation, minimising differences in key baseline variables, and improving statistical power.⁴⁶

Multiple measures will be taken to reduce bias: internet data capture, real-time validation and concealment of allocation; blinded assessment of events, and adjudication of events,

SAEs and neuroimaging; analysis by intention-to-treat; analyses adjusted for minimisation factors; adjustment for non-randomised treatment (e.g. statins, BP medications).

In the event that the website cannot be accessed, participants may be randomised by telephoning one of a series of emergency telephone numbers. These participants will be randomised without stratification or minimisation.

3.3 SELECTION AND WITHDRAWAL OF PARTICIPANTS

3.3.1 Recruitment

The initial approach will be from a member of the patient's usual care team (which may include the investigator or other members of the clinical research team).

The investigator or their nominee, e.g. from the research team or a member of the participant's usual care team, will inform the participant or their nominated representative (other individual or other body with appropriate jurisdiction), of all aspects pertaining to participation in the study.

If needed, the usual hospital interpreter and translator services will be available to assist with discussion of the trial, but the consent forms and participant information sheets may not be available printed in other languages. It will be explained to the potential participant that entry into the trial is entirely voluntary and that their normal treatment and care will not be affected by their decision. It will also be explained that they can withdraw at any time but attempts will be made to avoid this occurrence. In the event of their withdrawal it will be explained that their data collected so far cannot be erased and we will seek consent to use the data in the final analyses where appropriate.

3.3.2 Inclusion criteria

Adults at high risk of recurrent ischaemic stroke:

1. Age ≥ 50 years
2. Within 48 hours of ictus (24-48 hours if thrombolysed)
3. TIA with limb weakness and/or dysphasia lasting between 10 minutes and <24 hours with no residual symptoms and presenting with any of the following
 - a. ABCD2 score ≥ 4 , or
 - b. Crescendo TIA or
 - c. Already on dual antiplatelet therapy

Note: Neuroimaging is not necessary for transient ischaemic attack. Crescendo TIA is >1 TIA in one week and the onset time of last TIA is taken as time of ictus.

4. Ischaemic non cardioembolic stroke presenting with any of the following
 - a. Ongoing limb weakness and/or dysphasia of more than one hour duration
 - b. Resolved limb weakness of more than one hour duration with ongoing facial weakness
 - c. Ongoing isolated hemianopia of more than 1 hour duration with positive neuroimaging evidence to support the index event (e.g. ischaemic stroke in occipital lobe)
 - d. Resolved limb weakness and/or dysphasia between 24-48 hours after index event onset

Note: Neuroimaging is essential for ischaemic stroke to exclude intracranial haemorrhage and/or non stroke diagnosis

5. Informed consent from participant. If the participant is unable to give meaningful consent e.g. due to dysphasia, confusion, or reduced conscious level, proxy consent may be obtained from a relative, carer or legal representative .

.Exclusion criteria

1. Age < 50
2. Isolated sensory symptoms or vertigo/dizziness or facial weakness
3. Isolated hemianopia without positive neuroimaging evidence
4. Intracranial haemorrhage
5. Baseline neuroimaging showing parenchymal haemorrhagic transformation (PH I/II) of infarct, subarachnoid haemorrhage or other non ischaemic cause for symptoms
6. Presumed cardioembolic stroke (e.g. history or current AF, myocardial infarction within 3 months)
7. Participants with contraindications to, or intolerance of, aspirin, clopidogrel or dipyridamole.
8. Participants with definite need for treatment with aspirin, clopidogrel or dipyridamole individually or in combination (e.g. aspirin and clopidogrel for recent MI/acute coronary syndrome)
9. Participant has taken clopidogrel or dipyridamole after the index event but prior to randomisation (aspirin is allowed between ictus onset and randomisation)
10. Definite need for full dose oral (e.g. warfarin, dabigatran) or medium to high dose parenteral (e.g. heparin) anti-coagulation. NB Low dose heparin for DVT prophylaxis is allowed
11. Definite need for glycoprotein IIb-IIIa inhibitors
12. Received thrombolysis within the last 24 hours
13. No enteral access
14. Pre-morbid dependency (mRS > 2).
15. Severe high BP (BP > 185/110 mmHg).
16. Haemoglobin less than 10g/dL
17. Platelet count more than $600 \times 10^9/L$ or less than $100 \times 10^9/L$
18. White cell count more than $30 \times 10^9/L$ or less than $3.5 \times 10^9/L$
19. Major bleeding within 1 year (e.g. peptic ulcer, intracerebral haemorrhage).
20. Planned surgery during 3 month follow-up (e.g. carotid endarterectomy)
21. Concomitant STEMI or NSTEMI.
22. Stroke secondary to a procedure (e.g. carotid or coronary intervention)
23. Coma (GCS < 8)
24. Non-stroke life expectancy < 6 months
25. Dementia
26. Participation in another drug or devices trial concurrently or within 30 days. (participants may take part in observational studies or non-drug or devices trials)
27. Geographical or other factors that may interfere with follow-up e.g. no fixed address or telephone contact number, not registered with a GP, or overseas visitor.
28. Females of childbearing potential, pregnancy or breastfeeding

3.3.3 Removal of participants from therapy or assessments

Participants may be withdrawn from therapy or assessments either at their own request or at the discretion of the Investigator (e.g. for reasons of safety or new information becoming available on the trial medication or condition being treated). The participants will be made aware that this will not affect their future care. Participants will be made aware (via the information sheet and consent form) that should they withdraw the data collected to date cannot be erased and may still be used in the final analysis.

3.3.4 Informed consent / assent

All participants will provide written informed consent. The Informed Consent Form will be signed and dated by the participant before they enter the trial. If participants are not competent to consent, e.g. due to dysphasia or confusion, relatives will be invited to give consent. These approaches are standard practice in acute stroke trials. However, all attempts should be made to take further informed consent from the participants should their condition improve. A doctor knowledgeable about the trial will gain consent. Third party consent by an experienced, independent clinician would also be accepted in the event that no relatives were available. The Investigator will explain the details of the trial and provide a Participant / Relative/Independent Physician Information Sheet, ensuring that the individual providing consent has sufficient time to consider patient participation in the trial. The Investigator will answer any questions that the participant / relatives have concerning study participation.

Informed consent will be collected from each participant before they undergo any interventions (including physical examination and history taking) related to the study. The Investigator will keep the original, the participant will keep one copy, and a second will be retained in the participants's hospital records.

Should there be any subsequent amendment to the protocol, which might affect a participant's participation in the trial, continuing consent will be obtained using an amended consent form, which will be signed by the participant.

3.4 TRIAL TREATMENT AND REGIMEN

3.4.1 Intervention

The trial will compare intensive versus guideline antiplatelet therapy. Guidelines for secondary prevention of recurrent stroke may vary at local trial centres depending on local, national and international guidelines, and cost.

3.4.1.1 Intensive antiplatelet group

Participants in the intensive antiplatelet group will receive Asp+Dip+Clop triple therapy for 28-30 days (to cover the period of maximum risk of recurrence) along with standard 'best care' (including lifestyle advice, BP and lipid lowering). Clop will be given as a loading dose of 300 mg,¹² then 75 mg daily, Asp as a loading dose of 300 mg,²² then 75 mg daily, and Dip modified release 200 mg twice daily⁹ for 28-30 days.

3.4.1.2 Guideline antiplatelet group

Patients randomised to the guideline group will receive one of the following antiplatelet therapies depending on local policy and guidelines:

- A. For ischaemic strokes: Asp and Dip dual therapy or Clop monotherapy.
- B. For TIAs: Asp and Dip dual therapy or Clop monotherapy.

Clop will be given as a loading dose of 300 mg,¹² then 75 mg daily, Asp as a loading dose of 300 mg,²² then 75 mg daily, and Dip modified release 200 mg twice daily⁹ for 28-30 days.

3.4.1.3 Comparators

The trial will compare the following intensive versus guideline antiplatelet therapies.

- A. For ischaemic stroke:
 - 1) Asp+Clop+Dip : Clop: Asp+Dip (2:1:1)
 - 2) Asp+Clop+Dip : Clop (1:1)
 - 3) Asp+Clop+Dip : Asp+Dip (1:1)

B. transient ischaemic attack:

- 1) Asp+Clop+Dip : Clop : Asp+Dip (2:1:1)
- 2) Asp+Clop+Dip : Clop (1:1)
- 3) Asp+Clop+Dip : Asp+Dip (1:1)

All participating sites will choose what comparators they wish to use for ischaemic stroke and TIA separately (e.g A1/B1 or A2/B3, or A3/B3). Sites will only be allowed to randomise patients to the group that they have previously chosen. Sites can however change this group during the trial, but will need to inform the coordinating centre so that the computerised randomisation system can be reprogrammed.

3.4.1.4 Notes on treatment

- i. Dysphagic participants with enteral access may take crushed aspirin (or rectal aspirin), crushed or liquid dipyridamole (range 75 mg tds to 100mg qds), and crushed clopidogrel (if so randomised).
- ii. Participants having a headache on dipyridamole will have the dose weaned up from daily MR 200mg or standard release 75 mg od to MR 200 mg bd (as in PRoFESS⁴⁷). Fixed dose combinations of aspirin and dipyridamole can also be used.
- iii. At the discretion of the investigator, participants can take gastro-prophylaxis against upper gastrointestinal bleeding (proton pump inhibitor/histamine 2 receptor antagonist \pm H. pylori eradication) according to local practice and policy.⁴⁸
- iv. After the 30 day treatment period, participants will be expected to return to guideline antiplatelet therapy, such as combined aspirin and dipyridamole or clopidogrel as recommended by local, national or international guidelines. [Note: PRoFESS (aspirin/dipyridamole vs. clopidogrel) enrolled 8,113 (40%) of patients within 10 days, and ~1000 patients within 2 days of onset, so it is feasible to administer dipyridamole acutely and is apparently safe.].
- v. Study drugs may be stopped around procedures that become necessary after enrolment (however, this may constitute a protocol violation/deviation).

3.4.2 Baseline Measures

Pre-morbid modified Rankin Scale (mRS); stroke impairment (NIHSS); full blood count (part of routine clinical care); haemodynamics and ECG. Stroke type will be categorised according to modified TOAST criteria.⁴⁹

3.4.3 Follow-up

Face-to-face interview at 7 \pm 1 and 35 \pm 3 days. Central telephone follow-up will be performed at 90 \pm 7 days by an assessor blinded to outcome.

As stroke is the primary outcome, vascular events a key secondary outcome, and bleeding the main hazard, ascertaining these events is vitally important. All participants will be asked specifically about Serious Adverse Events (SAE) and Outcome events at every follow up. We will also triangulate this information from GPs and local researchers, especially between day 35 and 90. Such information will be obtained centrally by the coordinating centre from the GPs and by the local researchers from their hospital electronic systems.

3.4.4 Platelet Function

Platelet expression of P-selectin will be used to monitor platelet effects in participants. Blood will be taken from all participants at baseline & day 7 \pm 1, fixed (to allow batching of samples), posted to Nottingham using pre-purchased blood sample containers, and P-selectin measured using a standardised assay [Heptinstall; patent pending (PTC/GB2008/050169)] with blinding to participant and treatment identity. P-selectin has

been demonstrated to provide a robust means of identifying individual compliance with, and resistance to, aspirin, dipyridamole and clopidogrel; measurements will also be used to look for associations between successful platelet inhibition and clinical outcome. The analyses will be conducted at the Division of Cardiovascular Medicine at Queen's Medical Centre, Nottingham. All measurements are performed by flow cytometry and are subject to strict quality control

3.4.5 Additional Blood Samples

Tertiary questions in TARDIS include assessing the effects of the interventions on blood biomarkers and whether a participant's genotype alters response to the interventions. For example, the *CYP2C19* genetic variant is a major determinant of prognosis in young participants who are receiving clopidogrel treatment after myocardial infarction, and may be significant in ischaemic stroke.⁵⁰⁻⁵² Several blood biomarkers are surrogate markers of outcome, such as S-100.⁵³ However, whether they and other blood factors (to be identified during the course of the trial) are also markers of the efficacy of interventions has yet to be determined.

Centres should have appropriate storage facilities including access to a centrifuge and freezer. In addition to the full blood counts, the following blood samples are required for blood biomarkers and genetic analysis:

Genetics blood test sample

- 4mls EDTA. Frozen whole (i.e. no centrifugation)
- anytime from baseline to Day 35

Baseline:

- 4mls EDTA. Centrifuge to collect and freeze plasma.
- 6mls clotted sample. Centrifuge to collect and freeze serum

Day 7±1:

- 4mls EDTA. Centrifuge to collect and freeze plasma.
- 6mls clotted sample. Centrifuge to collect and freeze serum

(See table 3, page 24 for a tabulated summary of all blood samples)

If the centre concerned does not use blood bottles containing EDTA, then their bottles usually used for FBC samples is sufficient (this will contain appropriate anticoagulant). Blood samples should be anonymised and labeled with the centre number, participant number and initials (C999/9999/ZZ), day and date of sample (Day 7 or 35, dd/mm/yyyy), stored locally in a freezer at -20°C (or lower if possible at -60°C to -80°C) and accounted for using the TARDIS Blood Sample Freezer Log. The TARDIS Coordinating Centre will arrange transfer of blood samples to Nottingham for analysis. Blood samples will be destroyed once analysis is completed, this being dependent on the trial's completion date.

A separate consent form will allow the participant/relative to opt-in to the genetic sub-study. Participants may continue in the main trial even if they or their next-of-kin elect not to consent to the genetics sub-study. The participant or next-of-kin may request destruction of the genetic samples at any time after consent and prior to creation of an anonymised database. An important aim of the genetic analyses is to determine whether polymorphic differences in candidate genes explain resistance to antiplatelets (pharmacogenetic analysis). The exact genetic analyses to be performed are undefined at present and will depend on relevant scientific information available at the time of laboratory analysis and prior to sample destruction.

3.4.6 Scan Transfer and Storage

Baseline and subsequent clinical or research CT and/or MR brain scans should be sent electronically (ideally) using the secure internet web upload facility provided on the TARDIS

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website (www.tardistrial.org/). Scans should not be anonymised prior to upload as certain fields such as study date, birth date and sex are essential to ensure that the scan is matched to the correct participant. The upload facility will transfer data using RC4-MD5 (128 bit) cipher encryption and anonymise the DICOM header of the images automatically once the scan and participant have been matched. The DICOM header attributes that are anonymised are a subset of those specified in the 'Basic Application Level Confidentiality Profile' of the DICOM standard 3.15; namely the institution name, institution address, referring physician, referring physician's address, Patient name, patient identifier, date of birth, other patient id, other patient names and patient's address attributes.

If centres are unable to use the web upload facility, encrypted and anonymised scans can be copied on a CD/DVD and sent via recorded delivery to the TARDIS ICC. The password for de-encryption, site and participant number, participant initials and scan date should be communicated separately via email. The data will be unencrypted at the TARDIS ICC and uploaded to the database as described above.

If centres are unable to send the scans by the above methods, they will be advised to contact the TARDIS ICC, who will help them with the process. Under exceptional circumstances, for centres where the only method of transferring images is by films/hardcopies, centres will be advised to send anonymised films via recorded delivery. These will be digitised at the TARDIS ICC.

All digital brain image data will be stored on secure computer servers owned and maintained by the Information Services, University of Nottingham, with access restricted both physically (locked server rooms) and by password. Access for adjudication, analysis and archiving will be by login, password and PIN numbers.

Anonymised imaging data shall be adjudicated by trained neuroradiologists who may be based at the Coordinating Centre or elsewhere. The adjudication systems have been designed to ensure the highest levels of data security and participant confidentiality, and will be further enhanced if future technological advances permit it. The enhancements to the current system may include the use of e-Science and Grid technologies (e.g. NeuroGrid, www.neurogrid.ac.uk/) if they prove to be superior to current systems.

Reports from radiologists on clinical carotid imaging will also be collected (ultrasound, MRA or CTA). Reports on brain imaging and carotid imaging performed at local centres will be faxed to the TARDIS ICC.

3.4.7 Expected duration of participant participation

Participant participation and assessments are summarised in the table below (**table 3**):

	Day 0	Day 3±1	Day 7±1	Day 35±3	Day 90±*
Randomisation	+				
Safety assessments		+	+	+	+
Tolerability assessments		+	+	+	+
Bloods					
FBC	+		+	+	
P-Selectin	+		+		
Genetics/EDTA [†] sample	+				
Serum and plasma	+		+		
Clinical Efficacy					
Impairment (NIHSS)	+		+	+	
Function (mRS & BI)					+
Cognition, QoL & Mood					+

Table 3: *Day 90 assessment done by telephone questionnaire. †or anticoagulant provided in the hospital's usual FBC blood tubes. FBC, Full Blood Count; NIHSS, National Institutes of Health Stroke Scale; mRS, modified Rankin Score; BI, Bartel Index; QoL, Quality of Life

3.4.8 Co-enrolment into other studies

Uncoordinated co-enrollment of patients into two or more trials has the potential for introducing bias, e.g. when the treatments have a similar mechanism of action, potentially share adverse events or have common outcomes. Patients should not be enrolled into this trial if they are already in another drug or devices trial. Patients can be co-enrolled into non-drug or devices trials or observational studies.

3.4.9 Compliance

At each scheduled visit, compliance with the IMPs will be assessed on direct questioning or by reviewing medication charts. Patients stopping a drug because of adverse events will carry on with the remaining therapy and follow-up assessments with analysis by intention-to-treat.

3.4.10 Protocol Violations and Deviations

The study should be conducted in accordance with the approved protocol and that changes to that protocol will only be made to protect the safety, rights, or welfare of the subject.

3.4.10.1 Protocol Violation

A protocol violation is a major deviation from the trial protocol where a participant is enrolled in spite of not fulfilling all the inclusion and exclusion criteria, or where deviations from the protocol could affect the trial delivery or interpretation significantly.

The following baseline characteristics constitute a protocol violation

1. Randomisation > 48 hours from onset of symptoms
2. Participant less than 50 years of age
3. For ischaemic stroke:
 - a. No cranial imaging results available prior to randomisation
 - b. Isolated sensory symptoms, vertigo or dizziness or facial weakness as presenting symptoms of the index event
4. For TIAs:
 - a. Limb weakness and/or dysphasia lasting less than 10 minutes
 - b. ABCD2 score <4 and not a crescendo TIA and not on dual antiplatelet therapy
5. Failure to obtain appropriate consent prior to randomisation
6. Pre-morbid dependency (mRS) >2
7. Participant unable to swallow and does not have enteral access
8. Baseline cranial imaging shows parenchymal haemorrhagic transformation (PH I/II)
9. Subarachnoid haemorrhage
10. Intracerebral haemorrhage
11. On anticoagulation therapy except low dose low molecular weight heparin
12. Participant has taken dipyridamole or clopidogrel following the index event and prior to stroke randomisation
13. Thrombolysis less than 24 hours prior to randomisation
14. Presumed cardioembolic stroke or history of atrial fibrillation
15. Concomitant STEMI or NSTEMI
16. Baseline SBP reading >185 mm Hg or DBP > 110 mm Hg
17. Major bleeding within one year prior to randomisation
18. Planned surgery within the 3-month follow-up period

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19. Randomising event was secondary to a surgical procedure
20. Glasgow Coma Score < 8
21. Known history of dementia
22. Known probable life expectancy of less than 6 months
23. Unavailable for follow-ups
24. Female patient of childbearing potential, pregnant or breastfeeding at randomisation
25. Patient receiving treatment that they are not randomised to.

The following practice during the trial constitutes a 'protocol violation'

1. Subsequent randomisation into another drug or devices trial
2. Patient does not receive 5 days of randomised treatment in the first seven days and 16 days in the next 3 weeks
3. Failure to complete SAEs where appropriate
4. Failure to complete outcomes where appropriate
5. Follow-up assessments are performed (as opposed to submitted) outside the specified time as shown below:
 - a. 7-day follow-up: >7 days past the due date
 - b. 35-day follow-up: >7 days past the due date
 - c. Hospital event form: >30 days past the due date
 - d. 90-day follow up: >30 days past the due date

3.4.10.2 Protocol Deviation

A Protocol Deviation is a minor deviation from the protocol that affects the conduct of the trial in a minor way. This includes any deviation from the trial protocol that is not listed as a Protocol Violation. Examples of Deviations are given below but this is not exhaustive.

The following practice during the trial constitutes a 'protocol deviation'

1. Failure to switch to standard treatment following completion of treatment period
2. Patient receives more than 400mg daily of dipyridamole
3. Patient receives >75mg of aspirin or clopidogrel after Day 0
4. Non-receipt of Day 7 or Day 35 Full Blood Count
5. No blood pressure measurements at baseline, D7 or D35 follow-ups
6. Follow-up assessments are performed (as opposed to submitted) outside the specified time as shown below:
 - a. 7-day follow-up: >1day past the due date
 - b. 35-day follow-up: >3days past the due date
 - c. Hospital event form: >7days past the due date
 - d. 90-day follow-up: >7 days past the due date

3.4.10.3 Review of Protocol Violations and Deviations

Protocol Violations will be reviewed annually by both the Data Monitoring Committee (using unblinded data) and the Trial Steering Committee (with blinding to treatment assignment).

The list of protocol violations and deviations will be updated, as necessary, in a working practice document which will be uploaded and available on the trial website.

3.5 TRIAL MANAGEMENT

3.5.1 Sponsor

The University of Nottingham is the trial sponsor in the UK and will delegate responsibility

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for design and conduct of the trial to the Chief Investigator via our Sponsor/Chief Investigator agreement. The sponsor contact details are

Mr Paul Cartledge
Head of Research Grants and Contracts
Research Innovation Services
King's Meadow Campus,
Lenton Lane
Nottingham NG7 2NR UK

3.5.2 Coordinating Centre

The Stroke Trials Unit (STU), part of the University of Nottingham's Clinical Trials Unit (which has provisional registration), will co-ordinate the trial. STU will have overall responsibility for the conduct of the trial and will be responsible for provision of trial materials, collation and analysis of data and reporting of the final results. They will act as the International Coordinating Centre, UK National Coordinating Centre, the primary point of contact for UK centres, and the secondary point of contact for non-UK centres. The contact details are

Stroke Trials Unit
Division of Stroke Medicine
University of Nottingham
Clinical Science Building
City Hospital campus
Nottingham, NG5 1PBUK
Tel: +44 115 8230210
Fax: +44 115 8230273

3.5.3 Trial Steering Committee (TSC)

The TSC will provide overall supervision, as per their charter, and ensure that the trial is conducted in accordance with the principles of the ICH GCP and the relevant regulations. Any amendments to the trial will be agreed by the TSC. The TSC will provide advice to the investigators on all aspects of the trial. The composition of the TSC is given on the Trial website.

3.5.4 Data Monitoring Committee (DMC)

The Data Monitoring Committee (DMC) will monitor efficacy and safety as per their charter. As well as outcome measures, the DMC will also review recruitment, baseline data, balance in baseline factors between the treatment groups, completeness of data, compliance to treatment, co-administered treatments, and outcome by sub groups. They will also review all serious adverse events (both adjudicated and unadjudicated) and protocol violations. The DMC will usually meet at least yearly by teleconference; the chairman will receive 6 monthly updates from the statistician. The composition and charter of the DMC is given on the trial website (www.tardistrial.org).

3.5.5 Outcome and event adjudication

There will be 2 adjudication committees:

- Independent Events (vascular outcomes, SAE)
- Neuroimaging adjudication

4 STATISTICS

A medical statistician will support the TSC with analyses. An interim analysis will be done during the start-up phase, blinded to treatment allocation, to demonstrate feasibility of the trial.

4.1 Methods

Analysis will be performed using ordinal logistic regression for ordered categorical variables, binary logistic regression for binary outcomes, ANCOVA on continuous data and Kaplan-Meier and Cox proportional hazards regression on time to event data. Analyses will be adjusted for randomisation/minimisation factors.

Safety analyses will be reviewed 6 monthly during the start-up phase by the independent Data Monitoring Committee.

The effect of the intervention on the primary outcome will be performed within the following subgroups of subjects:

- a) By age - ≤ 75 years, > 75 years.
- b) By sex - male, female.
- c) By index event-stroke/TIA.
- d) By stroke sub-type - lacunar, posterior fossa, cortical.
- e) By stroke severity - severe, moderate/mild; NIHSS ≤ 10 , > 10 .
- f) By baseline systolic blood pressure - > 160 mmHg, 140-160 mmHg, < 140 .
- g) By treatment delay - > 24 hours, ≤ 24 hours.
- h) By patients enrolled into P-selectin substudy.
- i) By patients on antiplatelet therapy at randomisation - mono, dual
- j) Aspirin naïve vs aspirin.
- k) By heparin - none, unfractionated, LMWH.
- l) By number of TIAs in the last week.
- m) By thrombolysis - yes, no.
- n) By ABCD2 score - 4, > 4 .

Patients in the UK will be 'flagged' for death with the NHS Information Centre (Medical Research Information Service-MRIS) so that vital status can be obtained for all patients.

4.2 Sample size and justification

4.2.1 Start-up phase

The start-up phase was sized to assess safety, i.e. where intensive antiplatelet therapy *might be* hazardous compared to guideline therapy; the key concern for antiplatelet agents relates to bleeding. The sample size calculation⁵⁴ used assumptions based on data from our pilot trial of triple antiplatelet therapy.⁵⁵ Assuming bleeding rates for Asp+Dip was 15% and Asp+Dip+Clop was 30%, alpha 5%, power 90%, losses to follow-up 3%, total sample size = 320 rounded to 350. Analyses will, in reality, be performed using ordinal approaches to improve statistical power.⁴²

4.2.2 Main phase

The start-up phase informs the sample size calculation for the main phase which will assess the efficacy of intensive versus guideline therapy. Assuming the distribution in 5 level recurrent stroke/TIA outcome (stroke with mRS 6 = 0.1%/ mRS 2-5 = 0.7%/ mRS 0-1 = 1.53%/ TIA = 3.57%), odds ratio of 0.68, alpha 5%, power 90%, losses to follow-up 2%, treatment crossovers 5% the total sample size for the whole study is 4100.

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4.3 Definition of populations analysed

4.3.1 Safety Set

All randomised participants.

4.3.2 Intention-to-Treat (ITT) efficacy set

All participants in the Safety Set, and who took at least one treatment dose. Participants in the ITT will be defined prior to database lock.

4.3.3 Per Protocol Set (PPS) efficacy set

All participants in the ITT, and who are deemed to have no **protocol violations**. Participants in the PPS will be defined prior to database lock.

4.3.4 Analyses

All efficacy analyses will be assessed using the **ITT**; the robustness of the primary analyses will be assessed in the **PPS**. Safety summaries will be performed on the **Safety Set**. Major protocol deviations will lead to exclusion of a participant from the **PPS**.

4.4 Health economic analysis

The impact of intensive antiplatelet therapy with aspirin, dipyridamole and clopidogrel on quality of life will be assessed using the EuroQoL. A full health-economic analysis will only be performed after completion of the main phase of the trial.

5 ADVERSE EVENTS

5.1 Definitions

A Serious Adverse Event (SAE) is any adverse event occurring following study mandated procedures, having received the IMP that results in any of the following outcomes:

1. Death
2. A life-threatening adverse event
3. Inpatient hospitalisation or prolongation of existing hospitalisation
4. A disability / incapacity
5. A congenital anomaly in the offspring of a participant
6. Important medical events that may not result in death, be life-threatening, or require hospitalisation may be considered a serious adverse event when, based upon appropriate medical judgment, they may have been felt to jeopardize the participant and may require medical or surgical intervention to prevent one of the outcomes listed in this definition

All serious adverse events will be assessed for expectedness and causality:

A distinction is drawn between serious and severe AEs. Severity is a measure of intensity whereas seriousness is defined using the criteria above. Hence, a severe AE need not necessarily be serious.

5.2 Causality

Not related or improbable: a clinical event including laboratory test abnormality with temporal relationship to trial treatment administration which makes a causal relationship incompatible or for which other drugs, chemicals or disease provide a plausible explanation. This will be counted as "unrelated" for notification purposes.

Possible: a clinical event, including laboratory test abnormality, with temporal relationship to trial treatment administration which makes a causal relationship a reasonable possibility, but which could also be explained by other drugs, chemicals or concurrent disease. This will be counted as "related" for notification purposes.

Probable: a clinical event, including laboratory test abnormality, with temporal relationship to trial treatment administration which makes a causal relationship a reasonable possibility, and is unlikely to be due to other drugs, chemicals or concurrent disease. This will be counted as "related" for notification purposes.

Definite: a clinical event, including laboratory test abnormality, with temporal relationship to trial treatment administration which makes a causal relationship a reasonable possibility, and which can definitely not be attributed to other causes. This will be counted as "related" for notification purposes.

An AE whose causal relationship to the study IMP is assessed by the Chief Investigator or delegate as "improbable", "possible", "probable", or "definite" is an Adverse Drug Reaction.

With regard to the criteria above, medical and scientific judgment shall be used in deciding whether prompt reporting is appropriate in that situation.

5.3 Reporting of adverse events

Participants will be asked to contact the study site immediately in the event of any serious adverse event. All serious adverse events will be recorded and closely monitored until resolution, stabilisation, or until it has been shown that the study medication or treatment is not the cause. The Chief Investigator or delegate shall be informed immediately of any serious adverse events and shall determine seriousness and causality in conjunction with any treating medical practitioners. All SAEs will be reported to the Stroke Trials Unit, University of Nottingham.

In the event of a pregnancy occurring in a trial participant or the partner of a trial participant monitoring shall occur during the pregnancy and after delivery to ascertain any trial related adverse events in the mother or the offspring. Where it is the partner of a trial participant, consent will be obtained for this observation from both the partner and her medical practitioner. All serious adverse events will be recorded and reported to R&D and REC as part of the annual reports. SUSARs will be reported within the statutory timeframes to the MHRA and REC as stated below. The Chief Investigator shall be responsible for all adverse event reporting.

5.4 SUSARs

A serious adverse event that is either sudden in its onset, unexpected in its severity and seriousness or not a known side effect of the IMP *and* related or suspected to be related to the IMP is classed as Suspected Unexpected Serious Adverse Reaction and requires expedited reporting as per the clinical trials regulations.

All serious adverse events that fall or are suspected to fall within these criteria shall be treated as a SUSAR until deemed otherwise.

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The event shall be reported immediately of knowledge of its occurrence to the Chief Investigator.

The Chief Investigator will:

- Assess the event for seriousness, expectedness and relatedness to the study IMP
- Take appropriate medical action, which may include halting the trial and inform the Sponsor of such action
- If the event is deemed a SUSAR, shall, within seven days, complete the CIOMS form and send to the MHRA.
- Shall inform the REC using the reporting form found on the NRES web page within seven days of knowledge of the event
- Shall, within a further eight days send any follow-up information and reports to the MHRA and REC.
- Make any amendments as required to the study protocol and inform the ethics and regulatory authorities as required

5.5 Participant removal from the study due to adverse events

Any participant who experiences an adverse event may be withdrawn from the study at the discretion of the Investigator. Should the participant discontinue any trial medications due to, for example, an adverse event, they will remain in the study until the end of the trial at day 90 (± 7), as completeness of follow-up is essential. However, should they wish to do so, any participant is free to withdraw from the trial at any time and without giving reason.

6 ETHICAL AND REGULATORY ASPECTS

6.1 ETHICS COMMITTEE AND REGULATORY APPROVALS

The trial will not be initiated before the protocol, informed consent forms and participant and GP information sheets have received approval / favourable opinion from the Medicines and Healthcare products Regulatory Agency (MHRA), Research Ethics Committee (REC), and the respective National Health Service (NHS) Research & Development (R&D) department. Should a protocol amendment be made that requires REC approval, the changes in the protocol will not be instituted until the amendment and revised informed consent forms and participant and GP information sheets (if appropriate) have been reviewed and received approval / favourable opinion from the REC and R&D departments. A protocol amendment intended to eliminate an apparent immediate hazard to participants may be implemented immediately providing that the MHRA, R&D and REC are notified as soon as possible and an approval is requested. Minor protocol amendments only for logistical or administrative changes may be implemented immediately; and the REC will be informed.

The trial will be conducted in accordance with the ethical principles that have their origin in the Declaration of Helsinki, 1996; the principles of Good Clinical Practice, in accordance with the Medicines for Human Use Regulations, Statutory Instrument 2004, 1031 and its subsequent amendments and the Department of Health Research Governance Framework for Health and Social care, 2005.

6.2 RECORDS

6.2.1 Drug accountability

Hospitals/pharmacies should choose their own supplier for the trial medications. As is common with stroke trials, medication can be dispensed and kept on the relevant ward or department ready for use as soon as the patient is randomised. It may be kept as 'ward

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stock' or as separate trial medication according to the practices of the randomising hospital (see Section 1.1.3).

6.2.2 Case Report Forms

Each participant will be assigned a trial identity code number, allocated at randomisation, for use on CRFs other trial documents and the electronic database. The documents and database may also use their age. CRFs will be treated as confidential documents and held securely in accordance with regulations. The investigator will make a separate confidential record of the participant's name, date of birth, local hospital number and NHS number (UK patients), and Participant Trial Number (the Trial Recruitment Log), to permit identification of all participants enrolled in the trial, in case additional follow-up is required. CRFs shall be restricted to those personnel approved by the Chief or local Principal Investigator and recorded on the 'Trial Delegation Log.' All paper forms should be filled in using black ballpoint pen. Errors shall be lined out but not obliterated by using correction fluid and the correction inserted, initialled and dated.

6.2.3 Source documents

Source documents shall be filed at the investigator's site and may include but are not limited to, consent forms, current medical records, laboratory results and pharmacy records. A CRF may also completely serve as its own source data. Only trial staff as listed on the Delegation Log shall have access to trial documentation other than the regulatory requirements listed below.

6.2.4 Direct access to source data / documents

The CRF and all source documents, including progress notes and copies of laboratory and medical test results shall be made available at all times for review by the Chief Investigator, Sponsor's designee and inspection by relevant regulatory authorities (e.g., MHRA).

6.3 DATA PROTECTION

All trial staff and investigators will endeavour to protect the rights of the trial's participants to privacy and informed consent, and will adhere to the Data Protection Act, 1998. The CRF will only collect the minimum required information for the purposes of the trial. CRFs will be held securely, in a locked room, or locked cupboard or cabinet. Access to the information will be limited to the trial staff and investigators and relevant regulatory authorities (see above). Computer held data including the trial database will be held securely and password protected. Access will be restricted by user identifiers and passwords (encrypted using a one way encryption method).

Personal information (e.g. name and address of patients and secondary contacts) about trial participants will be held at local centres and will be passed onto the Coordinating Centre, Nottingham, UK and to National Coordinating Centres for centres situated outside the UK. This is necessary for the coordination and execution of the blinded 90 day follow up assessments, which will be carried out centrally for each country. Patient information will be held on a database in Nottingham but will be separated from all clinical information; the latter remain anonymous (identifiable only by initials, trial number and age). Computer data will be backed up regularly to an off-site secure repository (to enable disaster recovery). Personal patient information will be used only for the purposes of the TARDIS trial and will not be passed on to third parties. The personal patient information will be deleted at the end of the trial.

Trial paperwork will be anonymised, scanned and stored on a digital archiving system. This is with the exception of consent forms and patient details form. This will comply with the Data Protection Act and confidentiality rules, as outlined above.

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Where permissible, the TARDIS Coordinating Centres may use central databases to obtain additional follow-up information on patients enrolled into the trial. In England and Wales, this will involve use of the NHS Information Centre (MRIS), database. When information will be gathered on patients in this way, it will be clearly stated in the country specific patient/relative information sheets.

Information about the trial in the participant's medical records / hospital notes will be treated confidentially in the same way as all other confidential medical information.

7 QUALITY ASSURANCE & AUDIT

7.1 INSURANCE AND INDEMNITY

Insurance and indemnity for trial participants and trial staff is covered within the NHS Indemnity Arrangements for clinical negligence claims in the NHS, issued under cover of HSG (96)48. There are no special compensation arrangements, but trial participants may have recourse through the NHS complaints procedures.

The University of Nottingham has taken out an insurance policy to provide indemnity in the event of a successful litigious claim for proven non-negligent harm.

7.2 TRIAL CONDUCT

Trial conduct will be subject to systems audit of the Trial Master File for inclusion of essential documents; permissions to conduct the trial; Trial Delegation Log; CVs of trial staff and training received; local document control procedures; consent procedures and recruitment logs; adherence to procedures defined in the protocol (e.g. inclusion / exclusion criteria, correct randomisation, timeliness of visits); serious adverse event recording and reporting; drug accountability, pharmacy records and equipment calibration logs.

The Trial Coordinator, or where required, a nominated designee of the Sponsor, shall carry out a site systems audit at least yearly and an audit report shall be made to the Trial Steering Committee.

7.3 TRIAL DATA

Monitoring of trial data shall include confirmation of informed consent; source data verification; data storage and data transfer procedures; local quality control checks and procedures, back-up and disaster recovery of any local databases and validation of data manipulation. The Trial Coordinator, or where required, a nominated designee of the Sponsor, shall carry out monitoring of trial data as an ongoing activity.

Entries on CRFs will be verified by inspection against the source data. A sample of CRFs (10%) will be checked on a regular basis for verification of all entries made. In addition the subsequent capture of the data on the trial database will be checked. Where corrections are required these will carry a full audit trail and justification.

Trial data and evidence of monitoring and systems audits will be made available for inspection by the regulatory authority as required.

7.4 RECORD RETENTION AND ARCHIVING

In compliance with the ICH/GCP guidelines, regulations and in accordance with the University of Nottingham Research Code of Conduct, the Chief or local Principal Investigator will maintain all records and documents regarding the conduct of the study. These will be
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retained for at least 7 years or for longer if required. If the responsible investigator is no longer able to maintain the study records, a second person will be nominated to take over this responsibility.

The Trial Master File and trial documents held by the Chief Investigator on behalf of the Sponsor shall be finally archived at secure archive facilities at the University of Nottingham. This archive shall include all trial databases and associated meta-data encryption codes.

7.5 DISCONTINUATION OF THE TRIAL BY THE SPONSOR

The Sponsor reserves the right to discontinue this trial at any time for failure to meet expected enrolment goals, for safety or any other administrative reasons. The Sponsor shall take advice from the Trial Steering Committee and the Independent Data Monitoring Committee (IDMC) as appropriate in making this decision.

During the period of recruitment into the study, the trial statistician will perform interim analyses on major outcome events and supply these, in strict confidence, to the members of the Data Monitoring Committee, along with any other analyses that the committee may request. In the context of TARDIS, the balance between safety and efficacy will be considered.

With respect to safety the following outcomes in particular will initiate discussion for recommending early stopping or continuation of the study:

- The primary outcome ('shift' in modified Rankin Scale in patients having a recurrent stroke event or TIA)
- Combined outcome of fatal or non-fatal stroke or major bleeding
- The overall rate of symptomatic intracranial haemorrhage

With respect to efficacy, the committee will conduct formal interim analyses based on the following outcome.

Combined outcome of fatal or non-fatal stroke or major bleeding \pm event .

In making any decision, the committee will consider the overall internal and external evidence, the multiplicity of testing and the possibility that the trends in the data might be reversed with longer follow-up or increased recruitment.

In the light of these analyses, the IDMC will advise the Chairman of the Trial Steering Committee (TSC) and Sponsor (via the Chief Investigator) if, in their view, the randomised comparisons in TARDIS have provided both

- (i) "proof beyond reasonable doubt" that for all, or for some, specific types of patient, treatment is clearly indicated or clearly contraindicated, and
- (ii) evidence that might reasonably be expected to influence materially the patient management of the many clinicians who are already aware of the results of any other relevant trials.⁵⁶⁻⁵⁷

On the basis of information supplied by the IDMC, the TSC can then decide whether to modify intake to the study (or to seek extra data). Unless this happens, however, the TSC, the collaborators, and the central administrative staff (except the unblinded statistician) will remain ignorant of the interim results.

Further details and updates to the DMC charter will be made available via the TARDIS website (www.tardistrial.org). Investigators are advised to refer to the trial website for an up to date DMC charter.

If a trial is discontinued for any of the above reasons, participants will go back to receiving standard care from their GPs.

7.6 STATEMENT OF CONFIDENTIALITY

Individual participant medical information obtained as a result of this study are considered confidential and disclosure to third parties is prohibited with the exceptions noted above. Participant confidentiality will be further ensured by utilising identification code numbers to correspond to treatment data in the computer files. Such medical information may be given to the participant's medical team and all appropriate medical personnel responsible for the participant's welfare.

Data generated as a result of this trial will be available for inspection on request by the participating physicians, the University of Nottingham representatives, the REC, local R&D Departments and the regulatory authorities.

8 PUBLICATION AND DISSEMINATION POLICY

Data and results will be shared as follows:

8.1 Presentation

The main trial results will be presented to the investigators, and to funding bodies, and at major international and national scientific meetings, in the name of the trial and investigators i.e. 'TARDIS Investigators'.

8.2 Publication

The main results from the trial will be written by a 'Writing Committee' and published in quality peer-reviewed journal(s) in the name of the investigators, i.e. TARDIS Investigators. The writing committee will consist of as a minimum, the Chief Investigator, lead imaging and SAE adjudicators, statistical consultant and trial statistician.

Secondary publications will be published as 'Person(s), for the TARDIS Investigators', where the person(s) are those who conceived, designed, and analysed or interpreted the data, and/or wrote the paper for the publication.

Abstracts will be presented as 'TARDIS Investigators, person(s)', where the person(s) act as a contact point for the trial.

National and/or local investigators may present or publish data relating to their country or site once the main trial findings have been published. All papers will be approved by the TSC and all abstracts by the Chief Investigator.

8.3 Sharing of data

Anonymised subsets of data may be shared with other research groups and projects (e.g. Cochrane Collaboration, antithrombotic collaboration) once the main trial findings have been published, and following agreement by the Trial Steering Committee. A contract will be set up between the University of Nottingham (as represented by the Chief Investigator) and groups which are receiving the data.

9 USER AND PUBLIC INVOLVEMENT

The trial has been discussed with, and is supported by, the UK Stroke Research Network Prevention Clinical Studies Group, the Nottingham Stroke Users Research Committee. Their comments have been incorporated into the design. One member will be a member of the Trial Steering Committee.

10 STUDY FINANCES

10.1 Funding source

The start-up phase of the study is funded by The British Heart Foundation. Funding is being sought for the main phase from the United Kingdom Health Technology Assessment (HTA).

10.2 Participant stipends and payments

Participants will not be paid to participate in the trial. Travel or mileage/parking expenses will be offered for hospital visits.

11 SIGNATURE PAGE

Signatories to Protocol:

Chief Investigator:

(name)

PHILIP BATH

Signature:



Date:

22/12/11**Trial Statistician:**

(name)

CHERYL HOGG

Signature:



Date:

21/12/2011**Trial Pharmacist:**

(name)

Signature: _____

Date: _____

Appendices

Appendix A: Definitions

Bleeding Events

1. **Major bleed:**⁴⁵ All major bleeds will constitute a serious adverse event.
 - Fatal bleeding, and/or
 - Symptomatic bleeding in a critical area or organ, such as intracranial, intraspinal, intraocular, retroperitoneal, intraarticular or pericardial, or intramuscular with compartment syndrome, and/or
 - Bleeding causing fall in haemoglobin of 2 g/l (1.24 mmol/l) or more, or leading to transfusion of 2 or more units of whole blood or red cells.
2. **Moderate bleed:** Moderate bleeds may or may not constitute a serious adverse event depending on other criteria as determined by the investigator.
 - Not major, and
 - Bleeding causing fall in haemoglobin <2 g/l (1.24) mmol/l) and \geq 1g (0.62 mmol/l), and leading to no transfusion, or transfusion of only 1 unit of whole blood or red cells.
3. **Minor bleed:** Minor bleeds usually do not constitute a serious adverse event.
 - Not major or moderate, and
 - Comprising bruising, ecchymoses, gingival bleed or similar other type of bleeding
 - Fall in haemoglobin of less than 1g/l (0.62 mmol/l).

Other Clinical Events

4. **Stroke:** A clinical syndrome characterised by rapidly developing clinical symptoms and/or signs of focal (and at times global) loss of cerebral function with symptoms lasting \geq 24 hours or leading to death, with no apparent cause other than that of vascular origin'.⁵⁸
5. **TIA:** A sudden focal neurological deficit of the brain or eye, presumed to be of vascular origin and lasts less than 24 hours.

NB. TIAs and stroke usually present with 'negative' symptoms (e.g. loss of motor power, loss of speech) as opposed to symptoms that are 'positive' in nature such as parasthesia or limb jerking, which will usually have an alternative underlying cause.

6. **Recurrent Stroke:** A stroke defined as above occurring after the qualifying stroke **or** a progression of neurological symptoms or signs (increase in NIHSS score >4) in the same vascular territory as the index event.
7. **Neurological Deterioration:** An increase in NIHSS score by 4 points or more than the baseline value.
8. **Symptomatic Intracerebral Haemorrhage (SICH):** Any haemorrhage with neurological deterioration as defined above, or intracerebral haemorrhage leading to death. The haemorrhage must be the predominant cause of the neurological deterioration.⁵⁹
9. **Bleeding on CT/MRI head scans:** ⁶⁰⁻⁶¹
 - a. *Haemorrhagic Infarct (HI):* petechial infarction without space occupying effect.
 - i. HI1 - small petechiae
 - ii. HI2 - more confluent petechiae
 - b. *Parenchymal Haemorrhage (PH):* haemorrhage with mass effect.
 - i. PH1 - <30% of the infarcted area with mild space occupying effect
 - ii. PH2 - >30% of the infarcted area with significant space occupying effect.

Note: patients with PH should not be enrolled

10. **ABCD² Scoring Criteria** ⁶²⁻⁶³

A	Age \geq 60 years	1 point
B	Blood pressure \geq 140/90 mm Hg	1 point
C	Clinical features	
	Unilateral weakness	2 points
	Speech disturbance [§] without weakness	1 point
D	Duration	
	\geq 60 minutes	2 points
	10–59 minutes	1 point
D	Diabetes	
	Presence of diabetes mellitus*	1 point

§ Speech disturbance defined as either dysarthria or dysphasia or both

* Diabetes defined as requiring either oral medication or insulin

Note: patients with ABCD² <4 should not be enrolled

11. **Criteria for acute, evolving or recent Myocardial Infarction (MI):**⁶⁴ Either one of the following criteria satisfies the diagnosis for an acute, evolving or recent MI:

1. Typical rise and gradual fall (troponin) or more rapid rise and fall (CK-MB) of biochemical markers of myocardial necrosis with at least one of the following:

(a) ischaemic symptoms;

(b) development of pathologic Q waves on the ECG;

(c) ECG changes indicative of ischemia (ST segment elevation or depression); or

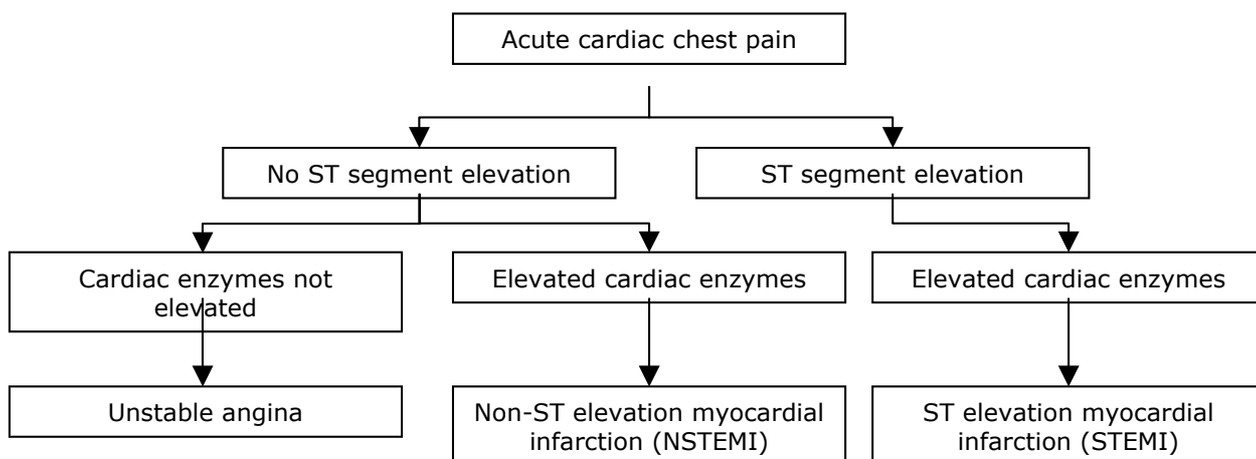
(d) coronary artery intervention (e.g., coronary angioplasty).

2. Pathologic findings of an acute MI.

12. **Unstable Angina**

Although there is no universally accepted definition of unstable angina, it has been described as a clinical syndrome between stable angina and acute myocardial infarction.

The diagram below will help distinguish between the types of acute coronary syndromes in patients presenting with acute cardiac chest pain:



Appendix B: The National Institutes of Health Stroke Scale (NIHSS)

All investigators should gain sufficient training and certification to measure NIHSS.

Administer stroke scale items in the order listed. Record performance in each category after each subscale exam. Do not go back and change scores. Follow directions provided for each exam technique. Scores should reflect what the patient does, not what the clinician thinks the patient can do. The clinician should record answers while administering the exam and work quickly. Except where indicated, the patient should not be coached (i.e., repeated requests to patient to make a special effort). (Please also see http://www.ninds.nih.gov/doctors/NIH_Stroke_Scale.pdf for pictures associated with this score)

1a. Level of Consciousness: The investigator must choose a response if a full evaluation is prevented by such obstacles as an endotracheal tube, language barrier, orotracheal trauma/bandages. A 3 is scored only if the patient makes no movement (other than reflexive posturing) in response to noxious stimulation.

0 = **Alert;** keenly responsive.

1 = **Not alert;** but arousable by minor stimulation to obey, answer, or respond.

2 = **Not alert;** requires repeated stimulation to attend, or is obtunded and requires strong or painful stimulation to make movements (not stereotyped).

3 = Responds only with reflex motor or autonomic effects or totally unresponsive, flaccid, and areflexic.

1b. LOC Questions: The patient is asked the month and his/her age. The answer must be correct - there is no partial credit for being close. Aphasic and stuporous patients who do not comprehend the questions will score 2. Patients unable to speak because of endotracheal intubation, orotracheal trauma, severe dysarthria from any cause, language barrier, or any other problem not secondary to aphasia are given a 1. It is important that only the initial answer be graded and that the examiner not "help" the patient with verbal or non-verbal cues.

0 = **Answers** both questions correctly.

1 = **Answers** one question correctly.

2 = **Answers** neither question correctly.

1c. LOC Commands: The patient is asked to open and close the eyes and then to grip and release the non-paretic hand. Substitute another one step command if the hands cannot be used. Credit is given if an unequivocal attempt is made but not completed due to weakness. If the patient does not respond to command, the task should be demonstrated to him or her (pantomime), and the result scored (i.e., follows none, one or two commands). Patients with trauma, amputation, or other physical impediments should be given suitable one-step commands. Only the first attempt is scored.

0 = **Performs** both tasks correctly.

1 = **Performs** one task correctly.

2 = **Performs** neither task correctly.

2. Best Gaze: Only horizontal eye movements will be tested. Voluntary or reflexive (oculocephalic) eye movements will be scored, but caloric testing is not done. If the patient has a conjugate deviation of the eyes that can be overcome by voluntary or reflexive activity, the score will be 1. If a patient has an isolated peripheral nerve paresis (CN III, IV or VI), score a 1. Gaze is testable in all aphasic patients. Patients with ocular trauma, bandages, pre-existing blindness, or other disorder of visual acuity or fields should be tested with reflexive movements, and a choice made by the investigator. Establishing eye contact and then moving about the patient from side to side will occasionally clarify the presence of a partial gaze palsy.

0 = **Normal.**

1 = **Partial gaze palsy;** gaze is abnormal in one or both eyes, but forced deviation or total gaze paresis is not present.

2 = **Forced deviation,** or total gaze paresis not overcome by the oculocephalic maneuver.

3. Visual: Visual fields (upper and lower quadrants) are tested by confrontation, using finger counting or visual threat, as appropriate. Patients may be encouraged, but if they look at the side of the moving fingers appropriately, this can be scored as normal. If there is unilateral blindness or enucleation, visual fields in the remaining eye are scored. Score 1 only if a clear-cut asymmetry, including quadrantanopia, is found. If patient is blind from any cause, score 3. Double simultaneous stimulation is performed at this point. If there is extinction, patient receives a 1, and the results are used to respond to item 11.

0 = **No visual loss.**

1 = **Partial hemianopia.**

2 = **Complete hemianopia.**

3 = **Bilateral hemianopia** (blind including cortical blindness).

4. Facial Palsy: Ask – or use pantomime to encourage – the patient to show teeth or raise eyebrows and close eyes. Score symmetry of grimace in response to noxious stimuli in the poorly responsive or non-comprehending patient. If facial trauma/bandages, orotracheal tube, tape or other physical barriers obscure the face, these should be removed to the extent possible.

0 = **Normal** symmetrical movements.

1 = **Minor paralysis** (flattened nasolabial fold, asymmetry on smiling).

2 = **Partial paralysis** (total or near-total paralysis of lower face).

3 = **Complete paralysis** of one or both sides (absence of facial movement in the upper and lower face).

5. Motor Arm: The limb is placed in the appropriate position: extend the arms (palms down) 90 degrees (if sitting) or 45 degrees (if supine). Drift is scored if the arm falls before 10 seconds. The aphasic patient is encouraged using urgency in the voice and pantomime,

but not noxious stimulation. Each limb is tested in turn, beginning with the non-paretic arm. Only in the case of amputation or joint fusion at the shoulder, the examiner should record the score as untestable (UN), and clearly write the explanation for this choice.

0 = **No drift;** limb holds 90 (or 45) degrees for full 10 seconds.

1 = **Drift;** limb holds 90 (or 45) degrees, but drifts down before full 10 seconds; does not hit bed or other support.

2 = **Some effort against gravity;** limb cannot get to or maintain (if cued) 90 (or 45) degrees, drifts down to bed, but has some effort against gravity.

3 = **No effort against gravity;** limb falls.

4 = **No movement.**

UN = **Amputation** or joint fusion, explain: _____

5a. Left Arm

5b. Right Arm

6. Motor Leg: The limb is placed in the appropriate position: hold the leg at 30 degrees (always tested supine). Drift is scored if the leg falls before 5 seconds. The aphasic patient is encouraged using urgency in the voice and pantomime, but not noxious stimulation. Each limb is tested in turn, beginning with the non-paretic leg. Only in the case of amputation or joint fusion at the hip, the examiner should record the score as untestable (UN), and clearly write the explanation for this choice.

0 = **No drift;** leg holds 30-degree position for full 5 seconds.

1 = **Drift;** leg falls by the end of the 5-second period but does not hit bed.

2 = **Some effort against gravity;** leg falls to bed by 5 seconds, but has some effort against gravity.

3 = **No effort against gravity;** leg falls to bed immediately.

4 = **No movement.**

UN = **Amputation** or joint fusion, explain: _____

6a. Left Leg

6b. Right Leg

7. Limb Ataxia: This item is aimed at finding evidence of a unilateral cerebellar lesion. Test with eyes open. In case of visual defect, ensure testing is done in intact visual field. The finger-nose-finger and heel-shin tests are performed on both sides, and ataxia is scored only if present out of proportion to weakness. Ataxia is absent in the patient who cannot understand or is paralyzed. Only in the case of amputation or joint fusion, the examiner should record the score as untestable (UN), and clearly write the explanation for this choice. In case of blindness, test by having the patient touch nose from extended arm position.

0 = **Absent.**

1 = **Present in one limb.**

2 = **Present in two limbs.**

UN = **Amputation** or joint fusion, explain: _____

8. Sensory: Sensation or grimace to pinprick when tested, or withdrawal from noxious stimulus in the obtunded or aphasic patient. Only sensory loss attributed to stroke is scored as abnormal and the examiner should test as many body areas (arms [not hands], legs, trunk, face) as needed to accurately check for hemisensory loss. A score of 2, "severe or total sensory loss," should only be given when a severe or total loss of sensation can be clearly demonstrated. Stuporous and aphasic patients will, therefore, probably score 1 or 0. The patient with brainstem stroke who has bilateral loss of sensation is scored 2. If the patient does not respond and is quadriplegic, score 2. Patients in a coma (item 1a=3) are automatically given a 2 on this item.

0 = **Normal;** no sensory loss.

1 = **Mild-to-moderate sensory loss;** patient feels pinprick is less sharp or is dull on the affected side; or there is a loss of superficial pain with pinprick, but patient is aware of being touched.

2 = **Severe to total sensory loss;** patient is not aware of being touched in the face, arm, and leg.

9. Best Language: A great deal of information about comprehension will be obtained during the preceding sections of the examination. For this scale item, the patient is asked to describe what is happening in the attached picture, to name the items on the attached naming sheet and to read from the attached list of sentences. Comprehension is judged from responses here, as well as to all of the commands in the preceding general neurological exam. If visual loss interferes with the tests, ask the patient to identify objects placed in the hand, repeat, and produce speech. The intubated patient should be asked to write. The patient in a coma (item 1a=3) will automatically score 3 on this item. The examiner must choose a score for the patient with stupor or limited cooperation, but a score of 3 should be used only if the patient is mute and follows no one-step commands.

0 = **No aphasia;** normal.

1 = **Mild-to-moderate aphasia;** some obvious loss of fluency or facility of comprehension, without significant limitation on ideas expressed or form of expression. Reduction of speech and/or comprehension, however, makes conversation about provided materials difficult or impossible. For example, in conversation about provided materials, examiner can identify picture or naming card content from patient's response.

2 = **Severe aphasia;** all communication is through fragmentary expression; great need for inference, questioning, and guessing by the listener. Range of information that can be exchanged is limited; listener carries burden of communication. Examiner cannot identify materials provided from patient response.

3 = **Mute, global aphasia;** no usable speech or auditory comprehension.

10. Dysarthria: If patient is thought to be normal, an adequate sample of speech must be obtained by asking patient to read or repeat words from the attached list. If the patient has severe aphasia, the clarity of articulation of spontaneous speech can be rated. Only if the patient is intubated or has other physical barriers to producing speech, the examiner should record the score as untestable (UN), and clearly write an explanation for this choice. Do not tell the patient why he or she is being tested.

0 = **Normal.**

1 = **Mild-to-moderate dysarthria;** patient slurs at least some words and, at worst, can be understood with some difficulty.

2 = **Severe dysarthria;** patient's speech is so slurred as to be unintelligible in the absence of or out of proportion to any dysphasia, or is mute/anarthric.

UN = **Intubated** or other physical barrier,
explain: _____

11. Extinction and Inattention (formerly Neglect): Sufficient information to identify neglect may be obtained during the prior testing. If the patient has a severe visual loss preventing visual double simultaneous stimulation, and the cutaneous stimuli are normal, the score is normal. If the patient has aphasia but does appear to attend to both sides, the score is normal. The presence of visual spatial neglect or anosagnosia may also be taken as evidence of abnormality. Since the abnormality is scored only if present, the item is never untestable.

0 = **No abnormality.**

1 = **Visual, tactile, auditory, spatial, or personal inattention** or extinction to bilateral simultaneous stimulation in one of the sensory modalities.

2 = **Profound hemi-inattention or extinction to more than one modality;** does not recognize own hand or orients to only one side of space.

Reference⁶⁵

Appendix C: Glasgow Coma Score

Eye movement

- 1 = None
- 2 = To pain
- 3 = To speech
- 4 = Spontaneous

Verbal response

- 1 = None
- 2 = Incomprehensible
- 3 = Inappropriate
- 4 = Confused
- 5 = Orientated

Motor response

- 1 = None
- 2 = Extension
- 3 = Flexor response
- 4 = Withdrawal
- 5 = Localises pain
- 6 = Obeys commands

Score out of 15 (range 3 – 15)

Reference⁶⁶

Appendix D: Modified Rankin Scale (mRS)

All investigators should gain sufficient training and certification to measure mRS.

- 0 No symptoms at all
- 1 No significant disability, despite symptoms; able to carry out all usual duties and activities
- 2 Slight disability; unable to carry out all previous activities but able to look after own affairs without assistance
- 3 Moderate disability; requiring some help, but able to walk without assistance
- 4 Moderately severe disability; unable to walk without assistance and unable to attend to own bodily needs without assistance
- 5 Severe disability; bedridden, incontinent and requiring constant nursing care and attention
- 6 Dead

Score 0 to 6 (range 0-6)

Reference ⁶⁷⁻⁶⁸

Appendix E: Barthel Index

Task	Criteria	Score
Bowels	Incontinent	0
	Occasional accident (once per week)	5
	Continent	10
Bladder	Incontinent, or catheterised and unable to manage alone	0
		5
	Occasional accident (maximum once per 24 hours)	10
	Continent	
Grooming	Needs help with personal care	0
	Independent face/hair/teeth/shaving (implements provided)	5
Toilet use	Dependent	0
	Needs some help, but can do something alone	5
	Independent (on and off, dressing, wiping)	10
Feeding	Unable	0
	Needs help cutting, spreading butter, etc.	5
	Independent	10
Transfer (bed to chair and back)	Unable, no sitting balance	0
	Major help (one or two people, physical), cab sit	5
	Minor help (verbal or physical)	10
	Independent	15
Mobility	Immobile	0
	Wheelchair independent, including corners	5
	Walks with help of one person (verbal or physical)	10
	Independent (but may use any aid: for example stick)	15
Dressing	Dependent	0
	Needs help but can do about half unaided	5
	Independent (including buttons, zips, laces, etc.)	10
Stairs	Unable	0
	Needs help (verbal, physical, carrying aid)	5
	Independent	10
Bathing	Dependent	0
	Independent (or in shower)	5

Score out of 100 (range 0-100)

Reference⁶⁹

Appendix F: EuroQOL

Group 1 ⁷⁰

I have no problems in walking about
 I have some problems in walking about
 I am confined to bed

Group 2

I have no problems with self care
 I have some problems with washing or dressing
 I am unable to wash or dress myself

Group 3

I have no problems performing my usual activities (e.g. work, study, housework, family or leisure activities)
 I have some problems performing usual activities
 I am unable to perform my usual activities

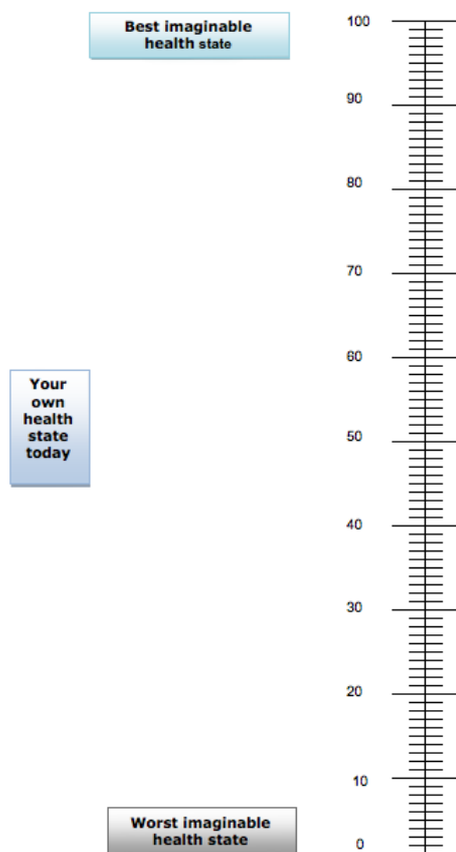
Group 4

I have no pain or discomfort
 I have moderate pain or discomfort
 I have extreme pain or discomfort

Group 5

I am not anxious or depressed
 I am moderately anxious or depressed
 I am extremely anxious or depressed

Health state today by visual analogue scale (best imaginable to worst imaginable)



Appendix G: Cognitive Testing

TICS-M⁷¹ – Adjusted for the TARDIS Trial

Please note that this test is designed for telephone use. In the event follow up is done in person the entire test must be completed verbally, i.e. the memory words must not be shown to the patient.

Question and Instructions Score

Orientation: Please ask them what day, date etc it is

7

Day

Date

Month

Season

Year

Age

Telephone Number (code+number)

Registration

10

I am going to read you a list of 10 words. Please listen carefully and try to remember them. When I am done, tell me as many as you can in any order. Ready?

Cabin

Pipe

Elephant

Chest

Silk

Theatre

Watch

Whip

Pillow

Giant

Attention and Calculation

6

Please take away 7 from 100. Now continue to take 7 away from what you have left over until I ask you to stop

93

86

79

72

65

Count backwards Please count back 20-1

No mistakes

Comprehension, Semantic and Recent Memory

5

What do people use to cut paper?

Scissors

What is the prickly green plant found in the desert?

Cactus

Who is the Prime Minister?

Correct surname

Who is the reigning monarch?

E, QE, QE2

What is the opposite direction to east? t

West

Language/Repetition

1

Please listen carefully and repeat No ifs ands or buts'

Score only if exactly right

Delayed Recall

10

Please repeat as many of the 10 words I asked you to remember earlier

Cabin

Pipe

Elephant	<input type="checkbox"/>
Chest	<input type="checkbox"/>
Silk	<input type="checkbox"/>
Theatre	<input type="checkbox"/>
Watch	<input type="checkbox"/>
Whip	<input type="checkbox"/>
Pillow	<input type="checkbox"/>
Giant	<input type="checkbox"/>

Total Score (1 point for each correct answer)

/39

Verbal Fluency

Letter

I'd like you to generate as many words as possible beginning with the letter 'P'. You have got a minute. Are you ready?

Write down each word and score 1 mark for each word. Do not score repetitions.

Animals

I'd like you to generate as many animals as possible, any kind of animal, beginning with any letter, it doesn't matter'. You've got a minute. Are you ready?

Write down each word and score 1 mark for each animal named. Do not score repetitions.

Total score _____

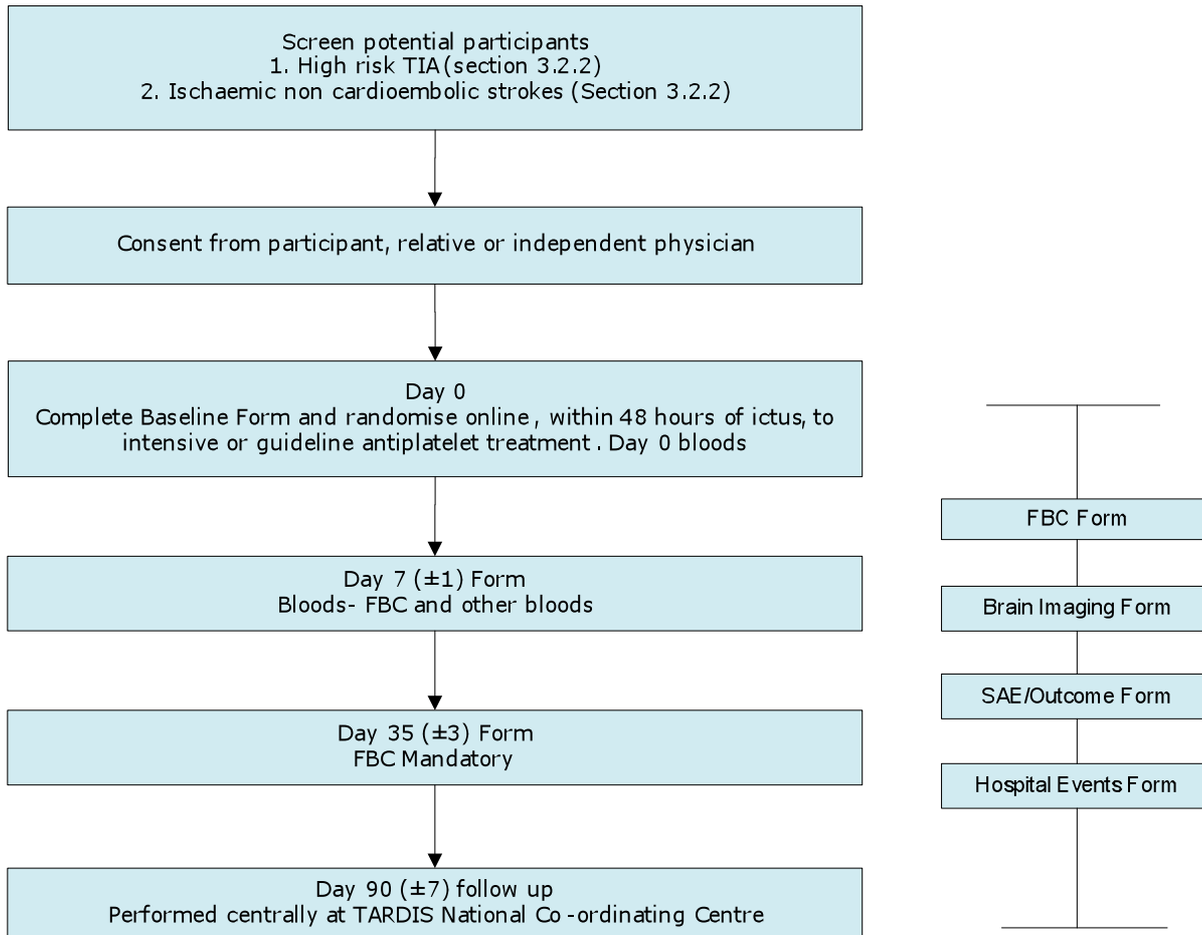
Appendix H: Zung Depression rating Scale (short)

	Seldom or never	Some of the time	Good part of the time	Most of the time
I feel down-hearted and blue	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Morning is when I feel best	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
I have trouble sleeping at night	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
I can eat as much as I used to	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
I get tired for no reason	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
I find it difficult to make decisions	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
I feel hopeful about the future	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
I feel that I am useful and needed	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
My life is somewhat empty	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
I still enjoy the things I used to do	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>

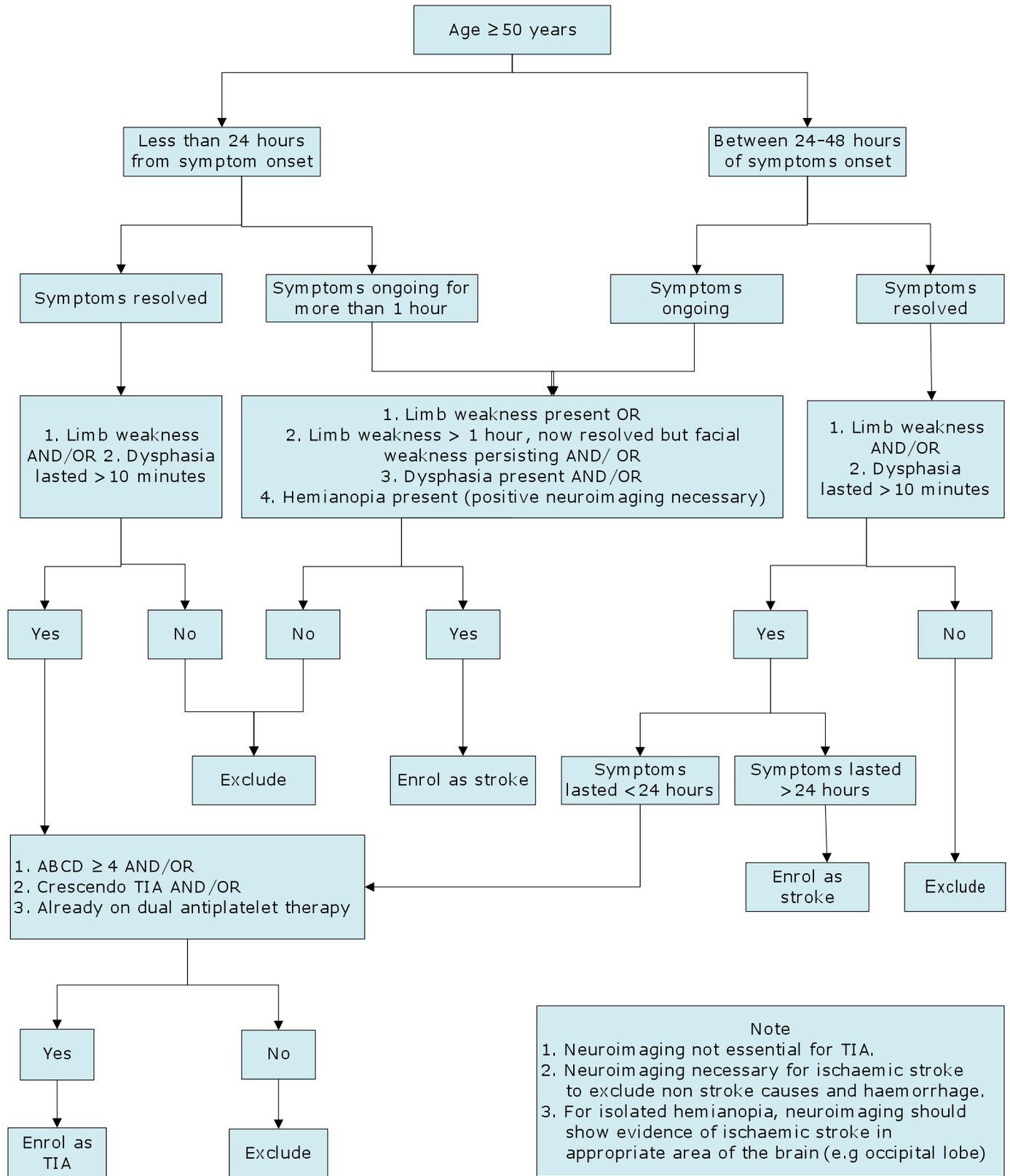
Reference⁴⁴Zung IDS Index = $100 \times \text{Total} / 40$

Depression => 70

Appendix I: Trial Flow



APPENDIX J: Trial inclusion flow chart



APPENDIX K: Sample Labels**CLOPIDOGREL LOADING DOSE** (taken on day 0, day of randomisation):**Eudract no: 2007-006749-42****TARDIS STUDY****4 x Clopidogrel 75mg tablets****Take Four tablets as a loading dose.**

Name.....Date.....

BN.....EXP.....

Clinical Trial use only

Investigator Prof P

Bath

KEEP OUT OF THE REACH OF CHILDREN

Do not store above 25⁰c

Pharmacy Dept, City Hospital Campus, NUH, Hucknall Rd, Nottm

NG5 1PB 0115 9691169.

Or

Eudract no: 2007-006749-42**TARDIS STUDY****1 x Clopidogrel 300mg tablet****Take one tablet as a loading dose.**

Name.....Date.....

BN.....EXP.....

Clinical Trial use only

Investigator Prof P

Bath

KEEP OUT OF THE REACH OF CHILDREN

Do not store above 25⁰c

Pharmacy Dept, City Hospital Campus, NUH, Hucknall Rd, Nottm

NG5 1PB 0115 9691169.

CLOPIDOGREL (days 1 to 30)**Eudract no: 2007-006749-42****TARDIS STUDY****30 x Clopidogrel 75mg tablets****Take ONE tablet DAILY.**

Name.....Date.....

BN.....EXP.....

Clinical Trial use only

Investigator Prof P

Bath

KEEP OUT OF THE REACH OF CHILDREN

Do not store above 25⁰c

Pharmacy Dept, City Hospital Campus, NUH, Hucknall Rd, Nottm

NG5 1PB 0115 9691169.

ASPIRIN LOADING DOSE (taken on day 0, day of randomisation):**Eudract no: 2007-006749-42****TARDIS STUDY****4 x Aspirin 75mg tablets****Take Four tablets as a loading dose.**

Name.....Date.....

BN.....EXP.....

Clinical Trial use only

Investigator Prof P

Bath

KEEP OUT OF THE REACH OF CHILDREN

Do not store above 25⁰c

Pharmacy Dept, City Hospital Campus, NUH, Hucknall Rd, Nottm

NG5 1PB 0115 9691169.

Or

Eudract no: 2007-006749-42**TARDIS STUDY****1 x Aspirin 300mg tablet****Take one tablet as a loading dose.**

Name.....Date.....

BN.....EXP.....

Clinical Trial use only

Investigator Prof P

Bath

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ASPIRIN (days 1 to 30)**Eudract no: 2007-006749-42****TARDIS STUDY****30 x ASPIRIN 75mg tablets****Take ONE tablet DAILY.**

Name.....Date.....

BN.....EXP.....

Clinical Trial use only

Investigator Prof P

Bath

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DIPYRIDAMOLE (days 0-30):**Eudract no: 2007-006749-42****TARDIS STUDY****60 x DIPYRIDAMOLE 200 mg tablets****Take two tablets daily.**

Name.....Date.....

BN.....EXP.....

Clinical Trial use only

Investigator Prof P

Bath

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REFERENCES

1. Albers GW, Caplan LR, Easton JD, Fayad PB, Mohr JP, Saver JL, et al. Transient ischemic attack--proposal for a new definition.[see comment]. *New England Journal of Medicine*. 2002;347:1713-1716
2. Johnston SC, Rothwell PM, Nguyen-Huynh MN, Giles MF, Elkins JS, Bernstein AL, et al. Validation and refinement of scores to predict very early stroke risk after transient ischaemic attack. *Lancet*. 2007;369:283-292
3. EAFT (European Atrial Fibrillation Trial) Study Group. Secondary prevention in non-rheumatic atrial fibrillation after tia or minor stroke. *Lancet*. 1993;342:1255-1262
4. Bath PMW, Zhao L, Heptinstall S. Current status of stroke prevention in patients with atrial fibrillation. *European Heart Journal*. 2005;7:C12-C18
5. Zhao L, Heptinstall S, Bath P. Antiplatelet therapy for stroke prevention. *British Journal of Cardiology*
Heart & Brain. 2005;12:57-60
6. Antithrombotic Trialists Collaboration. Collaborative meta-analysis of randomised trials of antiplatelet therapy for prevention of death, myocardial infarction, and stroke in high risk patients. *BMJ*. 2002;324:71-86
7. CAPRIE Steering Committee. A randomised, blinded, trial of clopidogrel versus aspirin in patients at risk of ischaemic events (caprie). *Lancet*. 1996;348:1329-1339
8. Ringleb PA, Bhatt DL, Hirsh AT, Topol EJ, Hacke W, for the CAPRIE Investigators. Benefit of clopidogrel over aspirin is amplified in patients with a history of ischemic events. *Stroke*. 2004;35:528-532
9. Diener HC, Cunha L, Forbes C, Sivenius J, Smets P, Lowenthal A. European stroke prevention study 2. Dipyridamole and acetylsalicylic acid in the secondary prevention of stroke. *J Neurological Sciences*. 1996;143:1-13
10. The ESPRIT Study Group. Aspirin plus dipyridamole versus aspirin alone after cerebral ischaemia of arterial origin (esprit): Randomised controlled trial. *Lancet*. 2006;367:1665-1673
11. Halkes PH, Gray LJ, Bath PM, Diener HC, Guiraud-Chaumeil B, Yatsu FM, et al. Dipyridamole plus aspirin versus aspirin alone in secondary prevention after tia or stroke: A meta-analysis by risk. *J Neurol Neurosurg Psychiatry*. 2008;79:1218-1223
12. Yusuf S, Fox KAA, Tognoni G, Mehta SR, Chrolavicius S, Anand S, et al. Effects of clopidogrel in addition to aspirin in patients with acute coronary syndromes without st-segment elevation. *New England Journal of Medicine*. 2001;345:494-502
13. Steinhubl SR, Berger PB, Mann JT, Fry ETA, DeLago A, Wilmer C, et al. Early and sustained dual oral antiplatelet therapy following percutaneous coronary intervention: A randomized controlled trial. *JAMA*. 2002;288:2411-2420
14. Bhatt DL, Fox KAA, Werner Hacke CB, Berger PB, Black HR, Boden WE, et al. Clopidogrel and aspirin versus aspirin alone for the prevention of atherothrombotic events. *The New England Journal of Medicine*. 2006;354:1706-1717
15. Diener HC, Bogousslavsky J, Brass LM, Cimminiello C, Csiba L, Kaste M, et al. Aspirin and clopidogrel compared with clopidogrel alone after recent ischaemic stroke or transient ischaemic attack in high-risk patients (match): Randomised, double-blind, placebo-controlled trial. *Lancet*. 2004;364:331-337
16. Bath PMW. Role of aspirin in match. *Lancet*. 2004;364:1662
17. Intercollegiate Stroke Working Party. National clinical guidelines for stroke. 2004
18. Leys D, Kwiecinski H, Bogousslavsky J, Bath PMW, Brainin M, Diener H-C, et al. Prevention. European stroke initiative *Cerebrovasc Dis*. 2004;17:15-29
19. European Stroke Organisation (ESO) Executive Committee, Committee EW. Guidelines for management of ischaemic stroke and transient ischaemic attack 2008. *Cerebrovascular Diseases*. 2008;25:457-507
20. Furie KL, Kasner SE, Adams RJ, Albers GW, Bush RL, Fagan SC, et al. Guidelines for the prevention of stroke in patients with stroke or transient ischemic attack: A

- guideline for healthcare professionals from the american heart association/american stroke association. *Stroke*. 2011;42:227-276
21. Sacco RL, Diener H-C, Yusuf S, Cotton D, Ounpuu S, Lawton W, et al. Aspirin and extended-release dipyridamole versus clopidogrel for recurrent stroke. *New England Journal of Medicine*. 2008;359:1238-1251
 22. International Stroke Trial Collaborative Group. The international stroke trial (ist); a randomised trial of aspirin, subcutaneous heparin, both, or neither among 19435 patients with acute ischaemic stroke. *Lancet*. 1997;349:1569-1581
 23. CAST (Chinese Acute Stroke Trial) Collaborative Group. Cast: Randomised placebo-controlled trial of early aspirin use in 20,000 patients with acute ischaemic stroke. *Lancet*. 1997;349:1641-1649
 24. Kennedy J, Hill MD, Ryckborst K, Eliasziw M, Demchuk AM, Buchan AM, et al. Fast assessment of stroke and transient ischaemic attack to prevent early recurrence (faster): A randomised controlled pilot trial. *Lancet Neurology*. 2007;6:961-969
 25. Dengler R, Diener HC, Schwartz A, Grond M, Schumacher H, Machnig T, et al. Early treatment with aspirin plus extended-release dipyridamole for transient ischaemic attack or ischaemic stroke within 24 h of symptom onset (early trial): A randomised, open-label, blinded-endpoint trial. *Lancet Neurol*. 2010;9:159-166
 26. Bath PM, Cotton D, Martin RH, Palesch Y, Yusuf S, Sacco R, et al. Effect of combined aspirin and extended-release dipyridamole versus clopidogrel on functional outcome and recurrence in acute, mild ischemic stroke: Profess subgroup analysis. *Stroke*. 2010;41:732-738
 27. Rothwell PM, Giles MF, Chandratheva A, Marquardt L, Geraghty O, Redgrave JN, et al. Effect of urgent treatment of transient ischaemic attack and minor stroke on early recurrent stroke (express study): A prospective population-based sequential comparison. *Lancet Neurology*. 2007;370:1432-1442
 28. Lavalley PC, Meseguer E, Abboud H, Cabrejo L, Olivot J, Simon O, et al. A transient ischaemic attack clinic with round-the-clock access (sos-tia): Feasibility and effects. *Lancet Neurology*. 2007;6:953-960
 29. Markus HS, Droste DW, Kaps M, Larrue V, Lees KR, Siebler M, et al. Dual antiplatelet therapy with clopidogrel and aspirin in symptomatic carotid stenosis evaluated using doppler embolic signal detection. The clopidogrel and aspirin for reduction of emboli in symptomatic carotid stenosis (caress) trial *Circulation*. 2005;111:2233-2240
 30. Wong KS, Chen C, Fu J, Chang HM, Suwanwela NC, Huang YN, et al. Clopidogrel plus aspirin versus aspirin alone for reducing embolisation in patients with acute symptomatic cerebral or carotid artery stenosis (clair study): A randomised, open-label, blinded-endpoint trial. *Lancet Neurol*. 2010;9:489-497
 31. Michelson AD, Cattaneo M, Eikelboom JW, Gurbel PA, Kottke-Marchant K, Kunicki TJ, et al. Aspirin resistance: Position paper of the working group on aspirin resistance. *Journal of Thrombosis and Haemostasis*. 2005;3:1309-1311
 32. Leonardi-Bee J, Bath PM, Bousser MG, Davalos A, Diener H-C, Guiraud-Chaumeil B, et al. Dipyridamole for preventing recurrent ischemic stroke and other vascular events: A meta-analysis of individual patient data from randomized controlled trials. *Stroke*. 2005;36:162-168
 33. Zhao L, Bath P, Heptinstall S. Effects of combining three different antiplatelet agents on platelets and leukocytes in whole blood in vitro. *British Journal Pharmacology*. 2001;134:353-358
 34. Scholz T, Zhao L, Temmler U, Bath P, Heptinstall S, Losche W. The gpiib/iiiia antagonist eptifibatide markedly potentiates platelet-leukocyte interaction and tissue factor expression following platelet activation in whole blood in vitro. *Platelets*. 2002;13:401-406
 35. Zhao L, Bath PMW, Fox S, May J, Judge H, Losche W, et al. The effects of gpIIb/IIIa antagonists and a combination of three other antiplatelet agents on platelet-leukocyte interactions. *Current Medical Research Opinion*. 2003;19:178-186
 36. Zhao L, Fletcher S, Weaver C, Leonardi-Bee J, May J, Fox S, et al. Effects of aspirin, clopidogrel and dipyridamole administered singly and in combination on platelet and

- leucocyte function in normal volunteers and patients with prior ischaemic stroke. *Thromb Haemost.* 2005;93:527-534
37. Zhao L, Gray LJ, Leonardi-Bee J, Weaver CS, Heptinstall S, Bath PM. Effect of aspirin, clopidogrel and dipyridamole on soluble markers of vascular function in normal volunteers and patients with prior ischaemic stroke. *Platelets.* 2006;17:100-104
 38. Sprigg N, Gray LJ, England T, Willmot MR, Zhao L, Sare GM, et al. A randomised controlled trial of triple antiplatelet therapy (aspirin, clopidogrel and dipyridamole) in the secondary prevention of stroke: Safety, tolerability and feasibility. *PLoS ONE [Electronic Resource]*. 2008;3:e2852
 39. Willmot M, Zhao L, Heptinstall S, Bath PMW. Triple antiplatelet therapy for secondary prevention of recurrent ischaemic stroke. *J Stroke Cerebrovasc Dis.* 2004;13:138-140
 40. Geeganage C, Wilcox R, Bath PM. Triple antiplatelet therapy for preventing vascular events: A systematic review and meta-analysis. *BMC Med.* 2010;8:36
 41. Hallas MJ, Dall M, Andries A, Andersen BS, Aalykke C, Hansen JM, et al. Use of single and combined antithrombotic therapy and risk of serious upper gastrointestinal bleeding: Population based case-control study. *British Medical Journal.* 2006;333:726
 42. Bath PMW, Geeganage C, Gray LJ, Collier T, Pocock SJ. Optimising the analysis of stroke prevention trials: Converting dichotomous vascular outcomes into ordinal measures. *Stroke.* 2008;In press
 43. Brooks R, with the EuroQol Group. Euroqol: The current state of play. *Health Policy.* 1996;37:53-72
 44. Zung WWK. A self-rating depression scale. *Archives of General Psychiatry.* 1965;12:63-70
 45. Schulman S, Kearon C, on behalf of the subcommittee on control of anticoagulation of the scientific and standardization committee of the international society on thrombosis and haemostasis. Definition of major bleeding in clinical investigations of antihemostatic medicinal products in non-surgical patients. *Journal of Thrombosis and Haemostasis.* 2005;3:592-694
 46. Weir CJ, Lees KR. Comparison of stratification and adaptive methods for treatment allocation in an acute stroke clinical trial. *Stat.Med.* 2003;22:705-726
 47. Diener H-C, Sacco RL, Yusuf S, for the Steering Committee and PROFESS Study Group. Rationale, design and baseline data of a randomized, double-blind, controlled trial comparing two antithrombotic regimens (a fixed-dose combination of extended-release dipyridamole plus asa with clopidogrel) and telmisartan versus placebo in patients with strokes: The prevention regimen for effectively avoiding second strokes trial (profess). *Cerebrovascular Diseases.* 2007;23:368-380
 48. Sung JJ. Combining aspirin with antithrombotic agents. *British Medical Journal.* 2006;333:726
 49. Adams HP, Bendixen BH, Kappelle LJ, Biller J, Love BB, Gordon DL, et al. Classification of subtype of acute ischemic stroke. Definitions for use in a multicenter clinical trial. *Stroke.* 1993;24:35-41
 50. Collet JP, Hulot JS, Pena A, Villard E, Esteve JB, Payo L, et al. Cytochrome p450 2c19 polymorphism in young patients treated with clopidogrel after myocardial infarction: A cohort study. *Lancet.* 2009;373:309-317
 51. Simon T, Verstuyft C, Mary-Krause M, Quteineh L, Drouet E, Meneveau N, et al. Genetic determinants of response to clopidogrel and cardiovascular events. *New England Journal of Medicine.* 2009;360:363-375
 52. Mega JL, Close SL, Wiviott SD, Shen L, Hockett RD, Brandt JT, et al. Cytochrome p-450 polymorphisms and response to clopidogrel. *New England Journal of Medicine.* 2009;360:354-362
 53. Abraha HD, Butterworth RJ, Bath PMW, Wassif WS, Garthwaite J, Sherwood RA. Serum s-100 protein, a prognostic marker of clinical outcome in acute stroke. *Annals of Clinical Biochemistry.* 1997;34:366-370
 54. Weaver CS, Leonardi-Bee J, Bath-Hexall FJ, Bath PMW. Sample size calculations in acute stroke trials: A systematic review of their reporting, characteristics, and relationship with outcome. *Stroke.* 2004;35:1216-1224

55. Sprigg N, Gray LJ, England T, Willmot MR, Zhao L, Sare GM, et al. A randomised controlled trial of triple antiplatelet therapy (aspirin, clopidogrel and dipyridamole) in the secondary prevention of stroke: Safety, tolerability and feasibility (isrctn 83673558). *PLoS One*. 2008;submitted
56. DAMOCLES study group. A proposed charter for clinical trial data monitoring committees: Helping them to do their job well. *Lancet*. 2005;365:711-722
57. Grant AM, Altman D, G, Babiker AB, Campbell MK, Clemens FJ, Darbyshire JH, et al. Issues in data monitoring and interim analysis of trials. *Health Technology Assessment*. 2005;9:1-237
58. Hatano S. Experience from a multicentre stroke register: A preliminary report. *Bulletin of the World Health Organisation*. 1976;54:541-553
59. Hacke W, Kaste M, Bluhmki E, Brozman M, Davalos A, Guidetti D, et al. Thrombolysis with alteplase 3 to 4.5 hours after acute ischemic stroke. *New England Journal of Medicine*. 2008;359:1317-1329
60. Hacke W. European cooperative acute stroke trial (ecass) (abstract). *Stroke*. 1994;25:542
61. Hacke W, Markku K, Fieschi C, von Kummer R, Davalos A, Meier D, et al. Randomised double-blind placebo-controlled trial of thrombolytic therapy with intravenous alteplase in acute ischaemic stroke (ecass ii). *Lancet*. 1998;352:1245-1251
62. Rothwell PM, Giles MF, Flossmann E, Lowelock CE, Redgrave JNE, Warlow CP, et al. A simple score (abcd) to identify individuals at high early risk of stroke after transient ischaemic attack. *Lancet*. 2005;366:29-36
63. Johnston SC, Rothwell PM, Nguyen-Huynh MN, Giles MF, Elkins JS, Bernstein AL, et al. Validation and refinement of scores to predict very early stroke risk after transient ischaemic attack. *Lancet*. 2007;369:283-292
64. Anonymous. Myocardial infarction redefined--a consensus document of the joint European society of cardiology/American college of cardiology committee for the redefinition of myocardial infarction. *European Heart Journal*. 2000;21:1502-1513
65. Brott T, Adams HP, Olinger CP, Marler JR, Barsan WG, Biller J, et al. Measurements of acute cerebral infarction: A clinical examination scale. *Stroke*. 1989;20:864-870
66. Teasdale G, Jennett B. Assessment of coma and impaired consciousness. A practical scale. *Lancet*. 1974;2:81-83
67. Rankin J. Cerebral vascular accidents in patients over the age of 60. 2. Prognosis. *Scottish Medical Journal*. 1957;2:200-215
68. Wade DT. *Measurement in neurological rehabilitation*. Oxford: Oxford University Press; 1992.
69. Mahoney FI, Barthel DW. Functional evaluation: The barthel index. *Maryland State Medical Journal*. 1965:61-65
70. Dorman PJ, Slattery J, Farrell B, Dennis MS, Sandercock PAG, United Kingdom Collaborators in the International Stroke Trial. A randomised comparison of the euroqol and short form-36 after stroke. *Br.Med.J*. 1997;315:461
71. de Jager CA, Budge MM, Clarke R. Utility of tics-m for the assessment of cognitive function in older adults. *International Journal of Geriatric Psychiatry*. 2003;18:318-324