TiMing of Intervention in patients with Acute **Coronary Syndromes**

Submission date	Recruitment status No longer recruiting	[] Prospectively registered	
27/11/2006		[_] Protocol	
Registration date	Overall study status Completed	Statistical analysis plan	
27/11/2006		[X] Results	
Last Edited 10/04/2019	Condition category Circulatory System	[] Individual participant da	

Plain English summary of protocol

Not provided at time of registration

Contact information

Type(s) Scientific

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Additional identifiers

EudraCT/CTIS number

IRAS number

ClinicalTrials.gov number NCT00552513

Secondary identifying numbers MCT-79654

Study information

Scientific Title

An International randomised trial of early versus delayed invasive strategies in patients with non-ST segment elevation acute coronary syndromes

Acronym

TIMACS

Study objectives

1. In patients with acute coronary syndromes (ACS), a strategy of routine early coronary angiography (less than 24 hours after randomisation) and intervention is superior to a strategy of delayed coronary angiography (more than 36 hours after randomisation) and intervention in preventing major cardiovascular events

2. In patients with ACS, a delayed invasive strategy will result in lower rates of major bleeding versus a strategy of early angiography and revascularisation

3. A strategy of early coronary angiography and revascularisation will be more cost effective than a strategy of delayed coronary angiography and revascularisation

Ethics approval required

Old ethics approval format

Ethics approval(s)

1. Research Ethics Board of the Hamilton Health Sciences/McMaster Health Sciences, Hamilton, Ontario, Canada, 26/05/2005

2. Comité de Ética em Pesquisa, Santa Casa de Belo Horizonte, Belo Horizonte, Brazil, 29/05/2006

3. Comité d'Ethique Médicale, Centre Hospitauer Regional De Huy, De Huy, Belgium, 12/10/2005 4. Ethics Committee of the Middle Slovak Institute of Cardiovascular Diseases, Banska Bistrica, Slovakia, 26/10/2006

5. University Clinical Emergency Hospital Mures Clinic of Cardiology, Targu Mures, Romania, 01 /11/2006

Study design

Therapeutic management strategy and procedures intervention type randomised parallel twoarmed multicentre multi-national trial with accessor blinding

Primary study design Interventional

Secondary study design Randomised controlled trial

Study setting(s) Hospital

Study type(s) Treatment

Participant information sheet

Not available in web format, please use the contact details below to request a patient information sheet

Health condition(s) or problem(s) studied

Acute coronary syndromes (unstable angina and non-ST segment elevation myocardial infarction)

Interventions

Early intervention: Coronary angiography and intervention as soon as possible (within 24 hours of randomisation).

Delayed intervention: Coronary angiography and intervention any time after 36 hours after randomisation.

Intervention Type

Procedure/Surgery

Primary outcome measure

The first occurrence of the composite death/myocardial (re-)infarction/stroke up to day 180.

Secondary outcome measures

1. The composite of death, myocardial (re-)infarction, stroke, refractory ischaemia or repeat revascularisation at 180 days

2. The first occurrence of any component of the composite of death myocardial infarction and refractory ischaemia until day 14, 30, 90 and at six months (day 180)

3. Stroke at 30 days and 180 days

4. Need for mechanical or pharmacological coronary revascularisation (i.e., percutaneous coronary intervention [PCI], coronary artery bypass graft [CABG], thrombolysis) at days 30, 90 and 180 days

Overall study start date

15/06/2005

Completion date

01/05/2008

Eligibility

Key inclusion criteria

1. Patients presenting or admitted to hospital with symptoms suspected to represent an acute coronary syndrome (unstable angina or MI without persistent ST elevation), i.e. clinical history consistent with new onset, or a worsening pattern, of characteristic ischaemic chest pain or ischaemic symptoms occurring at rest or with minimal activity (lasting longer than five minutes or requiring sublingual nitroglycerin for relief of the pain)

2. Able to randomise within 24 hours of the onset of the most recent episode of symptoms 3. At least two of the three following additional criteria:

3.1. Age more than or equal to 60 years, either sex

3.2. Troponin T or I or creatine kinase - myocardial bands (CK-MB) above the upper limit of normal for the local institution

3.3. Electrocardiogram (ECG) changes compatible with ischaemia (i.e., ST depression at least 1 mm in two contiguous leads or T wave inversion mroe than 3 mm or any dynamic ST shift or transient ST elevation)

4. Written informed consent dated and signed

Participant type(s)

Patient

Age group

Adult

Sex

Both

Target number of participants 4000

Key exclusion criteria

1. Age less than 21 years

2. Not a suitable candidate for revascularisation

3. Co-morbid condition with life expectancy less than six months

Date of first enrolment

15/06/2005

Date of final enrolment 01/05/2008

Locations

Countries of recruitment Argentina

Belgium

Brazil

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Bul	lgaria
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Canada

Chile

China

Czech Republic

France

Germany

Greece

India

Poland

Romania

Slovakia

Slovenia

Switzerland

United States of America

Study participating centre Hamilton Health Sciences Hamilton, Ontario Canada L8L 2X2

Sponsor information

Organisation Population Health Research Institute (PHRI) (Canada)

Sponsor details

McMaster University/Hamilton Health Sciences 237 Barton St East Hamilton, Ontario Canada L8L 2X2 +1 (0)905 527 4322 ext. 44555 crackbe@phri.ca

Sponsor type Research organisation

Website http://www.phri.ca

ROR https://ror.org/03kwaeq96

Funder(s)

Funder type Research organisation

Funder Name Canadian Institutes of Health Research

Alternative Name(s) Instituts de Recherche en Santé du Canada, Canadian Institutes of Health Research (CIHR), CIHR_IRSC, Canadian Institutes of Health Research | Ottawa ON, CIHR, IRSC

Funding Body Type Government organisation

Funding Body Subtype National government

Location Canada

Results and Publications

Publication and dissemination plan Not provided at time of registration

Intention to publish date

Individual participant data (IPD) sharing plan

IPD sharing plan summary Not provided at time of registration

Study outputs					
Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
<u>Basic results</u>				No	No
<u>Results article</u>	results	21/05/2009	10/04/2019	Yes	No