

## Form for participants who are not able to give written informed consent – Best Interest

Lay title:	MAnagement of Systolic blood pressure during Thrombectomy by Endovascular Route for acute ischaemic STROKE (MASTERSTROKE Trial							
Short title	MASTERSTROKE							
Locality:	Auckland City Hospita	Ethics committee ref.: 19NTB	163					
		Local Project number: A+ 817	3					
Lead investigator:	Dr Douglas Campbell	Contact phone number: 375709	)5					
PARTICIPANT INFORMATION - Investigator to complete								
Name of participant								
Participant <b>Number (if</b>	applicable)							
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What is the nature of the participant's <b>inability</b> to give own written informed consent?		The participant has suffered an acute ischaemic stroke and will be undergoing endovascular thrombectomy and is unable to provide their informed consent to be enrolled in the MasterStroke study to control Blood Pressure during clot retrieval. These patients may be confused and disoriented in a time critical period and thus not competent to make an informed choice and give informed consent.						
Note that as far as possible the wishes of the participant should still be taken into account in making the decision to be as to whether enroll them in this study								
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1. BEST INTEREST			<u> </u>					
Due to the patients inability to provide their own consent at this point in time and in accordance with NZ law and code of right 7 (4), the participant may be enrolled in the MasterStroke Study without signed consent, provided that their enrolment is clearly in their best medical interests. Progress to Step 2								
2. <b>PROCEDURE F</b>	OR TREATMENT WITHOU	T CONSENT						
Is there a family member(s) to consult regarding what the participant would have wanted done in this situation (or, if there is no suitable family member, a friend of the participant, or other person such as their GP who is familiar with their medical wishes)?  Whenever possible provide family with a brief overview of what is happening.								
If YES, fill in the three lin	nes immediately below be	fore completing the remainder of this form and then progress to Ste	p 3.					
Name of family member of friend consulted  Views of family/friend consulted:								

(please also document this in the patient clinical notes)								
picuse uiso document tins in the patient clinical notesy								
Date and time of discussion								
If NO, the participant may be treated without signed consent, provided that the treatment is clearly in their best medical								
interests. In that case progress to Step 3.								
3. INDEPENDENT CLINICIAN CONFIRMATION								
An agreement from an independent clinician (who is not directly involved in the study) should confirm that enrolment in the								
MasterStroke s	MasterStroke study to control Blood Pressure during clot retrieval is in the patient's best interests at this time.							
Please complete the details below of the name and signature of the independent clinician affirming study participation is in the								
participant's best interest and then proceed to Step 4.								
Name of independent clinician (please print)								
<u> </u>								
Signature		Date						
	INVESTIGATOR ASSESSMENT							
I believe that the inclusion of the participant in the research project would be in the best interests of the participant.								
In forming this view in relation to the best interests of the participant, I have taken into account:								
a) the wishes of the participant, so as far as they can be ascertained								
b) the nature and degree of any <b>significant benefits, discomforts and risks</b> to the participant in								
<ul> <li>b) the nature and degree of any significant benefits, discomforts and risks to the participant in participating in the research project</li> </ul>								
c) any other consequences to the participant if they do or don't participate in the research								
I have no reason to believe that including the participant in the research project would be against their wishes.								
	vestigator I will ensure that families members are ke							
the trial and when the patient has become stable that both the patient and family are given the opportunity to ask questions and offer ongoing agreement to continue in the study.as soon as reasonably practicable of:								
a) the participant's inclusion in the research project; and								
b)	b) the option to refuse consent for participation to be continued and withdraw the participant							
5,	without compromising the participant's ability to re	·						
	care.							
Name and signature of the investigator who is confirming participant eligibility								
radic and signature of the investigator who is commining participant enginnity								
Name of Trial Investigator (please print)								
Signature	Signature Date							

The investigator must also ensure that a copy of this document forms part of the participant's clinical records

## 5. **PATIENT REGAINS CAPACITY TO CONSENT**

If/when the participant subsequently regains the capacity to give written informed consent (e.g. post-enrolment); they should do so using the **Provision for follow-up**.