National Clinical Guideline Centre

Myocardial infarction with ST-segment elevation

The acute management of myocardial infarction with ST-segment elevation

Clinical guideline 167

Appendices A - H

July 2013

November 2020: NICE's original guidance on Myocardial infarction with ST-segment elevation was published in 2013. See the NICE website for the guideline recommendations and for the 2020 Acute coronary syndromes update. This document preserves evidence reviews and committee discussions from the 2013 guideline.

Commissioned by the National Institute for Health and Care Excellence











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Appendices

Appendix A: Scope

NATIONAL INSTITUTE FOR HEALTH AND CLINICAL EXCELLENCE

Centre for Clinical Practice

SCOPE

Clinical guideline title: Myocardial infarction with ST-segment-elevation: the acute management of myocardial infarction with ST-segment-elevation

Quality standard title: Management of acute coronary syndromes including myocardial infarction

1 Introduction

.1 Clinical guidelines

Clinical guidelines are recommendations by NICE on the appropriate treatment and care of people with specific diseases and conditions within the NHS. They are based on the best available evidence.

This scope defines what the guideline will (and will not) examine, and what the guideline developers will consider. The scope is based on the referral from the Department of Health.

.2 Quality standards

Quality standards are a set of specific, concise quality statements and measures that act as markers of high-quality, cost-effective patient care, covering the treatment and prevention of different diseases and conditions.

For this clinical guideline a NICE quality standard will be produced during the guideline development process, after the development of the clinical guideline recommendations.

This scope defines the areas of care for which specific quality statements and measures will (and will not) be developed.

The guideline and quality standard development processes are described in detail on the NICE website (see section 8).

2 Need for guidance

.3 Epidemiology

- a) ST-segment-elevation myocardial infarction (STEMI) is at one end of a spectrum of related conditions called acute coronary syndromes. The underlying common pathophysiology involves either the erosion or sudden rupture of an atheromatous plaque (cholesterol-rich material) within the wall of a coronary artery. This plaque erosion or rupture then stimulates blood clotting (thrombosis) within the affected coronary artery. Complete obstruction to blood flow is usually associated with the appearance of 'ST-segment-elevation' on the electrocardiograph the defining feature of STEMI. Occlusion of blood flow leads to heart muscle (myocardium) cell death that, without intervention, progressively worsens with time.
- b) Typically STEMI causes the onset of acute chest pain, although symptoms may include sweating, nausea and breathlessness. Symptoms may be atypical, particularly in women and people with diabetes. Cardiac arrhythmias may occur early in the onset of STEMI and may cause sudden death before the person is able to access emergency medical care. Certain groups of people, including women and those from ethnic minorities, may be slow to call for medical help.
- c) Although the incidence of STEMI has been declining over the past 20 years, it varies between regions of the UK and still averages around 750 cases per million people each year. Over the past 30 years in-hospital mortality following STEMI has fallen from around 20% to less than 5%, this has been attributed to various factors, including improved drug therapy and speed of access to effective treatments.

.4 Current practice

 a) The overriding concern in the management of STEMI is to rapidly and effectively restore coronary blood flow (reperfusion) because this limits the extent of heart muscle (myocardium) damage and reduces the likelihood of death or future heart failure. In the past, fibrinolysis (that is, reperfusion with fibrinolytic – or 'clot buster' – drugs) was the most common treatment. The treatment of choice now is mechanical reopening of the occluded artery by angioplasty and stent insertion (primary percutaneous coronary intervention [PPCI]).

- b) The Department of Health undertook a feasibility study (National Infarct Angioplasty Project) that reported in October 2008 and concluded that PPCI was both feasible and cost effective, and should become the treatment of choice for STEMI in England, although PPCI is more expensive than fibrinolytic therapy. Since 2009, cardiac networks have successfully implemented the new PPCI policy and by the end of 2011 it is estimated that 95% of the population in England and Wales will be covered by a PPCI care pathway. This PPCI strategy needs emergency access to specialist cardiac catheter laboratories and staff at all times.
- c) Fibrinolytic therapy is still offered a few people (5%) who live in remote rural surroundings and cannot access PPCI services within current recommended time frames.
- d) People may develop symptoms of STEMI and then call the emergency services or self-present to an emergency department. STEMI may also occur in someone already in hospital for a different reason, such as a surgical operation. Whatever the circumstances, care pathways should exist to ensure that PPCI is offered to all who may benefit, in a timely and efficient manner.
- e) The prime determinant of clinical benefit following reperfusion therapy for STEMI is the degree of myocardial salvage (a function of timeliness, effectiveness and maintenance of coronary reperfusion). Bleeding complications also play an important part in both morbidity and mortality if combinations of potent antiplatelet and antithrombin agents are used. After successful acute treatment, secondary prevention therapy, lifestyle modification and cardiac rehabilitation recommendations parallel those for

non-STEMI acute coronary syndromes, on which NICE recently produced guidance ('Unstable angina and NSTEMI', NICE clinical guideline 94, 2010).

f) This guideline will address the factors that influence the delivery of effective and timely coronary reperfusion treatment for people with STEMI.

3 Clinical guideline

.5 Population

.5.1 Groups that will be covered

- a) Adults (18 years or older) believed to be having spontaneous onset of STEMI (types 1 and 3 of the 'universal definition of myocardial infarction' categories).
- b) Adults with suggestive symptoms of spontaneous onset of STEMI, but whose electrocardiogram may be difficult to interpret because of the presence of left bundle branch block or permanent pacing.
- Where data exist, guidance will address differences between specific populations, such as older adults, women and people from ethnic minorities.
- d) Particular attention will be paid to people with STEMI who remain unconscious following resuscitation.

.5.2 Groups that will not be covered

- a) Children and young people (younger than 18 years).
 - Patients initially suspected as having STEMI once this diagnosis is excluded (for example, on cardiac catheterisation).
 - Patients once a diagnosis of STEMI has been excluded (for example, as a complication of coronary revascularisation).

.6 Healthcare settings

Primary, secondary and tertiary healthcare settings, including care from ambulance teams and other paramedical staff before admission to hospital.

.7 Management

.7.1 Key issues that will be covered

The diagnosis of STEMI will be considered to have been made once a patient is identified as having a suggestive clinical presentation and either ST-segment-elevation on the electrocardiograph or an electrocardiograph where interpretation is complicated by the presence of left bundle branch block or permanent pacing. The acute aspects of the following will be addressed, from symptom onset to the point of hospital discharge:

- Adjunctive pharmacotherapy (for example, antiplatelet and antithrombin agents).
- b) Time factors in relation to acute coronary reperfusion.
- The time interval from onset of STEMI beyond which fibrinolysis may be preferable to PPCI.
- d) Drug combinations administered before PPCI (facilitated PPCI).
- e) Timing and effectiveness of angiography or PCI following fibrinolytic therapy.
- f) Timing and effectiveness of PCI following failed fibrinolysis (rescue PCI).
- g) Procedural aspects of PPCI (for example, thrombus extraction).
- h) Guideline recommendations will normally fall within licensed indications; exceptionally, and only if clearly supported by evidence, use outside a licensed indication may be recommended. The guideline will assume that prescribers will use a drug's summary of product characteristics to inform decisions made with individual patients.

.7.2 Key issues that will not be covered

- Management of suspected brain injury in those with STEMI who have suffered cardiac arrest.
- Management of STEMI after hospital discharge, including postmyocardial infarction treatments (we will cross-refer to existing NICE guidance).

.8 Main outcomes

- a) Major cardiovascular events.
- b) Mortality.
- c) Stroke.
- d) Myocardial re-infarction.
- Reintervention at various time intervals (for example, in hospital, 30 days, and 1 year).
- Adverse events including bleeding complications.
- g) Length of hospital stay.
- h) Quality of life.

.9 Economic aspects

Developers will take into account both clinical and cost effectiveness when making recommendations involving a choice between alternative interventions. A review of the economic evidence will be conducted and analyses will be carried out as appropriate. The preferred unit of effectiveness is the quality-adjusted life year (QALY), and the costs considered will usually be only from an NHS and personal social services (PSS) perspective. Further detail on the methods can be found in 'The guidelines manual' (see Section 8).

4 Quality standard

Information on the NICE quality standards development process is available on the NICE website, see section 8.

.10 Mapped areas of care

The areas of care of a patient's journey that will inform the development of the quality statements are set out below (see 4.1.1). The content of the final quality standard statements may differ before and after consultation with stakeholders.

.10.1 Areas of care that will be considered

- a) Out of hospital presentation of acute chest pain
 - a. Assessment and ECG (see www.nice.org.uk/guidance/CG95)
 - Immediate management, pain relief, aspirin, and oxygen (see www.nice.org.uk/guidance/CG95)
- b) In-hospital assessment diagnosis
 - a. Clinical assessment and ECG (see www.nice.org.uk/quidance/CG95)
 - b. Troponin (see www.nice.org.uk/guidance/CG95)
- c) Management of STEMI (this guideline)
 - a. PPCI
 - b. Facilitated PCI
 - c. Rescue PCI
 - d. Fibrinolytic therapy
 - e. Angiography following fibrinolysis
 - f. Antiplatelet agents
 - g. Antithrombotic agents

- d) Management of unstable angina and NSTEMI
 - a. Antiplatelets (see www.nice.org.uk/guidance/CG94)
 - b. Antithrombotic agents (see www.nice.org.uk/guidance/CG94)
 - c. Angiography (see www.nice.org.uk/guidance/CG94)
- e) Discharge Planning (this guideline and CG48 currently scoping for update)
 - a. Cardiac rehabilitation
 - b. Initiation of secondary prevention

.10.2 Areas of care that will not be considered

- a) Adherence to secondary-prevention interventions after their initiation in hospital.
- b) Uptake of cardiac rehabilitation after discharge from hospital following acute coronary syndromes.

.11 Economic aspects

Developers will take into account both clinical and cost effectiveness when prioritising the quality statements to be included in the quality standard. The economic evidence will be considered, and the cost and commissioning impact of implementing the quality standard will be assessed.

5 Status

.12 Scope

This is the final scope.

.13 Timings

The development of the guideline recommendations and the quality standard will begin in August 2011.

6 Related NICE guidance

.13.1 NICE guidance that will be incorporated in or updated by the clinical guideline

The guideline will incorporate the following NICE guidance, subject to a technology appraisal review proposal agreement:

- Bivalirudin for the treatment of ST-segment-elevation myocardial infarction. NICE technology appraisal guidance 230 (2011). Available from www.nice.org.uk/guidance/TA230
- Guidance on the use of drugs for early thrombolysis in the treatment of acute myocardial infarction. NICE technology appraisal guidance 52 (2002). Available from www.nice.org.uk/guidance/TA52
- Ticagrelor for the treatment of acute coronary syndromes. NICE technology appraisal. Publication expected October 2011.

.14 Related NICE guidance

Published

- Hypertension (update). NICE clinical guideline 127 (2011). Available from www.nice.org.uk/guidance/CG127
- Stable angina. NICE clinical guideline 126 (2011). Available from www.nice.org.uk/guidance/CG126
- Prevention of cardiovascular disease. NICE public health guidance 25 (2010).
 Available from www.nice.org.uk/guidance/PH25

- Chest pain of recent onset. NICE clinical guideline 95 (2010). Available from www.nice.org.uk/guidance/CG95
- Unstable angina and NSTEMI. NICE clinical guideline 94 (2010). Available from www.nice.org.uk/guidance/CG94
- Clopidogrel and modified-release dipyridamole for the prevention of occlusive vascular events. NICE technology appraisal guidance 210 (2010). Available from www.nice.org.uk/guidance/TA210
- Cardiac resynchronisation therapy for the treatment of heart failure. NICE technology appraisal guidance 120 (2010). Available from www.nice.org.uk/guidance/TA120
- Prasugrel for the treatment of acute coronary syndromes with percutaneous coronary intervention. NICE technology appraisal guidance 182 (2009). Available from www.nice.org.uk/guidance/TA182
- Medicines adherence. NICE clinical guideline 76 (2009). Available from www.nice.org.uk/guidance/CG76
- Smoking cessation services in primary care, pharmacies, local authorities and workplaces, particularly for manual working groups, pregnant women and hard to reach communities. NICE public health guidance 10 (2008). Available from www.nice.org.uk/guidance/PH10
- Familial hypercholesterolaemia. NICE clinical guideline 71 (2008). Available from www.nice.org.uk/guidance/CG71
- Lipid modification. NICE clinical guideline 67 (2008). Available from www.nice.org.uk/guidance/CG67
- Drug-eluting stents for the treatment of coronary artery disease. NICE technology appraisal 152 (2008). Available from www.nice.org.uk/guidance/TA152
- MI: secondary prevention. NICE clinical guideline 48 (2007). Available from www.nice.org.uk/guidance/CG48
- Implantable cardioverter defibrillators (ICDs) for arrhythmias (2006). NICE technology appraisal 95 (2006). Available from www.nice.org.uk/guidance/TA95.
- Statins for the prevention of cardiovascular events. NICE technology appraisal guidance 94 (2006). Available from www.nice.org.uk/guidance/TA94
- Brief interventions and referral for smoking cessation in primary care and other settings. NICE public health guidance 1 (2006). Available from www.nice.org.uk/guidance/PH1

- Off-pump coronary artery bypass (OPCAB). NICE interventional procedure guidance 35 (2004). Available from <u>www.nice.org.uk/guidance/IPG35</u>
- Myocardial perfusion scintigraphy for the diagnosis and management of angina and myocardial infarction. NICE technology appraisal guidance 73 (2003).
 Available from www.nice.org.uk/guidance/TA73
- Guidance on the use of coronary artery stents. NICE technology appraisal guidance 71 (2003). Available from www.nice.org.uk/guidance/TA71
- Guidance on the use of drugs for early thrombolysis in the treatment of acute myocardial infarction. NICE technology appraisal guidance 52 (2002). Available from www.nice.org.uk/guidance/TA52
- Guidance on the use of glycoprotein IIb/IIIa inhibitors in the treatment of acute coronary syndromes. NICE technology appraisal guidance 47 (2002). Available from www.nice.org.uk/guidance/TA47

NICE guidance under development

NICE is currently developing the following related guidance (details available from the NICE website):

Hyperglycaemia in patients with acute coronary syndrome. NICE clinical guideline.
 Publication expected October 2011.

7 Further information

Information on the guideline development process is provided in:

- 'How NICE clinical guidelines are developed: an overview for stakeholders the public and the NHS'
- · 'The guidelines manual
- · 'Developing NICE quality standards: interim process guide'.

These are available from the NICE website (www.nice.org.uk/GuidelinesManual and www.nice.org.uk/aboutnice/qualitystandards). Information on the progress of the guideline and quality standards are also available from the NICE website (www.nice.org.uk).

Appendix B: Declarations of interest

B.1 Guideline development group members

B.1.1 Sotiris Antoniou

Item declared	Date	Expiry	Classification	Action taken
Chair of cardiac committee for the United Kingdom Clinical pharmacy Association	2005	Ongoing	Personal non- pecuniary	Declare and participate
Member NICE guideline development groups for Stable Angina and		Issued March	Expired	Expired
NSTEMI		2010		Declare and participate
Attended advisory board meeting for Chiesi Pharmaceuticals on integrated care pathways in ACS. Received honorarium.	26 August 2010	25 August 2011	Personal non-specific pecuniary	Expired
Attended presentation at Heart Rhythm Congress meeting sponsored by Sanofi Aventis.	2 October 2010	1 October 2011	Personal non-specific pecuniary	Declare and participate GDG1 and 2. Expired by GDG3
Attended advisory board meeting for Astra Zenica on ticagrelor. Received honorarium.	23 October 2010	22 October 2011	Personal specific pecuniary	GDG not discussing recommendations for ticagrelor (TA incorporated). Declare and participate GDG1 and 2. Expired by GDG3
Reimbursed for presentation on the role of pharmacy in health care (not specifically related to a drug) by Astra Zenica.	3 November 2010	2 November 2011	Personal specific pecuniary	GDG not discussing recommendations for ticagrelor (TA incorporated). Declare and participate GDG1 and 2. Expired by GDG3
Attended meeting on Ad Review (a drug used for imaging in heart failure). Honorarium received from GE Healthcare.	10 November 2010	9 November 2011	Personal non-specific pecuniary	Declare and participate GDG1 and 2. Expired by GDG3
Attended advisory board on ivabradine in Chronic Heart Failure for Servier. Received honorarium.	16 November 2010	15 November 2011	Personal non-specific pecuniary	Declare and participate GDG1 and 2. Expired by GDG3.
Attended advisory board on prasugrel for Lilly. Received honorarium.	23 November 2010	22 November 2011	Personal specific pecuniary	GDG not discussing recommendations for prasugrel (TA undergoing update).

Item declared	Date	Expiry	Classification	Action taken
				Declare and participate GDG1 and 2
Attended advisory board on HDL increasing prasugrel for Lilly. Received honorarium.	23 February 2011	22 February 2012	Personal specific pecuniary	GDG not discussing recommendations for prasugrel (TA undergoing update). Declare and participate.
Attended advisory board on anti Xa in Atrial Fibrillation (edoxaban - drug in development) for Daiichi Sankyo. Received honorarium.	3 March 2011	2 March 2012	Personal specific pecuniary	Re Daiichi Sankyo as manufacturer - GDG not discussing recommendations for prasugrel (TA undergoing update). Declare and participate.
Attended advisory board on dabigatan for atrial fibrillation. Received honorarium from Boehringer Ingelheim.	8 April 2011	7 April 2012	Personal specific pecuniary	GDG debating strategy only therefore declare and participate
Attended advisory board on prasugrel. Received honorarium from Lilly.	9 May 2011	8 May 2012	Personal specific pecuniary	GDG not discussing recommendations for prasugrel (TA undergoing update). Declare and participate.
Wrote an article for the journal 'Future Prescriber' on ticagrelor.	May 2011		Personal specific non- pecuniary	Declare and participate
Received reimbursement from Bayer for a presentation on the role of pharmacy and cardiac networks (not drug related)	29 June 2011	28 June 2012	Personal non-specific pecuniary	Declare ad participate
Attended advisory board on ticagrelor. Received honorarium from Astra Zeneca.	4 July 2011	3 July 2012	Personal specific pecuniary	GDG not discussing recommendations for ticagrelor (TA incorporated). Declare and participate
Reimbursed for analysis of risk stratification tools in NSTEMI by Daiichi Sankyo.	1 August 2011	31 July 2012	Personal specific pecuniary	Declare and participate
Received accommodation, travel and expenses to attend ESC congress in 2011 from Daiichi Sankyo.	27 August 2011	26 August 2012	Personal specific non- pecuniary	Declare and participate
Attended advisory board on dalcetrapib HDL agent. Received honorarium from Roche.	15 September 2011	14 September 2012	Personal non-specific pecuniary	Declare and participate

Item declared	Date	Expiry	Classification	Action taken
Attended UK antithrombotics meeting. No payment received.	13 October 2011	12 October 2012	Personal specific non- pecuniary	Declare and participate
Attended an advisory board on Integrated care pathway development for stable angina sponsored by Menarini who manufacture ranolazine. Honorarium received.	October 2012	September 2013	Personal, non-specific pecuniary	Declare and participate
Attended an advisory board on Heart failure sponsored by Servier who manufacture ivabradine. Honorarium received.	October 2012	September 2013	Personal, non-specific pecuniary	Declare and participate
Attend an advisory board by DS/Lilly partnership on prasugrel. Honorarium received.	17 December 2012	16 December 2013	Personal specific pecuniary	GDG not discussing recommendations for prasugrel (TA undergoing update). Declare and participate

B.1.2 Charles Deakin

	Date	Expiry	Classification	Action taken
Director, Prometheus Medical Ltd (http://www.prometheusmedical.co.uk) There is no direct link with STEMI management. Not actively engaged in sales or training (more as an advisory role).	24 August 2011	-	Personal non-specific pecuniary	Declare and participate
Divisional Medical Director, South Central Ambulance Service	24 August 2011	-	No conflict of interest	Declare and participate
Consultant in Cardiac Anaesthesia and Cardiac Intensive Care	24 August 2011	-	No conflict of interest	Declare and participate
Member of the Executive Committee, Resuscitation Council (UK)	24 August 2011	-	Personal non- pecuniary	Declare and participate
Board member, European Resuscitation Council	24 August 2011	-	Personal non- pecuniary	Declare and participate
Chair, Advanced Life Support Working Group, European Resuscitation Council	24 August 2011	-	Personal non- pecuniary	Declare and participate
Co-Chair, Advanced Life Support Working Group, International Liaison Committee on Resuscitation	24 August 2011	-	Personal non- pecuniary	Declare and participate
Royal College of Anaesthetists representative, Joint Royal Colleges Ambulance Liaison Committee (JRCALC)	24 August 2011	-	Personal non- pecuniary	Declare and participate
Guidelines Committee, Joint Royal Colleges Ambulance Liaison Committee(JRCALC)	24 August 2011	-	Personal non- pecuniary	Declare and participate

	Date	Expiry	Classification	Action taken
Honorary Civilian Consultant in Pre-Hospital Emergency Care to the British Army.	24 August 2011	-	No conflict of interest	Declare and participate
Deputy Chair (Chair as of January 2012), RNLI Medical & Survival Committee.	24 August 2011	-	Personal non- pecuniary	Declare and participate
Involved in developing the Position statement from JRCALC on paramedic airway management (see CV) - 30/06/11	24 August 2011	-	Personal non- pecuniary	Declare and participate
Various research papers and reviews on ambulance triage of patients with chest pain, management of cardiac arrest, and post-cardiac arrest management.	24 August 2011	-	Personal non- pecuniary	Declare and participate
Resuscitation Council (UK) – Research Grant 2007 – Lead applicant on grant to fund research into the safety of automatic chest compression devices.	24 August 2011	-	Non-personal pecuniary – non-healthcare manufacturer / industry related	Declare and participate
Resuscitation Council (UK) – Research Grant 2007 – Lead applicant to fund study of genetic polymorphism as a factor affecting survival from cardiac arrest. (Administers these funds)	24 August 2011	-	Non-personal pecuniary – non-healthcare manufacturer / industry related	Declare and participate
Resuscitation Council (UK) – Research Grant 2007 – Lead applicant to fund research a comparison of the efficacy of biphasic truncated exponential and rectilinear biphasic waveforms in cardioversion of atrial fibrillation. (Administers these funds)	24 August 2011	-	Non-personal pecuniary – non-healthcare manufacturer / industry related	Declare and participate
Resuscitation Council (UK) – Research Fellowship 2010 – A pilot prospective study of the role of cerebral oximetry in predicting outcomes in in-hospital cardiac arrest Principle Investigator. March 2010 (Administers these funds)	24 August 2011	-	Non-personal pecuniary – non-healthcare manufacturer / industry related	Declare and participate
Resuscitation Council (UK) – Research Fellowship 2010 – Principle Investigator. 'Hands on' Defibrillation – Is it safe? June 2010 (Administers these funds)	24 August 2011	-	Non-personal pecuniary – non- healthcare manufacturer / industry related	Declare and participate

B.1.3 Huon Gray

ltem declared	Date	Expiry	Classification	Action taken
President, British Cardiovascular Society (2003-5)	2003	2005	Expired	Not relevant
Lead (Cardiology), London Review of Cardiovascular Services, NHS London 2009-10.			Personal non- pecuniary	Declare and participate
Hon. Civilian Consultant Adviser (Cardiology) to the Army	2008	-	No conflict of interest	Not relevant
Clinical Advisor, NICE Clinical Guideline 95 (Unstable angina & non- ST elevation myocardial infarction) 2007-10		Issue March 2010	Expired	Expired Declare and participate
Chair, International Council (2008-date) & Member of Board of Trustees (2010-date), American College of Cardiology. Travel expenses paid by ACC for attendance at international & board meetings.	2008	-	Non-classifiable - ACC do not provide services to the NHS	Declare and participate
Co-chair of NIAP (with Roger Boyle, the National Director for Heart Disease at DH) from the project's inception to completion. Chaired meetings of the Steering Group, attended data monitoring meetings, and was principal clinical author on the two NIAP reports which appeared in 2008. I did not receive any reimbursement or payment for this work nor was I involved in allocating the £1 million of funds which the DH allocated to the Study. This was done by Civil Servants. I was also a co-author on the cost effectiveness paper which appeared in Heart [Heart 2010; 96:668-672]		2008	Personal non- pecuniary	Declare and participate
Advisory Board Member, Daiichi Sankyo; prasugrel for acute coronary syndromes (2 meetings – December 2010 & July 2011). Remuneration for each.	December 2010	July 2012	Personal specific pecuniary	GDG not discussing recommendations for prasugr (TA undergoing update).
				Deputy Chair wrote related NICE guidance chapter of guideline.
				Declare and participate
Attendance at an Advisory Board meeting for St Jude Medical who are discussing the introduction of their remote monitoring device for the management of heart failure patients in the UK.	24 November 2011	23 November 2012	Pecuniary non-specific	Declare and participate action at the discretion of the Guideline Lead

Item declared	Date	Expiry	Classification	Action taken
Seconded to the Department of Health as interim National clinical director for cardiovascular diseases.	12 March 2012		Non-specific pecuniary	Declare and participate

B.1.4 Robert Henderson

Item declared	Date	Expiry	Classification	Action taken
Sponsored by Edwards Life Science to attend EuroPCR Valve Live meeting. Reimbursed travel and accommodation costs only.	October 2010	October 2011	Expired by GDG2	No action required
Attended 'Cardiology and Diabetes at the Limits Conference' in Feb 2011; sponsored by Pfizer Ltd, F.Hoffman-La Roche Ltd, Novo Nordisk, AstraZenica South Africa, Medtronic Ltd, Saiichi-Sankyo/Lilly UK, Sanofi-Aventis, Lilly UK Ltd. Reimbursed travel and accommodation expenses only.	February 2011	February 2012	Within normal expenses of DOI policy	No action required
Member of council of British Cardiovascular Intervention Society Council member 2006-current Clinical standards lead 2009-current Member, British Cardiovascular Society clinical standards committee 2009-current Member, British Cardiovascular Society programme committee 2010-current			Personal non- pecuniary	Declare and participate No contribution to SH comments from BCIS because answering on behalf of GDG
Member of the steering committee of RITA-3 and co-author of publications			Personal non- pecuniary	Declare and participate
Member of NICE guideline development group for Unstable Angina/NSTEMI and clinical adviser to NICE Stable Angina guideline development group		Unstable Angina published March 2010, Stable Angina July 2011	Expired	No action required
Attended Euro-PCR meeting in Paris. Registration, travel and accommodation expenses paid by Edwards Lifesciences (heart valve manufacturer).	15-18 May 2012	18 th May 2013	Within normal expenses of DOI policy	No action required
Attended a one day meeting in London on TAVI on 15 June 2012. The meeting was sponsored by Edwards Lifesciences (heart valve manufacturer) and the company paid for travel and overnight	15 June 2012		Within normal expenses of DOI policy	No action required

Item declared	Date	Expiry	Classification	Action taken
Attended London PCR-valve conference (1-2 October). Edwards Lifesciences paid for travel, accommodation and registration but no honorarium or other financial benefit received.	1-2 October 2012		Within normal expenses of DOI policy	No action required
Attended TCT conference in Miami, USA (21-25 October). Biosensors (coronary stent manufacturer) paid for travel, accommodation and registration but I did not receive any other payment from the company.	21-25 October 2012		Within normal expenses of DOI policy	No action required

B.1.5 Jason Kendall

Item declared	Date	Expiry	Classification	Action taken
Authored a paper in the EMJ related to the management of STEMI in 2007: The Optimum Reperfusion Pathway for ST elevation myocardial infarction: development of a decision framework. Kendall JM. EMJ 2007; 24:52-56 Also published as a web-based interactive tool at: http://www.ddaccess.co.uk/europr/cdrom/index/htm		2007	Expired	No action required
Co-author of publication related to pre-hospital thrombolysis: Does pre-hospital thrombolysis increase the proportion of patients having an aborted myocardial infarction? Jackson L. Kendall JM, Castle N. EMJ 2009; 26:206-209	2009		Expired	No action required
Co-applicant on an NIHR-funded research project evaluating the cost effectiveness of diagnostic strategies for acute coronary syndromes (project based at the University of Sheffield).	2009	2011	Non-personal pecuniary	Declare and participate
Author of a BMJ Learning module on 'The Management of STEMI'.	2009	2010	Expired	Expired Declare and participate
Author of a Department of Health e-learning for Health (elfh) module on 'The Management of STEMI and its complications'.	2009	2010	Expired	Expired Declare and participate
Author of various e-learning modules on the College of Emergency Medicine's e-learning platform (ENLIGHTENme) related to Acute Coronary Syndromes (Pathophysiology, Clinical Assessment, STEMI, UA and NSTEMI)	2008 – on going		Personal non- pecuniary	Declare and participate
Member of NICE guideline development group CG95 on Chest Pain (2008-2010)		Issued March 2010	Expired	No action required
Steering Committee Member of STREAM Trial: Strategic Reperfusion Early after Myocardial Infarction Study. This is an international randomised control trial comparing PPCI with a contemporary lytic-based reperfusion strategy. (24/06/11) - no payment received in previous 12 months	24 June 2011		Personal non- pecuniary	Declare and participate
Attended an advisory board meeting on the diagnosis and treatment of pulmonary embolism, funded by Bayer who manufacture rivaroxaban which is one of the new oral anticoagulants used for the treatment of venous thromboembolic disorders.	September 2012	September 2013	Personal non-specific pecuniary	Declare and participate

B.1.6 Hugh McIntyre

Item declared	Date	Expiry	Classification	Action taken
Received research grant support from Takeda for a 2-year doctorate which HM supervised investigating the epidemiology of heart failure (to Jan 2011)	January 2009	January 2012	Non-personal pecuniary	Declare and participate
Regional acute lead for Heart Failure for the Enhancing Quality Programme run by NHS SE. One session per week reimbursed during secondment to April 2013.	April 2010	April 2013	Non-personal pecuniary	Declare and participate
Attended advisory board meeting for Novartis Pharmaceuticals on the topic of age-related muscular degeneration, therapy Lucentis. Honorarium received.	13 May 2011	12 May 2012	Personal non-specific pecuniary	Declare and participate
Attended advisory board meeting for Pfizer Pharmaceuticals on eplerenone in chronic heart failure (not yet licensed for chronic heart failure but licensed for post MI heart failure).	9 May 2011	8 May 2012	Personal non-specific pecuniary	Declare and participate
Honorarium received.				
Deputy Divisional Director of Medicine (Conquest Hospital East Sussex HealthcareTrust) recent establishment and ongoing developments in primary PCI service (no direct involvement).	24 August 2011		Personal non- pecuniary	Declare and participate
VAN Data Safety Monitoring Board member (comparison of bevacizumab and ranibizumab in macular degeneration)	24 August 2011		Personal non- pecuniary	Declare and participate
NICE Heart Failure GDG member (2008-10) and Topic Expert Group Chair (2010-11) – (reimbursed for role as Chair)	24 August 2011	2011	Expired	No action required
Principal investigator to international randomised control trials, including lipid lowering agent (HPS THRIVE), antiplatelet treatment for chronic cardiovascular disease (TRA2P), registry of patients treated for acute coronary syndrome (EPICOR), registry of patients with atrial fibrillation. Does not administer trial funds.	24 August 2011		Personal non- pecuniary	Declare and participate
Sub-investigator for study of antiplatelet agent in ACS (TAO study).	24 August 2011		Personal non- pecuniary	Declare and participate
European Society of Cardiology Annual meeting (from Servier). Travel and accommodation expenses only.	August 2011		Within normal expenses of DOI policy	No action required
Attended advisory board meeting for Boehringer Ingelheim Pharmaceuticals on dabigatran for anticoagulation in atrial fibrillation. Honorarium received.	12 September 2011	11 September 2012	Personal specific pecuniary	Q3, 5 & 6 - debating strategy only therefore declare and participate

tem declared	Date	Expiry	Classification	Action taken
Participated in consensus statement for heart failure meeting for Pfizer pharmaceuticals. Honorarium received.	16 September 2011	15 September 2012	Personal non-specific pecuniary	Declare and participate
Attended advisory board meeting for Servier Pharmaceuticals on vabradine. Honorarium received.	16 September 2011	15 September 2012	Personal non-specific pecuniary	Declare and participate
Attended Advisory Board meeting on rivaraoxaban (Bayer)	Nov 2011		Personal non-specific pecuniary	Declare and participate
Trust authorisation for two Charitable Funds (managed by East Sussex Healthcare Trust):	Declared January 2012	-	Non-personal pecuniary	Declare and participate
1. the Cardiovascular Research Fund - manages reimbursement from RCTs to support Research nurse funding; 2. and to the Hastings Heart Failure fund –supports MD student and heart failure nurses in continuing medical education, equipment and research.				
Attendance at an advisory board for Roche Pharmaceuticals regarding BNP in diagnosis of heart failure. Honorarium received.	13 March 2012	12 March 2013	Personal non-specific pecuniary	Declare and participate
Presented at a regional industry meeting on Chronic Heart Failure Pfizer pharmaceuticals) on 18th April (Honorarium received).	18 April 2012	17 April 2013	Personal non-specific pecuniary	Declare and participate
Presented at a regional educational meeting in London – Heart Failure New Challenges New solutions. Independent conference. Honorarium received from Servier.	26 April 2012	25 April 2013	Personal non-specific pecuniary	Declare and participate
Received educational grant from Pfizer to support conference attendance (travel, accommodation and registration only)	May 2012		Within normal expenses of DOI policy	No action required
Gave an expert cardiovascular briefing (Gilenya in multiple scleroisis) for Novartis. Honorarium received.	4 May 2012		Personal non-specific pecuniary	Declare and participate
Session chair for the 'from myocardial ischaemia to heart failure conference. (Independent conference – education grant from Servier) Honorarium received.	16 May 2012		Personal non-specific pecuniary	Declare and participate
Attended an advisory board for Novartis on LCZ (trial medication for neart failure not licensed) in Chronic Heart Failure. Honorarium received.	24 May 2012		Personal non-specific pecuniary	Declare and participate
Presented at an educational meeting on Heart Failure and received nonorarium from Pfizer.	13 June 2012		Personal non-specific pecuniary	Declare and participate
Presented at RCP Heart Failure Educational Update Lecture.	21 June 2012		Personal non-specific	Declare and participate

Item declared	Date	Expiry	Classification	Action taken
(Independent conference - education grant from Pfizer). Received honorarium.			pecuniary	
Advisory Board on Gilenya in multiple sclerosis for Novartis. Honorarium.	28 June 2012		Personal non-specific pecuniary	Declare and participate
Attended NICE acute heart failure stakeholder group	July 2012		Personal non- pecuniary	Declare and participate
Attended Department of Health cardiovascular disease long term conditions work stream.	July 2012		Personal non- pecuniary	Declare and participate
Presentation to Educational meeting (Pfizer)	July 17 2012		Personal non-specific pecuniary	Declare and participate
Presentation to Educational meeting (Servier)	July 24 2012		Personal non-specific pecuniary	Declare and participate
Presentation on the treatment of heart failure, 27 July 2012. Funded by Servier. ivabaradine.	July 2012		Personal non-specific pecuniary	Declare and participate
Educational grant to support one day development programme for local heart failure service. Pfizer pharmaceuticals reimbursed room hire, refreshments and facilitator.	Sept 2012	Sept 2013	Non-personal pecuniary	Declare and participate
Appointed chair of NICE Quality Standard Advisory Committee	2 September 2012		Personal non- pecuniary	Declare and participate
Attended American Heart Association annual meeting 3rd - 8th November and received accommodation expenses from Boehringer Ingelheim grants toward travel costs from Novartis, Servier and Pfizer.	3 – 8 November 2012		Within normal expenses of DOI policy	No action required
Participation on an advisory board for Novartis (regarding Relaxin for AHF).	28 November 2012	27 November 2013	Personal pecuniary non-specific interest	Declare and participate

B.1.7 Jim McLenachan

Item declared	Date	Expiry	Classification
Clinical lead for the development of PPCI in Leeds since 2004.			Personal non- pecuniary
Clinical lead for the development of PPCI in the West Yorkshire Cardiac Network since 2004.			Personal non- pecuniary

Action taken

Declare and participate

Declare and participate

Item declared	Date	Expiry	Classification	Action taken
Local investigator in the NIAP (National Infarct Angioplasty Project) in 2005.			Personal non- pecuniary	Declare and participate
Attended Lilly symposium on 17/11/10 and accepted the honorarium	17 November 2010	16 November 2011	Personal specific pecuniary	Declaration expires by date of Q4 (March 2012). Declare and participate.
The cardiology department at Leeds General Infirmary (in which I work) receives money from The Medicines Company to fund a research nurse post. Not administering any of these funds.	24 August 2011	-	Non-personal specific pecuniary	Declare and participate
Member (and past Honorary Secretary) of the British Cardiovascular Intervention Society and will attend the society's autumn meeting in October 2011. The society receives money from the British Cardiovascular Industry Association (BCIA) which is made up of a number of cardiac device and pharmaceutical companies. This money is then used to fund the meeting including overnight accommodation costs and travel expenses for invited speakers and chairmen.			Personal non- pecuniary	Declare and participate No contribution to SH comments from BCIS because answering on behalf of GDG
National Clinical Lead for PPCI with NHS Improvement, from September 2008 to date	September 2008	-	Personal non- pecuniary	Declare and participate
Gave presentation on 'Regional and National Roll-Out of PPCI' at a clinical cardiology meeting in York on 2/11/11. The meeting will be sponsored by the pharmaceutical company Lilly Ltd. No honorarium paid.	2 November 2011	1 November 2012	Personal non- pecuniary	Declare and participate
Attendance at ACI 2012 travel and expenses.	25/26 January 2012	24 January 2013	Within normal expenses of DOI policy	No action required

B.1.8 Francesco Palmer

Item declared	Date	Expiry	Classification	Action taken
None declared				No action required

B.1.9 Gerald Robinson

Item declared	Date	Expiry	Classification	Action taken
Member of the Birmingham, Sandwell & Solihull Cardiac and Stroke	29 July 2011	-	Personal non-	No action required

Item declared	Date	Expiry	Classification
Network Shadow Board. This is a voluntary position.			pecuniary

Action taken

B.1.10 Fiona Sayers

Item declared	Date	Expiry	Classification	Action
Member of the NICE guideline development group on the prevention of stroke and systemic embolism in patients with atrial fibrillation	24 August 2011	-	Personal non- pecuniary	Declar
Technology appraisal group member for rivaroxaban	24 August 2011	-	Personal non- pecuniary	Decla

Action taken	
Declare and participate	
Declare and participate	

B.1.11 David Smith (Wales)

Item declared	Date	Expiry	Classification	Action taken
Received sponsorship from Biosensors International (who make and sell coronary stents) to attend the TCT conference in Miami from October 22nd to 26 th 2012. I received sponsorship for registration, travel and accommodation costs but received no other monies.	22 – 26 October 2012	25 October 2013	Within normal expenses of DOI policy	No action required

B.1.12 LDR Smith

Item declared	Date	Expiry	Classification	Action taken
Attendance at European Society of Cardiology conference – registration, travel and expenses only reimbursed by Boehringer Ingleheim	30/31 August 2010	30 August 2011	Within normal expenses of DOI policy	No action required
Registration, travel and expenses only paid for by BCOS for BCIS Meeting and BCIS Clinical Standards Committee	7/8 October 2010	7 October 2011	Within normal expenses of DOI policy	No action required
Attendance at best practice in Cardiology (London) funded by PriMed, honorarium in addition to travel and accommodation received.	13 October 2011	12 October 2012	Personal non-specific	Declare and participate
Attendance at best practice in Cardiology (Manchester) funded by PriMed, honorarium in addition to travel and accommodation received.	10/11 November 2011	10 November 2012	Personal non-specific	Declare and participate
Accommodation, travel and expenses reimbursed by Boston	26/27 November	26 November	Within normal	No action required

Item declared	Date	Expiry	Classification	Action taken
Scientific to attend SWIG/IWEST in Bath	2010	2011	expenses of DOI policy	
Accommodation, travel and expenses reimbursed by Biosensors to attend meeting on PCI in the CGH in Basingstoke.	2/3 March 2011	2 March 2012	Within normal expenses of DOI policy	No action required
Accommodation, travel and expenses reimbursed by the Royal College of Physicians to attend a meeting on acute general medicine for physicians in Birmingham.	25/26 May 2011	25 May 2012	Within normal expenses of DOI policy	No action required
Accommodation, travel and expenses reimbursed by BCS to attend the BCS annual scientific sessions.	13-15 June 2011	14 June 2012	Within normal expenses of DOI policy	No action required
Member BCIS council			Personal non-	Declare and participate
			pecuniary	No contribution to SH comments from BCIS because answering on behalf of GDG
Member of the International Editorial board for Heart			Personal non- pecuniary	Declare and participate
Member of NIAP being one of 7 hospitals involved in study			Personal non- pecuniary	Declare and participate
Attended TCT conference in Miami, USA (21-25 October). Biosensors (coronary stent manufacturer) paid for travel, accommodation and registration but I did not receive any other payment from the company.	21-25 October 2012	24 October 2013	Within normal expenses of DOI policy	Declare and participate

B.1.13 Mark Whitbread

Item declared	Date	Expiry	Classification
Member of the JRCALC guidelines committee			Personal non- pecuniary
Member of NICE guideline development group for the UA/NSTEMI guideline		Issued March 2010	Expired

Action taken
Declare and participate
No action required

B.2 Invited experts

B.2.1 Keith Fox

Item declared	Date	Expiry	Classification	Action taken
Invited speaker for, and honoraria received from: Bayer, Johnson & Johnson, Astra Zeneca and Sanofi Aventis	September 2012	August 2013	Personal specific pecuniary	No action required – Keith Fox did not attend any GDG meetings and was not involved in discussing recommendations.
Grants received from Lilly and Bayer/Johnson & Johnson	Ongoing		Non-personal pecuniary	No action required – Keith Fox did not attend any GDG meetings and was not involved in discussing recommendations.

B.2.2 Tony Gershlick (Attended one session of GDG 10)

Item declared	Date	Expiry	Classification	Action taken
Support to attend meetings and financial remuneration for advisory boards for ABBOT Vascular, Metronic Corp and Boston Scientific.	September 2012	August 2013	Personal specific pecuniary	No action required – Professor Gershlick was not involved in discussing recommendations.
Remuneration for attendance at Steering Committee and TC re. trial management for the STREAM trial. Boehringer Ingelhiem.	Ongoing		Personal specific pecuniary	No action required – Professor Gershlick was not involved in discussing recommendations.

B.3 Technical team

B.3.1 Daria Bilan

Item declared	Date	Expiry	Classification	Action taken
None declared				No action required

B.3.2 Serena Carville (from October 2012)

Item declared	Date	Expiry	Classification	Action taken
None declared				No action required

B.3.3 Angela Cooper (from December 2011)

Item declared	Date	Expiry	Classification	Action taken
Co-author with Jane S Skinner on BMJ Clinical evidence reviews,	2011		No conflict	No action required
Secondary prevention of ischaemic cardiac events, Clinical Evidence				
2011; 08:2				

B.3.4 Martin Harker (from April 2012)

Item declared	Date	Expiry	Classification	Action taken
None declared				No action required

B.3.5 Taryn Krause (until October 2012)

Item declared	Date	Expiry	Classification	Action taken
None declared				No action required

B.3.6 Kate Lovibond (until August 2012)

Item declared	Date	Expiry	Classification	Action taken
None declared				No action required

B.3.7 Jill Parnham

Item declared	Date	Expiry	Classification	Action taken
Employment contract with the Royal College of Physicians	2001		No conflict	No action required No contribution to SH comments from RCP because answering on behalf of GDG
Accept NICE commissions			Non-personal pecuniary	Declare and participate
Various publications unrelated to STEMI			No conflict	No action required

B.3.8 Robert Pitcher (until January 2012)

Item dec	lared	Date	Expiry	Classification	Action ta
None dec	clared				No action

Action taken No action required

B.3.9 Carlos Sharpin

Item declared	Date	Expiry	Classification	Action taken
None declared				No action required

Appendix C: Review protocols

Table 1: Review protocol: Time to reperfusion

Review	What is the duration of PPCI-related time delay at which fibrinolysis becomes more clinically	
question	and cost effective compared to PPCI in people with STEMI and how is this modulated by	
	patient presentation delay and patient risk profile?	

Objectives To determine the duration of PPCI-related time delay at which fibrinolysis becomes more

clinically and cost effective compared to PPCI in people with STEMI.

Criteria Population:

Adults (≥ 18 years old) with STEMI to be managed by reperfusion therapy.

Intervention:

PPCI. Plus standard adjunctive pharmacotherapies (for example, antiplatelet and antithrombin agents).

Comparison:

Fibrinolytic therapy (in-hospital or pre-hospital; any agent). Plus standard adjunctive pharmacotherapies (for example, antiplatelet and antithrombin agents).

Outcomes (original study definitions will be used and recorded):

- Mortality (all-cause and cardiovascular specific)
- Non-fatal and all (non-fatal and fatal) stroke
- Non-fatal and all (non-fatal and fatal) myocardial reinfarction
- Heart failure
- · Minor and major bleeding
- Revascularisation (unplanned)
- Length of hospital stay
- Quality of life (report all, inc EQ-5D (EuroQol), SF-36 (Short Form 36), SF6D (Short Form 6-Dimensions), SF-12 (Short Form 12-Dimensions), RAND-36 (Research and Development Medical Outcomes Study Short Form-36))
- Time to equipoise between the treatments for outcomes listed above.

Study design:

RCTs that stratify people based on time to presentation or PPCI-related time delay Meta-regression analyses or other modelling techniques that assess impact of time to presentation or PPCI-related time delay on relative benefits of fibrinolysis and PPCI, based on : ≥10 RCTs or

UK registry data or

International registry data (n>100,000).

Population size and directness:

No limitations on sample size (unless registry data – n>100,000) Studies with indirect populations will not be considered.

Setting:

Pre-hospital Secondary care

Search See appendix F

Strategy

Review Strategy

Quality of life data

Collect all data for the stated quality of life measure, for meta-analysis and GRADE report only overall scores.

Appraisal of methodological quality

The methodological quality of each study will be assessed using adapted NICE checklists and GRADE. Limitations include the robustness of modelling technique (for example, use of patient/hospital level covariates, sensitivity analyses) and quality of input data (for example, patient/hospital/study-level data; number of studies; RCTs versus registry data; proportion of studies using streptokinase, stents and GPIs; conference abstracts versus full articles).

Data synthesis

We will record results of meta-regression analyses and other modelling techniques. We will also analyse each constituent RCT and where possible, meta-analyse data stratified by time to presentation.

Subgroups:

The following groups will be considered separately if data is present:

- People with diabetes
- People with renal dysfunction
- People aged over 70 years
- Ethnicity
- Gender
- MI location
- MI size.

Table 2: Review protocol: Facilitated PPCI

Table 2. Review protocol: Facilitated PPCI			
	Review question	What is the clinical and cost effectiveness of facilitated PPCI (fPPCI) compared to PPCI in people with STEMI?	
	Objectives	To compare the clinical and cost effectiveness of facilitated PPCI and PPCI, for the treatment of adults with STEMI.	
	Criteria	Population: Adults (≥ 18 years old) with STEMI eligible for PPCI.	

Intervention:

- In people already scheduled to have PCI. Facilitated PCI: use of pharmacological agents (GPIs, fibrinolytics or heparin OR a combination of any of these classes) before an anticipated PPCI to improve coronary patency
- pharmacological agents can be given any time before PPCI, before arrival in the cath lab, and given with the intention of facilitation (ambulance, hospital, accident and emergency)
- people must also be on a background of at least one oral antiplatelet agent, and may have been given an antithrombin
- older studies = aspirin with/without heparin
- newer studies = aspirin + clopidogrel (or other ADP antagonists) with/without heparin.

Comparison:

- In people with STEMI where the intended reperfusion strategy is PPCI. PPCI arm: PPCI with or without placebo or adjunctive therapy
- adjunctive therapy: GPIs
- adjunctive therapy drugs must be given in the cath lab or at time of PPCI, not before

- they should ideally should be the same drugs as given in the fPPCI arm
- newer studies give GPIs as part of standard PPCI therapy, in this case a suitable comparator fPPCI arm must be on combination therapy (GPI + fibrinolytic or heparin) OR they could be given the same drug but at an earlier time (ie. Studies comparing upstream/pre-cath lab administration fPPCI versus downstream/cath-lab administration fPPCI)
- people must also be on a background of at least one antiplatelet agent
- older studies = aspirin with/without heparin
- newer studies = aspirin + clopidogrel (or other ADP antagonists) with/without heparin)
- background antiplatelet therapy must be the same as in the fPPCI arm.

Additional intervention and comparison

• We will also look at studies that assess the effect of early versus later fPPCI.

Outcomes (original study definitions will be used and recorded):

- Mortality (all-cause and cardiovascular specific)
- Non-fatal and all (non-fatal and fatal) stroke.
- Non-fatal and all (non-fatal and fatal) myocardial reinfarction
- Major and minor bleeding. Note intracranial bleeding separately
- · Heart failure
- Repeat revascularisation (this includes TVR, repeat revascularisation, unplanned revascularisation and urgent revascularisation)
- Length of hospital stay
- Quality of life (report all, including EQ-5D (EuroQol), SF-36 (Short Form 36), SF6D (Short Form 6-Dimensions), SF-12 (Short Form 12-Dimensions), RAND-36 (Research and Development Medical Outcomes Study Short Form-36)).

Study design:

RCT

Additional inclusion / exclusion criteria including population size and directness:

- Studies with indirect populations will not be considered (population must be 100% STEMI or AMI).
- Only studies published after 1990 will be considered. This is to ensure that the extracted evidence is reflective of current practice, especially with regard to the widespread adoption of stenting in place of balloon angioplasty for PCI procedures over the last 15 years.
- Rescue PCI in either arm rather than scheduled PCI (this is after failed fibrinolysis, therefore not all people had PCI)
- Elective PCI in either arm rather than scheduled PCI (as this is indicative of not being STEMI population).
- Old meta-analyses (pre-2008).
- If, during sifting of the abstract lists, the abstract does not mention PCI occurring after fibrinolysis (as this is indicative of fibrinolysis NOT facilitated PCI).
- Outcomes stated are not post-PCI period just pre-PCI (as this means results are for the fibrinolysis and not the facilitated PCI).
- PCI must have been performed in at least 85% of people in each group.
- Exclude trials of n<60 where there are other larger trials using the same class of intervention (for example, GPI or fibrinolytic) and comparison.
- Exclude trials that did not use stents or <50% people received stents (as this would not be a good representation of current clinical practice). Also exclude studies if they do not mention the percentage of stent usage.

Setting:

- Pre-hospital (pharmacological component of facilitated PCI)
- Secondary care (pharmacological component of facilitated PCI; PCI).

Search strategy

See appendix F

Review strategy

Quality of life data

 Collect all data for the stated quality of life measure, for meta-analysis and GRADE report only overall scores.

Appraisal of methodological quality

The methodological quality of each study will be assessed using NICE checklists and GRADE.

Data synthesis of RCT data

Meta-analysis where appropriate will be conducted.

Subgroups:

The following groups will be considered separately if data is present (we will only consider subgroup analyses when people were stratified according to subgroups or the analyses were pre-specified in the study protocol):

- People with diabetes
- · People from black and minority ethnic groups
- People with renal dysfunction
- People of female gender
- People aged over 70 years (70 is an approximate cut-off age, record what age the study uses that is near to 75 years).

If heterogeneity is present among studies the following subgroups will be examined. in addition to those stated above:

- People receiving balloon angioplasty versus stenting
- People using GPIs administered in the catheterisation laboratory immediately before PCI versus selective GPIs versus no GPIs at this time-point.

Table 3: Review protocol: Radial versus femoral access for PPCI

What is the clinical and cost effectiveness of radial access compared to femoral access for coronary angiography and, if appropriate, follow-on PPCI in people with STEMI managed by PPCI? Objectives To determine whether it is more clinically beneficial to perform PPCI by radial artery access or femoral artery access. Criteria Population: Adults (≥ 18 years old) with STEMI eligible for PPCI. Intervention: Radial access to perform coronary angiography and PPCI (if clinically indicated); Plus standard adjunctive pharmacotherapies (for example, antiplatelet and antithrombin agents). Comparison:

Femoral access to perform coronary angiography and PPCI (if clinically indicated); Plus standard adjunctive pharmacotherapies (for example, antiplatelet and antithrombin agents).

Outcomes (original study definitions will be used and recorded):

- Mortality (all-cause and cardiovascular specific)
- Non-fatal and all (fatal and non-fatal) stroke

- Intracranial bleeding
- Non-fatal and all (fatal and non-fatal) myocardial reinfarction
- · Heart failure
- Repeat revascularisation
- · Access site crossover
- Inability to cross the lesion with a wire, balloon or stent during PCI
- Radiation exposure (X-ray time/fluoroscopic exposure) or procedural time if noted
- PCI procedural success
- Adverse events (major and minor bleeding)
- · Length of hospital stay
- Patient experience (pain)
- Vascular access site complications
- Quality of life (report all, including EQ-5D (EuroQol), SF-36 (Short Form 36), SF6D (Short Form 6-Dimensions), SF-12 (Short Form 12-Dimensions), RAND-36 (Research and Development Medical Outcomes Study Short Form-36)).

Study design:

RCT

Population size and directness:

- No limitations on sample size.
- Studies with indirect populations will not be considered.
- Only studies published after 1990 will be considered. This is to ensure that the extracted evidence is reflective of current practice, especially with regard to the widespread adoption of stenting in place of balloon angioplasty for PPCI procedures over the last 15 years.
- Where ≥3 RCTs (with a combined population of ≥500 people) deploy stents in≥50% of PPCI procedures (in which stenting is feasible) we will exclude studies where stents are deployed in <50% of PPCI procedures.
- Where <3 RCTs deploy stents in ≥50% of PPCI procedures (or the total population of RCTs deploying stents is <500 people) we will include all studies that began enrolling people after 1996.
- If <3 RCTs began enrolling people after 1996 (or the total population is <500 people), we will consider all studies published after 1990.

Setting:

Secondary care

Search strategy

See appendix F

Review strategy

Quality of life data

Collect all data for the stated quality of life measure, for meta-analysis and GRADE report only overall scores.

Appraisal of methodological quality

The methodological quality of each study will be assessed using NICE checklists and GRADE.

Data synthesis of RCT data

Meta-analysis where appropriate will be conducted.

Subgroups:

The following groups will be considered separately if data is present (we will only consider

subgroup analyses when people were stratified according to subgroups or the analyses were pre-specified in the study protocol):

- People with diabetes
- People with renal dysfunction
- BMI
- People aged > 70 years
- Gender
- Ethnicity

If heterogeneity is present among studies the following subgroups will be examined in addition to those stated above:

- Operator expertise
- People receiving GPIs versus people not receiving GPIs
- BMI.

Table 4: Review protocol: Thrombus extraction during PPCI

		·
Revie quest		What is the clinical and cost effectiveness of using thrombus extraction devices (catheter aspiration devices, mechanical thrombectomy devices) during PPCI compared with PPCI alone for the treatment of STEMI in adults?
Obje	ctives	To determine the clinical and cost effectiveness of thrombus extraction devices in people with STEMI managed by PCI.

Criteria Population:

Adults (≥ 18 years old) with STEMI managed by PPCI.

Intervention:

PPCI plus thrombus extraction device.

The devices will be sub-grouped:

- Thrombus aspiration devices (includes simple, manual, and aspiration devices)
- Mechanical thrombus extraction (includes mechanical and non-manual devices).

Comparator:

Standard PPCI.

Outcomes (original study definitions will be used and recorded):

- Mortality (all-cause and cardiovascular specific)
- Non-fatal and all (fatal and non-fatal) stroke.
- Non-fatal and all (fatal and non-fatal) myocardial reinfarction
- Heart failure
- Target vessel revascularisation
- Major and minor bleeding. Note intracranial bleeding separately.
- Length of hospital stay
- Quality of life (report all, inc EQ-5D (EuroQol), SF-36 (Short Form 36), SF6D (Short Form 6-Dimensions), SF-12 (Short Form 12-Dimensions), RAND-36 (Research and Development Medical Outcomes Study Short Form-36)).

Study design:

RCT

Population size and directness:

- No limitations on sample size.
- Studies with indirect populations will not be considered.

- Only studies published after 1990 will be considered. This is to ensure that the extracted evidence is reflective of current practice, especially with regard to the widespread adoption of stenting in place of balloon angioplasty for PCI procedures over the last 15 years.
- Where ≥3 RCTs (with a combined population of ≥500 people) deploy stents in≥50% of PCI procedures (in which stenting is feasible) we will exclude studies where stents are deployed in <50% of PCI procedures.
- Where <3 RCTs deploy stents in ≥50% of PCI procedures (or the total population of RCTs deploying stents is <500 people) we will include all studies that began enrolling people after 1996.
- If <3 RCTs began enrolling people after 1996 (or the total population is <500 people), we will consider all studies published after 1990.

Setting:

Secondary care

Search Strategy

See appendix F

Review Strategy

Quality of life data

Collect all data for the stated quality of life measure, for meta-analysis and GRADE report only overall scores.

Appraisal of methodological quality

The methodological quality of each study will be assessed using NICE checklists and GRADE.

Data synthesis of RCT data:

Meta-analysis where appropriate will be conducted.

Table 5: Review protocol: Culprit versus complete revascularisation

**Updated, see the 2020 evidence review **

Review question	What is the clinical and cost effectiveness of multivessel PCI compared to culprit-only PPCI in people with STEMI and multivessel coronary disease undergoing primary PCI (PPCI)?
Objectives	To compare the clinical and cost effectiveness of multivessel coronary artery, primary percutaneous revascularisation and culprit-only primary percutaneous revascularisation in people with STEMI and multivessel coronary disease.
Criteria	Population: Adults (≥ 18 years old) with STEMI and multivessel coronary disease.

Intervention:

Culprit vessel only PPCI. Plus standard adjunctive pharmacotherapies (for example, antiplatelet and antithrombin agents). Culprit vessel only PPCI defined as PPCI confined to culprit vessel lesions only.

Comparison:

- Multivessel PCI during the index procedure). Plus standard adjunctive pharmacotherapies
 (for example, antiplatelet and antithrombin agents). Multivessel PCI defined as PCI in which
 lesions in the culprit vessel as well as ≥ 1 non-culprit vessel were treated during the same
 procedure.
- Staged multivessel PCI. Plus standard adjunctive pharmacotherapies (for example, antiplatelet and antithrombin agents). Staged PCI defined as PCI confined to the culprit vessel only after which ≥ 1 non-culprit vessel were treated during planned secondary procedures. The timing of the secondary procedure is during the index hospitalisation for STEMI. The timing of staged PCI procedures was defined as reported in each study.

Outcomes (original study definitions will be used and recorded):

- Mortality (all-cause and cardiovascular specific)
- Non-fatal and all (fatal and non-fatal) stroke
- Non-fatal and all (fatal and non-fatal) myocardial reinfarction
- Heart failure
- Unplanned revascularisation
- Major and minor bleeding. Note intracranial bleeding separately
- Contrast-induced nephropathy (also note population that goes onto dialysis/renal replacement therapy)
- Length of hospital stay
- Radiation exposure (fluoroscopic time/X-ray time) or if not recorded, procedural time
- PCI procedural success
- Quality of life (report all, including EQ-5D (EuroQol), SF-36 (Short Form 36), SF6D (Short Form 6-Dimensions), SF-12 (Short Form 12-Dimensions), RAND-36 (Research and Development Medical Outcomes Study Short Form-36)).

Study design:

Cohort study > 500 people

RCT.

Population size and directness:

- No limitations on sample size for RCTs; limitation for cohort studies > 500 people.
- Studies with indirect populations will not be considered.
- Only studies published after 1990 will be considered. This is to ensure that the extracted
 evidence is reflective of current practice, especially with regard to the widespread adoption
 of stenting in place of balloon angioplasty for PCI procedures over the last 15 years.
- Where ≥ 3 RCTs (with a combined population of ≥ 500 people) deploy stents in≥ 50% of PCI procedures (in which stenting is feasible) we will exclude studies where stents are deployed in < 50% of PCI procedures.
- Where < 3 RCTs deploy stents in ≥\50% of PCI procedures (or the total population of RCTs deploying stents is <\500 people) we will include all studies that began enrolling people after 1996.
- If < 3 RCTs began enrolling people after 1996 (or the total population is < 500 people), we will consider all studies published after 1990.

Setting:

Secondary care

Search strategy

See appendix F

Review strategy

Quality of life data

Collect all data for the stated quality of life measure, for meta-analysis and GRADE report only overall scores

Appraisal of methodological quality

The methodological quality of each study will be assessed using NICE checklists and GRADE.

Data synthesis of RCT data

Meta-analysis where appropriate will be conducted.

Subgroups:

The following groups will be considered separately if data is present (we will only consider subgroup analyses when people were stratified according to subgroups or the analyses were

pre-specified in the study protocol):

- Gender
- · Black and minority ethnic groups
- People with diabetes
- People with renal dysfunction
- People aged over 70 years.

If heterogeneity is present among studies the following subgroups will be examined in addition to those stated above:

- People who are haemodynamically compromised versus people who are not
- People with cardiogenic shock versus no cardiogenic shock
- Stent use
- GPI use.

Table 6: Review protocol: Cardiogenic shock

Review question	In people with cardiogenic shock due to STEMI what is the clinical and cost effectiveness of early revascularisation compared to medical stabilisation?
Objectives	To compare routine early revascularisation with initial medical stabilisation in people with cardiogenic shock due to STEMI.

Criteria Population:

Adults (≥ 18 years old) with cardiogenic shock due to STEMI.

Intervention:

Early revascularisation (undefined time frame). Defined as immediate angiography and PPCI or CABG when indicated.

Comparison:

Initial medical stabilisation (with or without subsequent revascularisation).

Outcomes (original study definitions will be used and recorded):

- Mortality (all-cause and cardiovascular specific)
- Non-fatal and all (fatal and non-fatal) stroke.
- Non-fatal and all (fatal and non-fatal) myocardial reinfarction
- Heart failure
- Unplanned revascularisation
- Major and minor bleeding. Note intracranial bleeding separately
- Renal failure (use of dialysis)
- · Length of hospital stay
- Use of IABP (surrogate haemodynamic measures)
- Quality of life (report all, inc EQ-5D (EuroQol), SF-36 (Short Form 36), SF6D (Short Form 6-Dimensions), SF-12 (Short Form 12-Dimensions), RAND-36 (Research and Development Medical Outcomes Study Short Form-36)).

Study design:

RCT.

Population size and directness:

- No limitations on sample size.
- Studies with indirect populations will not be considered.
- Only studies published after 1990 will be considered. This is to ensure that the extracted evidence is reflective of current practice, especially with regard to the widespread adoption

of stenting in place of balloon angioplasty for PCI procedures over the last 15 years.

- Where ≥3 RCTs (with a combined population of ≥500 people) deploy stents in≥50% of PCI procedures (in which stenting is feasible) we will exclude studies where stents are deployed in <50% of PCI procedures.
- Where <3 RCTs deploy stents in ≥50% of PCI procedures (or the total population of RCTs deploying stents is <500 people) we will include all studies that began enrolling people after 1996.
- If <3 RCTs began enrolling people after 1996 (or the total population is <500 people), we will consider all studies published after 1990.

Setting:

Secondary care Emergency care.

Search Strategy

See appendix F

Review Strategy

Quality of life data

Collect all data for the stated QoL measure, for meta-analysis and GRADE report only overall scores.

Appraisal of methodological quality

The methodological quality of each study will be assessed using NICE checklists and GRADE.

Data synthesis of RCT data and subgroups:

Meta-analysis where appropriate will be conducted. The following groups will be considered separately or as subgroups if data is present (we will only consider subgroup analyses when people were stratified according to subgroups or the analyses were pre-specified in the study protocol):

- People with diabetes
- People with renal dysfunction
- People aged over 70 years (or as reported in the studies)
- Ethnicity
- Gender.

Table 7: Review protocol: People who remain unconscious after a cardiac arrest

Review	Does immediate angiography followed by PPCI where indicated improve outcomes of people
question	with presumed STEMI who are resuscitated but remain unconscious after a cardiac arrest?

Objectives To determine whether immediate angiography followed by PPCI where indicated improves outcomes in people who are resuscitated but remain unconscious after a cardiac arrest.

Criteria Population:

Adults (≥ 18 years old) who are resuscitated but remain unconscious after an out of hospital cardiac arrest due to STEMI.

Intervention:

Upon hospital entry, immediate transfer for coronary angiography followed by PPCI where indicated. Plus standard adjunctive pharmacotherapies (for example, antiplatelet and antithrombin agents) with or without therapeutic hypothermia.

Comparator:

Usual care. Defined as any care not involving immediate angiography followed by PPCI where indicated.

Outcomes (original study definitions will be used and recorded):

- Mortality (all-cause and cardiovascular specific)
- Non-fatal and all (fatal and non-fatal) stroke
- Non-fatal and all (fatal and non-fatal) myocardial reinfarction
- Heart failure
- Unplanned urgent target vessel revascularisation
- Major and minor bleeding; note intracranial bleeding separately
- Use of intra-aortic balloon pump
- · Length of hospital stay
- Neurologically intact survival at discharge (CPC score) or other measures of neurological disability
- Quality of life (report all, inc EQ-5D (EuroQol), SF-36 (Short Form 36), SF6D (Short Form 6-Dimensions), SF-12 (Short Form 12-Dimensions), RAND-36 (Research and Development Medical Outcomes Study Short Form-36)).

Study design:

RCTs and systematic reviews first. Prospective comparative observational prospective and retrospective studies if no RCTs are available.

Population size and directness:

No limitations on sample size.

Studies with indirect populations will not be considered.

Setting:

Pre-hospital

Hospital.

Search strategy

See appendix F

strategy Review

Quality of life data

strategy

Collect all data for the stated quality of life measure, for meta-analysis and GRADE report only overall scores.

Appraisal of methodological quality

The methodological quality of each study will be assessed using NICE checklists and GRADE.

Data synthesis of RCT data

Meta-analysis where appropriate will be conducted.

Subgroups:

The following groups will be considered separately if data is present (we will only consider subgroup analyses when people were stratified according to subgroups or the analyses were prespecified in the study protocol):

- People with diabetes
- People with renal dysfunction
- People aged over 70 years
- Ethnicity
- Gender.

If heterogeneity is present among studies the following subgroups will be examined in addition to those stated above:

People receiving GPIs versus people not receiving GPIs

• Balloon angioplasty versus stenting.

Table 8: Review protocol: Hospital volumes of PPCI

Review question	What is the impact of high volume versus low volume PPCI services on patient outcomes?
Objectives	To consider the clinical and cost effectiveness of volume on PPCI services in people with STEMI.
Criteria	Population: Adults (≥ 18 years old) with ST-segment elevation.

Prognostic factor:

Hospital PPCI volume (not rescue, facilitated, elective angiography).

Outcomes at following time intervals: in hospital, 30 days, 6 months, 1 year, longest follow-up (original study definitions will be used and recorded):

- Mortality (all-cause and cardiovascular specific)
- Non-fatal and all (non-fatal and fatal) stroke
- Non-fatal and all (non-fatal and fatal) myocardial reinfarction
- Heart failure
- Unplanned revascularisation (where information available we will record whether index lesion or not)
- · Major and minor bleeding
- Length of hospital stay
- Quality of life (report all, including EQ-5D (EuroQol), SF-36 (Short Form 36), SF6D (Short Form 6-Dimensions), SF-12 (Short Form 12-Dimensions), RAND-36 (Research and Development Medical Outcomes Study Short Form-36)).

Study design:

Prospective and retrospective observational data.

Population size and directness:

- Majority of the data collected post 1 January 1995
- Studies with >1000 participants
- Studies with indirect populations will not be considered
- Only data collected after 1995 will be considered with ≥50% of stent use in PCI procedures.
 This is to ensure that the extracted evidence is reflective of current practice, especially with regard to the widespread adoption of stenting in place of balloon angioplasty for PCI procedures over the last 15 years.

Setting:

Secondary care.

Search Strategy

See appendix F

Review Strategy

Quality of life data

Collect all data for the stated quality of life measure, for meta-analysis and GRADE report only overall scores.

Appraisal of methodological quality

The methodological quality of each study will be assessed using NICE checklists and GRADE.

Subgroups:

The following groups will be considered separately if data is present (we will only consider subgroup analyses when people were stratified according to subgroups or the analyses were pre-specified in the study protocol):

- Gender
- Black and minority ethnic groups
- People with diabetes
- People with renal dysfunction
- People aged over 70 years.

Table 9: Review	Review protocol: Pre-hospital versus in-hospital fibrinolysis What is the clinical and cost effectiveness of pre-hospital versus in-hospital fibrinolysis?
question	what is the chinical and cost effectiveness of pre-nospital versus in-nospital hornlolysis?
Objectives	To determine whether it is more beneficial to administer fibrinolysis before a patient arrives at hospital or in-hospital
Criteria	Population:
	Adults (≥ 18 years old) with STEMI to be managed by fibrinolysis.
	Intervention:
	Pre-hospital fibrinolysis (all agents). Plus standard adjunctive pharmacotherapies (for example, antiplatelet and antithrombin agents).
	Comparison:
	In-hospital fibrinolysis (all agents). Plus standard adjunctive pharmacotherapies (for example, antiplatelet and antithrombin agents).
	Outcomes (original study definitions will be used and recorded):
	Mortality (all-cause and cardiovascular specific)
	All-cause stroke (non-fatal and fatal)
	Intracranial bleeding
	Myocardial reinfarction (non-fatal and fatal)
	Heart failure
	Major and minor bleeding
	Subsequent revascularisation
	Length of hospital stay
	 Quality of life (report all, inc EQ-5D (EuroQol), SF-36 (Short Form 36), SF6D (Short Form 6-Dimensions), SF-12 (Short Form 12-Dimensions), RAND-36 (Research and Development Medical Outcomes Study Short Form-36)).
	Study design:
	RCTs.
	Population size and directness:

Setting:

- Pre-hospital
- Secondary care.

No limitations on sample size.

• Studies with indirect populations will not be considered.

Search See appendix F

Strategy	
Review	Quality of life data:
Strategy	 Collect all data for the stated QoL measure, for meta-analysis and GRADE report only overall scores.
	Appraisal of methodological quality:
	• The methodological quality of each study will be assessed using NICE checklists and GRADE.
	Data synthesis of RCT data and subgroups:
	Meta-analysis where appropriate will be conducted. If heterogeneity is present among studies the following subgroups will be examined:
	People with diabetes
	People with renal dysfunction
	• People aged > 70 years
	• Ethnicity
	• Gender.

Table 10: Review protocol: Use of antithrombin as an adjunct to fibrinolysis

Table 10: Review protocol: Use of antithrombin as an adjunct to fibrinolysis		
Does administration of anti-thrombin treatment at the same time as pre-hospital fibrinolysis improve outcomes compared to administration of pre-hospital fibrinolysis alone?		
To determine whether administration of anti-thrombin treatment at the same time as pre- hospital fibrinolysis improves outcomes compared to administration of pre-hospital fibrinolysis alone.		
Population:		
Adults (≥ 18 years old) with STEMI managed by pre-hospital fibrinolysis (reteplase or tenecteplase).		
Interventions:		
Unfractionated heparin plus antiplatelet therapy.		
LMWH plus antiplatelet therapy.		
Bivalirudin plus antiplatelet therapy.		
Fondaparinux plus antiplatelet therapy.		
Comparison:		
No pre-hospital anti-thrombin treatment.		
Outcomes (original study definitions will be used and recorded):		
Mortality (all-cause and cardiovascular specific)		
Non-fatal and all (fatal and non-fatal) stroke.		
Non-fatal and all (fatal and non-fatal) myocardial reinfarction		
Heart failure		
Unplanned revascularisation		
Major and minor bleeding. Note intracranial bleeding separately		
Length of hospital stay		
 Quality of life (report all, including EQ-5D (EuroQol), SF-36 (Short Form 36), SF6D (Short Form 6-Dimensions), SF-12 (Short Form 12-Dimensions), RAND-36 (Research and Development Medical Outcomes Study Short Form-36)). 		
Study design:		
RCT.		

Population size and directness:

No limitations on sample size

Studies with indirect populations will not be considered.

Setting:

Emergency care, secondary care.

Search Strategy

See appendix F

Review Strategy

Quality of life data

Collect all data for the stated quality of life measure, for meta-analysis and GRADE report only overall scores.

Appraisal of methodological quality

The methodological quality of each study will be assessed using NICE checklists and GRADE.

Data synthesis of RCT data

Meta-analysis where appropriate will be conducted.

Subgroups:

The following groups will be considered separately if data is present (we will only consider subgroup analyses when people were stratified according to subgroups or the analyses were pre-specified in the study protocol):

- People with diabetes
- People with renal dysfunction
- People aged over 70 years.

Table 11:	Review protocol: Rescue PCI
Review question	What is the clinical and cost effectiveness of rescue PCI, repeated fibrinolysis or conservative management compared to each other in people with STEMI who fail to reperfuse after fibrinolytic therapy?
Objectives	To compare the clinical and cost effectiveness of rescue PCI, repeated fibrinolysis and conservative management in people with STEMI who fail to reperfuse after fibrinolytic therapy.
Criteria	Population:
	Adults (≥ 18 years old) with STEMI who fail to reperfuse after fibrinolytic therapy (any agent).
	Intervention:
	Rescue PCI; defined as unplanned emergency PCI in people who fail to reperfuse after fibrinolysis. Plus standard adjunctive pharmacotherapies (for example, antiplatelet and antithrombin agents).
	Comparison:
	Repeat fibrinolysis (any agent) plus standard adjunctive pharmacotherapies (for example,

- antiplatelet and antithrombin agents) or
- Conservative management. Defined as standard medical therapy for MI without fibrinolysis or PCI.

Outcomes at following time intervals: in hospital, 30 days, 6 months, 1 year, longest followup (original study definitions will be used and recorded):

• Mortality (all-cause and cardiovascular specific)

- Non-fatal and all (non-fatal and fatal) stroke
- Non-fatal and all (non-fatal and fatal) myocardial reinfarction
- Heart failure
- Unplanned revascularisation (Where information is available we will record whether index lesion or not)
- · Major and minor bleeding
- · Length of hospital stay
- Quality of life (report all, including EQ-5D (EuroQol), SF-36 (Short Form 36), SF6D (Short Form 6-Dimensions), SF-12 (Short Form 12-Dimensions), RAND-36 (Research and Development Medical Outcomes Study Short Form-36)).

Study design:

RCT

Population size and directness:

- No limitations on sample size.
- Studies with indirect populations will not be considered.
- Only studies published after 1990 will be considered. This is to ensure that the extracted evidence is reflective of current practice, especially with regard to the widespread adoption of stenting in place of balloon angioplasty for PCI procedures over the last 15 years.
- Where ≥3 RCTs (with a combined population of ≥500 people) deploy stents in≥50% of PCI procedures (in which stenting is feasible) we will exclude studies where stents are deployed in <50% of PCI procedures.
- Where <3 RCTs deploy stents in ≥50% of PCI procedures (or the total population of RCTs deploying stents is <500 people) we will include all studies that began enrolling people after 1996.
- If <3 RCTs began enrolling people after 1996 (or the total population is <500 people), we will consider all studies published after 1990.

Setting:

Secondary care

Search Strategy

See appendix F

Review Strategy

Quality of life data

Collect all data for the stated quality of life measure, for meta-analysis and GRADE report only overall scores.

Appraisal of methodological quality

The methodological quality of each study will be assessed using NICE checklists and GRADE.

Data synthesis of RCT data

Meta-analysis where appropriate will be conducted.

Subgroups:

The following groups will be considered separately if data is present (we will only consider subgroup analyses when people were stratified according to subgroups or the analyses were pre-specified in the study protocol):

- Gender
- Black and minority ethnic groups
- People with diabetes
- People with renal dysfunction

- *People aged over 70 years (or age breakdown as reported in the study). If heterogeneity is present among studies the following subgroups will be examined in addition to those above:
- People receiving balloon angioplasty versus stenting
- People receiving GPIs with PCI versus no GPIs with PCI.

Table 12: Review protocol: Routine early angiography following fibrinolysis

Review	What is the clinical and cost effectiveness of routine early angiography following STEMI
question	successfully treated by fibrinolysis compared to routine deferred or selective angiography?
Objectives	To compare the clinical and cost effectiveness of routine early angiography following STEMI

Criteria Population:

Adults (≥ 18 years old) who were fibrinolysed (any agent) for STEMI.

treated by fibrinolysis to routine deferred or selective angiography.

Intervention:

Early routine angiography (and intervention if indicated) following fibrinolysis. Defined as planned angiography within 24 h after fibrinolysis was administered.

Comparison:

Conservative approach following fibrinolysis. Defined as either routine deferred (≥24 h after fibrinolysis administered) or selective angiography.

Outcomes at following time intervals: in hospital, 30 days and 1 year (or closest to 1 year)(original study definitions will be used and recorded):

- Mortality (all-cause and cardiovascular specific)
- Non-fatal and all (non-fatal and fatal) fatal stroke
- Non-fatal and all (non-fatal and fatal) myocardial reinfarction
- Unplanned revascularisation (Where information is available we will record whether index lesion or not)
- Major and minor bleeding. Intracranial bleeding recorded separately
- Length of hospital stay
- · Refractory ischaemia
- Heart failure
- Quality of life (report all, including EQ-5D (EuroQol), SF-36 (Short Form 36), SF6D (Short Form 6-Dimensions), SF-12 (Short Form 12-Dimensions), RAND-36 (Research and Development Medical Outcomes Study Short Form-36)).

Study design:

RCTs

Population size and directness:

- No limitations on sample size.
- Studies with indirect populations will not be considered.
- Only studies published after 1990 will be considered. This is to ensure that the extracted
 evidence is reflective of current practice, especially with regard to the widespread adoption
 of stenting in place of balloon angioplasty for PCI procedures over the last 15 years.
- Where ≥3 RCTs (with a combined population of ≥500 people) deploy stents in≥50% of PCI procedures (in which stenting is feasible) we will exclude studies where stents are deployed in <50% of PCI procedures.
- Where <3 RCTs deploy stents in ≥50% of PCI procedures (or the total population of RCTs deploying stents is <500 people) we will include all studies that began enrolling people after

1996.

• If <3 RCTs began enrolling people after 1996 (or the total population is <500 people), we will consider all studies published after 1990.

Setting:

Secondary care

Search Strategy See appendix F

Review Strategy

Quality of life data

Collect all data for the stated quality of life measure, for meta-analysis and GRADE report only overall scores.

Appraisal of methodological quality

The methodological quality of each study will be assessed using NICE checklists and GRADE.

Data synthesis of RCT data

Meta-analysis where appropriate will be conducted.

Subgroups:

The following groups will be considered separately if data is present (we will only consider subgroup analyses when people were stratified according to subgroups or the analyses were pre-specified in the study protocol):

- Gender
- Black and minority ethnic groups
- People with diabetes
- People with renal dysfunction
- People aged over 70 years.

If heterogeneity is present among studies the following subgroups will be examined in addition to those above:

- High risk versus low risk patients
- Mean time interval to angiography after fibrinolysis
- People receiving GPIs with PCI versus no GPIs with PCI.

Table 13: Health economic review protocol

Table 13. Health economic review protocol	
Review question	All review questions – health economic evidence
Objectives	To identify economic studies relevant to the review questions set out above.
Criteria	Populations, interventions and comparators as specified in the individual review protocols above. Must be a relevant economic study design (cost–utility analysis, cost–benefit analysis, cost–effectiveness analysis, cost–consequence analysis, comparative cost analysis).
Search strategy	An economic study search was undertaken using population-specific terms and an economic study filter – see Appendix F.
Review strategy	Each study is assessed using the NICE economic evaluation checklist which can be found in Appendix H of the NICE guidelines manual (2009). ⁸⁷
	Inclusion and exclusion criteria
	 If a study is rated as both 'Directly applicable' and 'Minor limitations' (using the NICE economic evaluation checklist) then it will be included in the guideline. An evidence table will be completed and it should be included in the economic profile.
	• If a study is rated as 'Partially applicable', 'Potentially serious limitations' or both then there

is discretion over whether it should be included. The health economist will make a decision based on the relative applicability and quality of the available evidence for that question, in discussion with the GDG if required. The ultimate aim is to include studies that are helpful for decision-making in the context of the guideline and the current NHS setting. Where exclusions occur on this basis this will be noted in the relevant section of the guideline with references.

- If a study is rated as 'Very serious limitations' then it will usually be excluded from the guideline. The health economist will make a decision based on the relative quality of the available evidence for that question, in discussion with the GDG if required.
- If a study is rated as 'Not applicable' then it will be excluded from the guideline. An evidence table will not be completed and it will not be included in the economic profile.

The following will also be excluded:

- unpublished reports unless submitted as part of a call for evidence
- abstract-only studies
- letters
- editorials and commentaries
- reviews of economic evaluations
- · articles not in English.

Where there is discretion

The health economist will be guided by the following hierarchies.

Setting:

- UK NHS
- OECD countries with predominantly public health insurance systems (for example, France, Germany, Sweden)
- OECD countries with predominantly private health insurance systems (for example, USA, Switzerland)
- non-OECD settings (always 'Not applicable').

Economic study type:

- cost-utility analysis (QALYs as outcome measure)
- other type of full economic evaluation (cost–benefit analysis, cost-effectiveness analysis, cost–consequence analysis)
- · comparative cost analysis
- non-comparative cost analyses including cost of illness studies (always 'Not applicable').

Year of analysis:

- Studies that are based on resource use and unit costs from more than 10 years ago will be downgraded in terms of applicability.
- Studies that are based on resource use and unit costs from more than 20 years ago will be judged 'Not applicable'.

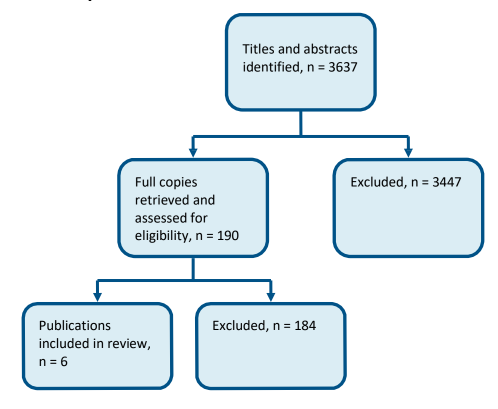
Quality and relevance of effectiveness data used in the economic analysis:

The more closely the effectiveness data used in the economic analysis matches with the
outcomes of the studies included in the clinical review the more useful the analysis will be
for decision-making for the guideline.

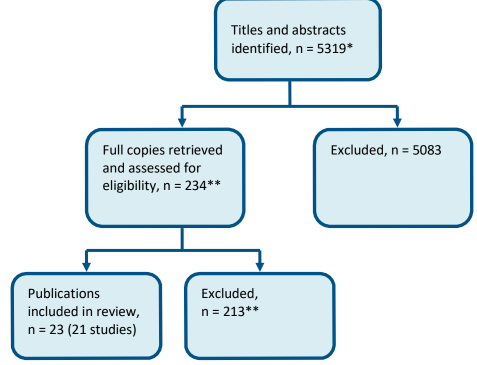
(a) Recent reviews will be ordered although not reviewed. The bibliographies will be checked for relevant studies, which will then be ordered

Appendix D: Clinical article selection

D.1 Time to reperfusion



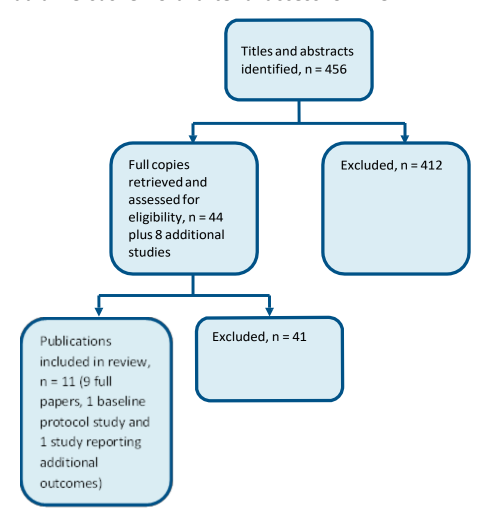
D.2 Facilitated PPCI



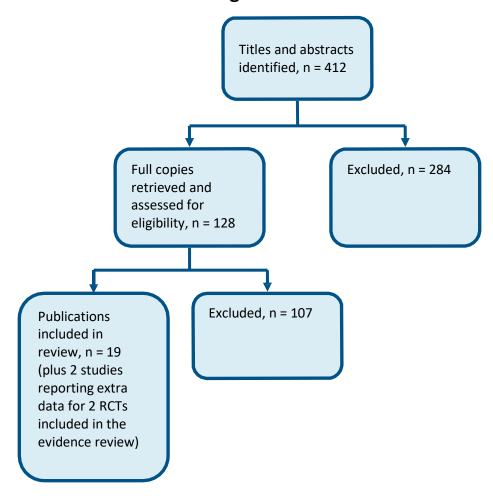
^{*}This includes n = 12 found by cross-referencing

^{**}Some studies retrieved from the search were duplicates so have not been included in the final (n = 174) number of excluded studies

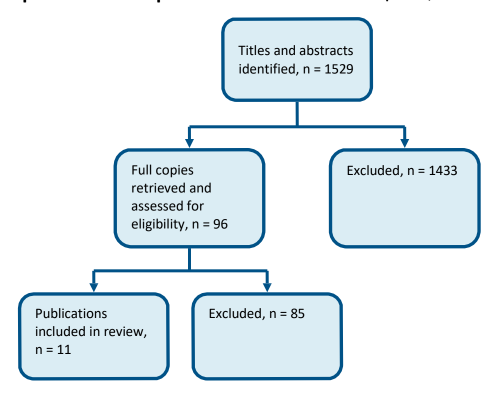
D.3 Radial versus femoral arterial access for PPCI



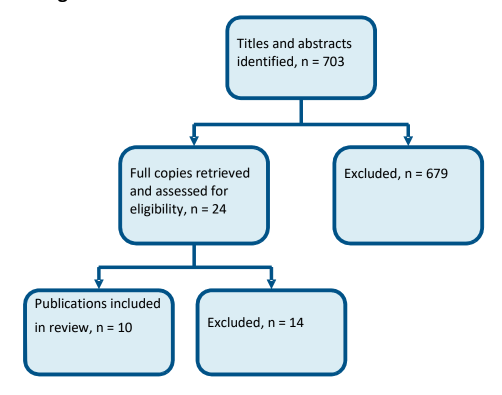
D.4 Thrombus extraction during PPCI



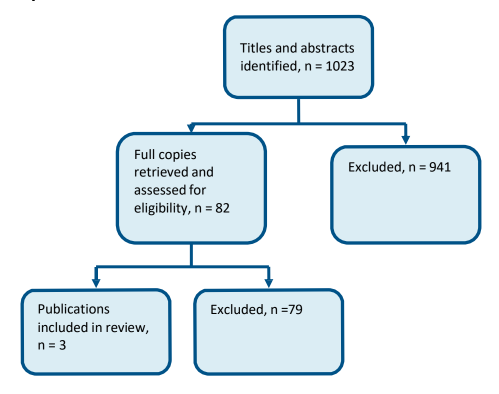
D.5 Culprit versus complete revascularisation **Updated, see 2020 evidence review**



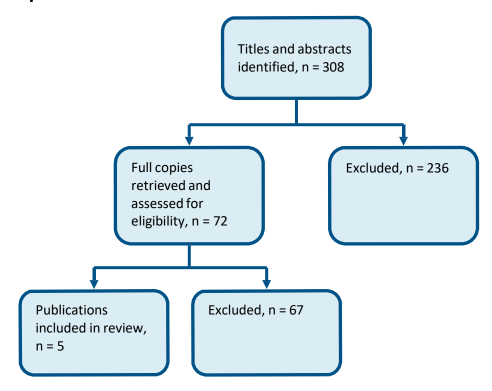
D.6 Cardiogenic shock



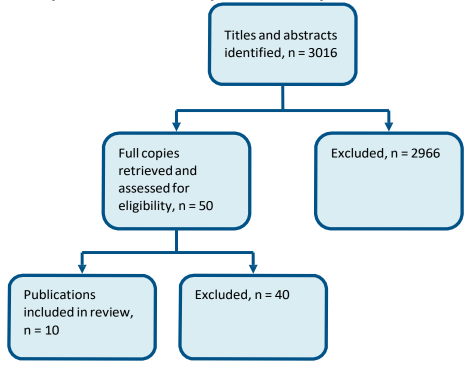
D.7 People who remain unconscious after a cardiac arrest



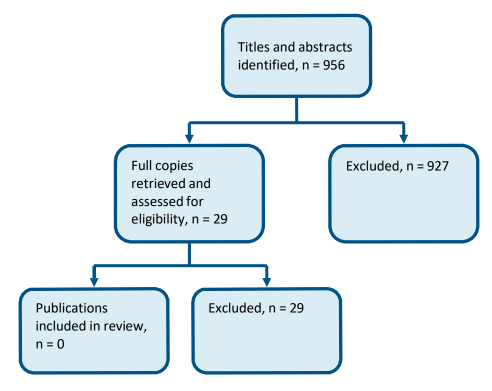
D.8 Hospital volumes of PPCI



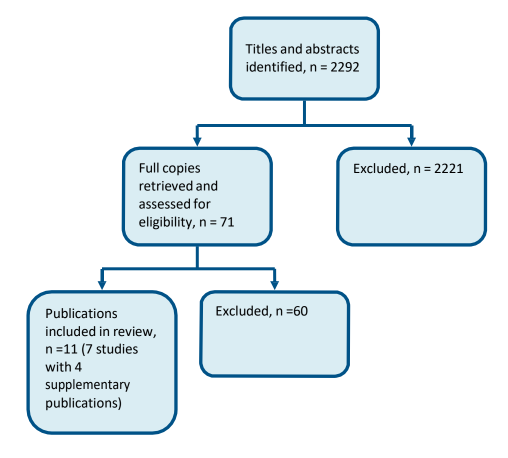
D.9 Pre-hospital versus in-hospital fibrinolysis



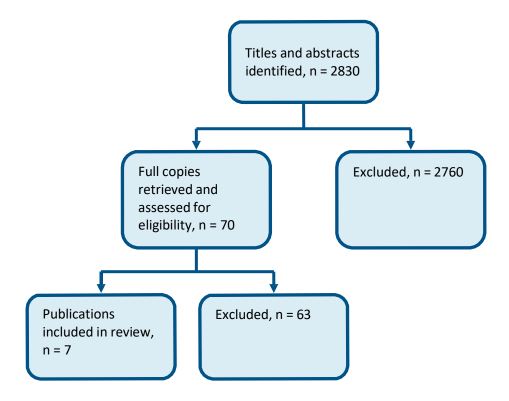
D.10 Use of antithrombin as an adjunct to fibrinolysis



D.11 Rescue PCI

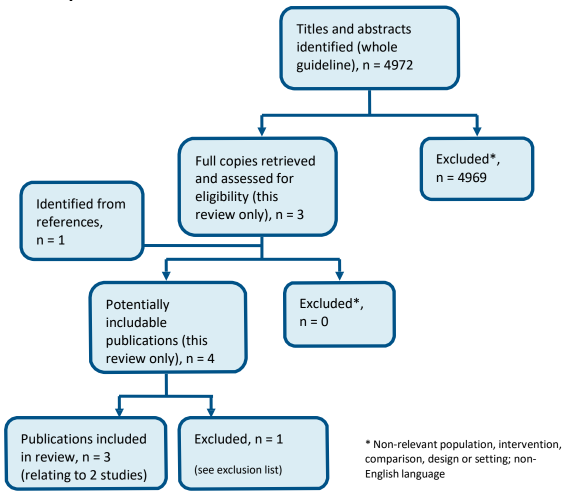


D.12 Routine early angiography following fibrinolysis

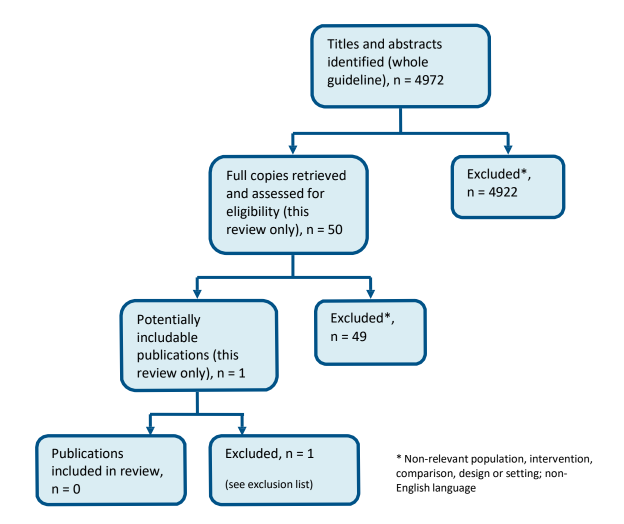


Appendix E: Economic article selection

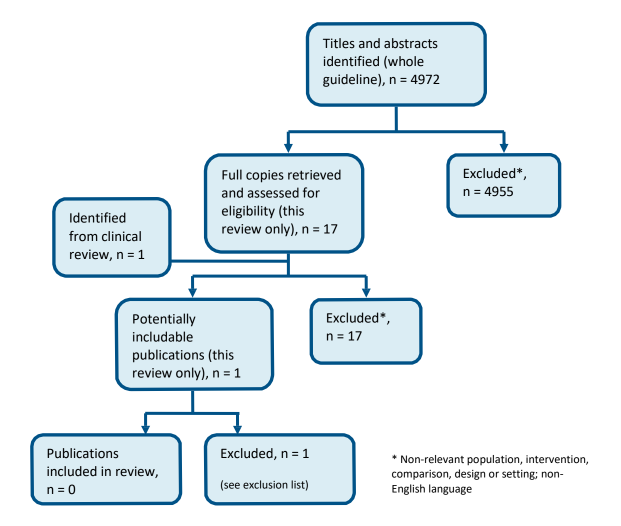
E.1 Time to reperfusion



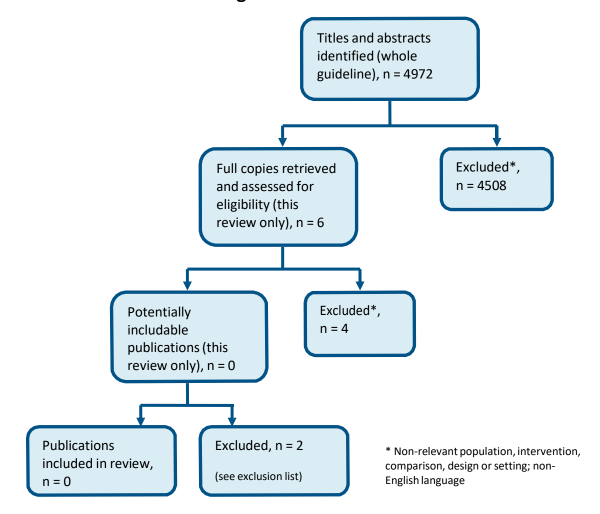
E.2 Facilitated PPCI



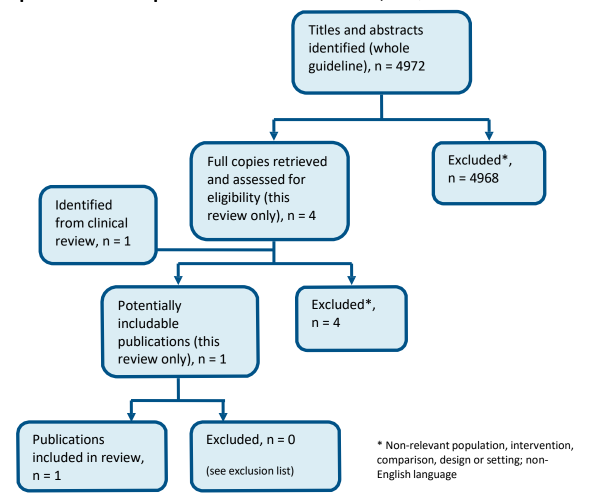
E.3 Radial versus femoral arterial access for PPCI



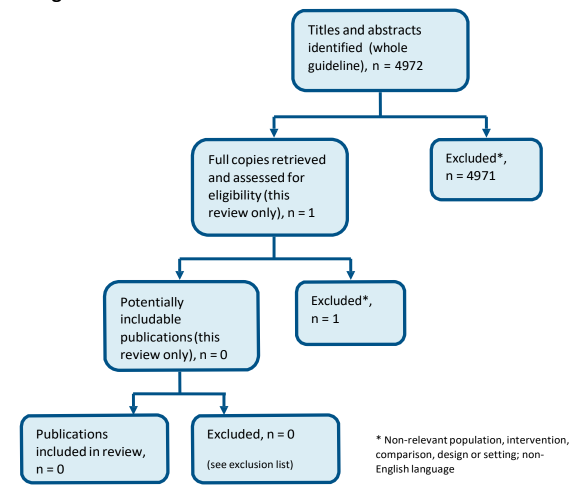
E.4 Thrombus extraction during PPCI



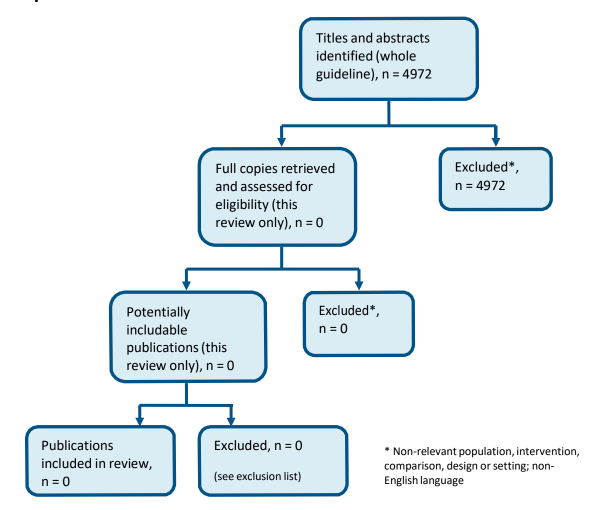
E.5 Culprit versus complete revascularisation **Updated, see the 2020 evidence review**



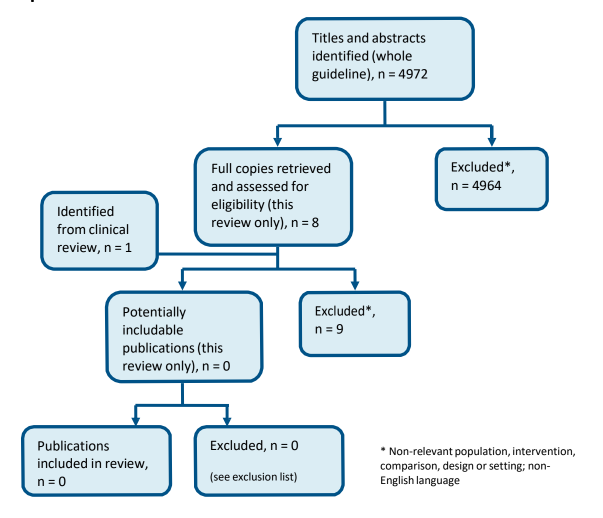
E.6 Cardiogenic shock



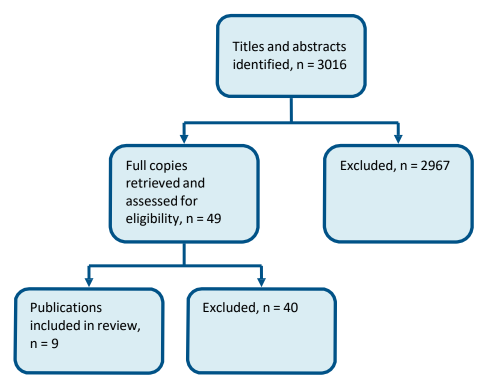
E.7 People who remain unconscious after a cardiac arrest



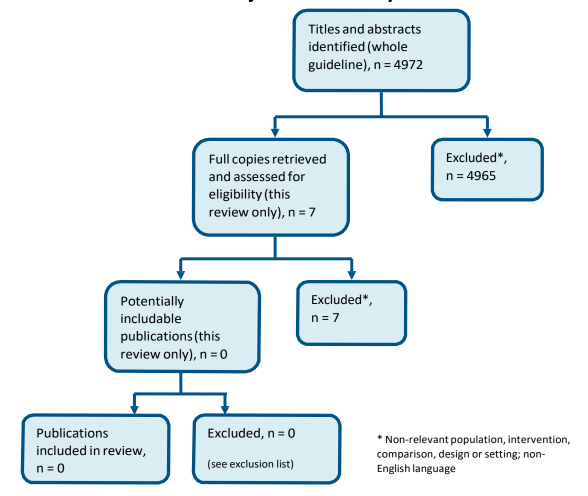
E.8 Hospital volumes of PPCI



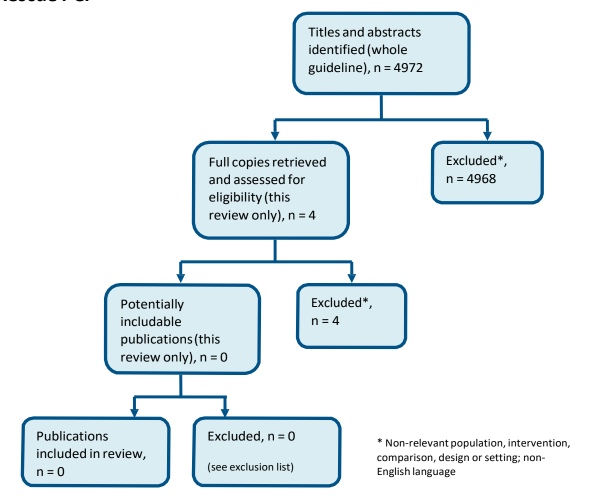
E.9 Pre-hospital versus in-hospital fibrinolysis



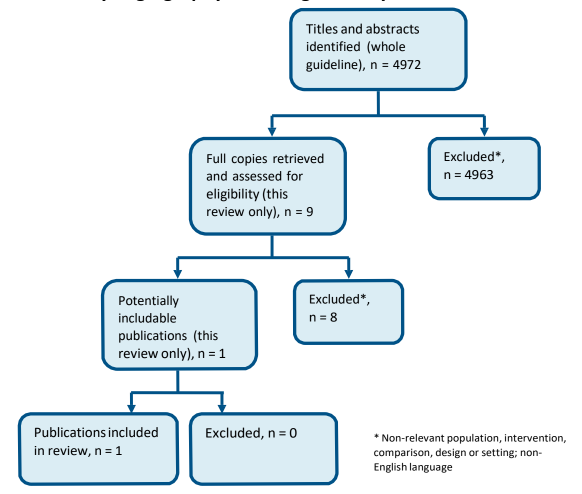
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Appendix F: Literature search strategies

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F.2 Introduction

Search strategies used for the STEMI guideline are outlined below and were run in accordance with the NICE guidelines manual (2009).⁸⁷

All searches were run up to 29 November 2012 unless otherwise stated. Any studies added to the databases after this date were not included unless specifically stated in the text. Journal web sites

were not searched in addition to the databases. Where possible searches were limited to retrieve material published in English.

Scoping searches

Scoping searches were conducted in January 2011 using the following websites and databases (listed below in alphabetical order). Browsing or simple search strategies were employed. The search results were used to provide information for scope development and project planning.

Guidelines	Website address	
Clinical Knowledge Summaries	www.cks.nhs.uk	
CMA Infobase (Canadian guidelines)	www.cma.ca/cpgs	
Guidelines International Network	www.g-i-n.net	
National Guidelines Clearinghouse	www.guideline.gov	
New Zealand Guidelines Group	www.nzgg.org.nz	
NHMRC (Australian Guidelines)	www.nhmrc.gov.au/guidelines/	
NICE Guidelines	guidance.nice.org.uk	
Scottish Intercollegiate Guidelines Network	www.sign.ac.uk	
TRIP Database	www.tripdatabase.com	
Reviews, clinical evidence sources, economic evaluations	Website address	
BMJ Clinical Evidence	clinicalevidence.bmj.com	
Cochrane Library (Systematic Reviews)	www.thecochranelibrary.com	
NHS Evidence	www.nelh.nhs.uk	
Other sources as agreed by reviewers	Website address	
British National Formulary (BNF)	www.bnf.org	
electronic Medicines Compendium (eMC)	www.medicines.org.uk	

Clinical searches

Searches for the **clinical reviews** were run in Medline (OVID), Embase (OVID) and the Cochrane Library. Searches were usually constructed using a PICO format: where population (P) terms were combined with Intervention (I) and sometimes Comparison (C) terms. Outcomes (O) are rarely used in search strategies for interventions. Search filters were also added to the search where appropriate.

Economic searches

Searches for the **health economic reviews** were run in Medline (OVID), Embase (OVID), the NHS Economic Evaluations Database (NHS EED), the Health Technology Assessment (HTA) database and the Health Economic Evaluation Database (HEED). HTA and NHSEED searches were carried out via the Centre for Reviews and Dissemination (CRD) interface. The HTA, NHS EED and HEED databases were accessed via the Wiley interface. Searches in these three databases were constructed using population terms only. For Medline and Embase a health economic filter (instead of a study type filter) was added to the standard population search strategy.

F.3 Population search strategies

F.3.1 Standard population search strategy

Medline search terms

1	exp *myocardial infarction/
2	myocardial infarct*.ti,ab.
3	(cardiac adj (infarct* or attack* or arrest* or event*)).ti,ab.
4	(stemi or st-segment or st segment or st-elevat* or st elevat*).ti,ab.
5	acute coronary syndrome/
6	acute coronary syndrome*.ti,ab,kw.
7	or/1-6

Embase search terms

1	myocardial infarct*.ti,ab.
2	(cardiac adj (infarct* or attack* or arrest* or event*)).ti,ab.
3	(stemi or st-segment or st segment or st-elevat* or st elevat*).ti,ab.
4	acute coronary syndrome/
5	acute coronary syndrome*.ti,ab,kw.
6	exp *heart infarction/
7	exp st segment elevation myocardial infarction/
8	or/1-7

Cochrane search terms

1	MeSH descriptor Myocardial Infarction explode all trees
2	(myocardial next infarct*):ti,ab
3	(cardiac next (infarct* or attack* or arrest* or event*)):ti,ab
4	(st-elevat* or st-segment* or stemi):ti,ab,kw
5	st next elevat*:ti,ab,kw
6	st next segment:ti,ab,kw
7	(("acute coronary") next syndrome*):ti,ab,kw
8	MeSH descriptor Acute Coronary Syndrome, this term only
9	(#1 or #2 or #3 or #4 or #5 or #6 or #7 or #8)

F.3.2 STEMI and Cardiogenic shock

Medline search terms

1.	exp *myocardial infarction/	
2.	myocardial infarct*.ti,ab.	
3.	(cardiac adj (infarct* or attack* or arrest* or event*)).ti,ab.	
4.	(stemi or st-segment or st segment or st-elevat* or st elevat*).ti,ab.	
5.	acute coronary syndrome/	
6.	acute coronary syndrome*.ti,ab,kw.	
7.	or/1-6	
8.	shock.ti,ab.	
9.	7 and 8	
10.	exp shock, cardiogenic/	

11.	(cardio* adj3 shock).ti,ab.
12.	9 or 10 or 11

Embase search terms

1.	myocardial infarct*.ti,ab.
2.	(cardiac adj (infarct* or attack* or arrest* or event*)).ti,ab.
3.	(stemi or st-segment or st segment or st-elevat* or st elevat*).ti,ab.
4.	acute coronary syndrome/
5.	acute coronary syndrome*.ti,ab,kw.
6.	exp *heart infarction/
7.	exp st segment elevation myocardial infarction/
8.	or/1-7
9.	shock.ti,ab.
10.	8 and 9
11.	exp cardiogenic shock/
12.	(cardio* adj3 shock).ti,ab.
13.	10 or 11 or 12

Cochrane search terms

1.	MeSH descriptor Myocardial Infarction explode all trees
2.	myocardial infarct*:ti,ab
3.	st next segment:ti,ab
4.	st next elevat*:ti,ab
5.	(cardiac near/3 (infarct* or attack* or arrest* or event*)):ti,ab
6.	stemi:ti,ab
7.	MeSH descriptor Acute Coronary Syndrome explode all trees
8.	(#1 or #2 or #3 or #4 or #5 or #6 or #7)
9.	shock:ti,ab,kw
10.	(#8 and #9)
11.	MeSH descriptor Shock, Cardiogenic explode all trees
12.	(cardio* next shock):ti,ab
13.	(#10 or #11 or #12)

F.3.3 STEMI and Cardiac arrest

Medline search terms

1.	exp *myocardial infarction/
2.	myocardial infarct*.ti,ab.
3.	(cardiac adj (infarct* or attack* or arrest* or event*)).ti,ab.
4.	(stemi or st-segment or st segment or st-elevat* or st elevat*).ti,ab.
5.	acute coronary syndrome/
6.	acute coronary syndrome*.ti,ab,kw.
7.	or/1-6
8.	heart arrest/
9.	((heart or cardiac) adj2 arrest*).ti,ab.
10.	((surviv* or resuscit* or postresuscit* or unconscious*) adj6 arrest*).ti,ab.
11.	or/8-10

12.	7 and 11
-----	----------

1.	myocardial infarct*.ti,ab.
2.	(cardiac adj (infarct* or attack* or arrest* or event*)).ti,ab.
3.	(stemi or st-segment or st segment or st-elevat* or st elevat*).ti,ab.
4.	acute coronary syndrome/
5.	acute coronary syndrome*.ti,ab,kw.
6.	exp *heart infarction/
7.	exp st segment elevation myocardial infarction/
8.	or/1-7
9.	*heart arrest/ or "out of hospital cardiac arrest"/
10.	((heart or cardiac) adj2 arrest).ti,ab.
11.	((surviv* or resuscit* or postresuscit* or unconscious*) adj6 arrest*).ti,ab.
12.	or/9-11
13.	8 and 12

Cochrane search terms

1.	MeSH descriptor Heart Arrest, this term only
2.	MeSH descriptor Out-of-Hospital Cardiac Arrest, this term only
3.	((heart or cardiac) next arrest*):ti,ab
4.	((unconscious or resuscit* or postresuscit* or surviv*) near/6 arrest):ti,ab
5.	(#1 or #2 or #3 or #4)

F.4 Study filter search terms

F.4.1 Systematic review (SR) search terms

Medline search terms

1	meta-analysis/
2	meta-analysis as topic/
3	(meta analy* or metanaly* or metaanaly*).ti,ab.
4	(reference list* or bibliograph* or hand search* or manual search* or relevant journals).ab.
5	((systematic* or evidence*) adj2 (review* or overview*)).ti,ab.
6	(search strategy or search criteria or systematic search or study selection or data extraction).ab.
7	(search* adj4 literature).ab.
8	(medline or pubmed or cochrane or embase or psychlit or psyclit or psychinfo or psycinfo or cinahl or science citation index or bids or cancerlit).ab.
9	cochrane.jw.
10	or/1-9

1	systematic review/
2	meta-analysis/
3	(meta analy* or metanaly* or metaanaly*).ti,ab.
4	((systematic or evidence) adj2 (review* or overview*)).ti,ab.

1	systematic review/
5	(reference list* or bibliograph* or hand search* or manual search* or relevant journals).ab.
6	(search strategy or search criteria or systematic search or study selection or data extraction).ab.
7	(search* adj4 literature).ab.
8	(medline or pubmed or cochrane or embase or psychlit or psyclit or psychinfo or psycinfo or cinahl or science citation index or bids or cancerlit).ab.
9	((pool* or combined) adj2 (data or trials or studies or results)).ab.
10	cochrane.jw.
11	or/1-10

F.4.2 Randomised controlled studies (RCTs) search terms

Medline search terms

1	randomized controlled trial.pt.
2	controlled clinical trial.pt.
3	randomi#ed.ab.
4	placebo.ab.
5	randomly.ab.
6	Clinical Trials as topic.sh.
7	trial.ti.
8	or/1-8

Embase search terms

1	random*.ti,ab.
2	factorial*.ti,ab.
3	(crossover* or cross over*).ti,ab.
4	((doubl* or singl*) adj blind*).ti,ab.
5	(assign* or allocat* or volunteer* or placebo*).ti,ab.
6	crossover procedure/
7	single blind procedure/
8	randomized controlled trial/
9	double blind procedure/
10	or/1-9

F.4.3 Cohort studies

Medline search terms

	epidemiologic studies/
1.	exp case control studies/
2.	exp cohort studies/
3.	cross-sectional studies/
4.	case control.ti,ab.
5.	(cohort adj (study or studies or analys*)).ti,ab.
6.	((follow up or observational or uncontrolled or non randomi#ed or nonrandomi#ed or epidemiologic*) adj (study or studies)).ti,ab.
7.	((longitudinal or retrospective or prospective or cross sectional) and (study or studies or review or analys* or cohort*)).ti,ab.

	8.	or/1-8
- 1	٥.	0.7 = 0

1.	clinical study/
2.	exp case control study/
3.	family study/
4.	longitudinal study/
5.	retrospective study/
6.	prospective study/
7.	cross-sectional study/
8.	cohort analysis/
9.	follow-up/
10.	cohort*.ti,ab.
11.	9 and 10
12.	case control.ti,ab.
13.	(cohort adj (study or studies or analys*)).ti,ab.
14.	((follow up or observational or uncontrolled or non randomi#ed or nonrandomi#ed or epidemiologic*) adj (study or studies)).ti,ab.
15.	((longitudinal or retrospective or prospective or cross sectional) and (study or studies or review or analys* or cohort*)).ti,ab.
16.	or/1-8,11-14

F.4.4 Health economic search terms

Medline search terms

1.	economics/
2.	value of life/
3.	exp "costs and cost analysis"/
4.	exp economics, hospital/
5.	exp economics, medical/
6.	economics, nursing/
7.	economics, pharmaceutical/
8.	exp "fees and charges"/
9.	exp budgets/
10.	budget*.ti,ab.
11.	cost*.ti.
12.	(economic* or pharmaco?economic*).ti.
13.	(price* or pricing*).ti,ab.
14.	(cost* adj2 (effective* or utilit* or benefit* or minimi* or unit* or estimat* or variable*)).ab.
15.	(financ* or fee or fees).ti,ab.
16.	(value adj2 (money or monetary)).ti,ab.
17.	or/1-16

1.	1. health economics/	
2.	exp economic evaluation/	
3.	exp health care cost/	

4.	exp fee/
5.	budget/
6.	funding/
7.	budget*.ti,ab.
8.	cost*.ti.
9.	(economic* or pharmaco?economic*).ti.
10.	(price* or pricing*).ti,ab.
11.	(cost* adj2 (effective* or utilit* or benefit* or minimi* or unit* or estimat* or variable*)).ab.
12.	(financ* or fee or fees).ti,ab.
13.	(value adj2 (money or monetary)).ti,ab.
14.	or/1-13

F.4.5 Quality of life search terms

Medline search terms

1.	exp quality-adjusted life years/	
2.	quality adjusted life.ti,ab.	
3.	exp "quality of life"/	
4.	exp "value of life"/	
5.	(qaly* or qald* or qale* or qtime*).ti,ab.	
6.	disability adjusted life.ti,ab.	
7.	daly*.ti,ab.	
8.	health status indicators/	
9.	(sf36 or sf 36 or short form 36 or shortform 36 or sf thirtysix or sf thirty six or shortform thirtysix or shortform thirty six or short form thirtysix or short form thirty six).tw.	
10.	(sf6 or sf 6 or short form 6 or shortform 6 or sf six or sfsix or shortform six or short form six).tw.	
11.	(sf12 or sf 12 or short form 12 or shortform 12 or sf twelve or sftwelve or shortform twelve or short form twelve).tw.	
12.	(sf16 or sf 16 or short form 16 or shortform 16 or sf sixteen or sfsixteen or shortform sixteen or short form sixteen).tw.	
13.	(sf20 or sf 20 or short form 20 or shortform 20 or sf twenty or sftwenty or shortform twenty or short form twenty).tw.	
14.	(eurogol or euro gol or eq5d or eq 5d).tw.	
15.	(hql or hqol or h qol or hrqol or hr qol).tw.	
16.	(hye or hyes).tw.	
17.	health* year* equivalent*.tw.	
18.	health utilit*.tw.	
19.	(hui or hui1 or hui2 or hui3).tw.	
20.	disutili*.tw.	
21.	rosser.tw.	
22.	(quality of wellbeing or quality of well being or quality of well-being or qwb).ti,ab.	
23.	willingness to pay.tw.	
24.	standard gamble*.tw.	
25.	(time trade off or time tradeoff or tto).tw.	
26.	or/1-25	

1.	quality adjusted life year/	
2.	quality of life/	
3.	(qaly* or qald* or qale* or qtime*).tw.	
4.	daly*.tw.	
5.	quality adjusted life.tw.	
6.	disability adjusted life.tw.	
7.	(sf36 or sf 36 or short form 36 or shortform 36 or sf thirtysix or sf thirty six or shortform thirtysix or shortform thirty six or short form thirtysix or short form thirty six).tw.	
8.	(sf6 or sf 6 or short form 6 or shortform 6 or sf six or sfsix or shortform six or short form six).tw.	
9.	(sf12 or sf 12 or short form 12 or shortform 12 or sf twelve or sftwelve or shortform twelve or short form twelve).tw.	
10.	(sf16 or sf 16 or short form 16 or shortform 16 or sf sixteen or sfsixteen or shortform sixteen or short form sixteen).tw.	
11.	(sf20 or sf 20 or short form 20 or shortform 20 or sf twenty or sftwenty or shortform twenty or short form twenty).tw.	
12.	(eurogol or euro gol or eq5d or eq 5d).tw.	
13.	(hql or hqol or h qol or hrqol or hr qol).tw.	
14.	(hye or hyes).tw.	
15.	health* year* equivalent*.tw.	
16.	health utilit*.tw.	
17.	(hui or hui1 or hui2 or hui3).tw.	
18.	disutili*.tw.	
19.	rosser.tw.	
20.	(quality of wellbeing or quality of well being or quality of well-being or qwb).ti,ab.	
21.	willingness to pay.tw.	
22.	standard gamble*.tw.	
23.	(time trade off or time tradeoff or tto).tw.	
24.	or/1-23	

F.4.6 Excluded study designs and publication types

The following study designs and publication types were removed from the retrieved results using the NOT operator.

Medline search terms

1	letter/
2	editorial/
3	news/
4	exp historical article/
5	Anecdotes as Topic/
6	comment/
7	case report/
8	(letter or comment*).ti.
9	or/1-8
10	randomized controlled trial/ or random*.ti,ab.
11	9 not 10
12	animals/ not humans/

13	exp Animals, Laboratory/
14	exp Animal Experimentation/
15	exp Models, Animal/
16	exp Rodentia/
17	(rat or rats or mouse or mice).ti.
18	or/11-17

	-
1.	letter.pt. or letter/
2.	note.pt.
3.	editorial.pt.
4.	case report/ or case study/
5.	(letter or comment*).ti.
6.	or/1-5
7.	randomized controlled trial/ or random*.ti,ab.
8.	6 not 7
9.	animal/ not human/
10.	nonhuman/
11.	exp Animal Experiment/
12.	exp Experimental Animal/
13.	animal model/
14.	exp Rodent/
15.	(rat or rats or mouse or mice).ti.
16.	or/8-15

F.5 Searches by specific questions with intervention terms

F.5.1 Time to reperfusion

What is the duration of PPCI-related time delay at which fibrinolysis becomes more clinically and cost effective compared to PPCI in people with STEMI and how is this modulated by patient presentation delay and patient risk profile?

Search constructed by combining the columns in the following table using the 'AND' Boolean operator, except for the exclusion filter which is combined with the search using the 'NOT' Boolean operator

Population	Intervention(s) / exposure(s)	Study filters used (Medline & Embase only)	Date parameters
STEMI	PCI terms AND fibrinolysis terms AND timing terms (line 24 below)	Exclusions	1990 to 29/11/12
STEMI	PCI terms AND fibrinolysis terms (line 23 below)	SRs, RCTs, Exclusions	1990 to 29/11/12

Medline search terms

1.	exp *stents/
2.	angioplasty/

3.	exp *angioplasty, balloon, coronary/
4.	*coronary angiography/
5.	exp *catheterization, peripheral/
6.	*myocardial reperfusion/
7.	percutaneous coronary intervention*.ti,ab.
8.	(peripheral adj3 catheter*).ti,ab.
9.	(coronary adj3 angiograph*).ti,ab.
10.	((heart or myocardi*) adj3 reperfusion).ti,ab.
11.	((primary or coronary or transluminal or balloon) adj3 angioplasty).ti,ab.
12.	coronary artery dilat*.ti,ab.
13.	stent*.ti,ab.
14.	or/1-13
15.	exp *thrombolytic therapy/
16.	((thrombolytic or fibrinolytic) adj1 therap*).ti,ab.
17.	(thrombolysis or fibrinolysis).ti,ab.
18.	fibrinolytic agents/
19.	or/15-18
20.	time factors/
21.	(delay* or timing or time).ti,ab.
22.	20 or 21
23.	14 and 19
24.	14 and 19 and 22

 percutaneous coronary intervention*.ti,ab. ((heart or myocardi\$) adj3 reperfusion).ti,ab. ((Primary or coronary or transluminal or balloon) adj3 angioplasty).ti,ab 	
3. ((Primary or coronary or transluminal or balloon) adj3 angioplasty).ti,ab	_
).
4. Coronary artery dilat\$.ti,ab.	
5. Stent\$.ti,ab.	
6. exp *stent/	
7. exp *transluminal coronary angioplasty/	
8. exp *percutaneous transluminal angioplasty/	
9. angioplasty/	
10. catheterization/	
11. heart muscle reperfusion/	
12. or/9-19	
13. fibrinolytic agent/	
14. exp *fibrinolytic therapy/	
15. ((thrombolytic or fibrinolytic) adj1 therap*).ti,ab.	
16. (thrombolysis or fibrinolysis).ti,ab.	
17. or/21-24	
18. (delay\$ or timing or time).ti,ab.	
19. time/	
20. 26 or 27	
21. 12 and 17 and 20	

1eSH descriptor Stents explode all trees
iestriuescriptor sterius explode all trees
1eSH descriptor Angioplasty, Balloon, Coronary explode all trees
1eSH descriptor Coronary Angiography explode all trees
neSH descriptor Catheterization, Peripheral explode all trees
leSH descriptor Myocardial Reperfusion explode all trees
percutaneous NEXT coronary NEXT intervention*):ti,ab
peripheral NEAR/3 catheter*):ti,ab
heart or myocardi*) NEAR/3 reperfusion):ti,ab
coronary or transluminal or balloon) NEAR/3 angioplasty):ti,ab
Coronary NEXT artery NEXT dilat*):ti,ab
tent*:ti,ab
1 OR #2 OR #3 OR #4 OR #5 OR #6 OR #7 OR #8 OR #9 OR #10 OR #11
neSH descriptor Thrombolytic Therapy explode all trees
1eSH descriptor Fibrinolytic Agents, this term only
thrombolytic or fibrinolytic) NEXT therap*):ti,ab
hrombolysis or fibrinolysis):ti,ab
13 OR #14 OR #15 OR #16 OR #17
1eSH descriptor Time Factors explode all trees
delay* or timing or time):ti,ab
18 or #19
12 and #17 and #20

F.5.2 Facilitated PPCI

What is the clinical and cost effectiveness of facilitated PPCI (fPPCI) compared to PPCI in people with STEMI?

Search constructed by combining the columns in the following table using the 'AND' Boolean operator, except for the exclusion filter which is combined with the search using the 'NOT' Boolean operator

Population	Intervention(s) / exposure(s)	Study filters used (Medline & Embase only)	Date parameters
STEMI	PCI terms AND Pharmacological agents terms (drugs used to facilitate PCI)	SRs, RCTs, Exclusions	1990 to 29/11/12

Medline search terms

1.	exp *stents/
2.	angioplasty/
3.	exp *Angioplasty, Balloon, Coronary/
4.	*Coronary Angiography/
5.	exp *Catheterization, Peripheral/
6.	*Myocardial Reperfusion/
7.	percutaneous coronary intervention*.ti,ab.
8.	(peripheral adj3 catheter\$).ti,ab.
9.	(coronary adj3 angiograph\$).ti,ab.

10.	((heart or myocardi\$) adj3 reperfusion).ti,ab.
11.	((primary or coronary or transluminal or balloon) adj3 angioplasty).ti,ab.
12.	Coronary artery dilat\$.ti,ab.
13.	Stent\$.ti,ab.
14.	or/1-13
15.	(Alteplase or Actilyse or Activase or Altepase or Activacin).ti,ab.
16.	(Reteplase or Rapilysin or Retavase or ecokinase).ti,ab.
17.	(Tenecteplase or Metalyse or TNKase).ti,ab.
18.	(Streptokinase or Streptase or Kabikinase or avelysin or celiase or kinalysin or plasmokinase or plasminokinase or stretodecase or zykinase).ti,ab.
19.	(abciximab or ReoPro or abcixi or centorx).ti,ab.
20.	(Eptifibatide or Integrilin or integrelin or intrifiban).ti,ab.
21.	(Tirofiban or Aggrastat or aggrastet).ti,ab.
22.	Tissue plasminogen activator/
23.	Fibrinolytic agents/
24.	Streptokinase/
25.	Platelet Glycoprotein GPIIb-IIIa Complex/
26.	(Glycoprotein* adj2 IIb*).ti,ab.
27.	or/15-26
28.	heparin/ or heparin, low-molecular-weight/ or dalteparin/ or enoxaparin/ or nadroparin/ or heparinoids/
29.	(Calciparine or Monoparin or Calcium Multiparin or Bemiparin or Zibor or Dalteparin or Fragmin or Enoxaparin or Clexane or Lovenox or Tinzaparin or Innohep or Antixarin or CY 222 or Embolex or monoembolex or Fragmin or Tinzaparin or Suleparoide or Ardeparin or Certoparin or Nadroparin or Parnaparin or Reviparin or Tedelparin).mp.
30.	or/28-29
31.	27 or 30
32.	14 and 31

1.	exp *stent/		
2.	Stent\$.ti,ab.		
3.	*angioplasty/		
4.	exp *percutaneous transluminal angioplasty/		
5.	exp transluminal coronary angioplasty/		
6.	((Primary or coronary or transluminal or balloon) adj3 angioplasty).ti,ab.		
7.	Coronary artery dilat\$.ti,ab.		
8.	*percutaneous coronary intervention/		
9.	percutaneous coronary intervention*.ti,ab.		
10.	catheterization/		
11.	heart muscle reperfusion/		
12.	((heart or myocardi\$) adj3 reperfusion).ti,ab.		
13.	or/1-12		
14.	(Alteplase or Actilyse or Activase or Altepase or Activacin).ti,ab.		
15.	(Reteplase or Rapilysin or Retavase or ecokinase).ti,ab.		
16.	(Tenecteplase or Metalyse or TNKase).ti,ab.		
17.	(Streptokinase or Streptase or Kabikinase or avelysin or celiase or kinalysin or plasmokinase or		

	plasminokinase or stretodecase or zykinase).ti,ab.
18.	(abciximab or ReoPro or abcixi or centorx).ti,ab.
19.	(Eptifibatide or Integrilin or integrelin or intrifiban).ti,ab.
20.	(Tirofiban or Aggrastat or aggrastet).ti,ab.
21.	(Glycoprotein* adj2 IIb*).ti,ab.
22.	Alteplase/
23.	Plasminogen activator/
24.	reteplase/
25.	tenecteplase/
26.	streptokinase/
27.	Abciximab/
28.	Eptifibatide/
29.	Tirofiban/
30.	fibrinogen receptor antagonist/
31.	or/14-30
32.	heparin/ or low molecular weight heparin/ or dalteparin/ or enoxaparin/ or nadroparin/ or heparinoid/
33.	(Calciparine or Monoparin or Calcium Multiparin or Bemiparin or Zibor or Dalteparin or Fragmin or Enoxaparin or Clexane or Lovenox or Tinzaparin or Innohep or Antixarin or CY 222 or Embolex or monoembolex or Fragmin or Tinzaparin or Suleparoide or Ardeparin or Certoparin or Nadroparin or Parnaparin or Reviparin or Tedelparin).mp.
34.	or/32-33
35.	31 or 34
36.	13 and 35

	sedicit terms
1.	MeSH descriptor Stents explode all trees
2.	MeSH descriptor Angioplasty, Balloon, Coronary explode all trees
3.	MeSH descriptor Coronary Angiography explode all trees
4.	MeSH descriptor Catheterization, Peripheral explode all trees
5.	MeSH descriptor Myocardial Reperfusion explode all trees
6.	(percutaneous NEXT coronary NEXT intervention*):ti,ab
7.	(peripheral NEAR/3 catheter*):ti,ab
8.	((heart or myocardi*) NEAR/3 reperfusion):ti,ab
9.	((coronary or transluminal or balloon) NEAR/3 angioplasty):ti,ab
10.	(Coronary NEXT artery NEXT dilat*):ti,ab
11.	Stent*:ti,ab
12.	#1 OR #2 OR #3 OR #4 OR #5 OR #6 OR #7 OR #8 OR #10 OR #11
13.	(Alteplase or Actilyse or Activase or Altepase or Activacin):ti,ab
14.	(Reteplase or Rapilysin or Retavase or ecokinase):ti,ab
15.	(Tenecteplase or Metalyse or TNKase):ti,ab
16.	(Streptokinase or Streptase or Kabikinase or avelysin or celiase or kinalysin or plasmokinase or plasminokinase or stretodecase or zykinase):ti,ab
17.	(abciximab or ReoPro or abcixi or centorx):ti,ab
18.	(Eptifibatide or Integrilin or integrelin or intrifiban):ti,ab
19.	(Tirofiban or Aggrastat or aggrastet):ti,ab
20.	(Glycoprotein* NEAR/2 IIb*):ti,ab

21.	MeSH descriptor Tissue Plasminogen Activator, this term only
22.	MeSH descriptor Fibrinolytic Agents, this term only
23.	MeSH descriptor Streptokinase explode all trees
24.	#13 OR #14 OR #15 OR #16 OR #17 OR #18 OR #19 OR #20 OR #21 OR #22 OR #23 OR #24
25.	MeSH descriptor Heparin explode all trees
26.	MeSH descriptor Dalteparin, this term only
27.	MeSH descriptor Enoxaparin, this term only
28.	MeSH descriptor Nadroparin, this term only
29.	(Calciparine or Monoparin or Calcium Multiparin or Bemiparin or Zibor or Dalteparin or Fragmin or Enoxaparin or Clexane or Lovenox or Tinzaparin or Innohep or Antixarin or CY 222 or Embolex or monoembolex or Fragmin or Tinzaparin or Suleparoide or Ardeparin or Certoparin or Nadroparin or Parnaparin or Reviparin or Tedelparin):ti,ab
30.	#25 OR #26 OR #27 OR #28 OR #29 OR #30
31.	#24 or 30
32.	#12 and #31

F.5.3 Radial versus femoral arterial access for PPCI

What is the clinical and cost effectiveness of radial access compared to femoral access for coronary angiography and, if appropriate, follow-on PPCI in people with STEMI managed by PPCI?

Search constructed by combining the columns in the following table using the 'AND' Boolean operator, except for the exclusion filter which is combined with the search using the 'NOT' Boolean operator

Population	Intervention(s) / exposure(s)	Study filters used (Medline & Embase only)	Date parameters
STEMI	PCI terms AND femoral or radial artery terms	SRs, RCTs, Exclusions	1990 to 29/11/12

Medline search terms

1.	exp *stents/
2.	exp *angioplasty, balloon, coronary/
3.	exp *coronary artery bypass/
4.	*coronary angiography/
5.	exp *catheterization, peripheral/
6.	*myocardial reperfusion/
7.	percutaneous coronary intervention*.ti,ab.
8.	(aortocoronary adj3 bypass).ti,ab.
9.	((coronary or heart) adj3 bypass*).ti,ab.
10.	(peripheral adj3 catheter*).ti,ab.
11.	(coronary adj3 angiograph*).ti,ab.
12.	((heart or myocardi*) adj3 reperfusion).ti,ab.
13.	((coronary or transluminal or balloon) adj3 angioplasty).ti,ab.
14.	coronary artery dilat*.ti,ab.
15.	stent*.ti,ab.
16.	or/1-15
17.	femoral artery/
18.	radial artery/

19.	(radial or femoral or (trans adj radial) or (trans adj femoral)).ti,ab.
20.	or/17-19
21.	16 and 20

percutaneous coronary intervention*.ti,ab.
(aortocoronary adj3 bypass).ti,ab.
((coronary or heart) adj3 bypass*).ti,ab.
(peripheral adj3 catheter*).ti,ab.
(coronary adj3 angiograph*).ti,ab.
((heart or myocardi*) adj3 reperfusion).ti,ab.
((coronary or transluminal or balloon) adj3 angioplasty).ti,ab.
coronary artery dilat*.ti,ab.
stent*.ti,ab.
exp *stent/
exp *transluminal coronary angioplasty/
exp *percutaneous transluminal angioplasty/
coronary artery bypass graft/
catheterization/
heart muscle reperfusion/
or/1-15
radial artery/
exp femoral artery/
(radial or femoral or (trans adj radial) or (trans adj femoral)).ti,ab.
or/17-19
16 and 20

1.	MeSH descriptor Stents explode all trees
2.	MeSH descriptor Angioplasty, Balloon, Coronary explode all trees
3.	MeSH descriptor Coronary Angiography explode all trees
4.	MeSH descriptor Coronary Artery Bypass explode all trees
5.	MeSH descriptor Catheterization, Peripheral explode all trees
6.	MeSH descriptor Myocardial Reperfusion explode all trees
7.	(aortocoronary near/3 bypass*):ti,ab
8.	((coronary or heart) near/3 bypass*):ti,ab
9.	(peripheral near/3 catheter*):ti,ab
10.	(coronary near/3 angiograph*):ti,ab
11.	((heart or myocardi*) near/3 reperfusion):ti,ab
12.	((coronary or transluminal or balloon) near/3 angioplasty):ti,ab
13.	(coronary next artery next dilat*):ti,ab
14.	(percutaneous next coronary next intervention*):ti,ab
15.	stent*:ti,ab
16.	(#1 or #2 or #3 or #4 or #5 or #6 or #7 or #8 or #9 or #10 or #11 or #12 or #13 or #14 or #15)
17.	MeSH descriptor Radial Artery explode all trees
18.	MeSH descriptor Femoral Artery explode all trees

19.	(radial or femoral or (trans next radial) or (trans next femoral)):ti,ab	
20.	(#17 or #18 or #19)	
21.	#16 and #20	

F.5.4 Thrombus extraction during PPCI

What is the clinical and cost effectiveness of using thrombus extraction devices (catheter aspiration devices, mechanical thrombectomy devices) during PPCI compared with PPCI alone for the treatment of STEMI in adults?

Search constructed by combining the columns in the following table using the 'AND' Boolean operator, except for the exclusion filter which is combined with the search using the 'NOT' Boolean operator

Population	Intervention(s) / exposure(s)	Study filters used (Medline & Embase only)	Date parameters
STEMI	PCI AND thrombectomy terms	SRs, RCTs, Exclusions	1990 to 29/11/12

Medline search terms

1.	exp heart catheterization/	
2.	exp stents/	
3.	myocardial reperfusion/	
4.	exp angioplasty/	
5.	exp myocardial revascularization/	
6.	percutaneous coronary intervention*.ti,ab.	
7.	((heart or cardiac) adj3 catheter*).ti,ab.	
8.	((heart or myocardi*) adj3 reperfusion).ti,ab.	
9.	((primary or percutaneous or coronary or transluminal or balloon) adj3 angioplasty).ti,ab.	
10.	(ppci or pci or ptca).ti,ab.	
11.	coronary artery dilat*.ti,ab.	
12.	stent*.ti,ab.	
13.	revasc*.ti,ab.	
14.	or/1-13	
15.	exp thrombectomy/	
16.	mechanical thrombolysis/	
17.	suction/	
18.	embolectomy/	
19.	(thrombectomy or thrombectomies or embolectomy or embolectomies).ti,ab.	
20.	(emboli* adj2 protect*).ti,ab.	
21.	thromboaspiration.ti,ab.	
22.	((thrombus or clot* or embol*) adj2 (remov* or extract* or aspirat*)).ti,ab.	
23.	((mechanical or manual) adj2 (clot disrupt* or thrombolysis or aspirat*)).ti,ab.	
24.	((catheter* or thrombo*) adj2 aspirat*).ti,ab.	
25.	(quickcat or thromcat or angiojet or x-sizer).ti,ab.	
26.	or/15-25	
27.	14 and 26	

1.	percutaneous coronary intervention*.ti,ab.		
2.	((heart or myocardi*) adj3 reperfusion).ti,ab.		
3.	((coronary or transluminal or balloon) adj3 angioplasty).ti,ab.		
4.	coronary artery dilat*.ti,ab.		
5.	stent*.ti,ab.		
6.	exp *stent/		
7.	exp *transluminal coronary angioplasty/		
8.	exp *percutaneous transluminal angioplasty/		
9.	catheterization/		
10.	heart muscle reperfusion/		
11.	or/1-10		
12.	artificial embolism/		
13.	exp thrombectomy/		
14.	coronary artery thrombosis/		
15.	exp percutaneous thrombectomy/		
16.	embolectomy/		
17.	(thrombus adj2 (aspirat* or extracti*)).ti,ab.		
18.	thrombectomy.ti,ab.		
19.	thromboaspiration.ti,ab.		
20.	(emboli* adj2 protect*).ti,ab.		
21.	embolectomy.ti,ab.		
22.	emboli#ation.ti,ab.		
23.	(export or pronto or diver or angiojet or (x adj sizer) or rescue).ti,ab.		
24.	or/12-23		
25.	11 and 24		

Cocinianc	Contrarie Search terms			
1.	MeSH descriptor Heart Catheterization explode all trees			
2.	MeSH descriptor Stents explode all trees			
3.	MeSH descriptor Myocardial Reperfusion, this term only			
4.	MeSH descriptor Angioplasty explode all trees			
5.	MeSH descriptor Myocardial Revascularization explode all trees			
6.	"percutaneous coronary intervention*":ti,ab			
7.	((heart or cardiac) near/3 catheter*):ti,ab			
8.	((heart or myocardi*) near/3 reperfusion):ti,ab			
9.	((percutaneous or primary or coronary or transluminal or balloon) near/3 angioplasty):ti,ab			
10.	(PPCI or PCI or PTCA):ti,ab			
11.	(revasc* or stent*):ti,ab			
12.	(coronary next artery next dilat*):ti,ab			
13.	(#1 or #2 or #3 or #4 or #5 or #6 or #7 or #8 or #9 or #10 or #11 or #12)			
14.	MeSH descriptor Thrombectomy explode all trees			
15.	MeSH descriptor Suction, this term only			
16.	MeSH descriptor Embolectomy, this term only			
17.	MeSH descriptor Mechanical Thrombolysis, this term only			
18.	(thrombectomy or thrombectomies or embolectomy or embolectomies or thromboaspiration or quickcat or thromcat or angiojet or x-sizer):ti,ab			

19.	(emboli* near/2 protect*):ti,ab
20.	((thrombus or clot* or emboli*) near/2 (remov* or extract* or aspirat*)):ti,ab
21.	((mechanical or manual) near/2 ("clot disrupt*" or thrombolysis or aspirat*)):ti,ab
22.	((catheter* or thrombo*) next aspirat*):ti,ab
23.	(#14 or #15 or #16 or #17 or #18 or #19 or #20 or #21 or #22)
24.	(#13 and #23)

F.5.5 Culprit versus complete revascularisation **Updated, see the 2020 evidence review**

What is the clinical and cost effectiveness of multivessel PCI compared to culprit-only PPCI in people with STEMI and multivessel coronary disease undergoing PPCI?

Search constructed by combining the columns in the following table using the 'AND' Boolean operator, except for the exclusion filter which is combined with the search using the 'NOT' Boolean operator

Population	Intervention(s) / exposure(s)	Study filters used (Medline & Embase only)	Date parameters
STEMI	PCI terms AND culprit or multivessel terms	SRs, RCTs, Observational study, Exclusions	1990 to 29/11/12

Medline search terms

1.	exp *stents/	
2.	percutaneous coronary intervention*.ti,ab.	
3.	((primary or coronary or percutaneous or transluminal or balloon) adj3 angioplasty).ti,ab.	
4.	coronary artery dilat*.ti,ab.	
5.	stent*.ti,ab.	
6.	(PPCI or PCI or PTCA).ti,ab.	
7.	exp angioplasty/	
8.	or/1-7	
9.	(culprit or non-culprit or nonculprit).ti,ab.	
10.	((infarct-related or infarct related or non-infarct-related) adj2 (artery or arteries)).ti,ab.	
11.	(complete adj2 revasc*).ti,ab.	
12.	((multivessel or multi-vessel or single-vessel or single vessel) adj3 (percutaneous coronary intervention* or PCI or stent* or revasc* or recanali* or angioplast*)).ti,ab.	
13.	or/9-12	
14.	8 and 13	

1.	percutaneous coronary intervention*.ti,ab.	
2.	coronary artery dilat*.ti,ab.	
3.	stent*.ti,ab.	
4.	exp *stent/	
5.	*percutaneous transluminal angioplasty/ or *angioplasty/ or *laser angioplasty/	
6.	exp *percutaneous coronary intervention/	
7.	(PPCI or PCI or PTCA).ti,ab.	
8.	((primary or percutaneous or coronary or transluminal or balloon) adj3 angioplasty).ti,ab.	
9.	or/1-8	

10.	(culprit or non-culprit or nonculprit).ti,ab.	
11.	((infarct-related or infarct related or non-infarct-related) adj2 (artery or arteries)).ti,ab.	
12.	(complete adj2 revasc*).ti,ab.	
13.	((multivessel or multi-vessel or single vessel or single-vessel) adj3 (percutaneous coronary intervention* or PCI or stent* or revasc* or recanali* or angioplast*)).ti,ab.	
14.	or/10-13	
15.	9 and 14	

1.	MeSH descriptor Stents explode all trees	
2.	MeSH descriptor Angioplasty explode all trees	
3.	stent*:ti,ab	
4.	(PCI or PCTA or PPCI):ti,ab	
5.	"percutaneous coronary intervention":ti,ab	
6.	(("coronary artery") next dilat*):ti,ab	
7.	((primary or coronary or percutaneous or transluminal or balloon) near/3 angioplasty):ti,ab	
8.	#1 or #2 or #3 or #4 or #5 or #6 or #7	
9.	(culprit or nonculprit or non-culprit):ti,ab	
10.	((infarct-related or "infarct related" or non-infarct-related) next (artery or arteries)):ti,ab	
11.	(complete next revasc*):ti,ab	
12.	((multivessel or multi-vessel or "single vessel" or single-vessel) near/3 (angioplas* or pci or stent* or "percutaneous coronary intervention" or recanali* or revasc*)):ti,ab	
13.	#9 or #10 or #11 or #12	
14.	#8 and #13	

F.5.6 Cardiogenic shock

In people with cardiogenic shock due to STEMI what is the clinical and cost effectiveness of early revascularisation compared to medical stabilisation?

Search constructed by combining the columns in the following table using the 'AND' Boolean operator, except for the exclusion filter which is combined with the search using the 'NOT' Boolean operator

Population	Intervention(s) / exposure(s)	Study filters used (Medline & Embase only)	Date parameters
STEMI OR cardiogenic shock terms	PCI terms OR revascularisation terms OR catheter terms OR bypass terms	SRs, RCTs, Exclusions	1990 to 29/11/12

Medline search terms

1.	exp *stents/
2.	*myocardial reperfusion/
3.	percutaneous coronary intervention*.ti,ab.
4.	((heart or myocardi*) adj3 reperfusion).ti,ab.
5.	coronary artery dilat*.ti,ab.
6.	stent*.ti,ab.
7.	((primary or coronary or percutaneous or transluminal or balloon) adj3 angioplasty).ti,ab.
8.	(PPCl or PCl or PTCA).ti,ab.

9.	exp myocardial revascularization/
10.	exp angioplasty/
11.	revasc*.ti,ab.
12.	CABG.ti,ab.
13.	((heart or coronary or aortocoronary or cardio*) adj2 bypass).ti,ab.
14.	(bypass adj2 (surg* or graft*)).ti,ab.
15.	exp *heart catheterization/
16.	((heart or coronary) adj3 catheter*).ti,ab.
17.	or/1-16

1.	percutaneous coronary intervention*.ti,ab.
2.	coronary artery dilat*.ti,ab.
3.	stent*.ti,ab.
4.	exp *stent/
5.	(PPCI or PCI or PTCA).ti,ab.
6.	((primary or percutaneous or coronary or transluminal or balloon) adj3 angioplasty).ti,ab.
7.	heart muscle revascularization/
8.	*heart muscle reperfusion/
9.	exp coronary artery surgery/
10.	*heart catheterization/
11.	((heart or coronary) adj3 catheter*).ti,ab.
12.	exp percutaneous coronary intervention/
13.	angioplasty/ or percutaneous transluminal angioplasty/
14.	revasc*.ti,ab.
15.	CABG.ti,ab.
16.	((heart or coronary or aortocoronary or cardio*) adj2 bypass).ti,ab.
17.	(bypass adj2 (surg* or graft*)).ti,ab.
18.	or/1-17

1.	MeSH descriptor Stents explode all trees
2.	MeSH descriptor Myocardial Reperfusion explode all trees
3.	MeSH descriptor Myocardial Revascularization explode all trees
4.	MeSH descriptor Angioplasty explode all trees
5.	("percutaneous coronary intervention"):ti,ab
6.	((heart or myocardial or coronary) near/3 reperfusion):ti,ab
7.	(("coronary artery" or coronary arteries") next dilat*):ti,ab
8.	stent*:ti,ab
9.	((primary or coronary or percutaneous or transluminal or balloon) near/3 angioplast*):ti,ab
10.	(PCI or PPCI or PTCA or CABG or revasc*):ti,ab
11.	(bypass near/2 (surg* or graft*)):ti,ab
12.	((heart or cardiac or coronary or aortocoronary or cardio*) next bypass):ti,ab
13.	MeSH descriptor Myocardial Infarction explode all trees
14.	MeSH descriptor Heart Catheterization explode all trees
15.	((heart or coronary) near/3 catheter*):ti,ab

16. (#1 or #2 or #3 or #4 or #5 or #6 or #7 or #8 or #9or #10 or #11 or #12 or #13 or	#14or #15)
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F.5.7 People who remain unconscious after a cardiac arrest

Does immediate angiography followed by PPCI where indicated improve outcomes of people with presumed STEMI who are resuscitated but remain unconscious after a cardiac arrest?

Search constructed by combining the columns in the following table using the 'AND' Boolean operator, except for the exclusion filter which is combined with the search using the 'NOT' Boolean operator

Population	Intervention(s) / exposure(s)	Study filters used (Medline & Embase only)	Date parameters
STEMI	PCI terms	SRs, RCTs, Exclusions	1990 to 29/11/12
AND cardiac arrest			

Medline search terms

1.	percutaneous coronary intervention*.ti,ab.
2.	((heart or myocardi*) adj3 reperfusion).ti,ab.
3.	((primary or percutaneous or coronary or transluminal or balloon) adj3 angioplasty).ti,ab.
4.	coronary artery dilat*.ti,ab.
5.	stent*.ti,ab.
6.	exp stents/
7.	exp angioplasty/
8.	myocardial reperfusion/
9.	exp myocardial revascularization/
10.	revasc*.ti,ab.
11.	(PPCI or PCI or PTCA).ti,ab.
12.	((heart or cardiac) adj3 catheter*).ti,ab.
13.	exp heart catheterization/
14.	coronary angiography/
15.	(coronary adj3 angiograph*).ti,ab.
16.	or/1-15

1.	percutaneous coronary intervention*.ti,ab.
2.	((heart or myocardi*) adj3 reperfusion).ti,ab.
3.	((primary or percutaneous or coronary or transluminal or balloon) adj3 angioplasty).ti,ab.
4.	coronary artery dilat*.ti,ab.
5.	stent*.ti,ab.
6.	exp *stent/
7.	exp *percutaneous coronary intervention/
8.	*percutaneous transluminal angioplasty/
9.	*angioplasty/
10.	*heart muscle reperfusion/
11.	heart muscle revascularization/
12.	revasc*.ti,ab.
13.	(PPCI or PCI or PTCA).ti,ab.
14.	((heart or cardiac) adj3 catheter*).ti,ab.

15.	*heart catheterization/
16.	angiocardiography/
17.	(coronary adj3 angiograph*).ti,ab.
18.	or/1-17

1.	MeSH descriptor Heart Catheterization explode all trees		
2.	MeSH descriptor Stents explode all trees		
3.	MeSH descriptor Myocardial Reperfusion, this term only		
4.	MeSH descriptor Angioplasty explode all trees		
5.	MeSH descriptor Myocardial Revascularization explode all trees		
6.	(percutaneous next coronary next intervention*):ti,ab		
7.	((heart or cardiac) near/3 catheter*):ti,ab		
8.	((heart or myocardi*) near/3 reperfusion):ti,ab		
9.	((percutaneous or primary or coronary or transluminal or balloon) near/3 angioplasty):ti,ab		
10.	(PPCI or PCI or PTCA):ti,ab		
11.	(revasc* or stent*):ti,ab		
12.	(coronary next artery next dilat*):ti,ab		
13.	(coronary next angiograph*):ti,ab		
14.	MeSH descriptor Coronary Angiography, this term only		
15.	(#1 or #2 or #3 or #4 or #5 or #6 or #7 or #8 or #9 or #10 or #11 or #12 or #13 or #14)		

F.5.8 Hospital volumes of PPCI

What is the impact of high volume versus low volume PPCI services on patient outcomes?

Search constructed by combining the columns in the following table using the 'AND' Boolean operator, except for the exclusion filter which is combined with the search using the 'NOT' Boolean operator

Intervention(s) /		Study filters used		
Population	exposure(s)	(Medline & Embase only)	Date parameters	
No population	PCI and volume terms	Exclusions	1990 to 29/11/12	

Medline and Embase search terms

1.	workload/
2.	((surg* or physician* or operat* or procedure* or hospital* or cardiol*) adj4 (volume* or workload* or caseload*)).ti,ab.
3.	1 or 2
4.	exp Angioplasty/
5.	percutaneous coronary intervention*.ti,ab.
6.	((primary or coronary or transluminal or balloon) adj3 angioplast*).ti,ab.
7.	or/4-6
8.	3 and 7

1.	MeSH descriptor Workload, this term only
2.	(surg* or physician* or operat* or procedure* or hospital* or cardiol*) near/4 (volume* or workload* or caseload*):ti,ab
3.	(#1 or #2)

4.	MeSH descriptor Angioplasty explode all trees
5.	("percutaneous coronary intervention" or "percutaneous coronary interventions"):ti,ab
6.	((primary or coronary or transluminal or balloon) near/3 angioplast*):ti,ab
7.	(#4 or #5 or #6)
8.	(#3 and #7)

F.5.9 Pre-hospital versus in-hospital fibrinolysis

What is the clinical and cost effectiveness of pre-hospital versus in-hospital fibrinolysis?

Search constructed by combining the columns in the following table using the 'AND' Boolean operator, except for the exclusion filter which is combined with the search using the 'NOT' Boolean operator

Population	Intervention(s) / exposure(s)	Study filters used (Medline & Embase only)	Date parameters
STEMI	(General fibrinolysis terms AND prehospital terms) OR Fibrinolysis agent terms	SRs, RCTs, Exclusions	All years up to 29/11/12

Medline search terms

 (tissue adj2 plasminogen activator*).ti,ab. (alteplase or actilyse or activase).ti,ab. (tenecteplase or metalyse or tnkase).ti,ab. (reteplase or rapilysin or retavase).ti,ab. duteplase.ti,ab. (monteplas* or cleactor).ti,ab. (streptokinase or streptase or kabikinase or mutose or heberkinasa).ti,ab. (urokinase adj2 plasminogen activator*).ti,ab. (urokinase or abbokinase or (syner adj kinase)).ti,ab. (saruplase or (pro adj urokinase) or rescupase).ti,ab. plasminogen streptokinase activator*.ti,ab. (anistreplas* or eminase).ti,ab. 	
 (tenecteplase or metalyse or tnkase).ti,ab. (reteplase or rapilysin or retavase).ti,ab. duteplase.ti,ab. (monteplas* or cleactor).ti,ab. (streptokinase or streptase or kabikinase or mutose or heberkinasa).ti,ab. (urokinase adj2 plasminogen activator*).ti,ab. (urokinase or abbokinase or (syner adj kinase)).ti,ab. (saruplase or (pro adj urokinase) or rescupase).ti,ab. plasminogen streptokinase activator*.ti,ab. 	
 (reteplase or rapilysin or retavase).ti,ab. duteplase.ti,ab. (monteplas* or cleactor).ti,ab. (streptokinase or streptase or kabikinase or mutose or heberkinasa).ti,ab. (urokinase adj2 plasminogen activator*).ti,ab. (urokinase or abbokinase or (syner adj kinase)).ti,ab. (saruplase or (pro adj urokinase) or rescupase).ti,ab. plasminogen streptokinase activator*.ti,ab. 	
 duteplase.ti,ab. (monteplas* or cleactor).ti,ab. (streptokinase or streptase or kabikinase or mutose or heberkinasa).ti,ab. (urokinase adj2 plasminogen activator*).ti,ab. (urokinase or abbokinase or (syner adj kinase)).ti,ab. (saruplase or (pro adj urokinase) or rescupase).ti,ab. plasminogen streptokinase activator*.ti,ab. 	
 (monteplas* or cleactor).ti,ab. (streptokinase or streptase or kabikinase or mutose or heberkinasa).ti,ab. (urokinase adj2 plasminogen activator*).ti,ab. (urokinase or abbokinase or (syner adj kinase)).ti,ab. (saruplase or (pro adj urokinase) or rescupase).ti,ab. plasminogen streptokinase activator*.ti,ab. 	
 (streptokinase or streptase or kabikinase or mutose or heberkinasa).ti,ab. (urokinase adj2 plasminogen activator*).ti,ab. (urokinase or abbokinase or (syner adj kinase)).ti,ab. (saruplase or (pro adj urokinase) or rescupase).ti,ab. plasminogen streptokinase activator*.ti,ab. 	
 (urokinase adj2 plasminogen activator*).ti,ab. (urokinase or abbokinase or (syner adj kinase)).ti,ab. (saruplase or (pro adj urokinase) or rescupase).ti,ab. plasminogen streptokinase activator*.ti,ab. 	
 (urokinase or abbokinase or (syner adj kinase)).ti,ab. (saruplase or (pro adj urokinase) or rescupase).ti,ab. plasminogen streptokinase activator*.ti,ab. 	
 10. (saruplase or (pro adj urokinase) or rescupase).ti,ab. 11. plasminogen streptokinase activator*.ti,ab. 	
11. plasminogen streptokinase activator*.ti,ab.	
12. (anistreplas* or eminase).ti,ab.	
13. tissue plasminogen activator/	
14. urokinase-type plasminogen activator/	
15. streptokinase/	
16. exp plasminogen activators/	
17. or/1-16	
18. fibrinolysis/	
19. thrombolytic therapy/	
20. (fibrinoly* or thromboly*).ti,ab.	
21. or/18-20	
22. (pre-hospital or prehospital).ti,ab.	
23. 21 and 22	
24. 17 or 23	

1.	(tissue adj2 plasminogen activator*).ti,ab.
2.	(alteplase or actilyse or activase).ti,ab.
3.	(tenecteplase or metalyse or tnkase).ti,ab.
4.	(reteplase or rapilysin or retavase).ti,ab.
5.	duteplase.ti,ab.
6.	(monteplas* or cleactor).ti,ab.
7.	(streptokinase or streptase or kabikinase or mutose or heberkinasa).ti,ab.
8.	(urokinase adj2 plasminogen activator*).ti,ab.
9.	(urokinase or abbokinase or (syner adj kinase)).ti,ab.
10.	(saruplase or (pro adj urokinase) or rescupase).ti,ab.
11.	plasminogen streptokinase activator*.ti,ab.
12.	(anistreplas* or eminase).ti,ab.
13.	tissue plasminogen activator/
14.	alteplase/
15.	urokinase/
16.	tenecteplase/
17.	plasminogen activator/
18.	reteplase/
19.	duteplase/
20.	monteplase/
21.	streptokinase/
22.	saruplase/
23.	anistreplase/
24.	or/1-23
25.	*fibrinolytic therapy/
26.	*fibrinolysis/
27.	(fibrinoly* or thromboly*).ti,ab.
28.	or/25-27
29.	(pre-hospital or prehospital).ti,ab.
30.	28 and 29
31.	24 or 30

1.	(tissue near/2 (plasminogen activator*)):ti,ab
2.	(alteplase or actilyse or activase):ti,ab
3.	(tenecteplase or metalyse or tnkase):ti,ab
4.	(reteplase or rapilysin or retavase):ti,ab
5.	duteplase:ti,ab
6.	(monteplas* or cleactor):ti,ab
7.	(streptokinase or streptase or kabikinase or mutose or heberkinasa):ti,ab
8.	(urokinase near/2 (plasminogen activator*)):ti,ab
9.	(urokinase or abbokinase or (syner next kinase)):ti,ab
10.	(saruplase or (pro next urokinase) or rescupase):ti,ab
11.	("plasminogen streptokinase activator*"):ti,ab
12.	(anistreplas* or eminase):ti,ab

MeSH descriptor Tissue Plasminogen Activator explode all trees
MeSH descriptor Urokinase-Type Plasminogen Activator explode all trees
MeSH descriptor Streptokinase explode all trees
MeSH descriptor Anistreplase explode all trees
MeSH descriptor Plasminogen Activators explode all trees
(#1 or #2 or #3 or #4 or #5 or #6 or #7 or #8 or #9 or #10 or #11 or #12 or #13 or #14 or #15 or #16 or #17)
MeSH descriptor Thrombolytic Therapy, this term only
MeSH descriptor Fibrinolysis, this term only
(fibrinoly* or thromboly*):ti,ab
(#19 or #20 or #21)
(pre-hospital or prehospital):ti,ab
(#22 and #23)
(#18 or #24)

F.5.10 Use of antithrombin as an adjunct to fibrinolysis

Does administration of antithrombin treatment at the same time as pre-hospital fibrinolysis improve outcomes compared to administration of pre-hospital fibrinolysis alone?

Search constructed by combining the columns in the following table using the 'AND' Boolean operator, except for the exclusion filter which is combined with the search using the 'NOT' Boolean operator

Population	Intervention(s) / exposure(s)	Study filters used (Medline & Embase only)	Date parameters
STEMI	Antithrombin agents OR prehospital terms	SRs, RCTs, Exclusions	All years up to 29/11/12

Medline search terms

1.	(reteplase or rapilysin or retavase).ti,ab.
2.	(tenecteplase or TNKase or metalyse).ti,ab.
3.	(pre-hospital or prehospital).ti,ab.
4.	or/1-3

Embase search terms

1.	reteplase/
2.	tenecteplase/
3.	(reteplase or rapilysin or retavase).ti,ab.
4.	(tenecteplase or TNKase or metalyse).ti,ab.
5.	(pre-hospital or prehospital).ti,ab.
6.	or/1-5

1.	(reteplase or rapilysin or retavase or tenecteplase or TNKase or metalyse):ti,ab	
2.	(pre-hospital or prehospital):ti,ab	
3.	(#1 or #2)	

F.5.11 Rescue PCI

What is the clinical and cost effectiveness of rescue PCI, repeated fibrinolysis or conservative management compared to each other in people with STEMI who fail to reperfuse after fibrinolytic therapy?

Search constructed by combining the columns in the following table using the 'AND' Boolean operator, except for the exclusion filter which is combined with the search using the 'NOT' Boolean operator

Population	Intervention(s) / exposure(s)	Study filters used (Medline & Embase only)	Date parameters
STEMI	(PCI terms OR fibrinolysis terms)	SRs, RCTs, Exclusions	1990 to 29/11/12
	AND rescue terms		

Medline search terms

1.	exp *stents/
2.	exp *angioplasty, balloon, coronary/
3.	percutaneous coronary intervention*.ti,ab.
4.	((rescue or emergency or unplanned) adj4 (percutaneous or angioplast*)).ti,ab.
5.	(rescue adj1 (PPCI or PCI)).ti,ab.
6.	exp *thrombolytic therapy/
7.	((thrombolytic or fibrinolytic) adj1 therap*).ti,ab.
8.	(thrombolysis or fibrinolysis).ti,ab.
9.	exp *tissue plasminogen activator/
10.	exp *urokinase-type plasminogen activator/
11.	exp *streptokinase/
12.	(alteplase or actilyse or reteplase or rapilysin or tenecteplase or metalyse or urokinase or syner-?kinase or streptokinase or streptase).ti,ab.
13.	or/1-12
14.	recurrence/
15.	retreatment/
16.	treatment failure/
17.	((fail* or after or repeat* or rescue) adj2 (reperfus* or treat* or therap* or thromb* or fibrino*)).ti,ab.
18.	or/14-17
19.	13 and 18

1.	exp *stent/	
2.	exp *transluminal coronary angioplasty/	
3.	exp *percutaneous transluminal angioplasty/	
4.	percutaneous coronary intervention*.ti,ab.	
5.	((rescue or emergency or unplanned) adj4 (percutaneous or angioplast*)).ti,ab.	
6.	(rescue adj1 (PPCI or PCI)).ti,ab.	
7.	exp *fibrinolytic therapy/	
8.	((thrombolytic or fibrinolytic) adj1 therap*).ti,ab.	
9.	(thrombolysis or fibrinolysis).ti,ab.	
10.	exp *tissue plasminogen activator/	

11.	exp *urokinase/
12.	exp *streptokinase/
13.	(alteplase or actilyse or reteplase or rapilysin or tenecteplase or metalyse or urokinase or syner-?kinase or streptokinase or streptase).ti,ab.
14.	or/1-13
15.	exp recurrent disease/
16.	exp treatment failure/
17.	exp retreatment/
18.	((fail* or after or repeat* or rescue*) adj2 (reperfus* or treat* or therap* or thromb* or fibrino*)).ti,ab.
19.	or/15-18
20.	14 and 19

1.	MeSH descriptor Stents explode all trees	
	- 	
2.	MeSH descriptor Angioplasty, Balloon, Coronary explode all trees	
3.	percutaneous next coronary next intervention*:ti,ab	
4.	((rescue or emergency or unplanned) near/4 (percutaneous or angioplast*)):ti,ab	
5.	(rescue next (PPCI or PCI)):ti,ab	
6.	MeSH descriptor Thrombolytic Therapy explode all trees	
7.	((thrombolytic or fibrinolytic) next therap*):ti,ab	
8.	(thrombolysis or fibrinolysis):ti,ab	
9.	MeSH descriptor Tissue Plasminogen Activator explode all trees	
10.	MeSH descriptor Urokinase-Type Plasminogen Activator explode all trees	
11.	MeSH descriptor Streptokinase explode all trees	
12.	(alteplase or actilyse or reteplase or rapilysin or Tenecteplase or metalyse or urokinase or	
	syner-kinase or Streptokinase or streptase):ti,ab	
13.	#1 or #2 or #3 or #4 or #5 or #6 or #7 or #8 or #9 or #10 or #11or #12	
14.	MeSH descriptor Recurrence explode all trees	
15.	MeSH descriptor Retreatment explode all trees	
16.	MeSH descriptor Treatment Failure explode all trees	
17.	((fail* or after or repeat* or rescue*) near/2 (reperfus* or treat* or therap* or thromb* or fibrino*)):ti,ab	
18.	#14 or #15 or #16 or #17	
19.	#13 and #18	

F.5.12 Routine early angiography following fibrinolysis

What is the clinical and cost effectiveness of routine early angiography following STEMI successfully treated by fibrinolysis compared to routine deferred or selective angiography?

Search constructed by combining the columns in the following table using the 'AND' Boolean operator, except for the exclusion filter which is combined with the search using the 'NOT' Boolean operator

Population	Intervention(s) / exposure(s)	Study filters used (Medline & Embase only)	Date parameters
STEMI	Fibrinolyis terms AND angiography terms	SRs, RCTs, Exclusion	1990 to 29/11/12

Medline search terms

1.	exp *thrombolytic therapy/
2.	exp *fibrinolytic agents/
3.	(antithrombic or antithrombotic or anti-thrombic or anti-thrombotic or thrombolytic or thrombolysis or fibrinolytic or fibrinolysis).ti,ab.
4.	or/1-3
5.	*coronary angiography/
6.	(angiogram or angiograph* or angioplasty or stent*).ti,ab.
7.	exp *angioplasty/
8.	exp *stents/
9.	or/5-8
10.	4 and 9

Embase search terms

1.	exp *fibrinolytic therapy/
2.	exp *fibrinolytic agent/
3.	(antithrombic or antithrombotic or anti-thrombic or anti-thrombotic or thrombolytic or thrombolysis or fibrinolytic or fibrinolysis).ti,ab.
4.	or/1-3
5.	exp *angiocardiography/
6.	exp *angioplasty/
7.	exp *stent/
8.	(angiogram or angiograph* or angioplasty or stent*).ti,ab.
9.	or/5-8
10.	4 and 9

1.	MeSH descriptor Thrombolytic Therapy explode all trees
2.	MeSH descriptor Fibrinolytic Agents explode all trees
3.	(antithrombic or antithrombotic or anti-thrombic or anti-thrombotic or thrombolytic or thrombolysis or fibrinolytic or fibrinolysis):ti,ab,kw
4.	(#1 or #2 or #3)
5.	MeSH descriptor Coronary Angiography explode all trees
6.	MeSH descriptor Angioplasty explode all trees
7.	MeSH descriptor Stents explode all trees
8.	(angiogram or angiograph* or angioplasty or stent*):ti,ab,kw
9.	(#5 or #6 or #7 or #8)
10.	(#4 and #9)

F.6 Economics search

F.6.1 General economic search

Search constructed by combining the columns in the following table using the 'AND' Boolean operator, except for the exclusion filter which is combined with the search using the 'NOT' Boolean operator.

Population	Intervention(s) / exposure(s)	Study filters used (Medline & Embase only)	Date parameters
STEMI OR PCI terms OR	none	Economic, Exclusions	Medline & Embase 2010 to 29/11/12
angioplasty terms (search terms below)			NHS EED and HEED all years to 29/11/12

Medline search terms

1.	exp *myocardial infarction/
2.	myocardial infarct*.ti,ab.
3.	(cardiac adj (infarct* or attack* or arrest* or event*)).ti,ab.
4.	(stemi or st-segment or st segment or st-elevat* or st elevat*).ti,ab.
5.	acute coronary syndrome/
6.	acute coronary syndrome*.ti,ab,kw.
7.	or/1-6
8.	*angiography/
9.	angiocardiography/
10.	*coronary angiography/
11.	(angiograph* or arteriograph* or angiocardiograph* or angiogram* or cardioangiograph* or angiocardiogram* or angio cardiograph* or coronarograph*).ti,ab.
12.	*myocardial revascularization/
13.	*myocardial reperfusion/
14.	((myocardi* or coronary or heart or cardiac) adj2 (revasculari* or reperfus*)).ti,ab.
15.	pci.ti,ab.
16.	ptca.ti,ab.
17.	exp angioplasty/
18.	blunt microdissection.ti,ab.
19.	angioplast*.ti,ab.
20.	((percutaneous or balloon or coronary or transluminal or primary) adj3 (dilation or dilatation or intervention*)).ti,ab.
21.	((coronary or drug-eluting or "bare metal") adj2 stent*).ti,ab.
22.	or/8-21
23.	7 or 22

1.	myocardial infarct*.ti,ab.	
2.	(cardiac adj (infarct* or attack* or arrest* or event*)).ti,ab.	
3.	(stemi or st-segment or st segment or st-elevat* or st elevat*).ti,ab.	
4.	acute coronary syndrome/	

5.	acute coronary syndrome*.ti,ab,kw.
6.	exp *heart infarction/
7.	exp st segment elevation myocardial infarction/
8.	or/1-7
9.	*Angiography/
10.	Angiocardiography/
11.	Coronary Angiography/
12.	Angiograph*.ti,ab.
13.	Arteriograph*.ti,ab.
14.	Angiocardiograph*.ti,ab.
15.	Coronary Arteriogra*.ti,ab.
16.	Angiogram*.ti,ab.
17.	Cardioangiograph*.ti,ab.
18.	Angiocardiogram*.ti,ab.
19.	Angio Cardiograph*.ti,ab.
20.	Coronary Arteriogra*.ti,ab.
21.	Coronarograph*.ti,ab.
22.	*Heart Muscle Revascularization/
23.	Angioplasty, Transluminal, Percutaneous Coronary/
24.	(Myocardial adj (revascularisation or revascularization)).ti,ab.
25.	PCI.ti,ab.
26.	Percutaneous coronary intervention.ti,ab.
27.	Percutaneous transluminal coronary angioplasty.ti,ab.
28.	PTCA.ti,ab.
29.	exp Angioplasty/
30.	Blunt Microdissection.ti,ab.
31.	((laser or patch) adj angioplasty).ti,ab.
32.	Percutaneous Transluminal Angioplasty.ti,ab.
33.	Transluminal Coronary Angioplasty.ti,ab.
34.	(Balloon adj3 coronary).ti,ab.
35.	(Balloon adj3 angioplasty).ti,ab.
36.	(Coronary adj2 stent*).ti,ab.
37.	or/9-36
38.	8 or 37

CRD search terms

1.	MeSH DESCRIPTOR Acute Coronary Syndrome IN NHSEED,HTA
2.	MeSH DESCRIPTOR Myocardial Infarction EXPLODE ALL TREES IN NHSEED, HTA
3.	MeSH DESCRIPTOR Coronary Thrombosis IN NHSEED, HTA
4.	(acute coronary syndrome*) OR (myocardial infarct*) OR (coronary NEAR thrombos*) IN NHSEED, HTA
5.	(heart NEAR infarct*) OR (heart NEAR attack) OR (heart NEAR arrest) OR (heart NEAR event) IN NHSEED, HTA
6.	(STEMI) OR (st-segment) OR (st segment) OR (st-elevation) OR (st elevation) IN NHSEED, HTA
7.	#1 OR #2 OR #3 OR #4 OR #5 OR #6
8.	MeSH DESCRIPTOR Angiography IN NHSEED,HTA

9.	MeSH DESCRIPTOR Angiocardiography IN NHSEED,HTA
10.	MeSH DESCRIPTOR Coronary Angiography IN NHSEED,HTA
11.	(Angiograph*) OR (Arteriograph*) OR (Angiocardiograph*) OR (Coronary Angiograph*) OR (Angiogram*) IN NHSEED, HTA
12.	(Cardioangiograph*) OR (Angiocardiogram*) OR (Angio Cardiograph*) OR (Coronary Arteriogra*) OR (Coronarograph*) IN NHSEED, HTA
13.	MeSH DESCRIPTOR Myocardial Revascularization EXPLODE ALL TREES IN NHSEED,HTA
14.	(Myocardial NEAR revascularization) OR (Myocardial NEAR revascularisation) OR (PCI) OR (Percutaneous coronary intervention) OR (Percutaneous AND Transluminal Coronary angioplasty) IN NHSEED, HTA
15.	MeSH DESCRIPTOR angioplasty EXPLODE ALL TREES IN NHSEED, HTA
16.	(PTCA) OR (Blunt Microdissection) OR (Laser NEAR angioplasty) OR (Patch NEAR angioplasty) OR (Percutaneous Transluminal Angioplasty) IN NHSEED, HTA
17.	(Transluminal Coronary Angioplasty) OR (Balloon NEAR coronary) OR (Balloon NEAR angioplasty) OR (coronary NEAR stent*) IN NHSEED, HTA
18.	#1 OR #2 OR #3 OR #4 OR #5 OR #6 OR #7 OR #8 OR #9 OR #10 OR #11 OR #12 OR #13 OR #14 OR #15 OR #16 OR #17

HEED search terms

IILLD Seal	en terms
1.	AX=Acute coronary syndrome or acute coronary syndromes
2.	Ax=myocardial infarction or myocardial infarct or myorcardial
3.	Ax=Coronary thrombosis
4.	Ax=Heart infarction or heart infarct or heart infarcts
5.	AX=Heart attack or heart attacks
6.	Ax=heart arrest or heart arrests
7.	Ax=heart event or heart events
8.	AX=STEMI
9.	AX=st-segment
10.	AX=st-elevation
11.	Ax=st segment
12.	Ax=st elevation
13.	CS=1 or 2 or 3 or 4 or 5 or 6 or 7 or 8 or 9 or 10 or 11 or 12
14.	Ax=angiography
15.	Ax=Angiocardiography
16.	Ax=Coronary angiography
17.	AX=Angiograph or angiographs
18.	AX=Arteriography or arteriograph or arteriographs
19.	AX=Angiogram or angiograms
20.	Ax=myocardial revascularization
21.	Ax=myocardial revascularisation
22.	Ax=PCI
23.	Ax=Percutaneous coronary intervention
24.	AX=Percutaneous transluminal coronary angioplasty
25.	Ax=PCTA
26.	AX=Angioplasty
27.	AX=Balloon and coronary
28.	AX=Balloon and Angioplasty

29.	Ax=Coronary and Stent
30.	Ax=coronary and stents
31.	CS=14 or 15 or 16 or 17 or 18 or 19 or 20 or 21 or 22 or 23 or 24 or 25 or 26 or 27 or 28 or 29 or 30
32.	CS=13 and 31

F.6.2 Quality of life reviews

These searches were carried out on Medline and Embase only.

Population	Intervention(s) / exposure(s)	Study filters used (Medline & Embase only)	Date parameters
STEMI	none	Quality of life, Exclusions	all years to 29/11/12

Appendix G: Clinical evidence tables

G.1 Time to reperfusion

Table 14: Asseburg 2006⁵

Reference/funding	Methods	Outcomes	RCTs
Asseburg C, et al. Assessing the effectiveness of primary angioplasty compared with fibrinolysis and its relationship to time delay: a Bayesian evidence synthesis. Heart. 2007; 93(10):1244-50. Funding: Unrestricted educational grant from Cordis Ltd; NHS	Updated previously published meta-analysis http://www.ncbi.nlm.nih.gov/pubmed/12517460 (Keeley et al 2003) by searching: Cochrane Controlled Trials Register, UK National Research Register, Medline, Embase, Database of Abstracts of Reviews of Effects, UK National Health Service Economic Evaluation Databases, and the Health Technology Assessment Database for English language RCTs published between 2002 and 2004 New meta-analysis was calculated by Bayesian statistical methods using random effects model (study-level data were used on an ITT basis). Meta-regression was used to estimate the relative treatment effect of PPCI and fibrinolysis as a function of the covariate PPCI-related time delay PPCI-related time delay was defined as the difference between time to balloon in PPCI and time to needle in fibrinolytic treatment, to avoid problems of different timing definitions across RCTs. Mean times (and SD) to treatment were preferred, when unavailable medians (and IQR) were used. A sensitivity analysis was undertaken to determine the effect of uncertainty around input parameters (prior distributions) and differences between fibrinolytic drugs.	Mortality, non-fatal strokes, non-fatal reinfarctions; all at 1 and 6 months PPCI-related time delay associated with equipoise between both reperfusion strategies Outcomes are presented as absolute probabilities of specific events and odds ratios (ORs; with 95% credible intervals (CrI)) as a function of the additional time delay associated with PPCI.	22 RCTs (n = 3760 PPCI; n = 3758 fibrinolysis) Zijlstra 1993; 1997 Ribeiro 1993 de Boer 1994; 2002 Berrocal 2003 Widimsky 2000; 2003 DeWood 1990 Grines 1993; 2002 Gibbons 1993 Ribichini 1998 Garcia 1999 GUSTO IIb 1997 Le May 2001 Bonnefoy 2002 Schomig 2000 Vermeer 1999 Kastrati 2002 Aversano 2002 Andersen 2003

Reference/funding	Methods	Outcomes

RCTs 1/22 used pre-hospital fibrinolysis 8/22 trials used streptokinase 14/22 used tPA 13/22 trials used stents 8/22 used GPIs

Table 15: Boersma 2006¹⁰

Reference/funding	Methods	Outcomes	RCTs
Boersma E, et al. Primary Coronary Angioplasty vs.	Based on patient-level data – All RCTs (n > 50) published between January 1990 and December 2002 were considered (non-English articles were not excluded).	30-day all-cause mortality, stroke and re-MI Outcomes are presented as	22 RCTs (n = 6763)
Angioplasty vs. Thrombolysis Group. Does time matter? A pooled analysis of randomized clinical crials comparing primary percutaneous coronary intervention and in-hospital fibrinolysis in acute myocardial infarction patients. Eur Heart J. 2006; 27(7):779-88. Funding: Boehringer ngelheim (the Netherlands)	They were identified by OVID MEDLINE and ISI Web of Science using a broad range of key words; References of identified papers and abstract listings of annual meetings of the American Heart Association, American College of Cardiology and European Society of Cardiology were also examined. Individual patient data from 22 RCTs were pooled, and multi-level logistic regression assessed the relationship among treatment, treatment delay, and 30-day mortality. Treatment delay was divided into 'presentation delay' (symptom onset to randomisation; categorised as 0–1, >1–2, >2–3, >3–6, >6 hours) and 'PPCI-related time delay' (median time from randomisation to PPCI minus median time to FT per hospital). PPCI-related time delay was calculated for each of the 153 hospitals and assigned to each patient within that hospital. PPCI-related time delay was then grouped into: 0–35, >35–50, >50–62, >62–79, and >79 minutes. At the patient level, age, gender, weight, diabetes mellitus, previous MI, prior revascularisation (PCI or CABG), anterior MI at presentation,	Outcomes are presented as OR and 95% CI Further analyses of how presentation delay affected PPCI benefit were performed in patient-, hospital- and study-level subgroups selected in advance (< 65 versus ≥ 65-years-old; gender; ± diabetes mellitus; ± previous MI; anterior versus nonanterior MI; systolic blood pressure (< 130 versus ≥ 130 mmHg); heart rate (< 70 versus ≥ 70 bpm), hospitallevel average annual PCI volume and study-level type of fibrinolytic agent used.	Zijlstra 1993; 1997 Ribeiro 1993 Grinfeld 1996 Akhras 1997 Kedev 1997 de Boer 2002 Widimsky 2000; 2003 Grines 1993; 2002 Gibbons 1993 Ribichini 1998 Garcia 1999 GUSTO IIb 1997 Le May 2001 Schomig 2000 Vermeer 1999 Kastrati 2002
	heart rate, systolic blood pressure, presentation delay, and study treatment (FT or PPCI) were considered fixed effects.		Aversano 2002 Andersen 2003
	At hospital level, the PPCI-related time delay and the average annual PCI volume (grouped into tertiles of its distribution: <10, 10−23, ≥24 PPCI/year) were considered.		Aoki 1997 0/22 used pre-hospital fibrinolysis

Reference/funding	Methods	Outcomes
	At the study level, the likelihood of PCI within 30 days after initial FT, use of stents, use of GPIs, type of fibrinolytic agent used (streptokinase, t-PA, or accelerated t-PA), single-centred versus multicentred trial, and the year of publication were considered. Two sensitivity analyses were performed: (i) impact of exclusion of 3 trials without patient data; (ii) fibrinolytic agent used.	

9/22 used streptokinase
1/22 used duteplase
12/22 used t-PA
10/22 used accelerated t-PA
10/22 used stents
6/22 used GPIs

Table 16: Kent 2001⁶³

Reference/funding	Methods	Outcomes	RCTs
Kent DM, et al. Balancing the benefits of primary angioplasty against the benefits of thrombolytic therapy for acute myocardial infarction: the importance of timing. Eff Clin Pract. 2001; 4(5):214-20. Funding: New England Medical Centre Research Fund Award	Based on study-level data from 10 RCTs included in a previously published meta-analysis (Weaver 1997). For each RCT this analysis calculated PPCI-related time delay (median 'door-to-balloon' time minus median 'door-to-needle' time) and survival benefit (30-day mortality after fibrinolytic therapy minus 30-mortality after PPCI) Linear meta-regression was used to assess the relationship between PPCI-related time delay and treatment benefit by estimating: 1) the decrease in benefit for each additional minute of delay in receipt of PCI and 2) the delay expected to lead to equipoise between PCI and fibrinolytic therapy. The magnitude and statistical significance of the relationship were estimated by weighting each RCT's results by the square root of the number of patients in that trial. A sensitivity analysis was performed by excluding the trials with the longest and the shortest PPCI-related time delay.	PPCI-related time delay associated with equipoise between both reperfusion strategies (Outcomes are presented as relative and absolute reduction in 30-day mortality, both as a function of the PPCI-related time delay) Reduction in mortality benefit for each additional 10 minutes of PPCI-related time delay.	2ijlstra 1993; 1997 Ribeiro 1993 DeWood 1990 Grines 1993 Gibbons 1993 Ribichini 1996 Garcia 1997 GUSTO IIb 1997 Grinfeld 1996

Table 17: Pinto 2006⁹⁰

Table 17. Fillto 2000			
Reference/funding	Methods	Outcomes	Registry
Pinto DS, et al. Hospital delays in reperfusion for ST-elevation myocardial infarction: implications when selecting a reperfusion strategy. Circulation. 2006; 114(19):2019-25. Funding: Genentech, Inc	Based on patient-level data from the National Registry of Myocardial Infarction (NRMI) 2, 3, and 4 NRMI 2–4 were voluntary, prospective registries that collected data from June 1994 to August 2003 on consecutive patients with documented STEMI (or LBBB on initial ECG) and < 12 hours after pain onset who received either fibrinolytic therapy or PPCI as initial reperfusion therapy. Patients transferred to an NRMI hospital for reperfusion were included, but patients transferred out of an NRMI hospital to a non-NRMI hospital were excluded because mortality data were unavailable. Patients with missing time-interval data were also excluded. Participating hospitals were required to manage ≥ 20 patients with STEMI (≥ 10 PPCI and ≥ 10 fibrinolysis). PPCI-related time delay was calculated by subtracting the median DN time from the median DB time at each hospital. For transfer patients, the point of reference to calculate DB and DN times was the first hospital arrival date/time. Hospitals were divided into 4 categories of PPCI-related time delays (< 60, 60–89, 90–120, and > 120 minutes). Then, the mean time delay within each of these 4 categories was calculated with the median PPCI-related time delay at each hospital. Hierarchical models that adjusted simultaneously for both patient-level risk factors and hospital-level covariates were used to evaluate the relationship between PPCI-related time delay, patient risk factors, and in-hospital mortality. Patient covariates included: treatment type (PPCI versus fibrinolysis), age, gender, race, diabetes mellitus, hypertension, angina, Killip class 2/3, Killip class 4, previous infarction, current smoking, stroke, pulse, systolic blood pressure, payer, pre-hospital delay, and discharge year. Hospital covariates included STEMI volume, PPCI volume, transfer-in rate, rural location, and status as a teaching hospital.	In-hospital mortality (before discharge) PPCI-related time delay associated with equipoise between both reperfusion strategies Relationship between DB-DN time delay and the mortality difference in patient subgroups stratified by age (< 65 versus ≥ 65 years), infarct location (anterior versus other), and time from symptom onset to hospital presentation (≤ 120 or > 120 minutes) Outcomes are presented as OR and 95% CI.	The selection criteria yielded 192,509 patients with STEMI and 645 hospitals eligible for analysis. PPCI was performed in 65,600 patients, and FT was administered to 126,909 patients (Fibrinspecific agents were administered in 92% (n = 117,256) of patients). > 65% of patients (n = 125,737) presented within 2 hours of symptom onset.

Table 18: Tarantini 2010¹¹⁰

Methods	Outcomes	RCTs
Based on study-level data – All RCTs ($n \ge 50$), published and unpublished (non- English articles were not excluded) comparing fibrin-specific fibrinolysis to PPCI.	30-day mortality risk leading to equipoise between	16 RCTs (n = 6281)
were searched (Jan. 1990–Dec. 2008) using a broad range of keywords. Also searched for abstracts in the New England Journal of Medicine, Circulation, European Heart Journal, Journal of the American College of Cardiology, and Heart. References of identified papers, relevant studies, and meta-analyses were additionally scanned. Oral presentations and expert slide presentations identified from www.theheart.org, www.tctmd.com, www.crtonline.com, www.clinicaltrialresults.org, www.esccardio.org, www.europcr.com, and www.acc.org were also examined. Weighted meta-regression was used to explore the relationship, (adjusted for prehospital time delay) between the mortality risk and the PPCI-related time delay	990—Dec. 2008) using a broad range of keywords. Also is in the New England Journal of Medicine, Circulation, hal, Journal of the American College of Cardiology, and Heart. ed papers, relevant studies, and meta-analyses were Oral presentations and expert slide presentations identified org, www.tctmd.com, www.crtonline.com, lts.org, www.esccardio.org, www.europcr.com, and so examined. ession was used to explore the relationship, (adjusted for presetween the mortality risk and the PPCI-related time delay lent 30-day mortality between PPCI and fibrin-specific ession analysis considered the absolute risk reduction as a inolytic mortality, time to treatment and PPCI-related time ession analysis as 30-day mortality minus PPCI 30-day arguments as a calculated as the difference between mean or median esse calculated as mean or median time from the symptom onset on or first medical contact) that was not influenced by the essay for baseline mortality risk. This rate can be considered as a enlying baseline risk and has the advantage of being a measure factors contributing to outcome occurrence.	DeWood 1992 Grines 1993; 2002; 2005 Gibbons 1993 Ribichini 1998 Garcia 1999 GUSTO IIb 1997 Le May 2001 Bonnefoy 2002 Schomig 2000 Vermeer 1999
fibrinolysis. A multiple linear regression analysis considered the absolute risk reduction as a linear function of fibrinolytic mortality, time to treatment and PPCI-related time delay. Mortality benefit was calculated as fibrinolysis 30-day mortality minus PPCI 30-day mortality.		Kastrati 2002 Aversano 2002 Andersen 2003 Armstrong 2006
DB time minus DN time. Presentation delay was calculated as mean or median time from the symptom onset to door (randomisation or first medical contact) that was not influenced by the allocated treatment. Because the outcome was mortality, the control (lytic arm) mortality rate was interpreted as a proxy for baseline mortality risk. This rate can be considered as a surrogate of the underlying baseline risk and has the advantage of being a measure that reflects multiple factors contributing to outcome occurrence.		fibrinolysis 0/16 stents 10/16 trials used stents 7/16 used GPIs
	Based on study-level data — All RCTs (n ≥ 50), published and unpublished (non-English articles were not excluded) comparing fibrin-specific fibrinolysis to PPCI. MEDLINE, CENTRAL, EMBASE, and the Cochrane Central Register of Controlled Trials were searched (Jan. 1990—Dec. 2008) using a broad range of keywords. Also searched for abstracts in the New England Journal of Medicine, Circulation, European Heart Journal, Journal of the American College of Cardiology, and Heart. References of identified papers, relevant studies, and meta-analyses were additionally scanned. Oral presentations and expert slide presentations identified from www.theheart.org, www.tctmd.com, www.crtonline.com, www.clinicaltrialresults.org, www.esccardio.org, www.europcr.com, and www.acc.org were also examined. Weighted meta-regression was used to explore the relationship, (adjusted for prehospital time delay) between the mortality risk and the PPCI-related time delay which leads to equivalent 30-day mortality between PPCI and fibrin-specific fibrinolysis. A multiple linear regression analysis considered the absolute risk reduction as a linear function of fibrinolytic mortality, time to treatment and PPCI-related time delay. Mortality benefit was calculated as fibrinolysis 30-day mortality minus PPCI 30-day mortality. PPCI-related time delay was calculated as the difference between mean or median DB time minus DN time. Presentation delay was calculated as mean or median time from the symptom onset to door (randomisation or first medical contact) that was not influenced by the allocated treatment. Because the outcome was mortality, the control (lytic arm) mortality rate was interpreted as a proxy for baseline mortality risk. This rate can be considered as a surrogate of the underlying baseline risk and has the advantage of being a measure	Based on study-level data – All RCTs (n ≥ 50), published and unpublished (non-English articles were not excluded) comparing fibrin-specific fibrinolysis to PPCI. MEDLINE, CENTRAL, EMBASE, and the Cochrane Central Register of Controlled Trials were searched (Jan. 1990–Dec. 2008) using a broad range of keywords. Also searched for abstracts in the New England Journal of Medicine, Circulation, European Heart Journal, Journal of the American College of Cardiology, and Heart. References of identified papers, relevant studies, and meta-analyses were additionally scanned. Oral presentations and expert slide presentations identified from www.theheart.org, www.tetmd.com, www.crtonline.com, www.clinicaltrialresults.org, www.esccardio.org, www.europcr.com, and www.acc.org were also examined. Weighted meta-regression was used to explore the relationship, (adjusted for prehospital time delay) between the mortality risk and the PPCI-related time delay which leads to equivalent 30-day mortality between PPCI and fibrin-specific fibrinolysis. A multiple linear regression analysis considered the absolute risk reduction as a linear function of fibrinolytic mortality, time to treatment and PPCI-related time delay. Mortality benefit was calculated as fibrinolysis 30-day mortality minus PPCI 30-day mortality. PPCI-related time delay was calculated as the difference between mean or median DB time minus DN time. Presentation delay was calculated as mean or median time from the symptom onset to door (randomisation or first medical contact) that was not influenced by the allocated treatment. Because the outcome was mortality, the control (lytic arm) mortality rate was interpreted as a proxy for baseline mortality risk. This rate can be considered as a surrogate of the underlying baseline risk and has the advantage of being a measure that reflects multiple factors contributing to outcome occurrence.

Table 19: Zjilstra 2002¹²²

Reference/funding	Methods	Outcomes	RCTs		
Zijlstra F, et al. Clinical characteristics and outcome of	Based on patient-level data from 9/10 RCTs included in a previously published meta-analysis (Weaver 1997) plus Akhras 1997 (identified subsequent to Weaver	Mortality, non-fatal reinfarction and total stroke at 30 days	10 RCTs (n = 2635; n = 1302 PPCI; n = 1333 fibrinolysis)		
patients with early (<2	meta-analysis)	Mortality and non-fatal	Zijlstra 1993; 1997		
h), intermediate (2-4 h) and late (>4 h) presentation treated by primary coronary angioplasty or thrombolytic therapy for acute myocardial infarction. Eur Heart J. 2002; 23(7):550-7.	The analysis examined the effects of presentation delay on outcomes after PPCI and fibrinolytic therapy using individual patient data on an ITT basis according to randomised groups. Time to presentation was measured from the outset of symptoms to randomisation in 6 RCTs and from	reinfarction at 6 months Both as a function of time to presentation	Ribeiro 1993		
			Grinfeld 1996		
			Grines 1993		
			Gibbons 1993		
			Ribichini 1998 Garcia 1999		
	symptom onset to hospital admission in 3 RCTs (unavailable in 1 RCT).		GUSTO IIb 1997 Akhras 1997		
	Patients were classified into 3 categories according to				
	the time delay from symptom onset to presentation				
Funding: None stated	(< 2 hours, 2–4 hours, ≥ 4 hours)		0/10 used pre-hospital fibrinolysis		
	Subgroups were investigated to allow multivariate analysis (lytic regimen; age [<50, 50–60, 60–70 and >70		5/10 trials used streptokinase		
	years); gender, diabetes, infarct location, prior MI, heart		5/10 used tPA		
	rate on admission [<65, 65–75, 75–85, and >85 beats		0/10 trials used stents		
	per minute] and SBP on admission [<115, 115–130, 130–150 and >150 mmHg].		0/10 used GPIs		

G.2 Facilitated PPCI

FINESSE^{41,42}

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
SG. Ellis, M Tendera, MA. de Belder, AJ. van Boven, P Widimsky et al., and FINESSE Investigators FINESSE. Facilitated PCI in patients with ST-	Design: RCT (20 sites – international: Europe, Argentina, Canada and USA) Enrolment: Aug 2002 – Dec 2006 Setting NO DETAILS GIVEN	n = 2452 Drop-outs (during the 90 days – not available for follow-up at 90 days): fPPCI: n = 8 (1%)	Inclusion criteria Adults ≥21 years Presented within 6 hours after the onset of signs and symptoms of cardiac ischaemia. Had ST-segment elevation suggestive of an acute MI eligible for fibrinolytic therapy or PPCI estimated time to diagnostic catheterisation was 1-4 hours after randomisation	fPPCI (n = 818) (Abciximab EARLY) Administere d in the emergency department . Abciximab: IV bolus 0.25	PPCI (Abciximab LATER) (n = 806) PPCI with abciximab initiated in the cardiac cath lab, immediately before PCI	90 days, 1 year For death and complications of MI: 90 days (maximum 100 days) after randomisation; 1 year (long-term follow-	1° Combined death (all causes), AND complications of MI [ventricular fibrillation (> 48 hours after randomisation), cardiogenic shock, HF (requiring hospitalisation or an emergency room visit through 90 days)].	Centocor and Eli Lilly
elevation myocardial infarction. N.Engl.J.Me d. 358	Randomisation Good: central randomisation (1:1:1 ratio)	Combinati on fPPCI: n = 15 (1.8%)	Exclusion criteria Low risk (<60 years and had localised inferior infarction) had risk factors for bleeding	mg / kg Combination fPPCI (n = 828)	Abciximab (an IV bolus of 0.25 mg /kg).	up)		
(21):2205- 2217, 2008. 1 year follow-up	Allocation concealment Unclear: central randomisation	PPCI: n = 13 (1.6%)	planned use of a direct thrombin inhibitor during PCI MI precipitated by a condition other than atherosclerotic coronary artery disease, recent (within 14 days)	Administere d in the emergency department	'Dummy' placebo medication s were administere d at all time		complications of MI through 90 days (as in the primary end pint)	
Ad J. van dummy (place	_	Drop-outs (at 1 year follow-up): fPPCI: 1.6%	use of fibrinolytic, administration of low-molecular–weight heparin (LMWH) within 24 hours PCI within 7 days	Reteplase + abciximab Reduced-	points to ensure that the study remained	Death (all causes through 90 days)		

Boven, P Widimsky, H R. Andersen, A Betriu, et al. and Investigators FINESSE. 1- year survival in a randomized trial of facilitated reperfusion: results from the FINESSE (Facilitated Intervention with Enhanced Reperfusion Sneed to Ston Events) trial. JACC Cardiovasc Interv 2 (10):909- 916, 2009.	to maintain the abciximab and reteplase bolus blind. Central adjudication (for cardiogenic shock, congestive HF and stroke – in all patients; and ST segment resolution – in of 50% of patients) by people blinded to treatment assignment Sample size calculation: NOT POWERED: aim was 3000 patients but recruitment terminated early due to slow recruitment and substantial cost overruns. Analysis	Combinati on fPPCI: 1.4% PPCI: 2.4%	known or suspected bleeding confirmed uncontrolled hypertension, and other contraindications to fibrinolytic treatment. If the protocol-specified ceiling dose of unfractionated heparin (40 U/kg, 3000U maximum) or an activated partial thromboplastin time (70 seconds) is exceeded before enrolment Demographics and baseline characteristics see below	reteplase: 2 5-U boluses separated by 30 minutes, for those <75 years old, or 1 5-U dose, for those ≥75 years old) Abciximab (IV bolus 0.25 mg / kg) Notes: IN PATIENTS AGED ≥75 YEARS: a novel regimen of reteplase 5U single bolus in combination with abciximab was studied.	blinded	ST-segment resolution (>70% from baseline as assessed at 60— 90 minutes after randomisation) Safety: Non- intracranial major or minor bleeding (TIMI) Intracranial haemorrhage (through discharge or day 7 whichever was sooner) Cardiogenic shock Congestive heart failure Stroke ST-segment
nanor -				was studied.		
ELLIC 2004	•			NOTE:		ST-segment
how shall I	aimed for was to			Before cathete	orisation	resolution
roforonco	detect a difference					
+hic3)	between combination fPPCI			aspirin: 81 - 32 250 to 500 mg	· ·	(see below for
SG. Ellis, P	and PPCI in			heparin:		definitions)
	primary end point			to 40 U /kg (m	aximum dose,	definitions)
Armstrong	(030/			, 51	,	

A Betriu, B Brodie, H Herrmann, et al., and FINESSE Investigators . Facilitated percutaneou s coronary intervention versus primary percutaneou s coronary intervention : design and rationale of the Facilitated Intervention n with Enhanced Reperfusion Speed to Stop Events (FINESSE)	ITT analysis: ITT for efficacy outcomes; As- treated analysis (treatment received) for safety outcomes		3000 U), with a target activated clotting time of 200 - 250 seconds At sites that participated in the low-molecular weight heparin sub-study, 0.5 mg /kg enoxaparin IV was administered, and 0.3 mg /kg was administered subcutaneous, with no target for the activated clotting time. Post-PCI — 0.125 μg/kg/minute abciximab (maximum dose, 10 μg / min) for 12 hours Stent type used — choice of stent at discretion of investigator PCI — 92% overall (91 and	
trial. Am.Heart J. 147 (4):E16,			92% in each group)	
2004.			Stents used – not mentioned	
		fPPCI (n = 818)	Combination fPPCI (n = 828)	PPCI (n = 806)
Mean age, yea	ars (SD)	61.9 (11.8)	62.6 (11.4)	62.5 (11.4)
Age <75 years	(%)	695 (85)	691 (83.5)	678 (84.1)
Age ≥75 years	(%)	123 (15)	137 (16.5)	128 (15.9)

Men (%)	602 (73.6)	609 (73.6)	599 (74.3)
Hypertension (%)	405 (49.5)	894 (47.6)	374 (46.4)
Hypercholesterolemia (%)	119 (14.5)	128 (15.5)	276 (34.2)
Diabetes mellitus (%)	249 (30.4)	291 (35.1)	133 (16.5)
Family history of CAD diagnosed at <55 years if age (%)	149 (18.5)	187 (22.9)	190(22.9)
Current smoker (%)	363 (44.4)	347 (41.9)	357 (44.3)
Previous MI (%)	82 (10.2)	80 (9.8)	104 (12.6)
Previous congestive heart failure (%)	13 (1.6)	9 (1.1)	12 (1.4)
Previous treatment (%)	NOT GIVEN	NOT GIVEN	NOT GIVEN
Anterior infarction (%)	370 (45.9)	403 (49.3)	40 (48.3)
*Activated clotting time			

Definitions of end points

Complications of MI: Complications are defined as resuscitated ventricular fibrillation occurring 48 hours after randomisation, rehospitalisation or emergency department visit for congestive heart failure, or cardiogenic shock.

Other end points not described further.

Outcomes

Outcomes at 90 days (unless specified)	fPPCI (n = 818)	Combination fPPCI (n = 828)	PPCI (n = 806)	Statistical significance / p value	p value (Kaplan-Meier)
Primary composite end point at 90 days, %	9.8	10.5	10.7	Combination versus fPPCI: HR 0.91 (95% CI 0.67 to 1.23)	Combination versus PPCI: p = 0.55 Facilitated versus PPCI: p = 0.86 Combination versus facilitated: p = 0.68
Complications of MI, %	7.4	7.5	9.0	NS differences (data not given)	
Death from all causes, %	5.2	5.5	4.5	NS differences (data not given)	
Ventricular fibrillation occurring > 48 hours after randomisation, %	0.6	0.2	0.4	NS differences (data not given)	
Cardiogenic shock, %	5.3	4.8	6.8	NS differences (data not given)	
Rehospitalisation or emergency room visit for	1.9	2.9	2.2	NS differences (data not given)	

congestive HF, %					
Heart failure - Index hospitalisation, n (%)	45 (5.5)	54 (6.5)	52 (6.5)	not given	not given
Recurrent MI	16 (2.0)	17 (2.1)	15 (1.9)	not given	not given
Any subsequent	111 (13.6)	111 (13.4)	111 (13.8)	not given	not given
revascularisation, n (%)	85 (10.4)	81 (9.8)	78 (9.7)	not given	not given
PCI CABG	26 (3.2)	31 (3.7)	37 (4.6)	not given	not given
SAFETY END POINTS (through discharge or day 7)	n = 814	n = 805	n = 795		
Non-intracranial TIMI	118 (14.5)	81 (10.1)	55 (6.9)	Combination versus PPCI	
bleeding, n (%)	39 (4.8)	33 (4.1)	21 (2.6)	(p < 0.001); fPPCI versus PPCI	
Major	79 (9.7)	48 (6.0)	34 (4.3)	(p < 0.05)	
Minor				Combination versus PPCI: p < 0.05	
				Combination versus fPPCI:	
				p < 0.05	
Stroke, n (%)	9 (1.1)	4 (0.5)	8 (1.0)	data not given	
Intracranial haemorrhage	5 (0.6)	0 (0)	1 (0.1)		
Ischaemic	4 (0.5)	4 (0.5)	7 (0.9)		
Fatal stroke, n (%)	3 (3.7)	0 (0)	0 (0)	data not given	
ST-segment resolution (>70%) in 60–90 minutes, %	43.9	33.1	31.0	p = 0.003 (combination versus PPCI)	
				p = 0.01 (combination versus fPPCI)	
TIMI flow grade 3 before PCI performed, %	32.8	14.1	12.0	p < 0.001(combination versus PPCI)	
				p < 0.001 (combination versus fPPCI)	
TIMI flow grade after PCI	not given	not given	not given	No substantial difference between groups (data not given)	
TIMI flow grade for ST- segment resolution at 180–	not given	not given	not given	No substantial difference between groups (data not given)	

240 minutes							
1 year follow-up							
Death from all causes at 1 year follow-up, %	7.4	6.3	7.0	NS differences (data not given)	Combination versus PPCI: $p = 0.603$ fPPCI versus PPCI: $p = 0.765$ Combination versus fPPCI: $p = 0.415$		
Median door to balloon time	(all patients): 2.2	hours (IQR 1.8 to	2.8)				
92% of patients underwent P	CI						
SUBGROUPS							
AGE 75: No intracranial haem	norrhages occurre	d in patients age	d ≥75 years				
Cardiac procedures							
n, %	fPPCI		Combination fPPCI	PPCI	n, %		
	(n = 818)		(n = 828)	(n = 806)			
Catheterisation	96%		93%	95%	Catheterisation		
PCI	92% overall each group)	(91 and 92% in		-	PCI		
Transfer and timing, and hos	Transfer and timing, and hospital stay						
No details given							

Table 20: ASSIST TRIAL⁶⁹

Reference	Study type	Patient characteristics	Intervention	Comparison	Outcome measures	Source of funding
LeMay M. et al. Primary	Design: RCT (prospective) Enrolment:	Number of patients: n = 400 Drop-outs: 1 patient lost to follow-up.	PPCI with heparin + eptifibatide (n = 201)	PPCI with heparin (n = 199) Note:	1° Death from any cause, recurrent MI, recurrent	Schering- Plough Canada Inc provided an
percutaneou s coronary angioplasty with and without eptifibatide in ST-	August 2005- March 2008 August 2005- March 2008 Setting ithout Hospital (3 Ottawa hospitals).	<12 hours of the onset of ischaemic chest discomfort ≥30 minutes ≥1mm (0.1mV) ST-elevation in 2 or more contiguous leads Exclusion criteria Active bleeding Stroke within 90 days Intracranial bleeding at any time Major surgery or trauma within 6 weeks Systolic blood pressure >200mmHg or of body weight bolus, followed (after 10 minute continuous infused of 2.0 microgram/kg/ ute with a second microgram/kg/ ute with a second microgram/kg. Eptifibatide initiated before catheterisation continued for 1 hours after PCI.	bolus, followed by (after 10 minutes) a continuous infusion of 2.0 microgram/kg/min	Before catheterisation: oral aspirin (160mg), oral clopidogrel (600mg) and IV unfractionated heparin (60U/kg up to max 4000U). 9 (4.5%) patient were given abciximab and 6 (3.0%) were given eptifibatide during PCI.	severe ischaemia at 30 days. Other: Stroke, congestive heart failure cardiogenic shock Length of follow-up: 30 days, 6 months	unrestricted grant and free distribution of eptifibatide, and Medtronic Canada Ltd provided free stents for the study.
segment elevation myocardial infarction. A safety end efficacy study of integrilin-	Randomisation: Yes. Randomly assigned in a 1:1 ratio to intervention or comparison group, in blocks of 10 at each site. Allocation concealment: Not mentioned / unclear.		bolus of 180 microgram/kg.			
facilitatyed versus primary percutaneou s coronary intervention in ST_segment elevation myocardial infarction (ASSIST)	Blinding: No. The intervention group received open-label drug. Corrected TIMI frame counts and Myocardial perfusion grades were evaluated by independent blinded investigators. Sample size calculation:	diastolic blood pressure >110mmHg Prolonged cardiopulmonary resuscitation PCI within 30 days Fibrinolytic agents within 7 days Any glycoprotein Ilb/Illa within 7 days Low-molecular weight heparin within 12 hours Coagulation disorder Current warfarin treatment Intolerance to aspirin or clopidogrel Other illness likely to result in death	Note: Before catheterisation : oral aspirin (160mg), oral clopidogrel (600mg) and IV unfractionated heparin (60U/kg up to max 4000U).	PCI performed in 184 (92.5%) patients. Stent insertion in 183 (92.0%) patients.		

difference between 15% and
5.0%, with a level of
significance of 0.05 and 90%
power using a χ2 test, was
determined to be 187 per
group. With an anticipated
loss to follow-up rate of 5%,
the minimum number of
patients required was 200
per group.

within 12 months
Pregnancy
Creatine > 200 micromol/L
Cardiogenic shock
Severe contrast allergy

PCI performed in 190 (94.5%) patients.

Stent insertion in 187 (93.0%) patients.

Demographics and baseline characteristics

see below

ITT analysis: Yes

Demographics and baseline characteristics

	Heparin + eptifibatide (n = 201)	Heparin (n = 199)
Age, years	60.4±12.1	60.6±11.8
Male gender	162 (80.6)	143 (71.9)
Hypertension	92 (45.8)	99 (49.8)
Diabetes mellitus	29 (14.4)	36 (18.1)
Current smoking	90 (44.8)	76 (38.2)
Hyperlipidimia	67 (33.3)	77 (38.7)
Prior MI	21 (10.5)	27 (13.6)
Prior angioplasty	16 (8.0)	16 (8.0)
Prior bypass surgery	10 (5.0)	8 (4.0)
Anterior MI	73 (36.3)	75 (37.7)
Heart rate, bpm	74±16	76±18
Systolic blood pressure, mmHg	130±23	134±24
Diastolic blood pressure	77±14	80±15
Killip class		
1	182 (90.6)	172 (86.4)
II	19 (9.5)	27 (13.6)
III or IV	0 (0.0)	0 (0.0)

BMI, KG/m ²	28.4±4.9	27.5±4.6
Eptifibatide started before initial catheterisation	189 (94.0)	0
Baseline creatine clearance, mL/min	93.3±35.2	87.3±31.1
Peak CK (creatine kinase), U/L	2009±1879	2047±1628

Data are presented as mean±SD or n (%)

Definitions of end points

Reinfarction:

Presence of recurrent ischaemic symptoms at rest, lasting at least 30 minutes and accompanied by any of the following:

New or recurrent ST-segment elevation of ≥1mm (0.1mV) in any contiguous leads

New left bundle branch block

Reelevation in serum creatine kinase level more than twice the upper limit of normal and at least more than 50% of the lowest level measured post-infarction.

Recurrent severe ischaemia:

Recurrent symptoms of ischaemia at rest associated with any of the following:

new ST-segment deviation (elevation at least 0.1mV or depression >0.05mV measured 80 ms after the J-point in at least 2 contiguous leads)

an episode of acute pulmonary edema, sustained ventricular arrhythmia, or haemodynamic instability

the need for urgent revascularisation within 30 days

recurrent myocardial infarction.

Effect Size / Outcomes

	Heparin + eptifibatide (n = 201)	Heparin (n = 199)	p value	Relative Risk (95% CI)
30 days				
Death	7 (3.5)	4 (2.0)	0.54	1.76 (0.51–6.11)
Reinfarction	3 (1.5)	1 (0.5)	0.62	3.00 (0.31–29.09)
Congestive heart failure	15 (7.5)	22 (11.1)	0.22	0.65 (0.33–1.29)
Stroke	0 (0.0)	1 (0.5)	0.50	
6 months				
Death	9 (4.6)	6 (3.1)	0.44	1.52 (0.53–4.34)
Reinfarction	4 (2.0)	2 (1.0)	0.69	2.01 (0.36–11.11)

0 1	45 (3.3)	24 (42 2)	0.40	0.50/0.00 4.45)
Congestive heart failure	15 (7.7)	24 (12.3)	0.12	0.59 (0.30–1.16)
Stroke	0 (0.0)	4 (2.0)	0.06	
Repeat target vessel revascularisation †	8 (4.0)	6 (3.0)	0.60	-
CABG	10 (5.0)	6 (3.0)	0.32	1.68 (0.60–4.73)
Repeat PCI of infarct-related artery	5 (2.5)	2 (1.0)	0.45	2.51 (0.48–13.11)
Repeat target vessel revascularisation †	8 (4.0)	4 (2.0)	0.38	1.76 (0.51–6.11)
Major bleeding	19 (9.5)	11 (5.5)	0.14	1.78 (0.83–3.85)
Major bleeding, non-CABG related	11 (5.5)	5 (2.5)	0.20	2.25 (0.77–6.59)
Minor bleeding	26 (12.9)	18 (9.1)	0.21	1.49 (0.79–2.82)
Minor bleeding, non-CABG related	25 (12.4)	18 (12.4)	0.27	1.43 (0.75–2.71)

Table 21: van't Hof et al. 2004¹¹⁶

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Outcome measures	Source of funding
AWJ. van't Hof, N Ernst, M Jan de Boer, R de	Design: RCT (multicentre - Netherlands)	n = 507 Drop-outs: 0 days: 3%	Inclusion criteria Presence of chest pain (>30 minutes) together with >0.2 mV (anterior	fPPCI (early tirofiban treatment – pre-hospital) (n = 251)	Placebo (later tirofiban treatment - in cath lab)	1° increase in incidence of TIMI grade 3	Educational I grant from Merck and
Winter, E Boersma, T Bunt, S Petronio, A.	Enrolment: June 2001-November 2002	(n = 15); 1 year: 4% (n = 18).	MI) or 0.1 mV (non- anterior MI) of ST- elevation in 2 contiguous ECG leads	IV bolus Tirofiban (10 microgram/kg) followed by maintenance infusion (0.15 mg/kg/min)	(n = 256) IV bolus and	flow at initial angiography prior to PCI	Co.
T. Marcel Gosselink, W Jap, F Hollak, JCA.	Setting Recruitment and randomisation in the	n = 15 patients withdrew	Ability to perform primary angioplasty within 6 hours after the onset of symptoms	Note:	infusion of placebo; tirofiban given	TIMI flow components	
Hoorntje, H Suryapranat a, JH Dambrink, F	ambulance or referring hospital; transported to tertiary hospital for	consent most often on the 2nd day of admission.	Exclusion criteria >80 years of age	Before transportation all patients received IV bolus 5000 IU unfractionated heparin + 250 mg aspirin	later (see 'note' below) until 24 hours post-PCI	Presence of thrombus fresh	
Zijlstra, and On-TIME study group.	centres were experienced cardiology centres).	ntres were Women perienced cardiology Treatm	Women <50 years of age IV. Treatment with fibrinolytic therapy in Emergency transportation			occlusion at initial angiography	
Facilitation of primary coronary angioplasty by early start of a	Randomisation: Blocks per institution. Randomised by selecting sealed study drug kits in		previous 24 hours On warfarin or acenocoumarol within last 7 days Contraindication to GPIs Severe heart failure or	performed after arrival of pt at the cath lab. After angiography, and before angioplasty, all patients received a second bolus of study drug IV (tirofiban in		pre-PCI myocardial blush grade	
glycoprotein 2b/3a inhibitor: results of	Allocation concealment: Not mentioned / unclear.		cardiogenic shock (Killip class III or IV) On haemodialysis	patients initially given placebo, and placebo in patients initially given tirofiban). After this		Other: Death Re-MI	
the ongoing tirofiban in myocardial infarction evaluation	Blinding: Double blind (mentioned, but details		Demographics and baseline characteristics see below	second bolus of drug, all patients were given openlabel tirofiban (maintenance infusion 0.15 microgram/kg/min)		Stroke Major bleeding (non- CABG related	

(On-TIME) trial. Eur.Heart J. 25 (10):837- 846, 2004.	not given of patient blinding). Outcome assessors blinded to treatment allocation for all angiographic parameters Sample size calculation: Yes: powered study. A sample size of 438 patients (80% power to show a significant difference in the incidence of the primary end point: increase in incidence of TIMI grade 3 flow at initial angiography prior to PCI). Trial recruited approx. 500 patients to allow for drop-outs and false-positive infarct diagnoses in the ambulance. ITT analysis: Yes		for 24 hours. Post-PCI all ts were treated with clopidogrel (300 mg loading dose followed by 75 mg daily for 1 month), aspirin, BB, statin therapy and ACEi. NOTE: Before catheterisation – all patients had heparin (5000U) and aspirin (500 mg) Post-PCI – antithrombin therapy of ticlopidine (250 mg bds, 4 weeks) and aspirin (100 mg bds, throughout study) Stent type used – different types of slotted-tube stents (evenly distributed between the study groups) PCI – 100% Stents used – 100%	major bleeding) Minor bleeding Composite (death/re- MI/stroke/maj or non-CABG related major bleeding) (see below for definitions) Length of follow-up: 30 days, 1 year	
		Early tirofib	oan (n = 251)	Later tirofiban (n = 256)	
Age, years; me		63 (10)		61 (11)	
Male gender,	%	79		80	
Hypertension	.%	10		11	
Diabetes, %		27		30	

Smoking (current or previous), %	62	68
Anterior MI, %	44	47
Previous MI, %	6	10
Previous CABG, %	2	2
Previosu PCI, %	5	6
Multivessel disease, %	58	53
Killip class >1, %*	17	15
TIMI, n(%) out of 243 and 244 total patients for each group		
respectively	104 (43)	82 (34)
2 or 3	107 (44)	143 (59)
0	32 (13)	19 (8)
1	58 (24)	46 (19)
2	46 (19)	36 (15)
3		
Treatment, %		
Angioplasty	88	90
CABG	3.6	2.3
Other	8.4	8.2
*defined as SBP <100 mmHg or HR >100/min		

defined as SBP < 100 mining of HK > 100/

Definitions of end points

Successful angioplasty: < 50% diameter stenosis and TIMI 3 flow of the IRV.

Presentation delay: Time from symptom-onset to infarct diagnosis (first ECG).

Major bleeding: Fall in Hb of ≥2.0 mmol/L and the need for transfusion of 2 or more units of blood, corrective surgery or both. OR bleeding that resulted in documented intra-cranial or retro-peritoneal haemorrhage

Minor bleeding: Fall in Hb ≥2.0 mmol/L without the need for a transfusion.

Recurrent MI: A new increase in creatine kinase (CK)-MB fraction of >3 times the upper limit of normal, whether accompanied by chest pain or ECG changes and present in 2 separated blood samples or not.

	Early tirofiban	Later tirofiban	p value
Presentation delay*	88 (58–137)	104 (55–155)	0.25
In-outdoor time†	26 (19–33)	25 (19–33)	0.49
Transportation delay	30 (17–47)	33 (18–46)	0.70

Door to angio time	25 (15–40)	24 (15–36)	0.29
Angio to balloon time	16 (12–21)	15 (12–20)	0.37
Ichaemic time**	196 (155–252)	199 (159–266)	0.48
Pre-treatment time††	79 (65–92)	15 (12–20)	-

Time delays, minutes; median (25-75% IQR)

Effect Size / Outcomes

After initial angiography, 89% patients underwent PCI and 3% were candidates for bypass surgery due to severe 3-vessel disease. Remainder of patients (8%) treated conservatively.

30 days:

Major bleeding in 44/492 (9%) patients, from which 25 events (57%) were blood transfusions related to the CABG procedure.

No differences in bleeding were observed between the groups.

No intracranial bleeding event occurred during treatments with tirofiban.

1 year:

Mortality no longer a difference between the groups

Combined incidence of death or re-MI was the same (7%) in both groups.

	Early tirofiban (n = 251)	Later tirofiban (n = 256)	p value
PCI success, % patients	90	91	-
Stents, % patients	72	74	0.71
At 30 days, n (%)	n = 245	n = 247	
Death	9 (3.7)	2 (0.8)	0.03
Re-MI	3 (1.2)	2 (0.8)	0.65
Stroke	0	1 (0.4)	1.00
Major bleeding*	11 (4.5)	8 (3.2)	0.47
Composite**	21 (8.6)	11 (4.4)	0.06
1 year, n (%)	n = 245	n = 244	

^{*}Defined as time from symptom onset to infarct diagnosis (1st ECG)

[†]Defined as time from infarct diagnosis to start of transportation

^{**}Defined as time from symptom onset to 1st balloon inflation

^{††}Defined as time between 1st active bolus of study drug and 1st balloon inflation

Death	11 (4.5)	9 (3.7)	0.66
re-MI	6 (2.4)	9 (3.7)	0.43
Death or re-MI	17 (7.0)	17 (7.0)	0.99
Post-PCI	n = 217	n = 228	
TIMI 3, n(%)	196 (90)	208 (91)	0.74

^{*}Non-CABG related major bleeding

Cardiac procedures

			In-hospital	
	TNK only n = 100)	Invasive strategy (n = 104)		
Cardiac catheterisation	Not reported	102 (within 24 h)	Not reported	Not reported
Revascularisation	60	89*	65	90*
Rescue PCI	14, at a median of 197 minute after randomisation (IQR 172–280 min).	38 (29 protocol-mandated; 6 ECG criteria)	Not reported	Not reported

^{*} Of these 9 patients received CABG

48% of the 91 patients undergoing PCI within 24 hours of randomisation in the invasive strategy arm received abciximab; 81% of these patients had TIMI 3 and 12% had TIMI 2 flow at the end of the procedure

Coronary stents were used in over 97% of all patients undergoing PCI

Transfer: There were no significant differences (in the primary end point) across the treatment groups in either setting (pre-hospital or in-hospital)

SUBGROUPS – TIMI3 flow (NS difference between the treatment groups)

AGE < 62: 22% (early) versus 11% (later) - NS AGE \leq 62: 16% (early) versus 12% (later) - NS MALE: 20% (early) versus 15% (later) - NS

FEMALE: 16% (early) versus 12% (LATER) - NS DIABETES (YES): 24% (early) versus 8% (later) - NS

DIABETES (NO): 18% (early) versus 16% (later) - NS

^{**}combined incidence of death/re-MI/stroke or major non-CABG-related major bleeding

Table 22: van't Hof et al. 2008 111,115,117

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Outcome measures	Source of funding
A W. J. van't Hof, J ten Berg, T Heestermans, T Dill, R C. F, W van Werkum, J H Dambrink, H S, G van Houwelingen, J Paul O, P Stella, E Giannitsis, C Hamm, and Ongoing Tirofiban In Myocardial infarction Evaluation (On-TIME). Prehospital initiation of tirofiban in patients with ST-elevation myocardial infarction undergoing primary angioplasty (On-TIME 2): a multicentre, doubleblind, randomised controlled trial. Lancet 372 (9638):537-546, 2008. A. W. van't Hof, C. Hamm, S. Rasoul, S. Guptha, J. F. Paolini, and J. M. Ten Berg.	Design: RCT (multicentre – 24 centres in Netherlands, Germany and Belgium) Enrolment: June 2006-November 2007 Setting: Recruitment and randomisation in the ambulance or referring hospital; transported to tertiary hospital for angiography and PCI (PCI centres were experienced cardiology centres). Randomisation: Random permuted blocks at each site;	n = 984 Drop-outs (acceptable): Clinical follow- up at 30 days: fPPCI – 4% and PPCI 3%; 1 year: 4.8% overall.	Inclusion criteria Age 21–85 years Symptoms of AMI (>30 minutes but <24 hours) and ST- elevation >1 mV in 2 adjacent ECG leads Exclusion criteria known severe renal dysfunction therapy resistant cardiogeniuc shock persistent severe HT contraindication to anticoagulation or increased risk of bleeding left bundle branch block pregnant or breastfeeding women life expectancy <1 year	fPPCI (early tirofiban treatment – pre-hospital) (n = 491) NOTE: given pre-hospital (ambulance or referring centre) Tirofiban – bolus 25 mg/kg; bail-out tirofiban could be given where needed Heparin – 5000U bolus given IV NOTE: Before catheterisation – aspirin IV (500 mg) clopidogrel orally (600 mg loading dose) UFH IV (5000 U)	PPCI (Placebo) (n = 493) IV bolus and infusion of placebo; tirofiban given post-PCI for 18 hours (see 'note' below) Heparin – 5000U bolus given IV	1°: extent of ST-segment deviation at 1 hour after PCI. Other: composite Death; Recurrent MI; Major and minor bleeding; stroke Length of follow-up In-hospital, 30 days, 1 year	Education I grant from Merck and Co.
Ongoing tirofiban in myocardial infarction evaluation (On-TIME) 2 trial: rationale and study design. EuroIntervention 3 (3):371-380, 2007.	stratified by intended place of recruitment Allocation		Demographics and baseline characteristics see below	Post-PCI – tirofiban infusion 0.15 microgram/kg/minute for 18 hours post-PCI			
	concealment: Unclear – assignment			Stent type used –DES			

			/ 250() 15145		
(methods)	given by each investigator		(approx. 25%) and BMS (approx 75%)		
	investigator		(approx 73%)		
and	Diadiaa		DCI 000/ in both manua		
	Blinding:		PCI – 99% in both groups		
J. M. Ten Berg, A. W. J.	Double blind				
Van 't Hof, T. Dill, T.			Stents used – 90% in		
Heestermans, J. W.	Sample size		both groups		
Van Werkum, A.	calculation:				
Mosterd, Houwelingen	Yes: powered study.				
G. Van, P. C. Koopmans, P. R. Stella,	A sample size of 814				
E. Boersma, and C.	patients (80% power for primary outcome)				
Hamm. Effect of Early,	ioi primary outcome;				
Pre-Hospital Initiation	. 				
of High Bolus Dose	ITT analysis:				
Tirofiban in Patients	Yes				
With ST-Segment					
Elevation Myocardial					
Infarction on Short-					
and Long-Term Clinical					
Outcome.					
J.Am.Coll.Cardiol. 55					
(22):2446-2455, 2010.					
(3 year results)			1		
%	fPPCI (early tirofiban – pre-hospit	tal) Place	ebo PPCI (n = 493)	p value	
	(n = 491)				
Age, years; mean	62	62		-	
Male gender	77	75		-	

%	fPPCI (early tirofiban – pre-hospital) (n = 491)	Place	bo PPCI (n = 493)	p value
Age, years; mean	62	62		-
Male gender	77	75		-
Diabetes	12	11		-
Hypertension	34	35		-
Current smoking	45	49		-
Hypercholesterolemia	29	25		-
Previous MI	9	8		-

Previous CABG	or PCI	12			9		-		
Killip Class >1		11			13		-		
TIMI flow (%)							0.069		
0		42			50				
1		14			11				
II		22			20				
III		22			20				
Critical time int	tervals, minutes; r	median (IQR)							
antithrombotic	pre-treatment a	nd study drug, to	angiography		55 (43–70)		-		
Effect Size									
Angioplasty									
Overall 99% (pe	eople had PCI and	l 90% in had stent	s (in both arms)						
Outcomes									
	In-hospital (aft	er procedure)		At 30 days			1 year		
	fPPCI (early			fPPCI (early	Placebo PPCI			fPPCI (early	p value Placebo PPCI
	tirofiban – pre-hospital)	Placebo PPCI (n = 493)		tirofiban – pre-hospital)	(n = 493)			tirofiban – pre-hospital)	(n = 493)
N (%)	(n = 491)	(,	p value	(n = 491)	(155)	p value	N (%)	(n = 491)	()
ST-segment			0.03	. ,		·	ST-segment	,	
recovery, %	286/436	264/440		-	-	-	recovery, %	286/436	264/440
Complete	(65.6)	(60.0)		-	-	-	Complete	(65.6)	(60.0)
Intermediate	10/436 (22.9)	104/440		-	-	-	Intermediate	10/436 (22.9)	104/440
None	50/436 (11.5)	(23.6)					None	50/436 (11.5)	(23.6)
		72/440 (16.4)							72/440 (16.4)
Normalised	160/451	132/455	-	-	-	-	Normalised ST	160/451	132/455
ST segment	(35.5)	(29.0)		4 (472 (0.2)	7/477/4.5	0.07	segment	(35.5)	(29.0)
Stroke	-	-	-	1/473 (0.2)	7/477 (1.5)	0.07	Stroke	-	-
Death	-	-	-	11/473 (2.3)	19/477 (4)	0.14	Death	-	-
Cardiac death	-	-	-	-	-	-	Cardiac death	-	-
Non-cardiac									

death

death

Recurrent MI	-	-	-	13/473 (2.7)	14/477 (2.9)	0.86	Recurrent MI	-	-
Major bleeding	-	-	-	19/473 (4.0)	14/477 (2.9)	0.36	Major bleeding	-	-
Minor bleeding	-	-	-	29/473 (6.1)	21/477 (4.4)	0.23	Minor bleeding	-	-
TIMI flow grade 0–2 or slow reflow	29/488 (5.9)	45/492 (9.1)	0.58	-	-	-	TIMI flow grade 0–2 or slow reflow	29/488 (5.9)	45/492 (9.1)
Urgent TVR	-	-	-	20/477 (4.2)	18/473 (3.8)	0.7614	Urgent TVR	-	-
	In-hospital (after procedure)	p value	At 30 days	p value	1 year	p value		In-hospital (after procedure)	p value
Cardiac proced	ures								
		fPPCI	(early tirofiban -	- pre-hospital)					
%		(n = 4	91)		%		p valu	e	
Coronary angio	gram	99%					-		
PCI		99%					-		
Transfer and tir	Transfer and timing, and hospital stay								
No details give	n								

Table 23: Kanakakis et al. 2009⁶²

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Outcome measures	Source of funding
J Kanakakis, J N. Nanas, E P. Tsagalou, G D. Maroulidis, S G. Drakos, A S. Ntalianis, P Tzoumele, E Skoumbourdis, P Charbis, S Rokas, and M Anastasiou- Nana. Multicenter randomized trial of facilitated percutaneous coronary intervention with low-dose tenecteplase in patients with acute myocardial infarction: the Athens PCI trial Cathatar Cardio vasc.Interv. 74 (3):398-405. 2009.	Design: RCT (Greece, 1 area, multiple hospitals) Enrolment: April 2005 – October 2006 Setting Patients were directly admitted or transferred to tertiary care hospital with PPCI facilities Randomisation: Not mentioned (just 'randomised') Allocation concealment: Not mentioned Blinding: Caccaccare only were blinded to treatment assignment	n = 284 Drop- outs/missing patients/ineligible: In-hospital: n = 0 (some deaths but this was the clinical outcome measure)	INCLUSION CRITERIA: Age ≥18 years <6hrs between symptom onset and randomisation ≥0.1 mV STE in ≥2 limb leads or ≥0.2 mV in 2 contiguous precordial leads or presence of left bundle branch block EXCLUSION CRITERIA expected arrival to cathlab <30 mins after randomisation SBP >180 mmHg or DBP >110 mmHg, or both, on multiple measurements use of GPIs within preceding 7 days major surgery, biopsy of parenchymal organ or severe trauma within 2 months head injury or trauma occurring after the onset of ongoing MI history of stroke. TIA, dementia or structural damage to the CNS ongoing treatment with oral anticoagulants, LMWH administered <12h before randomisation or platelet count <100.000 ul-1	fPCI (Tenecteplase) (n = 143) Tenecteplase – 10 mg (quarter of the standard dose) NOTE: Before catheteri patients had UFI 70 U/kg, IV) and 300 mg p.o). Her added in the cat and activated clc 250-300 sec. Immediately before all patients under received infusion (eptifibatide) in 180 ug/kg initial by 2ug/kg/min dinfusion for 21hr Patients with ste 300mg clopidogrel 75 mg received infusion for 21hr Patients with ste 300mg clopidogrel 75 mg received infusion for 21hr Patients with ste 300mg clopidogrel 75 mg received infusion for 21hr Patients with ste clopidogrel 75 mg received infusion for 21hr Patients with ste clopidogrel 75 mg received infusion for 21hr Patients with ste clopidogrel 75 mg received infusion for 21hr Patients with ste clopidogrel 75 mg received infusion for 21hr Patients with ste clopidogrel 75 mg received infusion for 21hr Patients with ste clopidogrel 75 mg received infusion for 21hr Patients with ste clopidogrel 75 mg received infusion for 21hr Patients with ste clopidogrel 75 mg received infusion for 21hr Patients with ste clopidogrel 75 mg received infusion for 21hr Patients with ste clopidogrel 75 mg received infusion for 21hr Patients with ste clopidogrel 75 mg received infusion for 21hr Patients with ste clopidogrel 75 mg received infusion for 21hr Patients with ste clopidogrel 75 mg received infusion for 21hr Patients with ste clopidogrel 75 mg received infusion for 21hr Patients with ste clopidogrel 75 mg received infusion for 21hr Patients with ste clopidogrel 75 mg received infusion for 21hr Patients with ste clopidogrel 75 mg received infusion for 21hr Patients with ste clopidogrel 75 mg received infusion for 21hr Patients with ste clopidogrel 75 mg received infusion for 21hr Patients with ste clopidogrel 75 mg received infusion for 21hr Patients with ste clopidogrel 75 mg received infusion for 21hr Patients with ste clopidogrel 75 mg received infusion for 21hr Patients with ste clopidogrel 75 mg received infusion for 21hr Patients with ste clopidogrel 75 mg received infusion for	sation – all H (single bolus, aspirin (150- parin could be h lab to reach potting time of ergoing PCI n of GPI the cath lab. bolus followed ose-adjusted rs ents received rel	Primary: TIMI flow grade 2 or 3 Secondary: composite of death, congestive HF and cardiogenic shock during hospitalisation Non-cerebral bleeding complications death congestive HF cardiogenic shock Length of follow-up: Inhospital	Not stated

Sample size calculation: Slightly underpowered study: 80% power using approximately 150 patients in each arm to demonstrate a 15% absolute increase in TIMI flow grade 2 or 3 (primary end point). ITT analysis: Not mentioned – it appears so (no dropouts but there were deaths!) – numbers given were always out of the total randomised		>10 mins of cardion resuscitation in preweeks pregnancy, lactation parturition in previous anticipated difficult vascular access participation in and or device investigat previous 30 days any other disorder expose the pt at incrisk inability or unwillin follow the protocol Demographics and characteristics see below	n or ous 30 cies with other drug cion in that would ordinate gness to			
	Escilitated BCI	 (tenecteplase) (n = 143)	PPCI (n = 1	41)	n va	alue
Age, yrs; mean (SD)	61 (12)	(tenectepiase) (ii – 145)	59 (12)	41)	0.16	
Age >75 yrs, n	13		11		0.34	
Male gender, %	92		89		0.23	
Killip Class, %	JL		03		0.23	
	75		76		0.63	3
/	17		16			
IV	8		8			
History of hypertension, %	54		50		0.58	8
History of diabetes, %	21		31		0.12	2

Anterior	46	44	0.25
Inferior	50	47	
Other	4	9	
SBP, mmHg; mean (SD)	125 (24)	127 (26)	0.83
Pre-PCI TIMI flow, n (%)			
0/I	54 (38)	81 (58)	0.0001
11/111	85 (59)	52 (37)	
Not measurable	4 (3)	8 (6)	
Previous PCI, %	16.7	23.2	0.35
Previous CABG, %	5	3	0.49
Previous MI, %	13	13	0.94
Onset of symptoms to hospital arrival	100 (60-169)	120 (30-240)	-
Onset of symptoms to tenecteplase	135 (90-228)	-	-
Onset of symptoms to 1st coronary angiogram (target lesion)	224 (172-306)	259 (172-368)	
Door to 1st coronary angiogram (target lesion)	108 (75-150)	99 (68-54)	
Onset of symptoms to 1st balloon	232(185-315)	275 (190-380)	
Door to 1st balloon	122 (91-175)	120 (89-175)	-
Tenecteplase to 1st balloon	121 (90-168)	.	
Before catheterisation			
tenecteplase	100	0	-
bolus heparin	100	100	1.00
During catheterisation			
GPI	91.2	93.5	0.55
Clopidogrel	88.2	90.4	0.85
After catheterisation			
clopidogrel or ticlopidine	88.2	90.4	0.85

Angioplasty								
Higher use of PCI in the fPCI group – 94% versus 88%								
Outcomes								
	Inhospital		p value					
n (%)	Facilitated PCI, tenecteplase	PCI						
	(n = 143)	(n = 141)						
Death	8 (6)	5 (3.5)	0.57					
Congestive HF	24 (17)	5 (3.5)	1.00					
Cardiogenic shock	12 (8)	11 (8)	0.91					
Intracranial haemorrhage	0	0	1.00					
Haemorrhagic stroke	0	0	1.00					
Primary (non-fatal) ischaemic stroke	1 (0.7)	0	0.93					
Unclassified stroke	0	0	1.00					
All strokes	1 (0.7)	0	0.92					
Major bleeding complications	8 (6)	5 (3.5)	0.55					
Death + HF + shock	39 (27)	34 (24)	0.59					
Patency / post-PCI TIMI flow								
0/1	13 (10)	13 (11)	NS					
11/111	122 (90)	52 (37)						
Not measurable	1 (3)	8 (6)						
Transfer: No details								

Table 24: LIPSIA-STEMI trial¹¹²

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Outcome measures	Source of funding
H. Thiele, I. Eitel, C. Meinberg, S. Desch, A. Leuschner, D. Pfeiffer, A. Hartmann, U. Lotze, W. Strauss, and G. Schuler. Randomized comparison of pre- hospitalinitia ted facilitated percutaneou s coronary intervention versus primary percutaneou s coronary intervention in acute myocardial infarction very early	Design: RCT (12 centres in 1 region of Germany) Enrolment: August 2006 – August 2009 Setting Ambulance to interventional centre for PCI Randomisation: Poor/unclear: sealed envelopes Allocation concealment: Good: contacted the study co-ordinating centre Blinding: No mention of	n = 162 Drop-outs/ missing data/ ineligible: Excluded due to no infarction: n = 1 (fPCI) and n = 3 (PPCI) None lost to follow-up at 30 days OVERALL acceptable losses (<20%): primary end point analysis fPCI - n = 12 (14.8%) missing and PPCI n = 10 (12%) missing. Secondary	Inclusion criteria: Peoplewith presence of ischaemic symptoms <3hours ST-segment elevation ≥0.1 mV in 2 or more extremity leads or ≥0.2 mV in 2 or more precordial leads Exclusion criteria: Contraindications to fibrinolysis (eg. previous stroke, active bleeding, history of major trauma or surgery <30 days, active peptic ulcer, neoplasms, uncontrolled HT >200 mmHg, chronic oral anticoagulation, cardiogenic shock and pregnancy) typical contraindications for MRI such as pacemakers, defibrillators and intracerebral metallic clips Demographics and baseline characteristics see below	fPCI (tenecteplase +clopidogrel background) (n = 81) NOTE: given pre-hospital Tenecteplase – weight-adjusted IV dose (as per ASSENT trial) During PCI: Heparin - dose calculated to achieve activated clotting time of 200-250 seconds GPIs – recommended at standard dose for 12h	PPCI (clopidogrel background) (n = 81) During PCI: Heparin - dose calculated to achieve activated clotting time of 200-250 seconds GPIs — recommended at standard dose for 12h	Primary: infarct size. Secondary: composite of death, reinfarctiona nd new congestive HF <30days after randomisation; post-hospital outcomes at 30 days follow-up: reinfarction; death; new HF; severe of life-threatening, moderate or minor bleeding; occurrence of ischaemic stroke Length of follow-up: Inhospital; 30 days	Supported in part by German Heart Research foundation and Boehringhe Ingelheim GmbH, Germany

after symptom onset: The LIPSIA- STEMI trial (Leipzig Immediate Prehospital Facilitated Angioplasty in ST- segment myocardial infarction). JACC: Cardiovascu I ar Intervention s 4 (6):605- 614, 2011.	physicians TIMI flow and clinical end points had blinded observers/adjudicati on committee Sample size calculation: Powered study: 80% power using at least 64 patients for infarct size outcome. ITT analysis: Yes	end point analysis fPCI — n = 1 (1%) and PPCI — n = 3 (4%) missing.				
%		fPCI (tenecteplase +clopidogrel background) (n = 80)	PPCI (clopi	idogrel background) (n = 78)	p value	
Age, yrs; medi	ian (IQR)	63 (54-73)	61 (53-72)		0.57	
Male gender		76	82		0.48	
Diabetes melli	itus	36	24		0.15	
Hypertension		66	64		0.91	
Current smoki	ing	41	48		0.43	
Hypercholeste	erolemia	40	26		0.12	
Previous MI		3	5		0.65	

Previous CABG	0		1		0.99	
Killip Class					0.36	
1	71		74			
2	14		18			
3	6		5			
4	9		3			
TIMI flow (%)					<0.001	
II	26.6		10.3			
III	44.3		24.4			
Concomitant medications						
ВВ	96		99		0.63	
ACEi	98		97		0.63	
Aspirin	99		100		0.99	
Clopidogrel	100		100		1.00	
Statins	98		97		0.63	
Aldosterone antagonists	6		4		0.74	
GPIs	29		88		<0.001	
Critical time intervals, mins; media	n (IQR)					
Symptom onset to first balloon infla	ition Symptom onset to first ba	alloon inflation	Symptom onset to first b	alloon inflation	Symptom onset	to first balloon inflation
Door to balloon	Door to balloon		Door to balloon		Door to balloon	
Effect size						
Angioplasty						
Overall 93% and 98% (fPCI and PPCi	respectively) people had PCI and	l 100% in each arı	m had stents			
Outcomes						
Inhospita	al (after procedure)		At 30 days			p value
fPCI	. , .		fPCI			fPCI (tenecteplase
(tenecte)	plase PPCI (clopidogrel		(tenectepla	se		+clopidogrei
N (%) +clopido		p value	+clopidogre		5)	background)

	background) (n = 80)	(n = 78)		background) (n = 80)		(n = 80)
ST-segment recovery,			0.18		ST-segment recovery,	
%	42.9	57.7		-	%	42.9
Complete	40.2	29.5		-	Complete	40.2
Intermediate	16.9	12.8		-	Intermediate	16.9
None					None	
Ischaemic stroke	-	-	-	1	Ischaemic stroke	-
Death	-	-	-	5 (6.1)	Death	-
Nonfatal reinfarction	-	-	-	5 (6.1)	Nonfatal reinfarction	-
New congestive HF	-	-	-	6 (7.4)	New congestive HF	-
Bleeding:	-	-	-		Bleeding:	-
severe/life threatening				2.5%	severe/life	
moderate				2.5%	threatening	
mild				4.9%	moderate	
					mild	
TIMI flow grade III (in	83.2	88.5	0.44	-	TIMI flow grade III (in	83.2
IRA), %					IRA), %	
Cardiac procedures						
						p value
%		fPCI (tenecteplase - background) (n = 80	•	PPCI (clopidogrel b	ackground) (n = 78)	%
Coronary angiogram		100%		-		Coronary angiogram
PCI		93%		98%		PCI
Transfer and timing, a	ınd hospital stay: No	details given				

Table 25: Mehilli et al. 2009^{81,99}

Reference	Study type	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Mehilli J. et	Design:	Number of patients:	Abciximab	Placebo	30 days; 1	1°	BRAVE-3
al. 2009	RCT (double	n = 800	(n = 401)	(n = 399)	year	Infarct size	was
	blind)	Drop-outs: None	PPCI	PPCI			supported
Abciximab in						Other:	by grants from
oatients with	Enrolment:	Inclusion criteria	Abciximab dose:	Dose:		Death, recurrent	Deutsches
icute ST-	June 2003 –	<24 hours of the onset symptoms	0.25 mg/kg,	Intravenous		MI, stroke,	Herzzentru
segment elevation	January 2008	Chest pain ≥ 20 min	bolus, followed	bolus of 70U		urgent	, Munich,
myocardial		≥ 0.1mV ST-elevation in 2 or more limb	by infusion of	heparin/kg		revascularisation,	Germany.
nfarction	Setting	leads, ≥ 0.2mV in 2 or more contiguous	0.125microgram/ kg/minute (max	followed by		in-hospital incidence of	
undergoing	Hospital (5 PCI	precordial leads	dose: 10	infusion of placebo for 12		major and minor	
primary	centres).		microgram/min)	hours.		bleeding	
percutaneou		Exclusion criteria	for 12 hours	nours.		complications	
coronary	Randomisation:	Received fibrinolytic therapy for the					
ntervention after	Yes. Computer-	index infarction	Note:				
clopidogrel	generated	Previous stroke (within last 3 months)	Before				
oading.	random sequence	Active bleeding or bleeding diatheses	catheterisation				
_	Allocation	Recent trauma or major surgery (during	: All patient				
and	Allocation concealment:	the last month)	received				
	Yes, sealed	Suspected aortic dissection	clopidgrel 600mg orally; aspirin				
S. Schulz, K.	envelopes.	Oral anticoagulation therapy with	bolus 500mg;				
A. Birkmeier,		coumarin derivatives (within the last 7 days)	heparin 60U/kg				
3.	Blinding:	Recent use of glycoprotein IIb/IIIa	(max dose:				
Ndrepepa,	Yes. Patients,	(within the last 14 days)	5000U)				
W. Moshage, Dotzer, K.	medical staff and	Severe uncontrolled hypertension	intravenous.				
Huber, J.	investigators	(>180mm Hg, unresponsive to therapy)	After reperfusion				
Dirschinger,	were blinded.	Relevant hematologic deviations	all patients were treated with				
M. Seyfarth,		(haemoglobin <100g/L or hematocrit	clopidogrel 75mg				
A. Schomig,	Sample size	<34%, platelet count < 100x109/L or >	twice daily for 3				
A. Kastrati,	calculation:	600 x109/L)	days and 75 mg/d				
and J.	The assumptions	Malignancies	thereafter for 30				

Mehilli. One- year clinical outcomes with abciximab in acute myocardial infarction: Results of the BRAVE-3 randomized trial. Clin.res.cardi ol. 99 (12):795- 802, 2010. (1 year results)	used for this purpose included an infarct size of 16.9% (SD, 13.9%) of the left ventricle in the placebo group and a 20% reduction with abciximab. Choosing a 2-sided \(\alpha\) level of 0.05 and power of 90%, we needed 353 patients with scintigraphic follow-up study in each group. We allowed for the possibility that not all patients would have a follow-up SPECT and included a total of 800 patients. ITT analysis: Yes	Prolonged cardiopulmonary resuscitation or cardiogenic shock >80 or <18 years of age Known or suspected pregnancy Allergy to study drugs Demographics and baseline characteristics see below	days and with aspirin 100mg twice daily indefinitely. All patients were sent to the cath lab for coronary angiography and PCI. The decision to perform a coronary intervention was at the discretion of the operator. The recommended intervention was coronary stenting.	
		Abciximab (n = 401)	Placebo (n = 399)	p value
Age, years		62.4±11.7	61.58±12.2	0.50
Women, n(%)		98 (24)	109 (27)	0.35
- , (,-)		· /	/	

Arterial Hypertension, n(%)	280 (70)	282 (71)	0.79
Hypercholesterolemia, n(%)	167 (42)	177 (44)	0.44
Diabetes mellitus, n(%)	76 (19)	65 (16)	0.32
Current smoking, n(%)	168 (42)	162 (41)	0.71
BMI, Kg/m ²	27.1±3.8	27.0±4.1	0.74
History of MI, n(%)	38 (10)	43 (11)	0.54
History of CABG surgery, n(%)	15 (4)	8 (2)	0.14
Infarct localisation, n(%)			0.81
Anterior	168 (42)	174 (44)	
Inferior	174 (43)	172 (43)	
Lateral	59 (15)	53 (13)	
Killip class, n(%)			0.89
T	304 (76)	307 (77)	
II	75 (19)	74 (19)	
III	15 (4)	11 (3)	
IV	7 (2)	7 (2)	
Arterial blood pressure, mm Hg			
Systolic	138±23	139±22	0.39
Diastolic	79±13	80±14	0.62
Heart rate, bpm	73±17	73±16	0.50
Peak CK (creatine kinase)-MB, U/L, U/L	258.9±251.5	265.5±240.5	0.72
Data are presented as mean±SD when ap	propriate		

Definitions of end points

Recurrent infarction

Diagnosis of recurrent infarction was based on the following criteria:

If the biomarkers of the index MI were still increasing or the peak had not been reached, the patients had to have both new ECG changes consistent with MI (new or reelevation of ST segments \geq 0.2 mV in \geq 2 contiguous precordial leads, \geq 0.1 mV in \geq 2 adjacent limb ECG leads, or development of new, abnormal Q waves considered distinct from the evolution of the index MI) and recurrent ischaemic discomfort lasting >20 minutes at rest or ischaemia-triggered haemodynamic instability;

If the biomarkers of the index MI were falling but still above the upper limit of normal, the patients had to have either an increase in creatine kinase-MB (creatine kinase) ≥50% over the nadir level or new ECG changes consistent with MI;

If the biomarkers of the index MI were normalised, the patients had to have a new increase in creatine kinase-MB (creatine kinase) ≥3 times the upper limit of normal. Stroke

The diagnosis of stroke required confirmation by computed tomography or magnetic resonance imaging of the head.

Bleeding

A bleeding complication was defined as major if it was intracranial or if clinically significant overt signs of haemorrhage were associated with a drop in haemoglobin of >5 g/dL (or, when haemoglobin was not available, an absolute drop in hematocrit of at least 15%).

Effect Size

	30 day results			1 year results		
n(%)	Abciximab (n = 401)	Placebo (n = 399)	n(%)	Abciximab (n = 401)	Placebo (n = 399)	n(%)
Death	13 (3.2)	10 (2.5)	0.53	27 (6.8)	16 (4)	0.09
Recurrent MI	6 (1.5)	6 (1.5)	0.99	12 (3.1)	11 (2.8)	0.83
IRA revascularisation	3 (0.8)	4 (1)	0.70	62 (16.3)	86 (22.3)	0.04
Stroke	1 (0.3)	1 (0.3)	1.0	3 (0.8)	1 (0.3)	0.32
TIMI major bleeding	7 (1.8)	7 (1.8)	0.99	-	-	-
TIMI minor bleeding	15 (3.7)	7 (1.8)	0.09	-	-	-
Repeat revascularisation of Target lesion	-	-	-	53 (13.9)	76 (19.7)	0.04
TIMI (post PCI, not 30 d	days or 1 year) - overall N	IS (p = 0.49)				
0	11 (3)	6 (1)	0.49 overall	-	-	-
1	4 (1)	7 (2)				
2	16 (4)	18 (5)				
3	370 (92)	368 (92)				
Repeat revascularisation of Target lesion	-	-	-	53 (13.9)	76 (19.7)	0.04

Table 26: Gabriel et al. 2006⁴⁷

	Studutuna	Number of	Dationt characteristics	Intervention	Comparison	Outcome	Source of
Reference	Study type	patients	Patient characteristics	Intervention	Comparison	measures	funding
H. Mesquita Gabriel, Joaquim A.	Design: RCT (prospective)	n = 80	Inclusion criteria: Clinical picture suggestive of	Early group (n = 36)	Later group (n = 38) Later described as	TFG 2 or 3	Not reporte d
Oliveira, Pedro Canas da Silva, J. Marques da Costa, and J.	Enrolment: August 2001- September 2002	Drop-outs: 6 in total did not receive PCI. 4 from	AMI of <12 hours duration Presence of an ST-segment elevation of >0.2mV in 2 or more adjunctive leads and suitability for angiographic	Early described as administration in the emergency room of abciximab bolus in MAI patients undergoing PCI.	administration in the catheterisation lab (CL) of abciximab bolus in MAI patients undergoing PCI.	Death Re-MI Urgent	u
A. C. da Cunha. Early	Setting Hospital (Santa	the early group and 2 from the	evaluation Exclusion criteria:	0.25 mg/kg abciximab bolus in the emergency room	0.25 mg/kg abciximab bolus in the CL (given	TVR	
administrati on of	Maria Hospital, Lisbon).	later group	Admitted >12 hours from onset	(given placebo in the	placebo in the	TIMI-	
abciximab			of symptoms	catherisation lab (CL)) Both groups: Pre-	emergency room) Both groups:	defined bleeding	
bolus in the emergency	Randomisation:		Had received fibrinolytic agent for the treatment of the	PCI	Pre-PCI		
department improves angiographic outcome after primary PCI as assessed by TIMI	Yes. Randomly assigned in a 1:1 ratio to intervention or comparison group. Allocation concealment:	tor t curr Expo Had up. Den chai	current episode Experienced cardiogenic shock, Had known bleeding diathesis Demographics and baseline characteristics see below	Recommended to received 250 mg aspirin and 5000 U bolus of heparin Other medications such as nitrates, morphine and beta-blockers used at the discretion of the emergency department cardiologist.	Recommended to received 250 mg aspirin and 5000 U bolus of heparin Other medications such as nitrates, morphine and betablockers used at the	Length of follow-up 30 days	
frame count: results of the early	Unclear. Blinding:			In the CL dose adjusted heparin was administered to achieve an activated clotting time (ACT) of ≥250 sec	discretion of the emergency department cardiologist.		
ReoPro administrati	2-way blinding.The study			Post PCI	In the CL dose adjusted		
on in myocardial infarction (ERAMI) trial.	packages were labelled in a way to preserve 2-way blinding.			In patients who received PCI coronary stenting was used liberally and 1 of 2 antithrombotics was associated to aspirin:	heparin was administered to achieve an activated clotting time (ACT) of ≥250 sec Post PCI		

Demographics and baseline characteristics	coronary patency at initial angiography with an overall type I error rate of 0.05 (2-sided). Sample size was calculated to be 78 patients. ITT analysis: No. Per protocol analaysis	at initial orally for \ge 24 day angiography with an overall type I followed by 250 m error rate of 0.05
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Demographics and baseline characteristics

	Early group (n = 36)	Later group (n = 38)
Age, years	60±12	63±16
Male gender	23 (64)	30 (79)
Hypertension	23 (64)	23 (61)
Diabetes	9 (25)	7 (18)
Smoker	19 (53)	16 (42)
Hypercholesterolemia	19 (53)	19 (50)
Prior MI	3 (8)	4 (11)
Anterior MI location	16 (44)	24 (63)
Treatment in emergency department		
Aspirin (250mg)	35 (97)	37 (97)
Heparin (5,000 U)	27 (75)	31 (82)
First angiography		

TFG 2 or 3	1	.1 (31)		10 (26)
TFG 3	4	(11)		3 (8)
TFG 2	7	' (19)		7 (18)
TFG 1	4	(11)		2 (5)
TFG 0	2	1 (58)		26 (68)
Data are presented as mean±	:SD or n (%)			
Effect Size / Outcomes				
30 days	Early group (n = 36)	Later group (n = 38)	p value	Details
Final angiography				
TFG 2 or 3	35 (97)	34 (90)	0.18	
TFG 3	32 (89)	29 (76)	0.08	
Death	4 (11%)	5 (13%)	0.78	Causes of death: heart failure (5); mechanical complications (2) (free wall rupture and ventricular septal defect); stroke (1) sudden death after discharge (1). groups not specified
Re-MI	0	1 (3%)	0.32	
Urgent target vessel revascularisation	1 (3%)	0	0.3	
TIMI-defined bleeding				
Significant	4 (11%)	2 (5%)	0.37	
Major	1 (3%)	0	0.51	
Minor	3 (8%)	2 (5%)	0.53	

Table 27: Bellandi et al. 2006⁸

Reference	Study type	Patient characteristics	Intervention	Comparison	Outcome measures	Source of funding
Bellandi, 2006	Design: RCT (prospective)	Number of patients: n = 55	Early group (n = 27)	Later group (n = 28)	TIMI flow grade	Not reporte
		Drop-outs: None.		All patients received	Stents	d
F Bellandi, M Maioli, M Leoncini, A	Enrolment: June 2003 – January 2004	Inclusion criteria: Presenting within 6 hours of	All patients received abciximab (0.25 mg/kg as a bolus followed by a 12 hour	abciximab (0.25 mg/kg as a bolus followed by a 12 hour infusion of 0.125 mcg/kg/min)	Death	
Toso, and R P. Dabizzi. Early	Setting	symptom onset with ST- segment elevation of more	infusion of 0.125 mcg/kg/min) in the emergency room	in the catheterisation lab after diagnostic angiography	Reinfarction	
abciximab administrati	Hospital.	than 1 mm in at least 2 contiguous leads of ECG	Afternoon de militation ell	After randomisation: all	Urgent TVR	
on in acute myocardial	Randomisation: Yes. Computerised	Exclusion criteria: Previous MI	After randomisation: all patients received heparin as a bolus of 70U/kg	patients received heparin as a bolus of 70U/kg (maximum 7000U) together with 250mg	Major bleeding	
infarction randomisation. treated with	Previous PCI	(maximum 7000U) togethe with 250mg aspirin	aspirin intravenously in the emergency room	Intra-cranial bleeding		
primary coronary intervention	Allocation concealment: Blinded envelopes	Previous coronary artery bypass Left bundle branch block	intravenously in the emergency room	If necessary additional boluses of heparin were	Length of follow-	
Int.J.Cardiol.	·	Bleeding diathesis	If necessary additional boluses of heparin were	administered to achieve an activated clotting time of	7 days, 1 month	
108 (1):36- Blinding: 42, 2006. No. Open label	Administration of fibrinolytic agents for the current episode Recent stoke	administered to achieve an activated clotting time of	200s.			
	Sample size	Uncontrolled hypertension	200s.	Before angiography: Intravenous injection of 740		
calculation: Not reported. ITT analysis: Yes		Recent surgery Oral anticoagulant therapy Known contraindications to	Before angiography: Intravenous injection of 740	to 1110 MBq of technetium- 99 m sestamibi.		
	therapy with abciximab, aspirin, clopidogrel or heparin	to 1110 MBq of technetium- 99 m sestamibi.	After PCI:			
		Demographics and baseline characteristics see below	After PCI: 24 hour infusion of 7U/kg per hour of heparin was	24 hour infusion of 7U/kg per hour of heparin was performed. The target activated partical-		

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Demographics and baseline characteristics				
	Early group (n = 27)	Later group (n = 28)		
Age, years	62.7±11.5	63.8±11		
Male gender	22 (81.5)	22 (78.6)		
Diabetes	6 (22.2)	7 (25)		
Hypertension	8 (29.6)	9 (32)		
Hyperlipidimia	13 (48)	13 (46.4)		
Current smoking	14 (51.8)	14 (50)		
Kilip class ≥	5 (18.5)	8 (28.6)		
Not-low-risk patients	15 (55.6)	15 (53.6)		
Infarct artery				
Left anterior descending	12 (44.4)	13 (46.4)		
Right coronary artery	12 (44.4)	13 (46.4)		
Circumflex artery	3 (11.2)	2 (7.2)		
Multivessel disease				

2 vessels	11 (40.7)	13 (46.4)			
3 vessels	3 (11.2)	3 ()10.7			
Symptom onset to admission (min)	129±63	122±95			
Door to abciximab (min)	32±14	55±34			
Door to balloon (min)	80±26	73±37			
Start of abciximab to balloon (min)	49±14	19±9			
Initial TIMI flow grade					
0	15 (55.6)	19 (67.9)			
1	1 (3.7)	2 (7.1)			
2	1 (3.7)	4 (14.3)			
3	10 (37)	3 (10.7)			
Data are presented as mean±SD or n (%)					

Effect Size / Outcomes				
Angiographic results	Early group (n = 27)	Later group (n = 28)	p value	Details
Final TIMI flow grade			0.37	
2	1 (3.7%)	4 (14.3%)		
3	26 (96.3%)	24 (85.7%)		
Direct stenting	8 (30%)	2 (7%)	0.07	
Multiple stents	4 (15%)	11 (39%)	0.08	
Number of stent	1.1±0.4	1.5±0.6	0.005	
1 month				
Death	1	1		Both cardiogenic shock
Reinfarction	0	0		
Urgent vessel revasculatisation	0	0		
Major bleeding (a decrease in haemoglobin >2g/dl with the need for transfusion	1	2		
Intra-cranial bleeding during treatment with abciximab	0	0		

Table 28: RELAX-AMI trial 79

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Outcome measures	Source of funding
M. Maioli, F. Bellandi, M.	Design: RCT (prospective)	n = 210	Inclusion criteria: Presenting within 12 hours of	Early group (n = 105)	Later group (n = 105)	Mortality	Not reporte
Leoncini, A. Toso, and R. P. Dabizzi. Randomized early versus late abciximab in acute myocardial infarction treated with primary coronary intervention (RELAX-AMI Trial). J Am Coll Cardiol 49 (14):1517- 1524, 2007.	Enrolment: June 2003 – March 2006	Drop-outs: 9 patients died before 1 month follow-up (3 from early group, 6 from later group.	symptom onset with ST- segment elevation of more than 1 mm in at least 2 contiguous leads of ECG	All patients received abciximab (0.25 mg/kg as a bolus followed by a 12 hour infusion of 0.125 mcg/kg/min) in the emergency room	as a bolus followed by a 12 o.125 mcg/kg/min) in the catheterisation lab after diagnostic angiography domisation: After randomisation: all patients received so a bolus of naximum 7000U) together with spirin 250mg aspirin ously in the	Recurrent MI Urgent PCI	d
	Setting Hospital (Italy).		Exclusion criteria: Previous MI Previous PCI			Coronary artery bypass grafting	
	Randomisation: Yes. Open label		Previous coronary artery bypass surgery Left branch bundle block Bleeding diathesis administration of fibrinolytic agents for the current episode	After randomisation: all patients received heparin as a bolus of 70U/kg (maximum 7000U) together with 250mg aspirin intravenously in the		Bleeding complications	
	randomisation.					Minor bleeding	
	Allocation concealment: Unclear.					Major bleeding Intracranial	
		Recent stroke Uncontrolled hypertension	emergency room	emergency room	bleeding		
	Blinding: Unclear. Blinded envelopes after open label randomisation. Sample size calculation:	Recent surgery Oral anticoagulant therapy	If necessary additional boluses of heparin were	If necessary additional boluses of heparin were administered to	TIMI flow rate		
		Known contraindications to therapy with abciximab, aspirin, clopidogrel or heparin	administered to achieve an activated clotting time of 200s.	achieve an activated clotting time of 200s.	Length of follow-up 1 month		
			Previous AMI likely to have pre-existent asynergies	After PCI:	After PCI: 24 hour infusion of		
	The sample size to detect a difference between 10% and 20% increase in TIMI flow grade 3.		Demographics and baseline characteristics see below	24 hour infusion of 7U/kg per hour of heparin was performed. The target activated	7U/kg per hour of heparin was performed. The target activated particalthromboplastin time		

With an α value of 0.05 and 80% power, was determined to be 95 per group. With regard to drop outs at least 105 patients were planned to be included in both groups. ITT analysis: No. Available case analysis		partical- thromboplastin time was between 1.5 and 2.0 times the control value. Patients routinely received aspirin (100 mg/day indefinitely) and clopidogrel (75 mg/day for at least 1 month PCI performed in all patients. Stent insertion in all patients.	was between 1.5 and 2.0 times the control value. Patients routinely received aspirin (100 mg/day indefinitely) and clopidogrel (75 mg/day for at least 1 month PCI performed in all patients. Stent insertion in all patients.
	Early group (n = 105)		Later group (n = 105)
Age, years	60.4±10		66±12
Age ≥75 years	17 (16)		26 (25)
Female gender	25 (24)		27 (26)
Hypertension	51 (49)		48 (46)
Diabetes mellitus	31 (29)		31 (29)
History of smoking	45 (43)		54 (51)
Hyperlipidimia	43 (41)		41 (39)
Low risk patients	36 (34)		43 (41)
Killip class ≥2	20 (21)		25 (26)
Infarct artery			
Right coronary artery	44 (42)		40 (38)
Circumflex artery	9 (9)		12 (11)
Left anterior descending	52 (49)		52 (49)
Main vessel	0		1 (1)

Multivessel disease				
2-vessel	32 (30)	40 (38)		
3-vessel	22 (21)	25 (24)		
Symptom onset to admission (min) (range)	85 (59–160)	105 (63–170)		
Door to abciximab (min) (range)	23 (15–37)	60 (40–88)		
Door to balloon (min) (range)	84 (67–107)	73 (59–104)		
Start of abciximab to balloon (min) (range)	55 (46–72)	14 (11–18)		
Initial TIMI flow grade				
0–1	63 (60)	88 (84)		
2	17 (16)	6 (6)		
3	25 (24)	11 (10)		
Data are presented as mean±SD or n (%)				

Effect Size / Outcomes

	Early group (n = 105)	Later group (n = 105)	p value	Details
1 Month				
Death	3	6		Early group: cardiogenic shock (1); sudden death (1); fatal repeat AMI (1)
				Later group: cardiogenic shock (4); free wall rupture (1); fatal repeat AMI (1)
Recurrent MI	2	4		
Repeated Urgent PCI	2	1	0.62	
Coronary artery bypass grafting	1	1		
Bleeding complications	9	6		
Minor bleeding	8	5		
Major bleeding	1	1		
Intracranial bleeding	0	0		
Final TIMI flow grade			NS	
0-1	4 (4)	5 (5)		
2	4 (4)	3 (3)		

3 97 (92) 97 (92)

Table 29: Dudek et al. 2010³⁵

s 30 (3):347- 353, 2010.			necessay to maintain an ACT optimal level 200–250 seconds PCI performed in all patients. Stent insertion in 86.4% patients.	optimal level 200– 250 seconds PCI performed in all patients. Stent insertion in 96.3% patients.	
		Early group (n = 24)		Later group (n = 27)	
Age, years		62±9		61±9	
Male sex %		75		81.5	
BMI (kg/m²)		27±3		27±4	
Diabetes mellit	cus %	16.7		11.1	
Arterial hypert	ension %	45.8		59.3	
Hyperlipidimia		41.7		33.3	
Smoking %				40.1	
SBP (mmHg)		143±27		138±25	
DBP (mmHg)		86±16		85±14	
Heart rate (bea	ats per min)	79±21		76±18	
Killip class I %		87.5		92.6	
Killip class II %		12.5		7.4	
Time from chest pain onset to clopidogrel administration (min)		165±88		189±99	
Time from clopidogrel administration to first device/balloon inflation (min)		93±27		106±31	
Time from chest pain onset to first device/balloon inflation (min)		252±95		285±98	
Time from chest pain onset to abciximab administration (min)		173±104		262±97	

Time from abciximab administration to first device/balloon inflation (min)	87±31	21±31				
Data are presented as mean±SD or %	Data are presented as mean±SD or %					
Effect Size / Outcomes	Effect Size / Outcomes					
	Early group (n = 24)	Later group (n = 27)				
30 days						
Death	1 (cardiac)	0				
Reinfarction	0	0				
Repeated revascularisation	0	0				
Intracranial bleeding	0	0				
Major bleeding	1	1				
Minor bleeding	1	1				
Total bleeding complications	12.5%	14.8%				
TIMI flow						
3	91.7%	88.9%				
2	0%	11.1%				
0 and 1	8.3%	0%				
Hospital stay Not reported						

Table 30: ASSENT-4 114

primary outcome, with a level of significance of 5% and 85% power using a χ2 test was determined to be 2000 per group. ITT analysis: No. Available case analysis	characteristics see below	the intervention. Glycoprotein was not allowed in the facilitated group except for bailouts but was allowed at the discretion of the investigator in the PPCI group. Instigators were encouraged to do coronary intervention on the culprit lesion but it was left to their decision. If stents were use a loading dose of 300mg clopidogrel was given followed by a maintenance dose of 75 mg PCI performed in 719 patients. Stent insertion in 667 patients.	Instigators were encouraged to do coronary intervention on the culprit lesion but it was left to their decision. If stents were use a loading dose of 300mg clopidogrel was given followed by a maintenance dose of 75 mg PCI performed in 762 patients. Stent insertion in 714 patients.		
		fPPCI (n = 829)	Standard PP	CI (n = 838)	
Age, years		61 ± 12.2	60 ± 12		
Age >75 years		98/829 (12)	106/837 (13)		
Women		193/829 (23)	189/838 (23)		
Weight (kg)		77.9 ± 14.7	77.7 ± 14.9		
Height (cm)		170.3 ± 8.2	169.8 ± 8.4		

Killip class		
1	753/829 (91)	773/834 (93)
11/111	66/829 (8)	53/834 (6)
IV	10/829 (1)	8/834 (1)
Congestive heart failure at randomisation	42/806 (5)	41/814 (5)
Heart rate (bpm)	74.3 ± 16.9	76.1 ± 17.1
Systolic blood pressure	133.7 ± 24.2	133.7 ± 22.2
Infarct location		
Anterior	403/828 (49)	389/837 (46)
Inferior	418/828 (50)	429/837 (51)
Other	7/828 (1)	19/837 (2)
Previous infarction	108/822 (13)	90/834 (11)
Previous congestive heart failure	7/824 (1)	13/834 (2)
Previous PCI	70/819 (9)	68/829 (8)
Previous coronary artery bypass graft	18/825 (2)	16/833 (2)
Hypertension	391/829 (47)	391/837 (47)
Diabetes	144/828 (17)	131/836 (16)
TIMI		
0	196/812 (24)	512/821 (62)
1	83/812 (10)	70/821 (9)
2	172/812 (21)	107/821 (13)
3	353/812 (43)	124/821 (15)
Not assessable	8/812 (1)	8/821 (1)
Symptom onset to randomisation (min) (median, Q1, Q3)	140 (90, 210)	135 (91, 210)
Symptom onset to unfractionated heparin bolus (min) (median, Q1, Q3)	150 (101, 220)	145 (100, 215)
Symptom onset to tenecteplase (min) (median, Q1, Q3)	153 (105, 225)	
Symptom onset to first balloon (min) (median, Q1, Q3)	263 (213, 339)	255 (200, 335)
Randomisation to first balloon (min) (median, Q1, Q3)	115 (94, 150)	107 (85, 140)
Tenecteplase to first balloon (min) (median, Q1, Q3)	104 (82, 135)	-

Effect Size / Outcomes			
After PCI / in hospital	fPPCI (n = 829 before and n = 719 after PCI)	Standard PPCI (n = 838 before and n = 763 after PCI)	Details
TIMI			
0	15/719 (2%)	13/763 (2%)	
1	15/719 (2%)	4/763 (1%)	
2	55/719 (8%)	68/763 (9%)	
3	631/719 (88%)	677/763 (89%)	
Not assessable	3/719 (<1%)	1/763 (<1%)	
Stroke			
Intracranial haemorrhage	8 (1%)	0	
Primary ischaemic stroke	5 (0.6%)	0	
Unclassified stroke	2 (0.2%)	0	
Total	15 (1.8%)	0	
In-hospital bleeding complications			
Major	46 (5.6%)	37 (4.4%)	
Minor	210 (25.3%)	159 (19%)	
Blood transufsions	48 (6.2%)	33 (4.2%)	
90 days			
Death	55/823 (7%)	41/831 (5%)	fPPCI: Reinfarction (4); cardiogenic shock (22); arrhythmia or sudden death (1); asystole or cardiac arrest (6); cardiac rupture or electromechanical dissociation (8); stroke of intracranial haemorrhage (8); other cardiac event (1); other non-cardiac event (5) Standard PPCI: Reinfarction (4); cardiogenic shock (17); arrhythmia or sudden death (3); asystole or cardiac arrest (5); cardiac rupture or electromechanical dissociation (5); other
	07/007/400/	75 (040 (00))	cardiac event (3); other non-cardiac event (3)
Congestive heart failure	97/807 (12%)	75/818 (9%)	
Shock	51/807 (6%)	39/817 (5%)	
Reinfarction	49/805 (6%)	30/820 (4%)	

Repeat target vessel revascularisation	53/805 (7%)
Rehospitalisation for congestive heart failure	15/807 (2%)
Rehospitalisation for shock	0/807 (0%)
Rehospitalisation for other cardiac reasons	83/806 (10%)
Stroke	
Intracranial haemorrhage	1 /829 (0.1%)
Primary ischaemic stroke	4 /829 (0.5%)
Unclassified stroke	2 /829 (0.2%)
Total	7/829

28/818 (3%)	
11/818 (1%)	
1/817 (<1%)	
90/819 (11%)	
1 /838 (0.1%) 0/838 0/838	
1/838	

Table 31: INTAMI-pilot trial 121

Study type	Number of patients	Patient characteristics	Intervention	Comparison	Outcome measures	Source of funding
Study type Design: RCT (prospective) Enrolment: October 2002- April 2004 Setting	n = 102 Drop-outs: 1 from the early group (1 DCM and 1 myocarditi	Inclusion criteria: >18 years old presented with an acute STEMI defined with angina or equivalent symptoms >30 minutes ST elevation >2 leads (>2 mm preccordial lead, < 1 mm limb	Early group (n = 53) Early eptifibatide, heparin and aspirin All patients received aspirin 50 mg i.v. and	Later group (n = 49) Heparin, aspirin and optional eptifibatide All patients received aspirin 50 mg i.v. and heparin 5.000 U i.v.	Outcome measures TIMI flow Death Reinfarction Repeat target vessel	Grant from ESSEX Pharma GbH, Munich, Germany
Hospital (3 Germany hospitals). Randomisation:	s). 2 from the later group (both myocarditi s)	mm precordial lead in posterior MI new or presumed new LBBB PCI planned	immediately followed but infusion 1.000 U/h – target aPTT 50–70 s.	followed but infusion 1.000 U/h – target aPTT 50–70 s.	revascularisation PCI different vessel	
Yes. Randomly stratified by centre in blocks of 10.		Exclusion criteria: Fibrinolytic therapy with 24 hours before randomisation oral anticoagulation with an	patients received a double bolus of 180 microgram/kg (10 minute interval)	and PCI with possible stent implantation were done according to the local	bypass Emergency CABG	
Allocation concealment: Yes: randomisation done in blinded		platelets < 100000 or known haemorrhagic diathesis stroke or TIA within 30 days evidence of an active	followed by infusion of 2.0 microgram/kg/minute > 12–24 hours. Catheter evaluation and PCI with possible stent implantation were done according to the local guidelines but within 3 hours after administration. The continuation of	3 hours after administration. The continuation of unfractionated heparin post	Severe bleeding complication	
Blinding: No. The trial was an open comparison.		gastrointestinal or urogenital bleeding major surgery within 6 weeks history or allergic reaction to eptifibatide severe renal or hepatic insufficiency		angiography or procedure was discouraged but left to the discretion of the investigator. Clopidogrel was started after PCI with stent, with a loading	Thrombocytopenia < 100.000 Length of follow-up: 60 minutes after PCI	
	Design: RCT (prospective) Enrolment: October 2002- April 2004 Setting Hospital (3 Germany hospitals). Randomisation: Yes. Randomly stratified by centre in blocks of 10. Allocation concealment: Yes: randomisation done in blinded envelopes. Blinding: No. The trial was an open	Design: RCT (prospective) Enrolment: October 2002- April 2004 Setting Hospital (3 Germany hospitals). Yes. Randomly stratified by centre in blocks of 10. Allocation concealment: Yes: randomisation done in blinded envelopes. Blinding: No. The trial was an open	Study typepatientsPatient characteristicsDesign:n = 102Inclusion criteria:RCT (prospective)>18 years oldDrop-outs:presented with an acuteEnrolment:1 from theSTEMI defined with angina or equivalent symptoms >30 minutesOctober 2002- April 2004group (1 DCM and 1 DCM and 1 ST elevation >2 leads (>2 mm precordial lead, < 1 mm limb lead) or ST depression > 1 the later group posterior MI (both myocarditi posterior MI new or presumed new LBBB PCI plannedRandomisation:yes. Randomly stratified by centre in blocks of 10.Exclusion criteria: Fibrinolytic therapy with 24 hours before randomisation oral anticoagulation with an INR >2 platelets < 100000 or known haemorrhagic diathesis stroke or TIA within 30 days evidence of an active gastrointestinal or urogenital bleeding	Design:	Design:	Design:

acute myocardial infarction (INTAMI) pilot trial. Eur.Heart J. 26 (19):1971- 1977, 2005.	calculation: Not reported ITT analysis: No. Available case analysis	angiography severe concomitant disease with life expectancy <1 year Demographics and baseline characteristics see below	procedure was discouraged but left to the discretion of the investigator. Clopidogrel was started after PCI with stent, with a loading dose of 300 mg and maintained with a dose of 75 mg daily for at least 30 days. PCI performed in 46 (87%) patients. Stent insertion in 73% patients.	maintained with a dose of 75 mg daily for at least 30 days. PCI performed in 46 (94%) patients. Eptifibatide was given in 42 (86%) of patients immediately before PCI or during PCI (immediately before procedure in 30 patients and during procedure in 12 patients) Stent insertion in 74% patients.	30 days	
		F (F2)		Later (n = 49)		
A ====		Early (n = 53)		61±11		
Age, years		61±13		66		
Male sex (%)		79		28±6		
BMI		29±7		74±17		
Pulse (b.p.m.)	/ 11.	80±21				
-	pressure (mmHg)	133±22		136±26		
	d pressure (mmHg)	77±13		79±17		
Anterior infar		43		39		
	sitive on admission (%)	21		28		
Killip class >1	(%)	16		14		
Smoker (%)		51		41		
Hyperlipidemi		55		67		
Diabetes melli	itus (%)	15		22		
Prior MI (%)		12		16		

Prior PCI (%)	4		18
Prior CABG (%)	0		2
Prior angina (%)	19		25
TIMI			
0/1 (%)	58.4		67.4
2 (%)	7.6		22.4
3 (%)	34		10.2
60 minutes after procedure	Early (n = 53)	Later (n = 49)	p value
TIMI			
0/1 (%)	7.7	2.1	
2 (%)	5.8	14.9	
3 (%)	86.5	83	
Death (n, %)	2 (3.8)	2 (4.1)	0.9
Reinfarction (n, %)	3 (5.7)	0	0.09
Repeat target vessel revascularisation (n, %)	2 (3.8)	1 (2)	0.6
PCI different vessel (n, %)	4 (7.6)	4 (8)	0.9
Coronary artery bypass (n, %)	3 (5.7)	5 (10.2)	0.4
Emergency CABG (n, %)	1/3 (1.9)	2/5 (4)	
Stroke (n, %)	0	0	
Severe bleeding complication (n, %)	2 (3.8)	2 (4.1)	0.9
Thrombocytopenia < 100.000 (n, %)	1 (2)	1 (2)	0.3

Table 32: Lee et al. 2003⁷⁰

Reference	Study type	Patient characteristics	Intervention	Comparison	Outcome measures	Source of fundin
Lee. et al.	Design:	Number of patients:	Early tirofiban, in the ER,	Later tirofiban,	1°	Grant
2003	RCT (prospective)	n = 100	before primary angioplasty	in the cath lab, after	Initial TIMI grade	from
		Drop-outs: None		diagnostic angioplasty, just	flow, corrected	Merck
Adjunctive	Enrolment:	·	(n = 50)	before primary angioplasty	TIMI frame	Co.
olatelet	July 1999 - December	Inclusion criteria:			counts, TIMI grade	
glycoprotein	2001	Clinical symptoms of AMI (acute	Notes:	(n = 50)	myocardial perfusion.	
llb/Illa		myocardial infarction) with the	(3 patients underwent PPCI		periusion.	
receptor inhibition	Setting	initial onset of chest pain in the	alone, 47received at least 1	Notes:	Other:	
with	Hospital	past 12 hours	stent after initial angioplasty)	(3 patients underwent PPCI	30-days major	
tirofiban		Deemed to be suitable	aligiopiasty)	alone, 47received at least 1	cardiac adverse	
before	Randomisation:	candidates for percutaneous revascularisation	Tirofiban dose: bolus 10	stent after initial	events.	
primary	Yes. Computerised	An ECG that demonstrated	microgram/kg over 3	angioplasty)		
angioplasty	randomisation	≥0.1mV ST-segment elevation	minutes, followed by 0.15		Length of follow-	
improves angiographic		in 2 or more contiguous leads	microgram/kg/minute per	Tirofiban dose: bolus 10 microgram/kg over 3	up	
outcomes.	Allocation	or documented new left	24 hours.	minutes, followed by 0.15	30 days	
Results of	concealment:	bundle-branch block.		microgram/kg/minute per		
the	Yes. Blinded		Other medications:	24 hours.		
TIrofiban	envelopes.	Exclusion criteria:	heparin bolus 70U/kg			
Given in the		Cardiogenic shock	followed by 5 U/kg/h.	Other medications:		
Emergency	Blinding:	Use of an intra-aortic balloon		heparin bolus 100U/kg with		
Room before	No. open-label.	pump (IABP)	If coronary stent was	a maintenance dose of 10		
Primary		Known bleeding diathesis	placed> clopidrogel (300mg orally, following by	U/kg/h.		
Angioplasty	Sample size calculation:		75 mg per day for at least 28	If coronary stent was		
(TIGER-PA)	study underpowered		days) or Ticlopidine (500 mg	placed>clopidrogel		
pilot trial	to show a significant	Demographics and baseline	orally followed by 250mg	(300mg orally, following by		
	difference in clinical	characteristics	twice per day for at least 28	75 mg per day for at least 28		
Circulation.	end points.	see below	days)	days) or Ticlopidine (500 mg		
2003;				orally followed by 250mg		
107:1497-	ITT analysis:			twice per day for at least 28		

Not mention unclear.	ned /				days).			
Demographics and baseline	characteristics							
		Early tirofiban (n = 50)			Late tirofiban (n = 50)	
Age, years		63.5±12.6				66.4±14.3		
Male gender, %		60				64		
Diabetes, %		24				24		
Hypertension, %		36				40		
Hyperlipidimia, %		32				32		
Previous CAD (coronary arto	ery disease), %	12				10		
Duration of CP (chest pain),	h	3.0±2.0				3.0±1.8		
Door to tirofiban, min		55.7±18.0				81.8±18.0		
Door to balloon, min		88.9±20.7		82.7±20.0				
Definitions of end points Bleeding: TIMI definition. Effect Size / Outcomes								
30 days	Early tirofib	an (n = 50)		Later tirofiban (n = 50	0)	р	value	
TIMI-defined bleeding, %								
Minor	10			6		N	IS	
Major	2			2		N	NS .	
30-day outcomes, %								
Death	2			2		N	NS .	
Re-MI (myocardial infarction)	0			2		N	IS	
Re-hospitalisation	4			6		N	IS	
Urgent TVR (target vessel revascularisation), %	0			2		N	NS	
NS=not significant								

Table 33: Emre 2006⁴³

Reference	Study type	Patient characteristics	Intervention	Comparison	Outcome measures	Source of funding
Emre et al.	Design:	Number of patients:	Early tirofiban	later tirofiban	1°	Not
2006	RCT	n = 66	(emergency room) Stenting	(cath lab) Stenting	Degree of MI salvage.	stated
Impact of	Enrolment:	Drop-outs: None	(n = 32)			
early	Not stated			(n = 34)	Other:	
tirofiban		Inclusion criteria:	Tirofiban bolus (10		30 day major	
administrati on on	Setting	Chest pain > 30 minutes	microgram/kg) followed by 0.15	Tirofiban bolus	adverse cardiac	
myocardial salvage in	Hospital. Consecutive patients	Presentation < 6 hours after the onset of symptoms	microgram/kg for 24 hours)	(10 microgram/kg)	events.	
patients	Randomisation:	ST-segment elevation of ≥1mm in	nours)	followed by 0.15 microgram/kg for	Length of follow-	
with acute myocardial	Yes, but details not given	≥2 contiguous leads.		24 hours)	up: 30 days	
infarction undergoing infarct- related	Allocation concealment: Not mentioned / unclear.	Exclusion criteria: Previous administration of fibrinolytic agents	Note: All patients received bolus 5000 U unfractionated			
artery stenting.	Blinding: Not mentioned / unclear.	Previous MI Previous percutaneous coronary intervention artery bypass graft	heparin during the procedure. And a loading dose of			
Cardiology 2006; 106:264-269	Sample size calculation: the study was powered to detect a	surgery Known bleeding diathesis or allergy to study drugs	clopidogrel 300mg and aspirin 325mg.			
100.204-203	20% difference in the myocardial salvage index (α 0.05 and β 0.80), with a minimum of 64 patients (32/arm). Study is underpowered to	Major surgery within 15 days Active bleeding Cardiogenic shock	Procedural success achieved in all patients (and all			
	show a significant difference.		patients given stents).			
	ITT analysis:	Demographics and baseline characteristics				
	Not mentioned / unclear.	see below				

Demographics and baseline characteristi	cs		
	Early tirofiban (n = 32)	Later tirofiban (n = 34)	p value
Age, years	58±10	59±12	ns
Male gender, %	81	82	ns
Anterior MI, %	13 (41)	15 (44)	ns
LVEF (Left ventricular ejection fraction)	48±11	49±11	ns
Symptom onset to presentation, min	118±52	122±48	ns
Smoking %	14 (44)	16 (47)	ns
Diabetes, %	7 (22)	8 (24)	ns
Hypercholesterolemia, %	15 (47)	15 (44)	ns
Systemic hypertension, %	15 (50)	17 (50)	ns
Door to tirofiban, min	18±4	52±10	0.004
Door to balloon, min	43±12	53±9	ns
Peak creatine kinase level, U/I, %	2,268±1,452	2,480±1,590	ns
Time to peak creating kinase level, h	8.6±5.6	8.8±4.4	ns
Initial TIMI, flow grade, %			
0	17 (53)	22 (64)	ns
1	2 (6)	3 (9)	ns
2	3 (10)	5 (15)	ns
3	10 (31)	4 (12)	0.04
30 day outcomes	Early tirofiban (n = 32)	Later tirofiban (n = 34)	p value
Death	0	0	ns
Recurrent MI	0	1 (3)	ns
Rehospitalisation	2 (6)	4 (12)	ns
TIMI-defined bleeding			
Minor	3 (10)	2 (6)	ns
Major	0	0	ns
Thrombocytopenia	1 (3)	0	ns

Table 34: Shen et al. 2008¹⁰⁰

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Outcome measures	Source of funding
Shen et al. 2008	Design: RCT , prospective	n = 172	INCLUSION CRITERIA Chest pain >30min Presented within 12 hours	early tirofiban (emergency room, upstream group) PPCI	later tirofiban (in cath lab, downstream group)	1° Occurrence rate of major	Grant from the Shanghai
Clinical benefits of adjunctive tirofiban therapy in patients with acute ST-segment elevation myocardial infarction undergoing primary percutaneous coronary intervention.	Enrolment: January 2005 – June 2006. Consecutive patients Setting Hospital. Randomisation: Yes, computer- generated random allocation system	Drop-outs: none	of symptoms New ST-segment elevation in at least 2 contiguous leads on electrocardiogram with the cut off points ≥0.2mV, with or without elevation of cardiac enzymes. EXCLUSION CRITERIA Cardiogenic shock Known bleeding diathesis	(n = 57) Tirofiban bolus (10 microgram/kg) followed by 0.15 microgram/kg/minute for 36 hours) Note: All patients received before procedure: aspirin 300 mg; clopidogrel 450 mg; unfractionated heparin 100 U/kg through the femoral arterial access sheaths.	PPCI (n = 57) Tirofiban bolus (10 microgram/kg) followed by 0.15 microgram/kg/minut e for 36 hours) Control group: PPCI alone (n = 58)	adverse cardiac events. Length of follow-up 30 days and 6 months	Science an Technolog Committee
Coronary Artery Disease 2008, 19; 271-277	Allocation concealment: Not mentioned / unclear. Blinding: Not mentioned / unclear. Sample size calculation: Not mentioned / unclear.		Demographics and baseline characteristics see below	All patients received after procedure: Clopidogrel (75 mg/day) for at least 9–12 months; aspirin (100 mg/day) infinitively; subcutaneous low molecular weight heparin for 7 days after the procedure. Procedural success achieved in all patients, and 99% stents - 1 person in control			

ITT analysis: Not mentioned unclear.	/	(intra-aortic counterpuls	ot have stent balloon aion because of mic instability).	
Demographics and baseline cha	aracteristics			
	Control (n = 58)	Early tirofiban	Later tirofiban	р
		upstream (n = 57)	Downstream (n = 57)	
Age, years	65.4±14.6	68.0±14.3	65.3±11.6	0.48
Male gender, %	86.2	75.4	82.5	0.32
Hypertension, %	65.5	73.3	68.4	0.63
Diabetes mellitus, %	27.2	31.6	21.1	0.44
Hypercholesterolemia, %	31.0	36.8	22.8	0.26
Current smoker, %	55.2	42.1	47.4	0.37
Previous PCI, %	5.2	5.3	7	0.89
Symptom-to-catheterisation time, h	6.5±2.7	6.7±3.6	5.6±2.9	0.17
Door-to-balloon time, min	75.3±25.9	79.9±26.7	68.9±24.3	0.08
Acute anterior MI, %	51.7	50.9	61.4	0.46
Acute inferior MI, %	43.1	40.4	33.3	0.54
Acute anterior+inferior MI, %	5.2	8.8	7.0	0.75
Sum of ST-segment elevation before procedure, mm	13.55±5.50	11.76±5.81	12.67±5.05	0.22
CK-MB before procedure, ng/ml	49.6±44.0	41.2±44.8	53.1±38.8	0.32
Tnl before procedure, ng/ml	10.8±9.2	9.8±8.5	10.8±16.9	0.89
Angiographic features				
No of implant stent	1.1±0.5	1.3±0.7	1.2±0.6	0.20
Effect Size				
Outcomes				
	Control	Early tirofiban	Later tirofiban	р
	(n = 58)	upstream (n = 57)	Downstream (n = 57)	

STEMI Clinical evidence tables

Death, %	5.2	3.5	5.3	0.88
nonfatal MI, %	6.9	0	0	0.02
TVR, %	3.4	0	0	0.14
LVEF	0.47±0.08	0.51±0.07	0.50±0.07	0.0008
Death, %	5.2	3.5	5.3	0.88
nonfatal MI, %	6.9	1.8	1.8	0.22
TVR, %	5.2	1.8	1.8	0.45
LVEF	0.54±0.07	0.59±0.06	0.57±0.07	<0.001
Minor bleeding	1.7	3.5	1.8	-
Major bleeding	3.4	5.3	8.8	-
Hospital stay, mean days (SD)	14.5 (6.5)	10.6 (5.4)	12.6 (4.7)	-
LVEF: left ventricular ejection fr	action. TVR: target vesse	l revascularisation		

Table 35: El Khoury et al. 2010³⁸

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow- up	Outcome measures	Source of funding
El Khoury. et al. 2010 Prehospital high-dose tirofiban in patients undergoing primary percutaneous intervention. The AGIR-2 study. Archives of cardiovascula r disease (2010) 103, 285-292	Design: RCT (multicentre, prospective) Enrolment: July 2007-July 2008. Out-of hospital patients managed by mobile intensive care units (MICU) staffed by a physicians. Setting MICU and Hospital Randomisation: Yes. Computergenerated random sequence. Allocation concealment: Yes. Scratch cards.	(n = 320) Drop-outs: 1	INCLUSION CRITERIA Presented within 12 hours after onset of symptoms of MI, ie: Characteristic pain lasting for at least 30 min Not responsive to nitrates Electrocardiographic ST-segment elevation ≥0.2mV in 2 or more contiguous precordial leads or 0.1mV for limb leads. EXCLUSION CRITERIA Haemorrhagic diathesis Pregnant Any allergy or contraindication to heparin, aspirin or tirofiban Suffered from severe renal or hepatic insufficiency Had major surgery within the past month Had any sign of cerebral ischaemic disease for <1 month or non-ischaemic disease whatever its date Received oral anticoagulant treatment, a fibrinolytic or a GP IIb/IIIa antagonist within the past 7 days Uncontrolled hypertension, severe conduction disorder or cardiogenic shock	Early tirofiban, in the ambulance (pre-hospital) PPCI (n = 164) Notes: Complete ST-segment resolution 60 minutes after the start of intervention: 52.6%. Stent placed in 112 (69.1%) of patients. Tirofiban dose: Bolus 25 microgram/kg in 3 min, followed by infusion of 0.15 microgram/kg per minute for	Later tirofiban, in the cath lab. PPCI (n = 156) Notes: Complete ST- segment resolution 60 minutes after the start of intervention: 55.4%. Stent placed in 113 (72.4%) of patients. Tirofiban dose: Bolus 25 microgram/k g in 3 min, followed by infusion of	In- hospital	In-hospital mortality Major bleeding In-stent thrombosis Stroke	Sponsor: Hospice Civil de Lyon, France. Merck Sharp Dhome and Iroko pharmaceut s provided the tirofiban free of charg to the sponsor.

Blinding: No. open-label. Sample size calculation: the initial sample size calculation was 300 patients with a 5%alpha risk and a 80% power to detect a 16% difference in the primary end point. The higher than expected rate of patency in the later tirofiban group lowered the power of the study to detect a difference. ITT analysis: Yes	If duration to transfer to the (entrance to the cath lab) exhour. Demographics and baseline characteristics see below		o.15 microgram/k g per minute for 18–24 hours. Other medications: clopidogrel 600mg		
	later tirofiban (cath lab) (n = 156)	early tirofiban (pre-hos (n = 164)	oital)	р	
Male gender	124 (79.5)	123 (75.0)		0.34	
Diabetes	17 (10.9)	18 (11.0)		0.98	
Hypertension	62 (39.7)	74 (45.1)		0.33	
Current smoker	56 (35.9)	61 (37.2)		0.81	
Dyslipidaemia	67 (43.0)	55 (33.5)		0.08	
Anterior MI	68 (43.6)	84 (51.2)		0.17	

Previous MI	24 (15.4)	15 (9.1)	0.09
Previous coronary artery bypass graft	6 (3.8)	2 (1.2)	0.13
Previous percutaneous coronary intervention	23 (14.7)	13 (7.9)	0.05
Killip class ≥2	13 (8.3)	19 (11.6)	0.33
Heart rate (beats/min)	75.4 ± 16.2	77.2 ± 22.2	0.42
Systolic arterial pressure (mmHg)	139 ± 28	141 ± 25	0.80
Cumulative ST-deviation on diagnostic electrocardiogram (mm)	12 ± 9	13 ± 8	0.25
Treatment delay, median [25–75%] (min)			
Onset of chest pain to MICU	98 [50–200]	104 [56–233]	0.30
MICU to cath lab	54 [45–69]	61 [54–74]	0.0002
Cath lab to first angiography	26 [15–35]	21 [15–35]	0.007
MICU to first angiography	83 [70–96]	85 [72–100]	0.46

Data presented as mean±SD or number (%) of patients otherwise stated. MICU: mobile intensive care unit arrival on site of intervention; Cath lab: admission to the catheterisation laboratory.

Definitions of end points

TIMI flow grade 2–3 of the infarct-related vessel at initial angiography.

ST-segment resolution 1 hour after percutaneous coronary intervention and peak serum troponin I concentration

Effect Size / Outcomes

	later tirofiban (cath lab) (n = 156)	early tirofiban (pre-hospital) (n = 164)	p
In-hospital mortality	5 (3.2)	9 (5.5)	0.26
Major bleeding	2 (1.3)	6 (3.7)	0.28
Stroke	1 (0.6)	2 (1.2)	Not reported
In-stent thrombosis	3 (1.9)	1 (0.6)	0.36
Data presented as number (%) of patients			

Table 36: Ohlmann et al. 2012 88

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow- up	Outcome measures	Source of funding
P Ohlmann, P Reydel, L Jacquemin, F Adnet, O Wolf, J Claude Bartier, A Weiss,et al. Prehospital Abciximab in ST-Segment Elevation Myocardial Infarction. Circulation: Cardiovascul ar Intervention s, 2012.	Design: RCT (11 centres in France serviced by 7 emergency ambulance services Enrolment: Jan 2005 – June 2009 Setting Ambulance to hospital Randomisation : Poor/unclear: stratified by centres, done according to manufacturer of treatment kits which had to be used consecutively	n = 256 Drop- outs/missing patients/inel igible: For clinical outcomes: n = 5 (2%) at 1 month and another n = 5 (4% cumulative)a t 6 months OVERALL acceptable losses (<20%)	INCLUSION CRITERIA: patients eligible for PPCI STEMI symptoms <6 hours ST-segment elevation 2 mm in V1 to V3 leads, or 1mm in remaining leads EXCLUSION CRITERIA Contraindications to anticoagulation increased risk of bleeding oral anticoagulation known hpersensitivity to study drugs pregnancy or breastfeeding presence of IV conduction abnormality (complete left or right bundle branch block) Demographics and baseline characteristics see below	EARLY abciximab (in ambulance) (n = 127) Abciximab – 0.25 mg/kg IV bolus additional bolus given if activated clotting time was < 150 seconds or 150–199 seconds NOTE: Before catheteris aspirin IV (250 m UFH (heparin) IV maximum 3000 U	g) (40 IU/kg,	Inhospital 30 days / 1 month 6 months	Primary: STR. Secondary: composite MACEs; death; nonfatal MI; coronary revasculari sation at 30 days and 6 months	Eli Lilly
	Allocation concealment: Not mentioned Blinding:			Post-PCI – 0.125 microgram abciximab infusio clopidogrel (300- dose) at discretio	on for 12 hours or 600-mg loading			

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow- up	Outcome measures	Source of funding
	Double blind							
				Stent type used discretion of inte				
	Sample size calculation:			discretion of file	erventionalist.			
	Powered study:			PCI – 93% (fPPCI) and 98% (PPCI)			
	83% power using at				,			
	least 240 patients for complete STR			Stents used – 10	0% in both groups	;		
	for complete 51K							
	ITT analysis: Yes							
Demographi	ics and baseline characte	eristics						
%		EARLY ab	ciximab	Placebo / later abciximal	b	p value		
		(in ambu	lance) (n = 127)	(in hospital cath lab) (n =	: 129)			
Age, yrs; me	ean (SD)	56.0 (11.9	9)	57.7 (12.7)		0.27		
Male gender		79		82		0.63		
Diabetes		9		11		0.84		
Hypertensio	n	38		37		1.0		
Current smo	king	51		53	(0.90		
Hypercholes	terolemia	38		39	(0.90		
Previous MI		5		5		1.0		
Previous CA	3G	0		0		1.0		
Previous PCI		4.7		4.7		1.0		
Previous stro	oke	1		0	(0.5		
Killip Class					(0.80		
1		92		91				
2		6		7				
3		0		2				
4		2		1				

TIMI flow (%)

Reference	Study type	Number of patients	Patient c	haracteristics	Interv	ention (Comparison	Length of follow- up	Outcome measures	Source of funding
0		49/109	(45)		67/117 (57.3)		0.	078		
1		9/109 (8.3)		9/117 (7.7)					
2		26/109	(23.9)		20/117 (17.1)					
3		25/109	(53.2)		21/117 (17.9)					
0–1		58/109	(53.2)		76/117 (65)		0.	08		
2–3		51/109	(46.8)		41/117 (35)					
Critical time (IQR)	intervals, minutes;	median								
Symptom on	set to balloon	190 (14	2–239)		198 (165–282)		0.	87		
Door (ambul	ance arrival) to bal	loon 105 (91	-125)		103 (90–122)		0.	91		
Arrival to ang	giography	25 (16–	35)		27 (20–35)		0.	14		
Bolus 1 to ba	illoon	7 (60–8	6)		74 (64–89)		0.	31		
Bolus 2 to ba	illoon	7 (4–12	.)		6 (3–14)		0.	92		
Effect Size		·	•		, ,					
Angioplasty	and808% (parly an	d later groups resp	actively) neor	ole had PCI and 100	% in each arm had	ctents				
Outcomes	andososo (earry and	a later groups resp	ectively) peop	ne nau i ci anu 100	70 III Cacii ai III iiau	3101113				
Outcomes	In-hospital (after procedure)		p value	At 30 days		p value	At 6 montl	ns	ру	value
N (%)	EARLY abciximab (in ambulance) (n = 127)	Placebo / later abciximab (in hospital cath lab) (n = 129)		EARLY abciximab (in ambulance) (n = 127)	Placebo / later abciximab (in hospital cath lab) (n = 129)		EARLY abciximab (in ambulance (n = 127)	Placeb later abcixim) (in hos cath la (n = 12	nab pital b)	
TIMI flow										
0	1/109 (0.9)	2/116 (1.7)	0.18	-	-	-	-	-	-	

		Number of						Length of follow-	Outcome	Source of
Reference	Study type	patients	Patien	characteristics	lı lı	ntervention	Comparison	up	measures	funding
2	7/109 (6.4)	11/116 (9.5)		-	-	-	-	-	-	
3	100/109	100/116		-	-	-	-	-	-	
0-1	(91.7)	(86.2)	0.45	-	-	-	-	-	-	
2–3	2/109 (1.8)	5/116 (4.3)		-	-	-	-	-	-	
	107/109 (98.2)	111/116 (95.7)								
Death	2 (2)	1 (1)	-	2 (2)	1 (1)	-	2 (2)	1 (1)	-	
MI	2 (2)	2 (2)	-	3 (2)	2 (2)	-	3 (2)	2 (2)	-	
TVR	2 (2)	4 (3)	-	2 (2)	4 (3)	-	9 (7)	11 (9)	-	
Cardiac proc	edures									
%		EARLY	abciximab		Placebo /	later abcixima	b	p value		
		(in am	bulance) (n	= 127)	(in hospit	al cath lab) (n =	= 129)			
Coronary ang	giogram	100%			-					
PCI		91%			88%					
Stents		100%								
Hospital stay										
No details giv	ven .									

Table 37: LIU 2012B

Reference	Study type	Patient characteristics	Intervention	Comparison	Outcome measures	Source of funding
Liu J, Fu X-H, Xue L, Wu W-L, Gu X-S, Li S-Q. Equilibrium	Xue L, Wu RCT (1 hospital centre in China) Li S-Q. Equilibrium radionuclide angiography for evaluating the effect of facilitated RCT (1 hospital centre in China) Enrolment September 2006 to September 2009 Randomisation Not described	Number of patients: n = 152 Drop-outs: n = 9 (fPPCI n=4, PCI n=5)	fPPCI with 50 mg reteplase before PCI Pre-PCI: All patients	PPCI	Congestive heart failure; TVR; mortality, cardiac mortality; IC haemorrhage;	Not stated
angiography for evaluating the effect of facilitated		Inclusion criteria Patients < 70 years with first AMI Onset within 6 hours ST elevation on contiguous ECG Transfer to PCI centre within 90 minutes	received aspirin 300mg and clopidogrel 300 mg as they were enrolled.		Length of follow-up: 1 week and 6 months	
s coronary intervention on	Allocation concealment Not described	Suitability for percutaneous revascularisation No prior use of t-PA Informed consent	Post-PCI: If stents were used then clopidogrel (75			
synchrony in patients with acute myocardial infarction.	blinded EDNA and ventricular phase analysis were carried out at 1 week and 6 months by 2 blinded independent observers. 2012; 26(4):928- Sample size calculation	Exclusion criteria Cardiogenic shock or severe heart failure Bleeding diathesis or recent stroke within 4 weeks	mg/day) and aspirin were			
Circulation Journal. 2012; 76(4):928- 935.		Recent surgery Previous inter-cranial or spinal surgery Neoplasm Severe hypertension	at discretion of attending physician.			
<i>3</i> 33.	Not described ITT analysis All randomised patients were included in ITT	Acute aortic dissection Contraindication for reperfusion therapy Serious arrhythmia or bundle branch block Demographics and baseline characteristics	PCI/STENTS: 96% / 97% PCI and 94% / 96% STENTS IN PPCI and fPPCI groups			
	comparisons of outcome	sas halau	respectively.			

Demographics and baseline characteristics (d	ata only reported in the patients ana	lysed, not the complete number randomise	ed)
	PPCI n = 71	fPPCI n = 72	p value
Age (years)	55.91 ± 9.26	58.34 ± 11.58	0.262
Male sex	56	60	0.101
Medical history, n (%)			
• Smoking	26(36.62)	23(31.94)	0.822
Diabetes mellitus	17 (23.94)	20 (27.78)	0.872
Pre-MI angina	24 (33.08)	22 (30.56)	0.574
Hypertension	40 (56.34)	44 (61.11)	0.608
Hyperlipidemia	21 (29.58)	24 (33.33)	0.720
Number of diseased vessels n(%)			
• 1	11 (42.8)	11 (44)	
• 2	6 (26.1)	8 (32)	
• 3	6 (26.1)	6 (24.0)	
OUTCOMES Complications and Outcomes of	the 2 PCI Groups; 6 months		
Parameter n (%)	PPCI n = 71	fPPCI n = 72	p value
New or worsening congestive heart failure	9 (12.68)	2 (2.78)	0.028
Recurrent ischaemia	7 (9.86)	2 (2.78)	0.043
Reinfarction	3 (4.23)	1 (1.39)	0.679
Urgent TVR	3 (4.23)	1 (1.39)	0.679
Cardiac death	4 (5.63)	1 (1.39)	0.161
Non-cardiac death	2 (2.82)	0	0.416
Intracranial haemorrhage	0	0	1.000
TIMI-defined bleeding			
• Minor	7 (9.86)	8 (11.11)	0.813
Major	0	0	1.000
•			
Access bleeding Transfusion	1 (1.41)	1 (1.39)	1.000

Table 38: ZORMAN 2002

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Outcome measures	Source of funding
Simona Zorman, Darko Zorman, and Marko Noc. Effects of abciximab pretreatment in patients with acute myocardial infarction undergoing primary angioplasty. Am.J.Cardiol. 90 (5):533-536, 2002.	Design RCT (1 hospital centre in Slovenia) Enrollment: June 1998 to June 2001 Randomisation: Not described (just says randomised) Allocation concealment: Not described Blinding: Not mentioned. Sample size calculation: Not mentioned ITT analysis: Yes as not mentioned and no drop-outs mentioned	n = 163 Drop-outs: n = not mentioned	 Inclusion criteria STEMI Admitted within 12 hours of symptom onset Exclusion criteria Not given Demographics and baseline characteristics see below 	GROUP A: fPPCI with abciximab / or early abciximab (0.25 mg/kg) followed by 12 hours of 0.125 microgram/kg/mi. Given immediately after entering emergency department and the initial aspirin and heparin bolus; before PCI. GROUP B: LATE abciximab (0.25 mg/kg) followed by 12 hours of 0.125 microgram/kg/mi. Given after angiography and before the angioplasty attempt (thus not true faciliatation) Pre-PCI: All patients received aspirin 250–200 (70 U/kg) immediately as arrived in endepartment. During/Post-PCI: 12 hours infusion of abciximab (0.125 microgram/mg/kg). PCI/STENTS: 93% / 100% PCI and 59% / 69% STENT PPCI groups respectively.	mergency	In-hospital: HF Bleeding Death 6 months: Death Length of follow-up: 6 months	Not stated

Demographics and baseline characteri	stics		
	fPPCI n = 56	PPCI n = 51	Late abciximab n = 56
Age, years (SD)	58 ± 13	63 ± 14	63 ± 11
Male sex, %	79	61	73
Medical history, n (%)			
• Smoking	20%	31%	22%
 Diabetes mellitus 	18%	29%	23%
• MI	11%	14%	14%
 Systemic Hypertension 	54%	61%	59%
Hyperlipidemia	55%	43%	52%
Killip class, %			
• 1	86	69	75%
• 2	9	10	16%
• 3	2	6	2%
• 4	3	15	7%
Outcome, n (%)	fPPCI n = 56	PPCI n = 51	Late abciximab n = 56
In-hospital complications			
Heart Failure	4 (7%)	15 (29%)	10 (18%)
Bleeding	16 (29%)	6 (12%)	11 (20%)
Death	0 (0%)	5 (10%)	4 (7%)
6-month follow-up			
Cumulatiive death	0 (0%)	7 (14%)	5 (9%)

G.3 Radial versus femoral arterial access for PPCI

Table 39: Gan et al. 2009⁴⁸

Reference	Study type	Number of patients	Patient characteristics	Interventi on	Comparison	Length of follow- up	Outcome measures	Source o
Gan L, Lib Q, Liuc R, Zhaoc Y, Qiuc J, Liao Y. Effectiveness and feasibility of transradial approaches for primary percutaneous coronary intervention in patients with acute myocardial infarction. Journal of Nanjing Medical University. 2009; 23(4):270-274.	Design RCT; 2 centres in China Enrolment June 2004 to July 2007 Randomisation Not detailed Allocation concealment Not detailed Blinding Patients and investigator s were not masked to treatment allocation. All other outcomes were as reported by the investigators.	n = 195 PPCI done in all patients Drop outs (at 6 month follow-up) Radial n = 9 Femoral n = 14 Crossover 1 patient in radial group required crossover to femoral group due to unsuccessful puncture of radial artery Operator expertise Not stated	Inclusion criteria Patients with STEMI recruited within 12 h of symptom onset Typical chest pain lasting > 30 min and < 12 h, nitrate losing efficacy, ST- segment elevation > 0.1 mV in limb leads or > 0.2 mV in 2 or adjacent chest leads Exclusion for transradial group Negative Allen test (these patients were switched to femoral group) Demographics and baseline characteristics see below Drug therapy All patients received 300 mg aspirin, 300 mg clopidogrel on diagnosis. 30000 IU heparin administered after sheath insertion. Additional heparin during procedure dependent upon patients body mass (100 IU/kg). GP IIb/Illa inhibitors were given dependent based on clinical need. After implantation of drug eluting	Radial access to perform coronary angiograph y and PPCI (if clinically indicated) (n = 90)	Femoral access to perform coronary angiography and PPCI (if clinically indicated) (n = 105)	In- hospital and 6 months	All-cause mortality Reinfarction Repeat revascularisati on CABG Hospital stay Angiographic procedural success access site complications Fluproscopy time (see below for definitions)	Language support from; Editorial Dept of the Journal of Nanjing Medical Universit

No	stents, patients were treated with 1000 IU/kg low molecular heparin twice a day for 5 to 7 days and 150 mg aspirin plus 75 mg aspirin plus 75 mg clopidogrel daily for 12 months			
Demographics and baseline characteristics				
Characteristics	Radial (n = 90)	Femoral (n = 105)		
Male/Female	73/17	84/21		
Age(years), mean(SD)	56.6(12.5)*	52.3(11.9)		
Smoking, n(%)	55(6.1)*	67(63.8)		
Diabetes, n(%)	25(27.8)*	31(29.5)		
Hypertension, n(%)	44(48.9)*	48(45.7)		
Hyperlipidaemia, n(%)	33(36.7)*	37(35.2)		
Prior MI, n(%)	8(8.9)*	11(10.5)		
Onset of symptoms to arrival (h), mean(SD)	4.3(2.1)*	4.8(2.2)		
Single vessel disease, n(%)	24(26.8)*	25(23.8)		
Multivessel disease, n(%)	66(73.3)*	80(76.2)		
Infarct related artery				
Left anterior descending artery, n(%)	47(52.2)*	57(54.3)		
Left circumflex, n(%)	15(16.7)*	19(18.1)		
Right coronary artery, n(%)	28(31.1)*	29(27.6)		
Compared with femoral group, $*p > 0.05$				
Definitions of operational data Cannulation time: time from patient arrival at cath lab to Reperfusion time: time from cannulation to balloon inflatorate procedural time: time from 1 st attempt to puncture.	ition			
Comparison of operation data between radial versus fe	moral groups			
Characteristics	Radial (n = 90)	Femoral (n = 105)		
Success of puncture, n(%)	89(98.9)*	105(100)		

Cannulation time (min), mean(SD)	3.15(1.56)*	2.86(0.97)	
Cannulation-to-balloon-infusion time (min), mean(SD)	18.56(4.37)*	17.75(3.21)	
Total procedural time (min), mean(SD)	29.75(4.38)**	27.89(3.95)	
Glycoprotein II b/II a inhibitor, n(%)	28(31.1)*	36(34.3)	
Final TIMI flow, n(%)			
TIMI 0	0	0	
TIMI 1	1	1	
TIMI 2	2	3	
TIMI 3	87(97.7)*	101(96.2)	
Angiographic procedural success, %	96.7*	96.2	
TIMI; thrombolysis in myocardial infarction, Compared with femoral group, *p > 0.05, **p < 0.05			

Definitions of outcomes

Angiographic procedural success: Residual obstruction < 20%, achieving TIMI flow of at least grade III and no major complications such as death, emergency surgical revascularisation or worsening of patients' clinical condition

Effect Size

Outcome	Radial	Femoral
6 month follow-up complete, n(%)	79(87.8)	88(83.8)
All-cause mortality at 6 months, n(%)	2(2.5)*	3(3.4)
All-cause mortality in-hospital, n(%)	2(2.2)*	3(2.9)
Reinfarction at 6 months, n(%)	1(1.3)*	0
Reinfarction in-hospital, n(%)	0	2(1.9)
CABD at 6 months, n(%)	0	0
CABD in-hospital, n(%)	0	0
Repeat revascularisation at 6 months, n(%)	2(13.3)*	2(9.5)
Access site complications in-hospital, n(%)	2(2.2)**	12(11.4)
Length of hospitalisation (days), mean(SD)	10.56(2.85)**	13.78(3.15)
Compared with femoral group, $*p > 0.05$, $**p < 0.05$		

Table 40: Hou et al. 2010⁵⁶

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow - up	Outcome measures	Source of funding
Hou L, Wei YD, Li WM, Xu YW. Comparative study on transradial versus transfemoral approach for primary percutaneous coronary intervention in Chinese patients with acute myocardial infarction. Saudi Medical Journal. 2010; 31(2):158- 162.	Design: RCT; 1 centre in China Enrolment: August 2005 to Sept 2008 Randomisation Not detailed Allocation concealment: Not detailed Blinding: Patients and investigator s were not masked to treatment allocation. All other outcomes were as reported by the investigators ITT analysis: Yes	n = 200 PPCI done in all patients Drop outs none Crossover 4 patients in radial group required crossover to femoral access (2 patients; severe subclavian artery tortuosity, 2 patients; radial artery tortuosity, artery tortuosity) Operator expertise 3 senior interventional cardiologists who had performed over 200 cases of radial PPCI	INCLUSION CRITERIA: Patients with Acute MI EXCLUSION Negative Allen test Non-palpable radial artery Cardiogenic shock Prior CABG Demographics and baseline characteristics see below Drug therapy Patients received 300 mg aspirin, 300 mg clopidogrel on diagnosis, and subcutaneous Fragmin (5000U) or FraxiParin (4100U) for all patients. Further 5000IU heparin given during procedure. GP Ilb/Illa inhibitors and stents were given during procedure dependent based on clinical need.	Radial access to perform coronary angiography and PPCI (if clinically indicated) (n =100)	Femoral access to perform coronary angiography and PPCI (if clinically indicated) (n = 100)	1 month	All-cause mortality Reinfarction Repeat revascularisatio n CABG Hospital stay Angiographic procedural success Access site complications Fluoroscopy time (see below for definitions)	None stated

Demographics and baseline characteristics					
Characteristics	Radial (n = 100)	Femoral (n = 100)	p value		
Age(years), mean(SD)	64.9(8.4)	66.2(7.7)	0.23		
Male, n(%)	72(72)	69(69)	0.64		
Hypertension, n(%)	42(42)	50(50)	0.25		
Diabetes, n(%)	22(22)	15(15)	0.20		
Smoker, n(%)	50(50)	42(42)	0.27		
Obesity, n(%)	23(23)	30(30)	0.26		
Hypercholesterolaemia, n(%)	35(35)	40(40)	0.47		
Three vessel disease, n(%)	22(22)	18(18)	0.48		
Killip class	0.89				
Class I, n(%)	60(60)	58(58)			
Class II, n(%)	30(30)	33(33)			
Class II, n(%)	10(10)	9(9)			
Infarct related artery					
Left anterior descending artery, n	44	50			
Left circumflex, n	8	13			
Right coronary artery, n	44	37			
Initial TIMI flow, n	0.40				
TIMI 0 to 1	72	68			
TIMI 2	20	18			
TIMI 3	8	14			
TIMI; thrombolysis in mycocardial infarction	on				
Comparison of operation data between r	adial versus femoral groups				
Characteristics	Radial (n = 100)	Femoral (n = 100)	p value		
Success of puncture, n	100	100			
Cannulation time (min), mean(SD)	2.5(0.6)	2.4(0.6)	0.24		
95% CI	2.2 to 2.4	2.1 to 2.3			
Reperfusion time (min), mean(SD)	16.4(1.7)	16.2(1.8)	0.42		

95% CI	16 to 16.7	15.8 to 16.6	
Total procedural time (min), mean(SD)	37.2(7.1)	35.7(8.1)	0.17
95% CI	35.8 to 38.6	34 to 34.3	
Fluoroscopy time (min), mean(SD)	11.2(2.0)	11.4(1.8)	0.14
	11.4 to 12.2	11.1 to 11.8	
Final TIMI flow, n	0.60		
TIMI 0 to 1	2	1	
TIMI 2	2	4	
TIMI 3	96	95	
Stents used, n	97	95	0.72
Tirofiban used, n	28	20	0.19
Angiographic procedural success*, n	96	97	1.00

TIMI; thrombolysis in myocardial infarction, Angiographic procedural success * = residual diameter stenosis < 30% with grade 3 coronary flow according to the classification of fibrinolysis in myocardial trial

Definitions of outcomes

Major access site bleeding: Haemoglobin loss ≥ 2 mmol/l, administration of blood transfusion, and needing vascular repair Minor access site bleeding: Hematoma formation not requiring specific therapy

Effect Size – 1 month follow-up

Outcome	Radial (n = 100)	Femoral (n = 100)	p value
All-cause mortality, n	4	5	1.0
Reinfarction, n	0	0	
Repeat revascularisation, n	0	0	
Vascular complications, n	3	11	< 0.01
Major bleeding, n	0	3	2.4
Minor bleeding (haematoma), n	2	6	2.8
Pseudoaneurysm, n	0	2	0.16
Artery occlusion without ischaemia, n	1	0	
Hospital stay (day)			
Mean(SD) 95% CI	8.6(1.8) 8.3 to 9.0	12.7(3.0) 12.1 to 13.3	< 0.001

Table 41: Li 2007⁷³

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Li WM, Li Y, Zhao JY, Duan YN, Sheng L, Yang BF et al. Safety and feasibility of emergent percutaneous coronary intervention with the transradial access in patients with acute myocardial infarction. Chin Med J (Engl). 2007; 120(7):598-600.	Design RCT; 1 centre in China Enrolment June 2004 to July 2006 Randomisation Not detailed Allocation concealment Not detailed Blinding: Patients and investigator s were not masked to treatment allocation. All other outcomes were as reported by the investigators. ITT analysis Yes	n = 370 PPCI done in all patients Drop outs None Crossover 3 patients in radial group required crossover to femoral and 2 patients in femoral group required crossover to radial group (reasons not given) Operator expertise Not stated	Inclusion criteria Acute MI within 12 h onset of chest pain (no further details given) Exclusion criteria for the transradial group Negative Allen test Aorto-arteritis Cardiogenic shock Non-palpable radial artery Severe tortuosity of radial arteries Body height > 150 cm Demographics and baseline characteristics see below Drug therapy All patients received aspirin and clopidogrel before PPCI, adjunctive bolus heparin was determined by body weight (70 to 100 IU/Kg).	Radial access to perform coronary angiography and PPCI (if clinically indicated) (n = 184)	Femoral access to perform coronary angiography and PPCI (if clinically indicated) (n = 186)	In- hospital	Angiograp hic procedural success	None stated

Baseline and clinical characteristics		
Baseline	Radial (n = 184)	Femoral (n = 186)
Male/Female	124/60	120/66
Age(years), mean(SD)	56.5(10.9)	55.4(12.8)
Body height (cm), mean(SD)	166(12.5)	165.8(13.1)
Diabetes, n(%)	37(20.1)	34(18.3)
Hypertension, n(%)	74(40.2)	78(41.9)
Hyperlipidaemia, n(%)	29(15.8)	31(16.7)
Smoker, n(%)	78(42.4)	80(43.0)
Clinical characteristics	Radial (n = 184)	Femoral (n = 186)
Anterior wall acute MI, n(%)	83(45.1)	80(43.0)
Inferior wall acute MI, n(%)	58(31.5)	63(33.9)
Lateral and posterior wall acute MI, n(%)	30(16.3)	32(17.8)
Right ventricular and inferior wall acute MI, n(%)	13(7.1)	11(5.9)
Culprit artery, n(%)		
• Left anterior descending artery, n(%)	82(44.6)	80(43.0)
• Left circumflex, n(%)	26(14.1)	29(15.6)
• Right coronary artery, n(%)	74(40.2)	77(41.4)
• Left anterior descending artery, n(%)	2(1.1)	0
Single vessel disease, n(%)	107(58.2)	107(57.5)
Multivessel disease, n(%)	77(41.8)	79(42.5)
Total occlusion, n(%)	76(41.3)	78(41.9)
Left ventricular ejection fracture, mean(SD)	48.42(8.48)	51.21(9.21)
No statistical difference between 2 groups for left ventri	cular ejection fracture and infarct location	
Procedural and angiographic details		
Angiographic procedural success: TIMI flow of at least gr	ade III	
Variables	Radial (n = 184)	Femoral (n = 186)
Cannulation time (min), mean(SD)	27.1(5.8)	26.5(7.0)
Total procedural time (min), mean(SD)	56.2(12.1)	58.4(15.1)

Stent implantation, n	200	202
Successful rates of puncture, %	98.4	98.0
Final TIMI III flow	94.8	94.2

Table 42: RADIAMI 2009²⁶

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow- up	Outcome measures	Source of funding
Chodór P, Krupa H, Kurek T, Sokal A, Swierad M, Was T et al. RADIal versus femoral approach for percutaneous coronary interventions in patients with Acute Myocardial Infarction (RADIAMI): A prospective, randomized, single-center clinical trial. Cardiology Journal. 2009; 16(4):332- 340.	Design RCT; 1 centre in Poland Enrolment April 2005 to June 2006 Randomisation Based on year of birth (radial group; even years, femoral group; odd years) Allocation concealment Not detailed Blinding Patients and investigator s were not masked to treatment	PPCI not completed 1 patient in femoral group due to inability of balloon passage through occluded area of artery Drop outs 1 due to PPCI not completed Crossover 4 patients in radial group required crossover to femoral group, 3 abnormal Allen test, 1 excessive tortuous radial artery. 1 pateint in femoral	Inclusion criteria Presence of MI defined as retrosternal pain lasting > 20 min, but < 12 h, resistant to nitroglycerin, and ECG changes; ST elevation of at least 1 mV in two neighbouring leads or new left bundle branch block, found in the qualifying ECG Age between 18 and 75 years Participation consent Exclusion criteria Age over 75 years Killip class III or IV Necessity of an intra-aortic balloon pumpplacement before the PPCI Necessity of an endocavitary stimulating electrode placement before the PPCI Height < 150 cm History of CABG if the infarction may be due to a closed venous or arterial bypass graft Demographics and baseline characteristics	Radial access to perform coronary angiography and PPCI (if clinically indicated) (n = 50)	Femoral access to perform coronary angiography and PPCI (if clinically indicated) (n = 50)	In- hospital	All-cause mortality Reinfarctio n Stroke, CABD Repeat revasculari sation Major bleeding, Fatal bleeding requiring operation, drop in Hb Intracrania I haemorrha ge Hospital stay Total radiograph ic contrast media used in	None stated

other outcomes were as reported by the investigators. ITT analysis Yes	crossed to radial due to atherosclerosis obliterations of the lower limb Operator expertise Conducted by physicians with many years experience of performing femoral access PPCI (300 to 400 PPCIs per year), who had performed at least 50 to 100 radial access PPCIs	Drug therapy Verapamil (5mg) after puncture of radial artery; dose was repeated in the case of a spasm, until reaching a total dose of 15 mg. Dependent on activated clotting time result heparin (70 U/kg) was administered. Fibrinolytic drugs and platelet glycoprotein Ilb/Illa receptor blockers were administered during the intervention based on clinical need. Heparin administration was continued after the intervention only in the presence of clinical indications. Abciximab was administered to a similar percentage of patients in both groups (44% versus 42%, radial and femoral respectively). Stents were given to all the patients who underwent PPCI.		PPCI procedure Fluoroscop y time Hematoma		
Characteristics	Entire study group (n	= 100)	Radial (n = 90)	Femoral (n = 105)	p value	
Age (years), mean(SD)	59.5(9.1)		59.9(9.4)	59.1(9.0)	NS	
Height (cm), mean(SD)	169.1(8,4)		167.8(7,5)	169.5(9.2)	NS	
Body weight (kg), mean(SD)	82.2(14.7)		79.5(11.7)	85.0(16.8)	NS	
Men, n(%)	68(68)		35(51.5)	33(48.5)	NS	
Diabetes, n(%)	15(15)		8(16)	7(14)	NS	
Smoking, n(%)	64(64)		34(68)	30(60)	NS	
Arterial hypertension, n(%)	47(47)		26(52)	21(42)	NS	
Hyperlipidemia, n(%)	19(19)		11(22)	8(16)	NS	
Past infarction, n(%)	11(11)		8(16)	3(6)	NS	

Family history*, n(%)	32(32)	16(32)	16(32)	NS
Killip class 1, n(%)	99(99)	50(100)	49(98)	NS
Killip class 2, n(%)	1(1)	0(0)	1(2)	NS
HR at admission (beats/min), mean(SD)	78.2(15.0)	78.3(3.7)	78.2(6.4)	NS
SBP at admission (mmHg), mean(SD)	135.2(29.2)	138.8(33.2)	131.6(24.5)	NS
DBP at admission mmHg), mean(SD)	78.4(17.9)	80.1(18.6)	76.7(17.1)	NS
Infarction location				
Anterior wall, n(%)	42(42)	21(42%)	21(42)	NS
Inferior wall, n(%)	54(54)	27(54)	27(54)	NS
Left bundle branch block, n(%)	0(0)	0(0)	0(0)	NS
Other, n(%)	4(4)	1(4)	2(4%)	NS
*Family history of coronary hear heart rate, NS; not significant	rt disease; **Mann-Whitney U te	est; MIN; mean infarction duratio	n; DBP; diastolic blood pressure;	SBP; systolic blood pressure; HR;
Angiographic data				
Outcome	Total population (n =100)	Radial (n = 50)	Femoral (n = 50)	p value
Outcome No. of pathologic vessels, n(%)	Total population (n =100)	Radial (n = 50)	Femoral (n = 50)	p value
	Total population (n =100) 36(37.5)	Radial (n = 50) 21(42.9)	Femoral (n = 50) 15(31.9)	p value
No. of pathologic vessels, n(%)				·
No. of pathologic vessels, n(%)	36(37.5)	21(42.9)	15(31.9)	NS
No. of pathologic vessels, n(%) 1 2	36(37.5) 40(41.7)	21(42.9) 19(38.8)	15(31.9) 21(44.7)	NS NS
No. of pathologic vessels, n(%) 1 2 3	36(37.5) 40(41.7)	21(42.9) 19(38.8)	15(31.9) 21(44.7)	NS NS
No. of pathologic vessels, n(%) 1 2 3 Infarct-related artery, n(%)	36(37.5) 40(41.7) 20(20.8)	21(42.9) 19(38.8) 9(18.4)	15(31.9) 21(44.7) 11(23.4)	NS NS NS
No. of pathologic vessels, n(%) 1 2 3 Infarct-related artery, n(%) LM	36(37.5) 40(41.7) 20(20.8) 0(0%)	21(42.9) 19(38.8) 9(18.4)	15(31.9) 21(44.7) 11(23.4)	NS NS NS
No. of pathologic vessels, n(%) 1 2 3 Infarct-related artery, n(%) LM LAD	36(37.5) 40(41.7) 20(20.8) 0(0%) 43(44.3)	21(42.9) 19(38.8) 9(18.4) 0(0) 21(42.9)	15(31.9) 21(44.7) 11(23.4) 0(0) 22(45.8)	NS NS NS NS
No. of pathologic vessels, n(%) 1 2 3 Infarct-related artery, n(%) LM LAD Cx Percutaneous coronary	36(37.5) 40(41.7) 20(20.8) 0(0%) 43(44.3) 12(12.4)	21(42.9) 19(38.8) 9(18.4) 0(0) 21(42.9) 9(18.4)	15(31.9) 21(44.7) 11(23.4) 0(0) 22(45.8) 3(6.2)	NS NS NS NS NS NS NS
No. of pathologic vessels, n(%) 1 2 3 Infarct-related artery, n(%) LM LAD Cx Percutaneous coronary intervention, n(%)	36(37.5) 40(41.7) 20(20.8) 0(0%) 43(44.3) 12(12.4)	21(42.9) 19(38.8) 9(18.4) 0(0) 21(42.9) 9(18.4)	15(31.9) 21(44.7) 11(23.4) 0(0) 22(45.8) 3(6.2)	NS NS NS NS NS

1	9(9.3)	5(10.2)	4(8.3)	NS
2	19(19.6)	8(16.3)	11(22.9)	NS
3	16(16.5)	10(20.4)	6(12.5)	NS
Final TIMI flow grade, n(%)				
0	1(1)	0(0)	1(2)	NS
1	0(0)	0(0)	0(0)	NS
2	9(9)	6(12)	3(6)	NS
3	90(90)	44(88)	46(92)	NS
Residual stenosis after the intervention, n(%)	1.7(10.7)	0.8(4.0)	2.6(14.7)	NS*
Abciximab, n(%)	43(43)	22(44)	21(42)	NS
Exposure time (min), n(%)	11.1(6.3)	10.9(5.6)	11.2(7.0)	NS
Contrast amount (ml , n(%)	198.7(45.7)	198.7(45.7)	197.7(72.0)	NS*
No. of stents per patient, n(%)	1.27(0.47)	1.28(0.45)	1.26(0.49)	NS
Stenting percentage, n(%)	99(99)	50(100)	49 (98)	NS

^{*}Mann-Whitney U test, LM; left mammary, LAD; left artery descending, Cx; circumflex artery, RCA; right coronary artery, TIMI; Thrombolysis In Myocardial Infarction, NS not significant

Time intervals during coronary angiography and PPCI

Time from admission to (door to), min (SD)	Total population (n = 100)	Radial (n = 50)	Femoral (n = 50)	p value
Arrival in the cath lab	35.7 (21.6)	37.8 (21.0)	33.7 (22.2)	NS
Sheath positioning	49.1 (22.9)	53.7 (21.9)	44.4 (23.1)	0.04
First contrast injection	56.0 (25.1)	50.2 (23.8)	62.3 (25.5)	0.02*
Balloon positioning	69.1 (27.9)	76.9 (25.9)	64.6 (26.9)	0.02*
Stent implantation (door to stent)	77.9 (27.2)	83.2 (26.3)	72.3 (27.3)	0.05
End of intervention	92.7 (28.7)	98.7 (26.8)	88.7 (30.1)	0.17
Arrival in the cath lab to sheath positioning time	13.6 (7.4)	15.7 (7.8)	11.4 (6.4)	0.0028
Sheath to injection time	6.6 (6.4)	8.6 (7.8)	4.5 (3.3)	0.008*
Injection to balloon time	15.1 (7.9)	15.6 (8.7)	14.6 (7.1)	NS

Balloon to stent time	8.0 (4.9)	7.3 (4.6)	8.7 (5.2)	0.21
Stent to end of intervention time	14.4 (10.6)	13.3 (8.6)	15.5 (12.4)	0.31
Procedure	56.8 (18.1)	58.3 (17.8)	55.1 (18.4)	0.38
*Mann-Whitney U test				

Angiographic procedural success: TIMI grade 3 flow rate was obtained and residual stenosis was lower than 30%

Major bleeding: Fatal bleeding, bleeding requiring blood transfusion, operation or resulting in a drop of haemoglobin count of more than 3 g/dl as well as any intracranial haemorrhage

Minor bleeding: all non-major bleeding

Procedure time: Period from the patient's arrival in the cath lab to the removal of the vascular sheath for radial group, and to the removal of the catheter from the sheath for group femoral group.

Effect size

Outcome	Entire study group	Radial (n = 50)	Femoral (n = 50)	p value	
All-cause mortality	1(1)	0(0)	1(2)	NS	
Myocardialinfarction	1(1)	1(2)	0(0)	NS	
Stroke	1(1)	1(2)	0(0)	NS	
Repeated revascularisation of the IRA	3(3)	1(2)	2(4)	NS	
Coronary artery bypass grafting	0(0)	0(0)	0(0)	NS	
Serious bleeding	10(9.3)	3(8.2)	17(14)	0.18	
Fatal bleeding	0(0)	0(0)	0(0)	NS	
Serious bleeding requiring transfusion	3(3%)	0(0)	3(6)	NS	
Serious bleeding resulting in haemoglobin level decrease of > 3 g/dl	7(7%)	3(6%)	4(8%)	NS	
Intracranial bleeding	0(0)	0(0)	0(0)	NS	
Hematoma > 5 cm	13(13)	5(10)	8(16)	0.37	
NS: not significant					

Table 43: RADIAMI II 2011²⁷

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Chodór P, Kurek T, Kowalczuk A, Swierad M, Was T, Honisz G et al. Radial vs femoral approach with StarClose clip placement for primary percutaneous coronary intervention in patients with ST- elevation myocardial infarction. RADIAMI II: A prospective, randomised, single centre trial. Kardiologia Polska. 2011; 69(8):763- 771.	Design RCT; 1 centres in Poland Enrolment Nov 2006 to Mar 2008 Randomisation Based on year of birth (radial group; even years, femoral group; odd years) Allocation concealment Not detailed Blinding Patients and investigator s were not masked to treatment allocation. All other outcomes were as reported by the investigators	n = 108 PPCI done in all patients Drop outs None Crossover 2 patients in radial group required crossover to femoral group, 1 patient in femoral group crossed over to radial (reasons not given) Operator expertise 3 physicians with 17 to 20 years of experience performed femoral	Inclusion criteria MI defined as retrosternal pain lasting between 20 min and 12 h and not relieved by nitroglycerine, accompanied by ECG changes in the form of ST elevation of at least 0.01 mV in two adjacent leads or new left bundle branch block Age between 18 and 75 years Participation consent Exclusion criteria Age over 75 years Killip class III or IV Necessity of an intra-aortic balloon pumping or temporary right ventricular pacing Placement before the PPCI Necessity of an endocavitary stimulating electrode Placement before the PPCI Height < 150 cm History of CABG Demographics and baseline characteristics see below Drug therapy	Radial access to perform coronary angiography and PPCI (if clinically indicated) (n = 49)	Femoral access and a StarClose device to perform coronary angiography and PPCI (if clinically indicated) (n = 59)	In- hospital	All-cause mortality Reinfarctio n Stroke CABD Repeat revasculari sation Serious bleeding Fatal bleeding requiring transfusion Serious bleeding requiring transfusion Serious bleeding requiring surgery Serious bleeding giving drop in Hb of > 3 g/dl Intracrania I haemorrha ge Fluoroscop y time	None stated

ITT analysis No	access PPCI, and several years experience in performing radial access PPCI	clotting tim heparin wa permitted clotting tim procedures of abcixima abciximab	ng the sheath, activated ne was determined and as administered in doses that the obtaining of activated ne of 350 to 450 s during a performed without the use ab, and 250 to 350 s when was used. Whether to use was a decision left to the		Total radiograph ic contrast media used in PPCI procedure hematoma > 5 cm
Demographics and baseline cha	aracteristics				
Characteristics	Entire study group	(n = 108)	Radial (n = 49)	Femoral (n = 59)	p value
Age (years), mean(SD)	59.6(10.0)		62.1(9.3)	57.6(10.3)	0.02
Height (cm), mean(SD)	168.0(7.8)		168.5(7.7)	167.5(8.0)	NS
Body weight (kg), mean(SD)	79.1(13.0)		81.0(14.4)	77.5(11.6)	NS
Men, n(%)	69(64)		32(65)	37(63)	NS
Diabetes, n(%)	20(19)		10(21)	10(17)	NS
Smoking, n(%)	72(67)		29(60)	43(73)	NS
Arterial hypertension, n(%)	42(39)		22(46)	20(34)	NS
Hyperlipidemia, n(%)	22(21)		12(25)	10(17)	NS
Prior MI, n(%)	12(11)		4(8)	8(14)	NS
Family history of early CVD, n(%)	44(41)		22(46)	22(37)	NS
Circulatory status on admission	1				
Killip class 1, n(%)	90(83)		40(82)	50(85)	NS
Killip class 2, n(%)	18(17)		9(18)	9(15)	NS
Mean duration of symptoms (min), mean(SD)	271.8(168.5)		252.2(181.3)	288.2(156.6)	NS
HR at admission (beats/min), mean(SD)	81.4(18.4)		80.3(19.5)	82.3(17.5)	NS
SBP at admission (mmHg), mean(SD)	141.5(22.9)		139.9(21.6)	142.9(24.1)	NS

DBP at admission (mmHg), mean(SD)	90.7(14.6)	90.8(14.4)	90.7(14.9)	NS
MI location				
Anterior wall, n(%)	40(37)	21(43)	19(32)	NS
Inferior wall, n(%)	61(56)	24(49)	37(63)	NS
Left bundle branch block, n(%)	7(13)	4(8)	3(5)	NS
Other, n(%)	40(37)	21(43)	19(32)	NS
CVD; cardiovascular disease, dia	stolic blood pressure; SBP; systoli	c blood pressure; HR; heart rate, N	S; not significant	
Comparison of angiographic an	d procedural data between radia	versus femoral groups		
Characteristics	Entire study group (n = 108)	Radial (n = 49)	Femoral (n = 59)	p value
1-vessel disease, n(%)	50(47)	21(44)	29(49)	NS
2-vessel disease, n(%)	43(40)	21(44)	22(37)	NS
3-vessel disease, n(%)	14(13)	6(12)	8(14)	NS
Infarct related artery, n(%)				
Left main stem, n(%)	0(0)	0(0)	0(0)	NS
Left anterior descending artery, n(%)	38(35)	21(43)	17(29)	NS
Circumflex artery, n(%)	14(13)	4(8)	10(17)	NS
Right coronary artery, n(%)	55(51)	24(49)	31(53)	NS
Initial TIMI flow, n(%)				
0	57(53)	22(45)	35(59)	NS
1	8(7)	4(8)	4(7)	NS
2	18(17)	13(27)	5(8)	0.012
3	25(23)	10(20)	15(25)	NS
Final TIMI flow, n(%)				
0	0(0)	0(0)	0(0)	NS
1	0(0)	0(0)	0(0)	NS
2	1(1)	0(0)	1(2)	NS
3	107(99)	49(100)	58(98)	NS

RS post-procedure < 20%, n(%)	108(100)	49(100)	59(100)	NS
Maximum activated clotting time (sec), mean(SD)	322.8(68)	304.8(64.8)	336(68)	0.025
Abciximab administration, n(%)	57(53)	25(51)	32(54)	NS
Fluoroscopy time (min), mean(SD)	7.0(3.0)	7.5(3.0)	6.9(3.0)	NS
Contrast material (ml), mean(SD)	163.4(43.7)	165.0(41.4)	162.0(46.0)	NS
Number of stents implanted				
1 stent, n(%)	70(65)	34(69)	36(61)	NS
2 stents, n(%)	31(29)	13(27)	18(31)	NS
3 stents, n(%)	5(5)	2(4)	3(5)	NS
Stenting ratio, %	98.2	100	94.9	NS
Successful placement of StarClose clip n(%), or Terumo band, n(%)		48(98.0)	55(93.2)	
RS; residual stenosis, NS; not sig	nificant			
Time intervals during coronary a	angiography and PPCI for radial a	nd femoral groups		
Time interval	Entire study group (n = 108)	Radial (n = 49)	Femoral (n = 59)	p value
Interval from cath lab arrival to	10.3(5.6)	11.2(6.5)	9.6(4.7)	NS
beginning of the procedure (min), mean(SD) (median; interquartile range)	(10.0; 5.0 to 13.0)	(10.0; 5.0 to 15.0)	(10.0; 6.0 to 10.0)	
Interval from beginning of	4.7(5.16)	5.8(6.8)	3.8(3.0)	NS
procedure to vascular sheath introduction (min), mean(SD) (median; interquartile range)	(3.0; 2.0 to 5.0)	(4.0; 2.0 to 5.0)	(3.0; 2.0 to 5.0)	
Interval from vascular sheath	5.4 (5.5)	5.4(3.9)	5.4 (6.6)	NS
introduction to first contrast injection (min), mean(SD) (median; interquartile range)	(4.5; 3.0 to 5.0)	(5.0; 3.5 to 6.0)	(4.0; 3.0 to 5.0)	
Interval from contrast injection	12.6(6.3)	11.9(4.9)	13.3(7.1)	NS

to balloon inflation (min), mean(SD) (median; interquartile range)	(11.0; 10.0 to 14.0)	(11.0; 10.0 to 13.0)	(11.5; 10.0 to 14.0)	
Interval from balloon inflation	7.3(5.7)	7.3(7.2)	7.3(4.2)	NS
to stent implantation (min), mean(SD) (median; interquartile range)	(9.0; 6.0 to 9.0)	(5.5; 4.0 to 8.0)	(6.5; 5.0 to 10.0)	
Interval from stent	11.0(7.2)	9.4(6.8)	12.6(7.3)	0.005
implantation to end of procedure (min)*, mean(SD) (median; interquartile range)	(10.0; 6.0 to 15.0)	(7.0; 5.0 to 13.00)	(10.0; 8.0 to 16.0)	
Total procedural time (from	50.2(20.2)	53.7 (20.6)	47.3(19.6)	NS
cath lab arrival to end of procedure) (min), mean(SD) (median; interquartile range)	(48.0; 40.0 to 58.0)	(50.0; 41.5 to 60.0)	(45.0; 40.0 to 56.0)	
				_

^{*}in cases of implantation of more than one stent, this is the interval from final stent implantation to the end of the procedure, NS; not significant

Angiographic procedural success: TIMI grade 3 flow rate was obtained and residual stenosis < 20%

Major bleeding complications: Serious bleeding that resulted in death or a need for blood transfusion or surgical intervention, caused haemoglobin level decrease by > 3 g/dl, and central nervous system bleedings.

Minor bleeding: all non-major bleeding

Procedure time: Period from the patient's arrival in the cath lab to the removal of the vascular sheath and placement of Terumo band dressing for radial group, and to the removal of the catheter from the sheath and VCD implantation for group femoral group.

Effect size

Outcome	Entire study population (n = 118)	Radial (n = 49)	Femoral (n = 59)	p value
All-cause mortality, n(%)	0(0)	0(0)	0(0)	NS
MI, n(%)	0(0)	0(0)	0(0)	NS
Stroke, n(%)	1(0.9)	0(0)	1(1.7)	NS
Repeated revascularisation of the IRA, n(%)	1(0.9)	1(2.0)	0(0)	NS
Coronary artery bypass	0(0)	0(0)	0(0)	NS

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grafting, n(%)				
PCI of a vessel other than the IRA , n(%)	2(1.9)	0(0)	2(3.4)	NS
Serious bleeding, n(%)	10(9.3)	4(8.2)	6(10.2)	NS
Serious bleeding resulting in death, n(%)	0(0)	0(0)	0(0)	NS
Serious bleeding requiring blood transfusion, n(%)		1(0)	1(0)	NS
Serious bleeding requiring surgery, n(%)	0(0)	0(0)	0(0)	NS
Serious bleeding resulting in haemoglobin level decrease of > 3 g/dl, n(%)	9(8.3)	3(6.1)	6(10.2)	NS
Intracranial bleeding, n(%)	0(0)	0(0)	0(0)	NS
Hematoma > 5 cm, n(%)	20(18.5)	8(16.3)	12(20.3)	NS
IRA: infarct-related artery, NS: n	ot significant			

Table 44: RIFLE-STEACS⁹³

						Long		
Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	th of follo w-	Outcome measures	Source of funding
		-				up		
Romagnoli E, Biondi-Zoccai G, Sciahbasi A, Politi L, Rigattieri S, Pendenza G, Summaria F, Patrizi R, Borghi A, Di RC, Moretti C, Agostoni P, Loschiavo P, Lioy E, Sheiban I, Sangiorgi G. Radial Versus Femoral Randomized Investigation in ST-Segment Elevation Acute Coronary Syndrome: The RIFLE- STEACS	Design: Multi-centre (4 centres) randomised, parallel group study Enrolment: 1,001 acute ST-segment elevation acute coronary syndrome patients undergoing primary/rescues percutaneous coronary intervention Randomisation: Based on a computer-generated random series and stratified by centre. Allocation concealment: Patients were randomised	Number of patients n = 1001 Drop-outs: None Crossover: None	Inclusion criteria: Patients with acute ST-segment elevation acute coronary syndrome patients undergoing primary/rescues percutaneous coronary intervention Exclusion criteria: Contraindication to either radial or femoral vascular access Recent stroke (within 4 weeks) Anti-coagulant therapy assumption with INR >2 Other severe bleeding diathesis Demographics and baseline characteristics: See below	Radial approach for PPCI	Femoral approach for PCI	w-		No extramu ral funding
(Radial Versus Femoral Randomized Investigation	according to opaque, numbered, sealed envelopes		Drug therapy: Procedural anticoagulation with administration of an un-					
in ST-Elevation Acute	Blinding: Patients and		fractionated heparin bolus at a dose of 70 UI/kg; ASA					

Coronary Syndrome) Study. Journal of the American College of Cardiologists. 2012 not blinded; end point adjudication was performed by a blinded central independent clinical event committee

Sample size calculation:
Assumptions for sample size analysis were based, for the control event rate, on NACE rates reported in the HORIZINS-AMI trial and pertinent metaanalyses.

ITT analysis: 1001 patients included in ITT analysis. No loss to follow-up.

plus a loading dose of clopidogrel 300 -600 mg; use of pre-procedural antithrombotic agents was left to the operators' discretion.

Demographics and baseline characteristics

- •			
Clinical characteristics n (%)	Femoral (n = 501)	Radial (n = 500)	p value
Age, yrs (median)	65 (55-77)	65 (56-75)	0.409
Female (%)	141 (28.1)	126 (25.2)	0.317
BMI, kg/m ²	26.6 (24-30)	27.2 (25-30)	0.140
LV ejection fraction	45.0 (40-50)	45.0 (40-52)	0.175
CK (GFR <60 ml/min/1.73m ²)	127 (25.3)	111 (22.2)	0.156
COPD	40 (8.0)	31 (6.2)	0.325
Peripheral arterial disease	68 (13.6)	75 (15.0)	0.529
Previous myocardial infarction	71 (14.2)	70 (14.0)	1.00

Previous CVA	22 (4.4)	19 (3.8)	0.750	
Previous revascularisation	52 (10.4)	65 (13.0)	0.2.2	
Previous PCI	45 (9.0)	60 (12.0)	0.123	
Previous CABG	12 (2.4)	7(1.4)	0.356	
Clinical characteristics n (%)	Femoral (n = 501)	Radial (n = 500)	p value	
Hypertension	309 (61.7)	299 (59.8)	0.561	
Hypercholesterolemia	199 (39.7)	218 (43.6)	0.223	
Smoking	191 (38.1)	210 (42.0)	0.221	
Family history of CAD	81 (16.2)	96 (19.2)	0.215	
Diabetes	122 (24.4)	115 (23.0)	0.656	
Single vessel disease	265 (52.9)	279 (55.8)	0.374	
Double vessel disease	149 (29.7)	136 (27.2)	0.401	
Triple vessel disease	80 (16.0)	79 (15.8)	1.00	
Killip I	330 (65.9)	348 (69.6)	0.224	
Killip II	108 (21.5)	102 (20.4)	0.670	
Killip III	28 (5.6)	24 (4.8)	0.670	
Killip IV	35.(7.0)	26 (5.2)	0.290	
Results: 30 day outcome n (%)	Femoral (n = 501)	Radial (n = 500)	p value	
NACE	105 (21.0)	68 (13.6)	0.003	
MACE	57 (11.4)	36 (7.2)	0.029	
Cardiac death	46 (9.2)	26 (5.2)	0.020	
Stroke	3 (0.6)	4 (0.8)	0.725	
MI	7 (1.4)	6 (1.2)	1.000	
Target lesion revascularisation	9 (1.8)	6 (1.2)	0.604	
Stent thrombosis	9 (1.8)	6 (1.2)	0.604	
Non-CABG bleeding	61 (12.2)	39 (7.8)	0.026	
Access site related bleeding	34 (6.8)	13 (2.6)	0.002	
Non-access site related	27 (5.4)	26 (5.2)	1.000	

Results: n (%)	Femoral (n = 501)	Radial (n = 500)	p value
Fatal bleeding	3 (0.6)	3 (0.6)	0.684
Hospital stay, days (range)			
Total hospital stay	6 (5-8)	5 (4-7)	0.0008
Intensive coronary care unit	4 (3-5)	3 (2-4)	<0.001
Cardiology ward	3 (1-4)	2 (1-4)	0.472

Table 45: RIVAL^{59,60}

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Jolly SS, Yusuf S, Cairns J, Niemela K, Xavier D, Widimsky P et al. Radial versus femoral access for coronary angiography and intervention in patients with acute coronary syndromes (RIVAL): a randomised, parallel group, multicentre trial. Lancet.	Design RCT (multicentre; 158 hospitals in 32 countries) Enrolment June 2006 to Nov 2010 Randomisation 1:1 randomisation, variable sizes (2, 4 and 6) stratified by centre Allocation concealment 24h computerised central automated voice response system (at the Population Health Research institute)	n = 7021 Drop-outs: n = 29 (radial) n = 47 (femoral) Crossover: n = 245 (7%) Radial to femoral after failed radial access due to; radial spasm (n = 80), radial artery loop (n = 20), subclavian tortuosity (n = 31), other data unavailable n = 32(2%) Femoral to radial	Inclusion criteria Patients with STEMI Patients with ACS with or without ST segment elevation Patients with UA or NSTEMI an invasive approach was planned intent to perform same- sitting coronary angiography and PPCI during index hospitalisation suitable candidate for either radial or femoral artery PPCI the interventional cardiologist was willing to proceed with either radial or femoral access (and had expertise for both, including at least 50 radial procedures for coronary angiography or intervention within the	Radial access to perform coronary angiography and PPCI (if clinically indicated) (n = 3507) Note: For both groups, the use of an arterial vascular closure device is allowed at the discretion of the treating	Femoral access to perform coronary angiography and PPCI (if clinically indicated) (n = 3514) Note: For both groups, the use of an arterial vascular closure device is allowed at the	48 h 30 days	All-cause mortality Reinfarction Stroke Repeat revascularisati on Non-CABG- related major bleeding within 30 days Angiographic procedural success Major vascular access site complications at 48 h	Sanofi- Aventis, Populati on Health Researc Institute Canadia n Network for Internati (CANNe CTIN)

2011; 377(9775):1 409-1420. RIVAL trial methods Jolly SS, Niemela K, Xavier D, Widimsky P, Budaj A, Valentin V et al. Design and rationale of the radial versus femoral access for coronary intervention (RIVAL) trial: a randomized comparison of radial versus femoral access for coronary angiography or intervention in patients with acute coronary syndromes. American Heart	Blinding Patients and investigators were not masked to treatment allocation, but a masked central committee adjudicated the primary outcome and its components and stent thrombosis. All other outcomes were as reported by the investigators. Sample size calculation Lower than expected overall event rate for the primary outcome, so sample size was increased from 4000 to 7000 (provide 80% power to detect 25% RR reduction with control event rate of 6% and 30% risk reduction with control event rate of 4.5%) ITT analysis All randomised patients were included in ITT comparisons of outcome (regardless of whether crossed over)	after failed femoral access due to iliac tortuosity (n = 10), peripheral vascular disease (n = 9), other data unavailable Operator expertise Each operator had performed ≥ 50 radial procedures within the previous year	previous year) dual circulation of the hand was intact as assessed by Allen test Exclusion criteria <18 years Active bleeding or significant increased risk of bleeding (severe hepatic insufficiency, current peptic ulceration, proliferative diabetic retinopathy) uncontrolled hypertension carcinogenic shock severe peripheral vascular disease precluding a femoral approach previous coronary bypass surgery with use of >1 internal mammary artery Previously entered in the study Investigational treatment (drug or drug device) within the previous 30 days Demographics and baseline characteristics see below Drug therapy Antithrombotic regimen (including glycoprotein Ilb/Illa inhibitors) used for	physician	discretion of the treating physician	(see below for definitions)	
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Journal. 2011; 161(2):254.	PPCI was at the discretion of treating physician, (see Demographics and baseline characteristics). Stents were used in 95% of patients in both radial and femoral groups.	
Demographics and baseline characteristics		
	Radial (n = 3507)	Femoral (= 3514)
Age (years), mean (SD)	62 (2)	62 (12)
	(14.4% >75 years)	(15.1% >75 years)
Male sex, n (%)	2599 (74.1)	2561 (72.9)
Diagnosis at admission, n (%)		
Unstable angina	1544 (44.3)	1606 (45.7)
NTSTEMI	998 (28.5)	905 (25.8)
STEMI	995 (27.2)	1003 (28.5)
Medical history, n (%)		
Present smoker	1083 (30.9)	1097 (31.2)
Diabetes mellitus	781 (22.3)	722 (20.5)
MI	658 (18.8)	622 (17.7)
PPCI	431 (12.3)	408 (11.6)
Hypertension	2118 (60.4)	2076 (59.1)
CABG	79 (2.3)	75 (2.1)
Peripheral vascular disease	91 (2.6)	82 (2.3)
Antithrombotic treatment in-hospital, n(%)		
Aspirin	3479 (99.2)	3489 (99.3)
Clopidogrel	3368 (96.0)	3358 (95.6)
Low Mwt heparin	1806 (51.5)	1819 (51.8)
IV unfractionated heparin	1168 (33.3)	1110 (31.6)
Fondaparinux	383 (10.9)	381 (10.8)

Bivalirudin		76 (2.2)		109 (3.1)	
Glyc IIb/IIIa inhibitor		887 (25.3)		844 (24.0)	
Other in-hospital medicat	ions, n(%)				
PPIs		1050 (29.9)		1097 (31.2)	
BBs		3104 (88.5)		3130 (89.1)	
ACEs		2546 (72.6)		2539 (72.3)	
ARBs		377 (10.7)		386 (11.0)	
Statins		3309 (94.4)		3289 (93.6)	
CCBs		655 (18.7)		623 (17.7)	
Procedural complications	and outcomes and patient p	reference			
Outcome	Radial (n = 3507)	Femoral (n = 3514)	HR	95% CI	p value
Major vascular complications at 30 days					
Large haematoma, n(%)	42(1.2)	106(3.0)	0.40	0.28-0.57	< 0.0001
Pseudoaneurysm needing closure, n(%)	7(0.2)	23(0.6)	0.30	0.13-0.71	0.006
Arteriovenous fistual, n(%)	0(0)	5(0.1)			
Ischaemic limb needing closure, n(%)	1(0)*	0(0)			
PPCI complications#					
Abrupt closure, n(%)	12(0.5)	11(0.5)	1.11	0.49 to 2.51	0.81
No reflow, n(%)	21(0.9)	31(1.3)	0.69	0.40 to 1.20	0.19
Dissection with reduced flow, n(%)	30(1.3)	25(1.1)	1.22	0.72 to 2.07	0.46
Coronary perforation, n(%)	5(0.2)	4(0.2)	1.27	0.34 to 4.37	0.72
Catheter thrombus, n(%)	2(0.1)	2(0.1)	1.01	0.14 to 7.21	0.99
Stent thrombosis, n(%)‡	16(0.7)	26(1.2)	0.63	0.34 to 1.17	0.14
Definite, n(%)	8(0.4)	16(0.7)	0.51	0.22 to 1.19	0.12

Probable, n(%)	8(0.4)	11(0.5)	0.74	0.30 to 1.84	0.52
PPCI procedural time (min), median(IQR)	35(22,50)	34(22,50)			0.62
Fluoroscopy time (min), median(IQR)†	9.8(5.8,15.0)	8.0(04.5, 13.0)			< 0.0001
Length of stay in hospital (days), median(IQR)	4(3,7)	4(3,7)			0.18
Persistent pain at access site for > 2 weeks, n(%)	87/3378(2.6)	104/3392(3.1)	0.84	0.63 to 1.12¶	0.22
Patient prefers radial next procedure, n(%)	2962/3282(90.2)	1629/3210(50.7)	8.99	7.86 to 10.28¶	< 0.0001

^{*}Related to iliac thrombosis secondary to intra-aortic balloon pump inserted via femoral site. As a proportion of patients having PPCI; n =2311 in radial group and n = 2349 in femoral group. As a proportion of patients receiving a stent; n = 2197 in femoral group and n = 2243 in femoral time. Fluoroscopy times added to case reports and available for 2850 patients in the radial group and 2890 patients in the femoral group. Odds ratio (95%CI)

Angiographic procedural success

- Failure: no success at dilating attempted lesion(s) and/or failure to cross/dilate/not attempted
- Partial success: one of ≥2 attempted lesions was successfully dilated and procedure performed but >50% residual or TIMI flow <3 or failure
- Full success: lesions(s) attempted was successfully dilated with <50% residual or TIMI 3 flow

Major bleeding: Bleeding that was: 1. fatal, 2. resulted in transfusion of 2 or more units of red blood cells or equivalent whole blood, 3. caused substantial hypotension or with the need for inotropes, 4. needed surgical intervention, 5. caused severely disabling sequelae, 6. was intracranial and symptomatic or intraocular and ked to significant visual loss, or 7. led to a drop in Hb of at least 50 g/l

Minor bleeding: Bleeding events that did not meet the criteria for major bleeding and required transfusion of 1 unit of blood or modification of the drug regiment (ie. cessation of antiplatelet or antithrombotic therapy)

Major vascular access site complications: Included pseudoaneurysms needing closure, large haematoma (as judged by investigator), arteriovenous fistula, or and ischaemic limb needing surgery; these complications were classed as a major bleeding event or a minor bleeding event only if they also met the above definitions of major or minor bleeding

Effect Size

Total combined populations; hazard ratios (log rank) and 95%CI

STEMI Clinical evidence tables

Outcome	Radial (n = 3507)	Femoral (n = 3514)	HR	95% CI	p value	
Non-CABG major bleeding at 30 days, n(%)	24 (0.7)	33 (0.9)	0.73	0.43 to 1.23	0.23	
Non-CABG major bleeding at 48 h, n(%)	11 (0.3)	18 (0.5)	0.61	0.29 to 1.30	0.20	
All-cause mortality at 30 days, n(%)	44 (1.3)	51 (1.5)	0.86	0.58 to 1.29	0.47	
All-cause mortality at 48 h, n(%)	9 (0.3)	15 (0.4)	0.60	0.26 to 1.37	0.23	
Reinfarction at 30 days, n(%)	60 (1.7)	65 (1.9)	0.92	0.65 to 1.31	0.65	
Reinfarction at 48 h, n(%)	29 (0.8)	31 (0.9)	0.94	0.56 to 1.56	0.80	
Stroke at 30 days, n(%)	20 (0.6)	14 (0.4)	1.43	0.72 to 2.83	0.30	
Stroke at 48 h, n(%)	7 (0.2)	6 (0.2)	1.17	0.39 to 3.48	0.78	
Angiographic procedural success Angiographic procedural success*	2204 (95.4%)	2235 (95.2%)	1.01	0.95 to 1.07	0.83	
Access site crossover	265 (7.6)	70 (2.0)	3.82	2.93 to 4.97	<0.0001	
Major vascular complications	49 (1.4)	131 (3.7)	0.37	0.27 to 0.52	<0.0001	
Minor bleeding	100 (2.9)	118 (3.4)	0.84	0.65 to 1.10	0.21	
Non-CABG TIMI major bleeding	19 (0.5)	19 (0.5)	1.00	0.53 to 1.89	1.00	
CABG related bleeding	48 (1.4)	48 (1.4)	1.00	0.67 to 1.49	1.00	
Non-CABG-related blood transfusions	39 (1.1)	45 (1.3)	0.87	0.56 to 1.33	0.51	
All blood transfusions	99 (2.8)	98 (2.8)	1.01	0.76 to 1.33	0.95	
*As a proportion of patient	s who had PPCI: n = 2311 (ra	dial group) and n = 2349 (fen	noral group)			
NSTE-ACS versus STEMI patients; hazard ratios (log rank) and 95%CI						

Outcome	Total	Radial (n/total population)	Femoral (n/total population)	HR	95% CI	p value
Non-CABG major blee	eding at 30 days, n(%)					
NSTE-ACS	5063	16/2552(0.6)	24/2511(1.0)	0.66	0.35 to 1.23	0.19
STEMI	1958	8/955(0.8)	9/1003(0.9)	0.92	0.36 to 2.39	0.87
Overall	7021	24/3507(0.7)	33/3514(0.9)	0.73	0.43 to 1.23	0.23
All-cause mortality at	30 days, n(%)					
NSTE-ACS	5063	32/2552(1.2)	19/2511(0.8)	1.66	0.94 to 2.92	0.082
STEMI	1958	12/955(1.3)	32/1003(3.2)	0.39	0.20 to 0.76	0.006
Overall	7021	44/3507(1.3)	51/3514(1.5)	0.86	0.86 to 1.29	0.47
Major vascular compl	ications at 30 days, n(%)					
NSTE-ACS	5063	37/2552(1.4)	96/2511(3.8)	0.38	0.26 to 0.55	< 0.0001
STEMI	1958	12/955(1.3)	35/1003(3.5)	0.36	0.19 to 0.70	0.002
Overall	7021	49/3507(1.4)	131/3514(3.7)	0.37	0.27 to 0.52	< 0.0001
Access site crossover,	n(%)					
NSTE-ACS	5063	214/2552(8.4)	54/2511(2.2)	3.94	2.92 to 5.31	< 0.0001
STEMI	1958	51/955(5.3)	16/1003(1.6)	3.32	1.89 to 5.82	< 0.0001
Overall	7021	265/3507(7.6)	70/3514(2.0)	3.82	2.93 to 4.97	< 0.0001
STEMI patients						
Outcome	Radial	Fem	noral	RR (95%CI)		
	(n/population)	(n/p	opulation)			
Stroke	5/955	4/10	003	1.31 (0.35 to 4.87)		
Reinfartion	11/955	18/	1003	0.64 (0.30 to 1.35)		
Minor bleeding	33/955	22/:	1003	1.58 (0.93 to 2.38)		

Table 46: TEMPURA 2003⁹⁵

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Saito S,	Design	n = 149	Inclusion criteria	Radial access to	Femoral access	In-	All-cause	None stated
Saito S, Tanaka S, Hiroe Y, Miyashita Y, Takahashi S, Tanaka K, Satake S. Comparative study on transradial approach vs. transfemoral approach in primary stent imlplantation for patients with acute myocardial infarction: results of the test for myocardial infarction by prospective unicenter		patients			•	follow-up	measures	funding
unicenter randomization for access		to control and engage	conditions Culprit vessel was previous					
sites (TEMPURA) trial.		several types of guiding catheters	CABG Culprit vessel not identified					
Cathetarizatio n and cardiovascular interventions.		into LAD due to tortuous	Culprit vessel too small by extreme tortuosity and /or					

59:26-33 (2003)	iliac artery and enlarged aortic arch PCI Operator expertise Not stated	artery or of smaller vessel (< 2.5 mm diameter by visual estimate) Demographics and baseline characteristics see below Drug therapy Heparin given after arterial puncture (men; 6000 units, women; 5000 units). Any fibrinolytic agents were not given before or after PPCI. GP IIb/IIIa inhibitors were not given as not licensed in Japan. All patients received stents. Once daily aspirin (162 mg or more) and ticlopidine (200 mg) were started as soon as possible after stent implantation and continued for > 4 weeks			
Baseline characteristics					
	Radial (n	= 77)	Femoral (n = 72)	p value	
Male, n(%)	62(80.5)		59(81.9)	0.824	
Age, mean(SD)	66(12)		67(10)	0.872	
Diabetes, n(%)	19(24.7)		19(26.4)	0.810	
Hypertension, n(%)	38(49.4)		38(52.8)	0.676	
Hyperlipidemia, n(%)	21(27.3)		17(23.6)	0.608	
Smoking, n(%)	30(39.0)		39(54.2)	0.063	

Prior MI, n(%)	5(6.5)	6(8.3)	0.668
Cerebrovascular disease, n(%)	2(2.6)	6(8.3)	0.121
Killip classification, n(%)			0.222
• 1	48(62.3)	44(61.1)	
• 2	16(20.8)	20(27.8)	
• 3	11(14.3)	4(5.6)	
• 4	2(2.6)	4(5.6)	
Symptom onset to arrival (hr), mean(%)	3.1(2.4)	3.3(2.8)	0.670
Number of diseased arteries, n(%)			0.538
• Single	57(74.0)	48(66.7)	
• Double	14(18.2)	15(20.8)	
• Triple	6(7.8)	9(12.5)	
Culprit artery, n(%)			0.918
 Left anterior descending artery 	37(48.1)	37(51.4)	
Left circumflex	9(11.7)	6(8.3)	
 Right coronary artery 	29(37.7)	27(37.5)	
 Left anterior descending artery 	2(2.6)	2(2.8)	
Initial TIMI flow			0.492
• 0	52(76.6)	56(77.8)	
• 1	4(5.2)	7(9.7)	
• 2	7(9.1)	6(8.3)	
• 3	7(9.1)	3(4.2)	
Absence of collateral	52(67.5)	50(69.4)	0.802
Left ventricular ejection fraction			
<0.40, n(%)	3(3.9)	6(8.3)	0.256
Left ventricular end-diastolic			
Pressure (mm Hg), mean (SD)	25(9)	21(9)	0.016
TIMI; thrombolysis in mycocardial			

infarction

Comparison of operation data between radial versus femoral groups						
	Radial group (n = 77)	Femoral group (n = 72)	p value			
IABP support, n(%)	7(9.1)	7(9.7)	0.895			
Temporal pacing, n(%)	4(5.2)	6(8.3)	0.444			
Guideline catheters used, n(SD)	1.1(0.4)	1.1(0.3)	0.307			
Fluoroscopy time (min), mean(SD)	15.1(7.6)	16.1(7.9)	0.500			
Total procedure time (min), mean (SD)	44(18)	51(21)	0.033			
Onset of the end of PPCI (hr), mean(SD)	4.3(2.2)	4.7(3.0)	0.366			
Total amount of radiographic contrast media used (ml), mean(SD)	180(61)	186(66)	0.579			
Stent diameter (mm), mean(SD)	3.3(0.4)	3.2(0.4)	0.650			
Stents given, n(%)	77(100)	72(100)				
Number of stents, n(SD)	1.4(0.6)	1.3(0.6)	0.760			
Total stent length (mm), mean(SD)	19.8(8.0)	19.4(7.3)	0.763			
Final TIMI flow, n(%)			0.624			
• 0	1(1.3)	0				
• 1	0	0				
• 2	2(2.6)	2(2.8)				
• 3	74(96.1)	70(97.2)				
Reference vessel diameter (mm), mean(SD)						
• Pre	2.99(0.87)	3.10(0.74)	0.797			
• Post	3.18(0.55)	3.25(0.55)	0.826			
Minimum lumen diameter (mm), mean (SD) post	3.11(0.85)	3.10(0.50)	0.249			
Post diameter stenosis, %(SD)	7.3(11.6)	6.6(9.1)	0.358			
Peak CK (IU/I), mean (SD)	3170(2192)	3256(2123)	0.807			

Peak CRP(mg/d), mean(SD)	3.6(4.0)	4.2(4.6)	0.448		
Success of procedure, %	96.1	97.1	0.939		
IABP; Intra-aortic balloon pump, CRP;	IABP; Intra-aortic balloon pump, CRP;				
C-reactive protein, CK; creatine kinase					

Definitions:

- Procedure times: Time from entry of patient into the catheterisation laboratory to arterial sheath removal within the laboratory in patients of the radial group and that to the removal of catheters from the arterial sheath in those of the femoral group
- Angiographic procedural success: Achievement of TIMI grade 3 flow at end of PPCI
- Restenosis: increase in % diameter stenosis to ≥ 50%
- Major bleeding: bleeding requiring blood transfusion and/or surgical repair or cerebral bleeding

Effect size

In-hospital					
Outcome	Radial	Femoral	p value		
N	77	72			
All-cause mortality, n(%)	4(5.2)	6(8.3)	0.444		
Reinfarction, n	0	0			
Repeat revascularisation, n(%)	0	0			
Crossover to opposite arm, n(%)	0	1(1.5)	0.300		
Major bleeding, n(%)	0	2(3.0)	0.141		
Excluding patients with in-hospital death					
N	73	66			
Hospital stay, day(SD)	5.7(4.9)	7.4(0.95)	0.204		
Success at day 3 discharge, n(%)	43(58.9)	32(48.5)	0.218		
6 month follow-up angiogram					
N	44	43			
Reference vessel diameter (mm), mean(SD)	3.17(0.61)	3.08(0.53)	0.480		
Minimum lumen diameter (mm), mean(SD)	2.19(0.76)	1.94(0.92)	0.189		

Diameter stenosis, %(SD)	31(21)	39(27)	0.153
Binary restenosis, n(%)	13(29.5)	14(33.3)	0.705
9 month follow-up			
N	73	66	
All-cause mortality, n(%)	0	1(1.5)	0.629
MI, n(%)	2(2.7)	1(1.5)	0.291
Repeat revascularisation, n(%)	13(17.8)	15(22.7)	0.351

Table 47: Brasselet 2007¹⁴

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Brasselet C, Tassan S, Nazeyrollas P, Hamon M, Metz D. Randomised comparison of femoral versus radial approach for percutaneous coronary intervention using abciximab in acute myocardial infarction: results of the FARMI trial. Heart. 2007; 93(12):1556- 1561.	Design RCT; recruited either from the emergency unit of institution, mobile intensive care units, or referred from other hospitals to institution for emergent PCI, French study. Enrolment Jan 2004 to Sept 2005 Randomisation Not detailed Allocation	n = 114 PCI done in all patients Drop outs None Crossover Radial group; 7 (12%) required conversion to femoral access despite primary arterial cannulation success.	 Acute coronary syndrome with ST segment elevation associated with sustained chest pain Recruited either from the emergency unit of institution, mobile intensive care units, or referred from other hospitals to institution for emergent PCI. Emergent PCIs were defined as primary, rescue or facilitated PCI as follows: PPCI was defined as mechanical coronary recanalisation without previous thrombolysis or pretreatment by glycoprotein IIb/IIIa (GpIIb/IIIa) inhibitors; rescue PCI was defined as mechanical coronary recanalisation when thrombolytic treatment had failed; facilitated PCI was defined as mechanical coronary recanalisation after successful thrombolytic treatment, assessed by clinical and electrocardiographic criteria 	Radial access to perform coronary angiography and PCI (if clinically indicated) (n = 57)	Femoral access to perform coronary angiography and PCI (if clinically indicated) (n = 57)	In- hospital	All-cause mortality Major bleeding Minor bleeding Angiograp hic procedural success, Fluoroscop y time n = 114	None stated

Blinding Patients and investigator s were not masked to treatment allocation. All other outcomes were as reported by the investigators. ITT analysis Yes	approach failures resulted from technical difficulties (ie, inability to selectively catheterise	 Killip state > 2 or cardiogenic shock) Need for an intra-aortic balloon pump or temporary pacemaker History of a coronary artery bypass graft Intolerance to abciximab Demographics and baseline characteristics see below Drug therapy Before PCI, patients were pretreated by either an intravenous bolus of heparin as follows: unfractionated heparin 50 IU/kg with an upper limit of 4000 IU in patients > 75 years old, or low molecular weight heparin (enoxaparin) 30 mg intravenously and 1 mg/kg subcutaneously in patients aged > 75 years, and a bolus of aspirin (250 mg intravenously). This protocol was used irrespective of arterial access since it was started by emergency units, mobile intensive care units or hospitals referring patients to institution for emergent PCI. When complementary PCI was required, abciximab was conventionally given (ie, a 0.25 mg/kg bolus followed by a 0.125 mg/kg/min infusion during 12 hours). After completion of PCI, subcutaneous enoxaparin (100 IU/kg) was injected twice a day at the most during the first 72 hours, if necessary. All patients received oral clopidogrel (300 mg), followed by 75 mg daily for 1 year, plus 75–300 mg/day oral 		
Patients and investigator s	failures resulted from technical difficulties	 Need for an intra-aortic balloon pump or temporary pacemaker History of a coronary artery bypass graft 		
to treatment allocation. All other outcomes were as reported by the	inability to selectively catheterise the coronary	Demographics and baseline characteristics see below Drug therapy		
	anatomical variations of aortic roots, painful brachial artery	either an intravenous bolus of heparin as follows: unfractionated heparin 50 IU/kg with an upper limit of 4000 IU in patients > 75 years old, or low molecular weight heparin (enoxaparin) 30 mg intravenously and 1 mg/kg subcutaneously in patients aged > 75 years, and a bolus of aspirin (250 mg intravenously). This protocol was used		
	expertise	started by emergency units, mobile intensive care units or hospitals referring patients to institution for emergent PCI. When complementary PCI was required, abciximab was conventionally given (ie, a 0.25 mg/kg bolus followed by a 0.125 mg/kg/min infusion during 12 hours). After completion of PCI, subcutaneous		
		a day at the most during the first 72 hours, if necessary. All patients received oral clopidogrel (300 mg), followed by 75 mg		

	aspirin.			
Demographics and baseline characte	eristics			
Characteristics	Radial (n = 57)	Femoral (n = 57)	p value	
Age (years), mean(SD)	60(12)	58(13)	0.27	
Men, n(%)	49(86.0)	47(82.5)	0.80	
Diabetes, n(%)	12(21.1)	9(15.8)	0.48	
Hypertension, n(%)	30(52.6)	17(29.8)	0.01	
Current smoker, n(%)	42(73.7)	45(78.9)	0.66	
Hypercholesterolaemia, n(%)	28(49.1)	22(38.6)	0.26	
Systolic blood pressure (mm Hg), mean(SD)	123(21)	123(25)	0.93	
Delay: pain onset-cath-lab (min) , mean(SD)	409(305)	358(14)	0.33	
LVEF (%), mean(SD)	46(9)	49(11)	0.18	
Myocardial infarction topography, n(%)			
Anterior	- 31(54.4)	26(45.6)	5.3	
• Posterior	23(40.4)	27(47.4)		
• Lateral	3(5.3)	5(8.8)		
Angiographic and procedural charact	teristics			
Diffusion of the coronary artery disea	se			
One-vessel disease, n(%)	27(47.4)	30(52.6)		
Two-vessel disease, n(%)	16(28.1)	20(35.1)	0.22	
• Three-vessel disease, n(%)	14(24.6)	7(12.3)		
Target lesion, n(%)				
Left anterior descending coronary	29(50.9)	26(45.6)		
Left circumflex	8(14.0)	8(14.0)	0.83	
Right coronary	20(35.1)	23(40.4)		
Pre-PCI TIMI flow grade, n(%)				
• 0	27(47.4)	31(54.4)		

• 1	5(8.8)	5(8.8)	0.87
• 2	11(19.3)	9(15.8)	
• 3	14(24.6)	12(21.1)	
Coronary angiography duration (min), mean(SD)	17(8)	13(6)	< 0.01
Crossover, n(%)	7(12.3)	1(1.8)	0.03
Indication of PCI, n(%)			
Primary PCI	26(45.6)	32(56.1)	
Rescue PCI	28(49.1)	20(35.1)	0.65
Facilitated PCI	3(5.3)	5(8.8)	
Stents (n), mean(SD)	1.15(0.36)	1.28(0.61)	0.22
Direct stent implantation, n(%)	29(50.9)	27(47.4)	0.87
Stent with predilatation, n (%)	25(43.9)	27(47.4)	
Angiographic success of PCI, n(%)	52(91.2)	55(96.5)	0.43
Duration of PCI (min), mean(SD)	28(14)	26(18)	0.72
Delay: pain onset to TIMI 3 flow (min), mean(SD)	450(46)	381(31)	0.22
Contrast medium for PCI (ml), mean(SD)	97(57)	91(47)	0.45
Overall fluoroscopy duration (min), mean(SD)	13(9)	8(6)	< 0.01

Thrombolysis in myocardial infarction (TIMI) major bleeding; Heamoglobin drop of > 50 g/I, or intracranial haemorrhage or cardiac tamponade

Thrombolysis in myocardial infarction (TIMI) major bleeding; Heamoglobin drop of > 30 g/l but < 50 g/l, with bleeding from a known sight or spontaneous gross haematuria, haemoptysis or haematemesis

Groin haematoma: Local induration of > 4 cm diameter

Ecchymosis: Cutaneous bruise or induration of > 4 cm diameter or both

Effect size

Outcome Radial (n = 57) Femoral (n = 57) p value

STEMI Clinical evidence tables

In-hospital all-cause mortality, n(%)	3(5.3)	3(5.3)	NS
Ischaemic complication due to in-stent thrombosis, n(%)	4(7.0)	4(7.0)	NS
Duration of hospitalisation (days), mean(SD) 7.5(0.4) 0.59	7.2(0.5)	7.5(0.4)	NS
TIMI major bleeding, n(%)	3(5.3)	3(5.3)	NS
TIMI minor bleeding, n(%)	0(0)	1(1.8)	NS
Haematoma	2(3.5)	11(19.3)	0.05
Ecchymosis	6(10.5)	9(15.8)	NS
Transfusion, n(%)	1(1.8)	0(0)	NS
NS; not significant			

G.4 Thrombus extraction during PPCI

Table 48: AIMI 2006²

Reference	Study type	Number of patient	s Intervention		Outcome measures	Source of funding
Ali, Cox, Dib.et al. Rheolytic thrombectomy with percutaneous coronary intervention for infarct size reduction in acute myocardial infarction. Journal of the American college of Cardiology. Vol. 48, No. 2. 2006	Prospective multicentre RCT US and Canada 2001–2004	n = 480 Timing of randomisation: after diagnostic angiogra confirmed vessel sires > 2.0 mm Length of follow-ur 30 days	sephy Inclusion: > 18 years of age segment elevation of > 1 m contiguous leads within V1 MI (new ST-segment elevations)	e; anterior MI (new ST- mm in at least 2 to V6) or large inferior tion of >1 mm in 2 nd a VF); presented et; and reference in diameter ction fraction <35%, blood pressure <80 opic support), ent with GP IIb/IIIas, s 6 weeks, or history of	Primary end point infarct size. 30-day MACE (death, new C wave MI, emergency CABG, target vessel revascularisation, stroke or stent thrombosis)	
Baseline characteristics:						
Characteristic			chanical thrombus extraction (n = 0) (%)	No thrombectomy	(n = 240) (%) p	value
Age (years), mean		60		59.9	0	.92
Male, n (%)		182	2 (75.8)	178 (74.2)	0	.75
Diabetes, n (%)s		40	(16.7)	28 (15.8)	0	.81
Hypertension (requiring me	dication, n (%)	103	3 (42.9)	101 (42.1)	0	.93
Dyslipidemia requiring med	ication, n (%)	53	(22.1)	61 (25.4)	0).52
Smoker in past year, n (%)		106	5 (44.2)	108 (45.0)	O).71

Prior coronary artery disease, n (%)	42 (17.5)	33 (13.8)	0.26
Prior stroke, n (%)	7 (2.9)	10 (4.2)	0.62
Rescue after failed fibrinolysis, n (%)	34 (14.2)	32 (13.3)	0.90
Prior percutaneous intervention in target vessel	10 (4.2)	8 (3.3)	0.81
Time from symptom onset to emergency departmentt (h), mean (SD)	2.4 (3.3)	2.5 (3.2)	0.76
Time from emergency department to randomisation (h), mean (SD)	2.7 (4.3)	2.5 (3.2)	0.61
Total time to procedure (min), mean (SD)	75.4 (30.9)	59.6 (26.8)	< 0.001
Before or during the procedure	228 (95.0)	226 (94.2)	0.84
Number of major coronary arteries with ≥50% stenosis, n (%)			
1	112 (46.7)	116 (48.3)	
2	86 (35.8)	78 (32.5)	
3	39 (16.3)	44 (18.3)	
Stents, n (%)			
None	16 (6.3)	13 (5.3)	
1	154 (64.5)	173 (72.1)	
2	57 (23.8)	45 (18.8)	
3 or more	13 (5.4)	9 (3.8)	
Crossover to RT treatment, n (%)		6 (0.25)	
Angiographic inclusion criteria: Did not require angiographica	lly visible thrombus.		

Angiographic inclusion criteria: Did not require angiographically visible thrombus.

Concomitant therapy: All patients received 325 mg oral aspirin, clopidogrel 300 mg loading dose, clopidogrel 75 mg daily (cont. 4 weeks after) (ticlopidine 500mg loading dose and 250 mg twice a day dose given in event of intolerance to clopidogrel), unfractionated heparin during procedure to achieve activated clotting time >250 s. Use of beta-adrenergic blockers, nitroglycerine, and ACE-I before and after procedure at discretion of individual physician.

Results:	Mechanical thrombus extraction, n = 240	No thrombectomy, n = 240	p value
MACE, n (%)	16 (6.7)	4 (1.7)	0.01
All-cause mortality, n (%)	11 (4.6)	2 (0.8)	0.02
Q-wave MI, n (%)	0	0	-
Stroke, n (%)	4 (1.7)	2 (0.8)	0.69
Emergent CABG, n (%)	0	0	-

TLR/SAT, n (%)	5 (2.1)	1 (0.4)	0.22
Bleeding: requiring transfusion, n (%)	16 (6.7)	17 (7/1)	1.00
Bleeding: GI/GU, n (%)	11 (4.6)	15 (6.3)	0.55
Bleeding: retroperitoneal, n (%)	3 (1.3)	6 (2.5)	0.50
Bleeding: pericardial haemorrhage/tamponade, n (%)	2 (0.8)	0	0.50
Cardiogenic shock, post-procedure, n (%)	0	1 (0.4)	1.00
Ventricular fibrillation: pre-procedure, n (%)	12 (5.0)	5 (2.1)	0.34
Ventricular fibrillation: during/post-procedure, n (%)	7 (2.9)	3 (1.3)	
Ventricular tachycardia: non-sustained, n (%)	28 (11.7)	19 (7.9)	0.22
Ventricular tachycardia: sustained, n (%)	3 (1.3)	4 (1.7)	1.00

Table 49: Antoniucci 2004³

Reference	Study type	Number of patients	Intervention	Outcome measures	Source of funding
Antoniucci, Valenti, et al. Comparison of rheolytic thrombectomy before direct infart artery stenting versus direct stenting alone in patients undergoing Percutaneous coronary intervention for acute myocardial infarction. The American Journal of Cardiology. Vol. 93, 2004 15; 93(8):1033-5.	RCT Data collected November 2002 – June 2003 Italy	n = 100 Sample size calculation to detect 80% power and type 1 error of 0.05, 50 people in each group. Timing of randomisation: after angiography Length of follow up: 1 month	Mechanical thrombus extraction; anjojet rheolytic thrombectomy as an adjunct to conventional PCI or to conventional PCI alone. Inclusion Chest pain > 30 min and ST-segment elevation of ≥ 1 mm in at least 2 contiguous leads Exclusion Prior MI, administration of FT therapy, LBBB or ventricular pacing on baseline ECG preventing analysis of ST-segment changes, IRA diameter < 2.5 mm, inability to give informed consent	All-cause mortality Reinfarction Stroke Target vessel revascularisation Major bleeding	Not specified

Baseline characteristics:	Thrombectomy (n = 50)	No thrombectomy (n = 50)	p value
Age (years), mean (SD)	63 (13)	66 (12)	0.251
Men, n (%)	41 (82)	39 (78%)	0.617
Current smoker, n (%)	19 (38%)	14 (28)	0.288
Hypertension, n (%)	18 (36)	19 (38)	0.836
Cholesterolemia > 200 mg/dl, n (%)	23 (46)	24 (48)	0.841
Diabetes mellitus, n (%)	9 (18)	8 (16)	0.790
Angina pectoris, n (%)	10 (20)	4 (8)	0.084
Anterior wall acute myocardial infarction, n (%)	17 (34)	23 (46)	0.221
Cardiogenic shock, n (%)	3 (6)	6 (12)	0.295
Infarct coronary artery: left anterior descending, n (%)	17 (34)	23 (46)	
Infarct coronary artery: right, n (%)	26 (52)	21 (42)	
Infarct coronary artery: circumflex, n (%)	7 (14)	6 (12)	
Multivessel coronary disease, n (%)	15 (30)	20 (40)	0.295
Preprocedural TIMI grade flow 0–1, n (%)	38 (76)	40 (80)	0.629
Time-to-treatment, h, mean (SD)	3.9 (2)	4.4 (2.8)	0.295
Infarct artery stenting, n (%)	49 (98)	49 (98)	1.000
Direct stenting, n (%)	47 (94)	41 (82)	0.065
Multiple stents, n (%)	12 (24)	13 (26)	0.817
Stent length (mm), mean (SD)	20.7 (9.2)	21.1 (20.1)	0.848
Rheolytic thrombectomy, n (%)	48 (96)	4 (8)	< 0.001
Abciximab administration, n (%)	49 (98)	49 (98)	1.000
Intra-aortic balloon counterpulsation	3 (6)	5 (10)	0.461
Angiographic inclusion criteria:			

Concomitant therapy: All patients received abciximab if not contraindicated before the procedure to achieve activated clotting time of 200 to 300 seconds. Routine aspirin (325 mg/day indefinitely) and ticlopidine (500mg/day for 1 month) or clopidogrel (75 mg/day for 1 month).

Results:					
Outcome	Mechanical thrombus extraction (n = 50)	No thrombectomy (n =50)	p Value		
All-cause mortality, n	0	0			
Reinfarction, n	0	0			
Target vessel revascularisation, n	0	0			
Major bleeding requiring blood transfusion, n	0	1	0.315		
Disabling stroke, n	1 (2)	0	0.315		

Table 50: Beran 2002⁹

Reference	Study	Number of patients	Patient characteristics	Intervention	Comparison	Outcomes	Source of funding
Beran G, Lang I, Schreiber W, Denk S, Stefenelli T, Syeda B, Maurer G, Glogar D, Siostrzonek P. Intracoronary thrombectomy with the X-sizer catheter system improves epicardial flow and accelerates ST-segment resolution in patients with acute coronary syndrome: a prospective, randomized,	RCT Austria Randomisation 1:1 ratio Allocation concealment Not stated Blinding Investigators blinded to outcome	n = 66 Initially 66 patients were randomised, and 5 were subsequentl y excluded Length of follow-up: 30 days.	Inclusion criteria Chest pain for > 30 minutes and ST- segment elevation > 1 mm in 2 or more contiguous leads on the 12- lead ECG, rescue PCI was defined as PCI within 12 hours after failed systemic fibrinolysis. patients with unstable angina included if presented with recurrent chest pain at rest associated with ischaemic ST- segment or T-wave changes Exclusion criteria Not stated Demographics and baseline characteristics see below Drug therapy	n = 30 Mechanical thrombus extraction; X-sizer catheter system (EndiCOR Medical Inc) consists of a dual-lumen catheter shaft connected to a handheld control module.3 Two catheter sizes, 1.5 mm (7F compatible) and 2- mm (8F compatible)	n = 31 Standard PPCI	All-cause mortality Target vessel revasculari sation	Not stated

controlled study. Circulation. 2002 May 21; 105(20):2355- 60.	Patients with STEMI received unfractionated heparin, acetylsalicylic acid (blockers, and analgesics unless contraindicated, at operation; patients with unsurangina were pretreated with low-molecular-weight hasa, and nitroglycerin	(ASA), beta perators stable	
Baseline characteristics	No thrombectomy (n = 31)	Mechanical thrombus extraction (n = 30)	p value
Age (years), mean (SD)	53.9 (10.0)	55.0 (9.9)	NS
Male, n (%)	24 (77)	22 (73)	NS
STEMI, %	74	77	NS
Risk factors, %			
• Diabetes	13	17	NS
 Hypertension 	36	53	
Rescue PCI	10	23	
 Current smoking 	55	57	
Baseline angiographic and procedural da	ta		
Target vessel, %			
 Left anterior descending coronary artery 	10	9	NS
 Left circumflex coronary artery 	9	3	
 Right coronary artery 	12	18	
Results			
One month	No thrombectomy (n = 31)	Mechanical thrombus extraction (n = 30)	
All-cause mortality, n	1	2	
Target vessel revascularisation, n	1	0	

Table 51: Bulum 2012¹⁷

Reference	Study	Number of patients	Patient characteristics	Intervention	Comparison	Length of outcome	Outcomes	Source
Bulum J, Ernst A, Strozzi M. The impact of successful manual thrombus aspiration on in-stent restenosis after primary PCI: angiographic and clinical follow-up. Coronary Artery Disease. 2012; 23:487-491.	RCT Croatia Randomisation 1:1 ratio; no other details were reported Allocation concealment Not reported Blinding Not reported Intention to treat Yes	n = 60	Inclusion criteria Symptoms suggesting acute myocardial ischaemia >20 minutes, the onset of symptoms <12 hours previously, and ST-segment elevation of > 0.1 mV in 2 or more contiguous ECG leads. Exclusion criteria Rescue PCI after failed fibrinolysis, cardiogenic shock, triple-vessel disease, significant left main stem stenosis, previous PCI of an infarct-related artery, previous coronary artery bypass grafting, and known existence of a concomitant disease with life expectancy less than 6 months. Demographics and baseline characteristics see below Drug therapy Before PPCI Aspirin (300 mg) and clopidogrel (600 mg), weight adjusted dose of unfractionated heparin, and glycoprotein IIb/IIIa inhibitor (eptifibatide) at the discretion of the operator. After PPCI Beta-blockers, high-dose lipid-lowering agents, angiostensin-converting enzyme inhibitors or angiostensin-II receptor	Thrombus aspiration with Export Aspiration catheter	Standard PPCI	6 months	Primary Referent vessel diameter, minimal lumen diameter (MLD), lesion length, and perentage diameter stenosis (acute gain, and late lumen loss) Secondary Incidence of major adverse cardiocerebro vascular events (death, reinfarction, stroke, and the need for TLR)	None stated

	antagonists, and w-3-acid ethyl esters				
Demographics and baseline characteristics:					
Characteristics	Thrombus aspiration PPCI (n = 30) Standard PPCI (n =	= 30)		
Age (years), mean (SD)	54.3 (8.6)	58.5 (9.7)			
Men, n (%)	25 (83)	22 (73.3)			
Hypertension, n (%)	12 (40)	13 (43.3)			
Diabetes, n (%)	3 (10)	3 (10)			
Current smoker, n (%)	16 (53.3)	15 (50)			
Hyperlipdaemia, n (%)	25 (83.3)	26 (86.7)			
Family history of CAD, n (%)	14 (46.7)	11 (36.7)			
Angiographic data and procedural results					
Number of stents					
1, n (%)	19(63.6)	18(60.0)			
2, n (%)	9(30.0)	9(30.0)			
3, n (%)	1(3.9)	3(10.0)	3(10.0)		
4, n (%)	1(3.9)	0	0		
Left anterior descending, n (%)	14 (46.7)	11 (36.7)	11 (36.7)		
Left circumflex, n (%)	4 (13.3)	8 (26.7)	8 (26.7)		
Right coronary, n (%)	12 (40.0)	11 (36.7)	11 (36.7)		
Results at 6 month follow-up					
	Thrombus aspiration PPCI (n = 30)	Standard PPCI (n = 30)	p value		
All-cause mortality, n	0	0	1		
Reinfarction, n	0	0	1		
Target vessel revascularisation, n	8	5	0.347		

Table 52: EXPORT 2008²⁵

Reference	Study type	Number of patients	Intervention	Length of follow-up	Outcome measures	Source of funding
Chevalier B, Gilard M, et al. Systematic primary aspiration in acute myocardial Percutaneous intervention: a multicentre randomised controlled trial of the export aspiration catheter. EuroIntervention. 2008: 222-228.	RCT France, Austria, Italy, India Randomised using computerised telephone system.	n = 249 Inclusion Consecutive patients > 18 years old, with 2\mm or more of ST-segment elevation in 2 or more contiguous leads, with visual reference vessel diameter ≥\2.5 mm, and TIMI flow 0 or 1 before placing the wire in the infarct-related artery Exclusion Cardiogenic shock, cardiac arrest at any time before the intervention, pre- catheterisation with lytic agents, with GPI Ilb/Illa, or with pacemakers; a current medical condition with expected survival less than a year, and current participation in other investigations	Thrombus aspiration using export aspiration catheter by stenting versus conventional stenting without thrombectomy	30 days	All-cause mortality Myocardial reinfarction	Not stated
		Data collected May 2005 – April 2007				

Baseline characteristics:

	No thrombectomy (n = 129)	Thrombus aspiration PPCI (n = 120)	p value
Age (years), (SD)	61.2 (12.9)	59.2 (12.8)	0.18
Male (%)	81.4	80.8	0.91
Prior Q-wave MI (% patients)	8.5	9.2	0.86
Prior non-Q-wave MI (%)	2.3	1.7	1.00
Prior CABG (%)	0.0	0.8	0.48
Prior non-target vessel PCI (%)	2.3	3.3	0.71
Prior target vessel PCI (%)	3.1	5.0	0.53
Smoking (%)	35.7	42.5	0.18
Diabetes (%)	13.2	16.7	0.44

NIDDM	94.1	65.0	0.05
IDDM	5.9	35.0	
Hypertension (%)	44.2	41.7	0.69
Hypercholesterolaemia (%)	41.9	36.7	0.40
Family history of cardiovascular disease (%)	25.6	32.5	0.23
Killip class ≥ 2 (% patients)	10.9	11.6	0.69
Symptom onset to randomisation, min			
Mean	271.4 (197.6	321.7 (413.5	0.53
Median (range)	219 (145–362.5)	225 (149–333)	

Angiographic inclusion criteria: The choice of PCI including stent treatment was at the investigator's discretion, with permissible options including predilatation followed by stention, direct stenting, and stenting with postdilatation.

Concomitant therapy: Aspirin, heparin, clopidogrel, and GPI IIb/IIIa inhibitors at investigator's discretion, and administered according to standard hospital procedure.

Results

	No thrombectomy (n = 129)	Thrombus aspiration PPCI (n = 120)	p value
Predilatation performed (%)	55.8	28.3	< 0.001
Postdilatation performed (%)	18.6	8.3	0.02
Number of stents used, mean (SD)	1.17 (0.58)	1.28 (0.58)	0.08
Procedure time, mean (SD)	34.5 (21.5)	36.7 (18.0)	0.08
Median (IQR)	30 (20–45)	32 (25–45)	
Bailout performed (% patients)	14.7	5.8	0.02
No thrombectomy (n = 129)	Thrombus aspiration PPCI (n = 120)	57.1 (4)	
Rescue distal protection	0	0	
Rescue aspirations, , % (n)	42.1 (8)	42.9 (3)	
30-day cardiac mortality, n	5	3	
Myocardial reinfarction, n	1	2	

Table 53: De Luca 2006³⁰

Reference	Study type	Number of par	tients	Intervention	Length of follow-up	Outcome measures	Source of funding
De Luca L, Sardella G, Davidson C, et al. Impact of intracoronary aspiration thrombectomy during primary angioplasty on left ventricular remodelling in patients with anterior ST- elevation myocardial infarction. Heart. 2006, 92: 951-957	RCT	Inclusion >18 years with anterior STEMI identifiable thr on IRA at coror angiography. Exclusion Coronary bypa grafting, 3 vess coronary arter disease, severe heart disease, grade 2 or 3 flo time of initial angiography, o unsuccessful P defined as no anterograde flo >50% residual in the IRA.	I and rombus nary ass sel ry e valvar TIMI bow at	Thrombus aspiration with Diver CE aspiration thrombectomy catheter (Invatex, Brescia, Italy).	6 months	All-cause mortality Chronic heart failure Myocardial infarction	Not stated
Baseline characteristics:							
			No throm	nbectomy (n = 38)	Thrombus aspirat	ion PPCI (n = 38)	
Age (years)			64.6 (12.5	5)	66.7 (14.1)		
Men, n (%)			21 (55.3)		27 (71)		
Hypertension, n (%)		19 (50)		15 (39.5)			
Diabetes, n (%)			7 (18.4)	9 (23.7)			
Smoking, n (%)			10 (26.3)		7 (18.4)		

Family history of CAD, n (%)	14 (36.8)	5 (13.1)*
Cholesterol (mmol/l), n (%)	4.32 (0.39)	4.17 (0.28)*
Triglycerides (mmol/l), n (%)	1.41 (0.29)	1.37 (0.44)
Renal failure, n (%)	3 (7.9)	5 (13.1)
Killip class III, n (%)	11 (28.9)	8 (21)
Previous PCI, n (%)	4 (10.5)	7 (18.4)
Symptoms to balloon (h), n (%)	7.6 (1.8)	7.2 (1.9)
Medication at follow-up:		
ACE inhibitors/ARBs, n (%)	31 (81.6)	36 (94.7)
Aldosterone blockers, n (%)	22 (57.9)	19 (50)
Beta-blockers, n (%)	27 (71)	31 (81.6)
Statins, n (%)	36 (94.7)	32 (84.2)
Aspirin, n (%)	37 (97.4)	37 (97.4)
Data are made (CD) ar number (0/)		

Data are mean (SD) or number (%)

Angiographic inclusion criteria:

Concomitant therapy: Aspirin 300 mg orally and heparin 8000 IU intravenously before the procedure and with abciximab as a 0.25 mg/kg bolus and 0.125 ug/kg/min intravenous infusion immediately before the revascularisation and continued for 12 hours. Heparin for 48 hours, aspirin 100mg/day, and ticlopidine (250 mg orally twice a day for at least 4 weeks) or clopidogrel (loading does of 300 mg followed by 75 mg/day for at least 4 weeks. Other adjunctive pharmacotherapy was administered at the discretion of the operator.

Results:

6 months	No thrombectomy (n = 38)	Thrombus aspiration PPCI (n = 38)	
Mortality, n	2	0	
New onset MI, n	0	1	
Hospitalisation for chronic heart failure, n	3	2	
MACE, n	4	3	

^{*}p < 0.05 between conventional and thrombectomy groups

Table 54: DEAR-MI 2006¹⁰²

Reference	Study type	Number of patients	Intervention	Length of follow-up	Outcome measures	Source of funding
Silva-Orrego P, Colombo P, Bigi R, Thrombus aspiration before primary angioplasty improves myocardial reperfusion in acute myocardial infarction. The DEAR-MI (Dethrombosis to enhance acute reperfusion in myocardial infarction) study. Journal of the American College of Cardiology, Vol. 48, No. 8, 2006.	RCT Italy Data collected March 2004 to June 2005 12% dropout due to protocol exclusions	n = 148 Inclusion: continuous chest pain ≥ 30 min, ST-segment elevation > 0.1mV (0.2 mV in case of anterior leads) in ≥3 contiguous leads on 12-lead ECG, and technical feasibility for primary angioplasty independently of initial TIMI flow or angiographic evidence of intraluminal thrombus in the culprit artery. Exclusion: cardiogenic shock, previous myocardial infarction or CABG, bundle branch block or pacemaker induced rhythm on admission ECG, and contraindication to GP IIb/IIIa inhibitors.	Standard PPCI with stenting and abciximab versus thrombus aspiration using Pronto extraction catheter (Vasc.solutions, Minneapolis, Minnesota) plus standard PPCI.	30-days	Mortality, myocardial reinfarction, left ventricular failure, Target vessel revascularisation.	Not stated
Baseline characteristics:		Thrombus aspiration PPCI	(n = 74) No th	ombectomy (n = 74	l) p Valu	a
Age, years (SD)		57.3 (13)	58.9 (1		0.472	
Male, n (%)		62 (84)	56 (76)		0.314	
Diabetes, n (%)		16 (21)	11 (15)		0.465	
Hypertension, n (%)	28 (37)		32 (46)		0.349	
Dyslipidemia, n (%)		26 (34)	18 (25)	0.309	
Smoking, n (%)		38 (54)	43 (60)	0.571	
Killip class >1, n (%)		8 (11)	4 (5)		0.297	
Anterior infarction, n (%)		32 (42)	38 (51		0.355	

Ejection fraction, n (%)	53 (7)	51 (9)		0.133
Ischaemic time, min (SD)	206 (115)	199 (124)		0.722
Target vessel				
Left anterior descending coronary artery, n (%)	32 (43)	38 (51)		0.420
Left circumflex coronary artery, n (%)	7 (10)	10 (14)		0.623
Right coronary artery, n (%)	35 (47)	26 (35)		0.191
CAD extension				
1 vessel, n (%)	36 (49)	36 (49)		0.876
2 vessels, n (%)	23 (31)	24 (32)		0.965
3 vessels, n (%)	15 (20)	14 (19)		0.958
TIMI flow				
0/1, n (%)	60 (81)	54 (73)		0.657
2, n (%)	12 (16)	16 (22)		0.646
3, n (%)	2 (3)	4 (5)		0.822
Glycoprotein IIb/IIIa inhibitors, n (%)	100	100		
Stenting, n (%)	73 (99)	72 (97)		
Direct stenting, n (%)	52 (70)	18 (24)		< 0.0001
Procedural time (min), mean (SD)	57 (19)	53 (21)		0.363
Results:				
	Thrombus aspiration, n = 74		No thrombus aspiration,	n = 74
In-hospital mortality, n	0		0	
In-hospital myocardial reinfarction, n	0		0	
In-hospital left ventricular failure, n	0		1	
In-hospital target vessel revascularisation, n	1		0	

Table 55: EXPIRA 2010⁹⁶

Reference	Study	Number of patients and Patient characteristics	Intervention	Compari son	Length of outcome	Outcomes	Source of funding
Sardella G, Mancone M, Canali E, Di RA, Benedetti G, Stio R et al. Impact of thrombectomy with EXPort Catheter in Infarct-Related Artery during Primary Percutaneous Coronary Intervention (EXPIRA Trial) on cardiac death. American Journal of Cardiology. 2010; 106(5):624-629.	RCT Italy Randomisation 1:1 manner Allocation concealment Not stated Blinding Single blind Intention to treat Unclear	n = 175 Inclusion criteria First STEMI within 9 h from symptoms onset ischaemic chest pain > 30 min and ST-segment elevation ≥ 2mm in ≥ 2 contiguous electrocardiographic leads, infarct-related artery 2.5 mm diameter, identifiable lesion, thrombus score ≥ 3, TIMI flow grade O to 1, and age > 18 years Exclusion criteria Previous PCI on infarct-related artery, previous CABG, cardiogenic shock, 3-vessel disease, left main disease, severe valvular heart disease, fibrinolysis, fPPCI, contraindication to glycoprotein Ilb/IIIa inhibitors administration Demographics and baseline characteristics see below Drug therapy Before PPCI Patients treated with 300 mg aspirin, heparin to maintain clotting time > 250 seconds, abciximab (0.25 mg/kg followed by intravenous infusion at 0.125 microgram/kg/minute) Subsequently patients received heparin for 48 h, aspirin 100 mg/day, clopidogrel (300 mg loading dose, followed by 75 mg/day for ≥ 12 months) Adjunctive therapy at operator discretion; standard therapies after PPCI; beta blockers, lipid lowering drugs, ACE or ARB inhibitors	Thrombus aspiration with Export Medronic PCI (EM-PCI)	Standard PPCI	2 years	All-cause mortality, Myocardial reinfarction, Target vessel revascularisati on	None stated

Demographics and baseline characteristics:			
Characteristics	No thrombectomy (n = 88)	Thrombus extraction (n = 87)	p value
Age (years), mean (SD)	64.6 (12.5)	66.7 (14.1)	0.298
Men, n (%)	48.0 (51.1)	57 (64.7)	0.218
Hypertension, n (%)	43 (41.4)	59.0 (67.2)	0.021
Diabetes, n (%)	16.0 (18.4)	21 (23.8)	0.459
Smoking, n (%)	23.0 (26.4)	43.0 (48.8)	0.003
Cholesterol mg/dl), mean (SD)	167 (15)	161 (11)	0.002
Renal failure, n (%)	7 (8.0)	7 (7.9)	1
Killip class III, n (%)	25.0 (28.7)	17.0 (19.3)	0.160
Symptom to balloon (h), mean (SD)	6.1 (1.8)	6.2 (0.9)	0.642
LVEF, n (%)	40.7 (9.3)	42.0 (10.5)	0.192
MV coronary disease, n (%)	16.0 (18.4)	21.0 (23.8)	0.459
Direct stenting, n (%)	2.0 (2.3)	67.0 (76.2)	0.0001
Drug eluting stent, n (%)	53.0 (60.9)	49.0 (55.7)	0.540
Location of infarct –related coronary artery			
Left anterior descending, n (%)	38.0 (43.7)	38.0 (43.2)	1
Left circumflex, n (%)	20.0 (23.0)	22.0 (25.0)	0.859
Right coronary, n (%)	29.0 (33.3)	28.0 (31.8)	0.872
Results at 2 year follow-up			
	Thrombus aspiration PPCI (n = 88)	No thrombectomy (n = 87)	p value
All-cause mortality, n	0	6	0.0001
Reinfarction , n	0	1	0.999
Target vessel revascularisation, n	4	5	0.651

Table 56: INFUSE-AMI 2012¹⁰⁶

Reference	Study type	Number of patients	Intervention	Length of follow-up	Outcome measures	Source of funding
Stone G, Maehara A, Witzenbichler R, et al. Intracoronary abciximab Ind aspiration Ihrombectomy in patients With large anterior Impocardial infarction. The NFUSE-AMI randomised Irial. JAMA. 2012; 307 17): 1817-1826.	RCT USA Multicentre, single-blind Data collected November 2009- December 2011 Powered for infarct size at 30-days	Inclusion: >18 years, symptoms consistent with STEMI longer than 30 minutes' duration and 1 mm or greater of ST-segment elevation in 2 or more contiguous leads in V1-V4, or new LBBB, with anticipated symptom onset to device time of 5 hours of less Exclusion: contraindications to study medications or contrast; prior MI, bypass graft surgery or LAD stenting; planned surgery necessitating antiplatelet agent interruption, contraindication to cMRI; known creatine clearance less than 30 mL/min/1.73 m2, dialysis, platelet count less than 100000 or > 7000000 cells/mm3, or haemoglobin level less than 10 g/dL; recent major bleeding, bleeding diathesis, or current warfarin use; history of intracranial disease; ischaemic stroke or transient ischaemic attack within 6 months or any permanent neurologic defect; pre-randomisation cardiogenic shock or cardiopulmonary resuscitation; prior fibrinolysis or IIb/IIIa inhibitor for the present admission; and any comorbid conditions likely to interfere with protocol compliance or associated with less than 1-year survival.	Thrombus aspiration performed with 6 F Export Catheter (Medtronic), PPCI for anterior STEMI, PCI performed using standard techniques with bare metal or drugeluting stent implantation at operator discretion Patient randomised to either: a) thrombus aspiration followed by intracoronary bolus abciximab, (b) thrombus aspiration without abciximab, (c) intracoronary plus abciximab without aspiration, or (d) no abciximab and no aspiration	30 day	MACE: All-cause mortality Myocardial reinfarction New onset severe heart failure, or rehospitalisation for heart failure. MACCE: all-cause mortality, reinfarction, stroke, or clinically driven Target vessel revascularisation.	Atrium medical

Baseline characteristics:		
	Thrombus aspiration PPCI + no abciximab (n = 111)	No aspiration PPCI + no abciximab (n = 112)
Age, median (IQR), years	62.0 (53.0–73.0)	62.5 (52.5–71.0)
Male, n/total (IQR)	85/111 (76.6)	81/112 (72.3)
Body mass index, median (IQR) ^a	26.8 (24.3–30.5)	26.6 (24.0–28.7)
Killip class, n/total (%)		
T	82/110 (74.5)	90/112 (80.4)
II	13/110 (11.8)	13/112 (11.6)
III	0/110	2/111 (1.8)
Hypertension n/total (%)	39/111 (35.1)	36/112 (32.1)
Hyperlipidemia n/total (%)	18/111 (1632)	14/112 (12.5)
Diabetes mellitus n/total (%)	19/110 (17.3)	8/112 (7.1)
Prior myocardial infarction n/total (%)	1/110 (0.9)	0/112
Prior Percutaneous coronary intervention n/total (%)	3/111 (2.7)	3/112 (2.7)
Cigarette smoking, current n/total (%)	46/109 (42.2)	55/112 (49.1)
Symptom to hospital arrival, median (IQR), min	107.0 (66.5–152.5)	98.0 (67.0–136.0)
Hospital arrival to first device, median (IQR), min ^b	42.0 (30.0–61.0)	46.5 (34.0–70.5)
Symptom onset to first device, median (IQR)	151 (117–205)	160 (126–217)
Infarct artery lesion location ^c :		
Proximal left anterior descending n/total (%)	68/111 (61.3)	74/112 (66.1)
Mid left anterior descending n/total (%)	47/111 (39.6)	48/112 (42.9)
Left ventricular ejection fraction, median (IQR), % ^d	40.0 (38.0–50.0)	40.0 (31.0–50.0)
a , , , , , , , , , , , , , , , , , , ,	_	

^a calculated as weight in kilograms divided by height in meters squared

Angiographic inclusion criteria: infarct lesion to be located in the proximal or mid LAD with visually assessed TIMI 0–2 flow, and absence of excessive tortuosity, diffuse disease, heavy calcification, or significant left main disease.

Concomitant therapy: 324 [sic] aspirin orally, or 250 to 500 mg intravenously; clopidogrel, 600 mg, or prasugrel, 60 mg. Patients undergoing PCI received procedural anticoagulation with bivalirudin (intravenous bolus 0.75 mg/kg plus infusion of 1.75 mg/kg per hour, discontinued at procedure end. Following procedure, all patients

^b Balloon angioplasty, local drug delivery, or aspiration

^c Some patients had both proximal and mid left anterior descending lesions

^d From contrast left venticulography during the index procedure

were treated with aspirin indefinitely and with clopidogrel or prasugrel for at least 1 year. Discharge medicines included aspirin in 99% of patients, clopidogrel in 66.4%, prasugrel in 31.8%, statins in 97.7%, beta-blockers in 96.6% and angiotensin-converting enzyme inhibitors or receptor blockers in 94.1%, with no significant differences between the groups.

Results:

30-day outcomes ^a	Aspiration thrombectomy (n = 229)	No thrombectomy ^c (n = 223)	p value
All-cause mortality, n (%)	7 (3.1)	6 (2.7)	0.81
Reinfarction, n (%)	1 (0.5)	2 (0.9)	0.55
New-onset severe heart failure, n (%)	8 (3.5)	9 (4.1)	0.77
Rehospitalisation for heart failure, n (%)	0	2 (0.9)	0.15
Stroke, n (%)	0	1 (0.5)	0.31
Clinically driven total vascular revascularisation, n (%)	1 (0.5)	4 (1.8)	0.17
HORIZONS-AMI major bleeding, n (%)	9 (4.0)	10 (4.6)	0.79
TIMI major bleeding, n (%)	2 (0.9)	4 (1.8)	.40
TIMI minor bleeding, n (%)	1 (0.5)	2 (0.9)	0.55
GUSTO severe bleeding, n (%)	9 (4.0)	10 (4.5)	0.77
GUSTO moderate, n (%)	2 (0.9)	1 (0.5)	0.58
GUSTO mild bleeding, n (%)	1 (0.4)	4 (1.8)	0.17
Any blood product transfusion	2 (0.9)	3 (1.4)	0.64
Thrombocytopenia, in-hospital ^d , n/total (%)	1/186 (0.5)	3/189 (1.6)	0.62

^aData are Kaplan-Meier estimates

^cPooled, either with or without intracoronary abciximab

d<100 000 cells/mm3 in patients with a baseline platelet count > 150 000 cells/mm³ (n = 384)

Table 57: ITTI 2012⁷⁵

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow- up	Outcome measures	Source of funding
Liu CP, Lin MS, Chiu YW, Lee JK, Hsu CN, Hung CS, Kao HL. Additive benefit of glycoprotein llb/Illa nhibition and adjunctive chrombus aspiration during primary coronary ntervention: results of the nitial Thrombosuction and Tirofiban nfusion (ITTI) crial. nternational lournal of Cardiology. 2012; 156(2):174-179.	Design RCT (5 hospital centres in Taiwan) Enrolment Not stated Randomisation 2 x 2 factorial design Allocation concealment Not stated Blinding Not stated Sample size calculation Based on previous studies a sample size of 22 patients in each group ware required to achieve the power of 0.80 for the primary end point of MBC 3 rate ITT analysis All randomised patients were included in ITT comparisons of outcome (regardless of whether crossed over) ad baseline characteristics:	n = 100 2 x 2 factorial study and 2 2 arms relevant to the review (47 patients) Drop- outs: n = 0 Crossove r: n = 2	Inclusion criteria Patients > 18years with STEMI undergoing PPCI within 12 hours of symptom onset Exclusion criteria <18 years cardiogenic shock history of bleeding tendency hepatic or renal insufficiency major operation within 6 weeks contraindication to tirofiban use Demographics and baseline characteristics see below Drug therapy All patients received aspirin (300 mg loading followed by 100 mg daily), and clopidogrel (300 mg loading followed by 75 mg daily) and unfractionated heparin 100 IU /kg before the procedure	Thrombus aspiration with thrombuster device	Standard PPCI	6 months	MACE All-cause mortality Reinfarction Target lesion revascularisati on Stroke	Not stated

Characteristics	No thrombectomy (n = 23)	Thrombus extraction (n = 24)
Age, year (SD)	57 (13)	62 (9)
Male, n (%)	20 (87)	21 (87.5)
Smoker, n (%)	13 (56.5)	10 (40)
Hypertension, n (%)	13 (56.5)	14 (58.3)
Diabetes mellitus, n (%)	5 (21.7)	7 (29.2)
Hypercholesteremia, n (%)	7 (30.4)	4 (16.7)
Previous PCI, n (%)	0 (0)	1 (4.2)
Body-mass index, n (%)	25 (2)	25 (3)
Cardiogenic shock, n (%)	2 (8.7)	0 (0)
Killip class III or IV, n (%)	2 (8.7)	0 (0)
Systolic blood pressure (mmHg)	144 (31)	148 (26)
Diatolic blood pressure (mmHg)	82 (18)	83 (18)
Heart rate (bpm), n (%)	85 (24)	80 (13)
Number of disease vessels 1 2 3	11 (47.8) 6 (26.1) 6 (26.1)	8 (33.3) 10 (41.7) 6 (25.0)
Infarct related vessel	0 (20.1)	0 (23.0)
Left anterior descending artery	12 (52.2)	15 (62.5)
Results	No thrombectomy (n = 23)	Thrombus extraction (n = 24)
All-cause mortality (6 months), n	0	1
Reinfarction (6 months) , n	3	0
Target lesion revascularisation (6 months), n	3	1
Stroke (6 months) , n	0	1

Table 58: JETSTENT 2010⁸³

Reference	Study	Number of patients and Patient characteristics	Intervention	Comparison	Length of outcome	Outcomes	Source of funding
Migliorini, Stabile, Rodriguez et al. Comparison of angioJet rheolytic thrombectomy before direct infarct artery stenting with direct stenting alone in patients with acute myocardial infarction. The JETSTENT Trial. Journal of the American College of Cardiology. 2010: 56(16) 1298-306	RCT (multicentre, international trial) Italy, Germany Poland, Argentina, Randomisation Randomised to thrombectomy before direct stenting or direct stenting alone was performed after coronary angiograpy and wiring of the infarct artery. The TIMI study thrombus score was used for the assessment of thrombus dimensions. Patients with angiographic evidence of TIMI thrombus grade 3−5 after infarct artery wiring could be randomised if the reference diameter of the infarct artery was ≥ 2.5mm on visual assessment. Angiographic criteria for exclusion from randomisation included: 1)TIMI thrombus grade < 3; infarct artery reference diameter < 2.5 mm on visual assessment; previous stenting of the infarct artery; and inability to identify the infarct artery	Inclusion criteria Patients with ST-segment elevation AMI were considered eligible for the study without restriction based on age or clinical status on presentation, patients with cardiogenic shock were eligible Acute MI diagnosed according minutes and M 12 hours and 2) at least 2 contiguous leads or branch block Exclusion criteria 1) Thombolysis for current AMI; 2) Major surgery <6 weeks; 3) Stroke <30 days or any history of haemorrhagic stroke; 4) Comorbidities with expected survival <1 year; and participation in another study Demographics and baseline characteristics see below	Mechanical thrombus aspiration; angioJet rheolytic thrombectomy system consists of a drive unit console, a disposable pump set and a 4-F disposable catheter. The single-pass anterograde thrombectomy technique was used. Multivessel intervention was allowed only in patients with cardiogenic shock; otherwise, procedures in noninfarct arteries were performed after the 1-month scintigraphy	In both arms, direct stenting was attempted in all cases using baremetal stents. The stent type choice was at the operator's discretion. If the stent failed to directly cross the lesion, predilation was performed using an balloon to risk of before stent. Pressure and were at the	1, 6 and 12 months Baseline and 30- min post procedur e electroc a rdiogram undertak en	All-cause mortality Myocardial infarction Target vessel revascularisation Stroke	Medrad Interventic al/Possis (Minneapos, Minnesota Provided financial support ar thrombect my devices only (not involved in the manageme t, collectio or analysis of data

out by computer-generated
sequence and assignments
were provided by a
centralised telephone system

Allocation concealment

Blinding

All clinical events were adjudicated by an independent clinical event committee blinded to treatment allocation after review of original source documentation

Drug therapy

Patients received 325mg of aspirin orally, or 250mg intravenously at the emergency room, and a loading dose of 600mg of clopidogrel before or immediately after the procedure. All patients received abciximab unless contraindicated. Abciximab was administered immediately before or during the procedure as a bolus of 0.25 mg/kg body weight followed by a 12 hour infusion at a rate of 0.125 microgram/kg/minute. Heparin was given as an initial bolus of 70U/kg and additional boluses were administered during the procedure to achieve an activated clotting time of 200 to 250 seconds. After the procedure, patients were treated with aspirin (100 to 325 mg daily indefinitely) and clopidogrel (75 mg daily for 6 months). Other drugs such as beta blockers, angiotensinconverting enzyme inhibitors and statins were used in accordance with standard and recommended practice.

the operator according to the characteristi c of the stent and lesion

Baseline characteristics			
	Mechanical thrombus aspiration (n = 256)	No thrombectomy (n = 245)	p value
Age, years (range)	63 (12.3 (24–94)	64.3 (11.5 (34–91)	0.208
Male, n (%)	195 (76)	199 (81)	0.168
Hypertension, n (%)	120 (47)	116 (47)	0.916
Dyslipidemia, n (%)	77 (30)	85 (35)	0.270
Diabetes mellitus, n (%)	36 (14)	37 (15)	0.742
Previous myocardial infarction, n (%)	10 (3.9)	12 (4.9)	0.588
Anterior myocardial infarction, n (%)	101 (39)	91 (37)	0.595
Cardiogenic shock, n (%)	7 (2.7)	13 (5.3)	0.142
Previous PCI, n (%)	11 (4.3)	10 (4.1)	0.904
Previous CABG, n (%)	1 (0.4)	1 (0.4)	0.975
Symptoms to admission (min), median (interquartile ranges)	125 (85–221)	135 (86–227)	0.853
ST-segment elevation (mm), mean (SD)	3.98 (2.49)	4.02 (2.69)	0.886
Baseline angiographic characteristics			
Infarct artery			
• Left anterior descending artery, n (%)	107 (42)	91 (37)	0.483
• Right coronary artery, n (%)	112 (44)	120 (49)	
• Circumflex artery, n (%)	37 (14)	34 (14)	
Reference vessel diameter (mm), median (interquartile ranges)	29.4 (2.67–3.24)	2.91 (2.62–3.25)	0.67
Multivessel disease, n (%)	114 (44)	95 (39)	0.192
Pre-wiring TIMI flow grade 0–1, n (%)	212/254 (83.5)	203/242 (83.9)	0.899
Post-wiring TIMI flow grade 0–1, n (%)	142/231 (61.5)	129/222 (58.1)	0.465
TIMI thrombus grade post-wiring			
• 1–2, n (%)	3 (1.4)	3 (1.4)	0.64
• 3, n (%)	73 (32.5)	80 (37.4)	

• 4, n (%)	83 (37.4)	79 (36.9)	
• 5, n (%)	63 (28.4)	52 (24.3)	
Procedural characteristics (values are m	edian (interquartile range), n (%) or mean	(SD).	
Procedural time (minutes, median (interquartile range)	59.5 (44.7–70)	46 (35–60)	< 0.001
Stent per patient, mean (SD)	1.26 (0.54)	1.40 (0.73)	0.022
Multiple stenting, n (%)	58 (23)	72 (30)	0.079
Abciximab, n (%)	249 (97)	239 (98)	0.841
Procedural success *, n (%)	237 (92.6)	229 (93.6)	0.696
* Residual stenosis < 30% and TIMI flow	grade 3 by operator's assessment.		
Results			
1 month	Mechanical thrombus aspiration	No thrombectomy	(n = 245)
	(n = 256)		
All-cause mortality, n (%)	4 (1.6)	7 (2.9)	
Myocardial infarction, n (%)	2 (0.8)	3 (1.2)	
Target vessel revascularisation, n (%)	2 (0.8)	6 (2.5)	
Stroke, n (%)	0 (0)	1 (0.4)	
Stroke, n (%) 6 months	0 (0)	1 (0.4)	
	0 (0) 7 (2.8)	1 (0.4) 11 (4.5)	
6 months			
6 months All-cause mortality, n (%)	7 (2.8)	11 (4.5)	
6 months All-cause mortality, n (%) Myocardial infarction, n (%)	7 (2.8) 2 (0.8)	11 (4.5) 3 (1.2)	

TIMI: thrombolysis in myocardial infarction

Table 59: Kaltoft et al. 2006⁶¹

Reference	Study type	Number of pa	tients	Intervention	Length of follow-up	Outcome measures	Source of funding
Kaltoft A, Bottcher M, Nielsen S, et al. Routine thrombectomy in Percutaneous coronary intervention for acute ST- segment elevation myocardial infarction: a randomised, controlled trial. Circulation 2006, 114:40-47	Data collected 2004–2005 Randomisation using varying block sizes stratified by sex and diabetes.	but < 12 h and elevation ≥2 n Exclusion crite branch block, infarction with fibrinolytics tr coronary bypa stenosis, need ventilation, ar	aptoms lasting > 30 min I cumulative ST-segment InV in ≥2 contiguous leads Pria were left bundle- acute myocardial Inin the previous 30 days, eatment, previous Iss surgery, left main stem I for mechanical Ind sever heart failure Intra-aortic balloon pump	Thrombus aspiration versus no thrombectomy	30 days	All-cause mortality Reinfarction Stroke.	Boston Scientific, Denmark.
Baseline characteristics:			Thrombectomy (n = 108)		No thrombecto		
Age, y, mean (SD)			65 (11)		63 (13)	illy (II – 107)	
Male, n (%)			82 (76)		86 (80)		
Diabetes mellitus, n (%)			9 (8)		6 (6)		
Current smoker, n (%)			59 (55)		69 (64)		
Hypertension*, n (%)			33 (31)		22 (21)		
Hypercholesterolemia*			10 (9)		10 (9)		
Prior myocardial infarction,	n (%)		14 (13)		11 (10)		
Symptom onset to first ballo	oon inflation (minute	es) mean	242 (171–321)		208 (155–329)		
ST-segment elevation imme	ediately before PCI, i	mV, n (%)	0.65 (0.35–1.3)		0.7 (0.3–1.1)		
Systolic blood pressure befo	ore PCI, mmHg, n (%)	133 (114–150)		130 (117–155)		
Killip class ≥			7 (6)		(4)		
* Requiring medication, n (9	%)						

Abciximab administered, n (%)	104 (96)	100 (93)	0.37
Stent implanted, n (%)	103 (95)	104 (97)	1.0
Device success, n (%)	96 (89)	-	

Concomitant therapy: Before intervention: aspirin 300 mg orally or intravenously, clopidogrel 300 mg orally, and unfractionated heparin 100000 IE intravenously. During the intervention, all treated with abciximab. Post intervention: infusion of abciximab for 12 hours and aspirin 75 mg/d and clopidogrel 75 mg/day for 12 months.

Results:

	Thrombectomy (n = 108)	No thrombectomy (n = 107)	p value
Total procedure time (min), mean (IQR)	39 (29–48)	29 (23–38)	< 0.0001
All-cause mortality, n	0	1	
Reinfarction, n	0	1	
Disabling stroke, n	2	0	

Table 60: Liistro 2009⁷⁴

Reference	Study	Number of patients and Patient characteristics	Intervention	Comparison	Length of outcome	Outcomes	Source
Liistro F, Grotti S, Angioli P, Falsini G, Ducci K, Baldassarre S et al. Impact of thrombus aspiration on myocardial tissue reperfusion and left ventricular functional recovery and remodeling after primary	RCT Italy Randomisation 1:1 manner, computer generated random series of numbers. And block randomisation (blocks of 10 patients) Allocation concealment	Inclusion criteria Symptoms suggesting acute myocardial ischaemia lasting > 30 minutes, the onset of symptoms < 12 hours previously, and ST-segment elevation of > 0.1 mV in 2 or more leads on the ECG, patients without contraindication to the use of platelet glycoprotein IIb/IIIa inhibitors. Exclusion criteria Rescue PCI after fibrinolysis, previous MI, the absence of an optimal echocardiographic apical view, the known existence of a disease resulting in a life expectancy of < 6 months, and the lack of	Thrombus aspiration with Export Aspiration catheter; 6 F catheter	Standard PPCI	6 months	Cardiac mortality Target vessel revascularisati on Heart failure	None stated

angioplasty. Circulation Cardiovascular	Physicians unaware of block randomisation	informed consent, patients with poor images	
Interventions.	Blinding	Demographics and baseline characteristics	
2009; 2(5):376- 383.	Single bind, outcomes blinded to operator	see below	
		Drug therapy	
	Intention to treat	Before PPCI	
	Yes	Aspirin (a loading dose of 500 mg), heparin (70 IU/kg), clopidogrel (a loading dose of 600 mg), glycoprotein IIb/IIIa inhibitor abciximab with an intravenous procedural bolus of 0.25 mg/kg followed by a continuous intravenous infusion of 0.125 microgram/kg/minute for 12 hours and postprocedural infusion without heparin	
Demographics ar	nd baseline characteristics:		
Characteristics		No thrombectomy (n = 88)	Thrombus extraction (n = 87)
Age (years), mear	n (SD)	65 (11)	64 (11)
Men, n (%)		43 (77)	43 (64.7)
Hypertension, n (%)	43 (41.4)	59.0 (67.2)
Diabetes, n (%)		16.0 (18.4)	21 (23.8)
Current smoker, i	n (%)	23.0 (26.4)	43.0 (48.8)
Hyperlipdaemia,	n (%)	167 (15)	161 (11)
History of CAD, n	(%)	7 (8.0)	7 (7.9)
Killip class ≥ III, n	(%)	25.0 (28.7)	17.0 (19.3)
Symptom to ballo	oon (min), mean (SD)	6.1 (1.8)	6.2 (0.9)

Symptom to door (min), mean (SD)

Door to balloon (min), mean (SD)

MV coronary disease, n (%)

Direct stenting, n (%)

Stented patients, n (%)

21.0 (23.8)

67.0 (76.2)

49.0 (55.7)

16.0 (18.4)

53.0 (60.9)

2.0 (2.3)

Location of infarct–related coronary artery			
Left anterior descending, n (%)	38.0 (43.7)	38.0 (43.2)	1
Left circumflex, n (%)	20.0 (23.0)	22.0 (25.0)	0.859
Right coronary, n (%)	29.0 (33.3)	28.0 (31.8)	0.872
Results at 2 year follow-up			
· ·			
	Thrombus aspiration PPCI (n = 88)	No thrombectomy (n = 87)	p value
All-cause mortality, n	•	No thrombectomy (n = 87)	p value 0.0001
	88)		·

Table 61: Napodano 2003⁸⁵

Reference	Study type	Number of patients		Intervention	Length of follow-up		tcome asures	Source of funding
Napodano M, Pasquetto G, Sacca S. Intracoronary thrombectomy improves myocardial reperfusion in patients undergoing direct angioplasty for acute myocardial infarction. Journal of the American College of Cardiology. Vol. 42. No. 8. 2003.	RCT Italy Randomisati 1:1 manner	Inclusion: continuous chest pain for at le 30 minutes and within 12 hours of onset pain, and ST-segment elevation ≥1 mm (mV) in 2 or more contiguous leads, angiographic evidence of intraluminal thrombus in the IRA, TIMI flow ≤2, or ≥70 diameter stenosis. Exclusion: presence of LBBB or pacemake induced rhythm at admission ECG, left m stem lesions, IRA diameter <2.5 mm.	of 0.1 0% er-	Mechanical thrombus extraction with X- sizer catheter	30-days	mo hea No reir Stro Sev Tar	cause rtality Chronic art failure, nfatal nfarction oke rere bleeding get vessel ascularisation	Not reporte d
Baseline characteristics:								
	Mechanical thrombus extraction (n = 46)		No	No thrombectomy (n = 46)			p value	
Age (years), mean (SD)		61.3 (10.8)	63.0	63.6 (11.7)		0.33		
Male, mean 82.6		71.	71.7		0.32			
Smokers								

Current, %	45.6	34.8	0.39
Former, %	20.5	30.4	0.34
Hypertension, %	60.9	65.2	0.38
Dyslipidemia, %	50.0	52.1	1.0
Diabetes, %	13.0	13.0	1.0
Previous MI, %	17.4	6.5	0.19 [¥]
Anterior MI, %	39.1	43.5	0.83
Time from onset of symptoms to hospital presentation (min), mean (SD)	202.9 (204.9)	165.7 (134.7)	0.54*
Time from hospital presentation to angioplasty (min), mean (SD)	35 (12)	38 (15)	0.82
Multivessel disease, %	52.2	41.3	0.46
Killip class, mean (SD)	1.5 (1.0)	1.5 (0.9)	0.83*
Killip class IV, %	8.7	8.7	1.0
HR > 100 beats/min, %	19.6	13.0	0.57
BP < 100 mm Hg, %	15.2	15.2	1.0
*Wilcoxon rand-sum test, ^ categorical da	ata are presented as frequency values and	were compared by chi-square, [¥] Fisher ex	act test
Stents, %	93.5	91.3	1.0
Intraaortic balloon pump, %	10.9	10.9	1.0
Baseline TIMI flow, mean(SD)	1.46(1.24)	1.48(1.31)	0.93
TIMI flow final, mean(SD)	2.91(0.35)	2.89(0.53)	0.82
Concomitant therapy			
GP IIIb/IIa inhibitors, %	43.4	41.3	1.0
In-hospital	Mechanical thrombus extraction (n = 46)	No thrombectomy (n = 46)	p value
All-cause mortality, %	6.5	6.5	1.0*
Chronic heart failure, %	10.9	21.7	0.17^
Nonfatal reinfarction, %	2.2	2.2	1.0*
Stroke, %	0	0	1.0*
Severe bleeding, %	2.2	2.2	1.0*

Target vessel revascularisation, %	0	0	1.0*		
30 days					
All-cause mortality, %	6.5	6.5	1.0*		
Reinfarction, %	4.3	4.3	0.2*		
Stroke, %	0	0	0*		
Target vessel revascularisation, %	0	0	1.0*		
* Data compared using Fischer exact test, or ^ chi-square					

Table 62: PIHRATE 2010³⁴

D D M	DCT		Intervention	Comparison	outcome	Outcomes	funding
Burzotta F, Gasior M, Witkowski A, Horvath IG et al. Thrombus aspiration followed by direct stenting: a novel strategy of primary percutaneous coronary intervention in ST- segment elevation myocardial infarction. Results of the Polish- Italian-Hungarian RAndomized ThrombEctomy Trial (PIHRATE Trial).	RCT Poland Hungary Italy Randomisation 1:1 ratio Allocation concealment Not stated Blinding Investigators blinded to outcome	Inclusion criteria Patients with first STEMI with 6 hours from chest pain onset, ≥ 2 mm ST elevation in at least 2 contiguous leads and , ≥ 3 mm ST elevation in a lead, occluded infarct related artery Exclusion criteria Prior MI, CABG, cardiogenic shock, treated with fibrinolysis before admission to cath lab Demographics and baseline characteristics see below Drug therapy	n = 100 Thrombus aspiration with DIVER 6 F compatible catheter	n = 96 Standard PPCI	In-hospital: All-cause mortality Reinfarction Target vessel revascularisa tion Heart failure 6 months All-cause mortality Reinfarction	In-hospital: All-cause mortality Reinfarction Target vessel revascularisa tion Heart failure 6 months All-cause mortality Reinfarction	Not stated

Patients received aspirin (325 mg), clopidogrel loading dose, unfractionated heparin (70 U/kg), GPPI IIIb/IIa inhibitors at operators discretion					
Baseline characteristics					
	No thrombectomy (n = 96)	Thrombus aspiration PPCI (n = 100)	p value		
Age (years), mean (SD)	58.8 (10.3)	58.8 (10.3)	NS		
Male, %	80	81.7	NS		
Risk factors, %					
• Diabetes	13	9.6	NS		
Hypertension	58.0	53.7	NS		
Current smoking	63.7	62.0	NS		
Prior angina	4.3	13.0	0.033		
Baseline angiographic and procedural dat	ta				
Target vessel, %					
 Left anterior descending coronary artery 	39.6	39.0	NS		
 Left circumflex coronary artery 	12.5	11.0			
Right coronary artery	47.9	50.0			
Results					
In-hospital	No thrombectomy (n = 96)	Thrombus aspiration PPCI (n = 100)			
All-cause mortality, n	3	3			
Reinfarction, n	1	0			
Target vessel revascularisation, n	1	2			
Heart failure, n	6	10			
6 months					
All-cause mortality, n	3	3			
Reinfaction, n	1	0			

Table 63: REMEDIA 2005¹⁹

Reference	Study	Number of patients and Patient characteristics	Intervention	Comparison	Length of outcome	Outcomes	Source of funding
Burzotta F, Trani C, Romagnoli E, Mazzari MA, Rebuzzi AG, De VM et al. Manual thrombus- aspiration improves myocardial reperfusion: the randomized evaluation of the effect of mechanical reduction of distal embolization by thrombus- aspiration in primary and rescue angioplasty (REMEDIA) trial. Journal of the American College of Cardiology. 2005; 46(2):371-376.	RCT (single centre) Italy Randomisation Undertaken after enrolment and before coronary angiography 1:1 randomisation using a computer generated random series of numbers Allocation concealment Not stated Blinding Cardiologist analysing ECG was blinded to procedural and clinical data. Otherwise not stated ITT analysis used	Inclusion criteria All patients within 12 hours of onset of STEMI referred for primary or rescue PCI to a catheterisation laboratory were entered for the study. Exclusion criteria No angiographic exclusion criteria. Demographics and baseline characteristics see below Drug therapy All patients were treated by heparin (initial weight adjusted intravenous bolus then further boluses administered with the aim of obtaining an activated clotting time of 250 to 300s in patients treated with abciximab and >300s in the remaining subjects) and with double antiplatelet therapy with aspirin and clopidogrel (loading dose of 300mg followed by 75 mg/day) for at least 4 weeks. Unless contraindicated, abciximab (0.25 mg/kg bolus plus infusion of	PCI with thrombus-aspiration n = 50, 1 had no significant culprit lesion, 1 had surgical coronary anatomy, therefore 48 eligible for procedure 1 failure to cross the lesion with the guidelines, 1 death during PCI 2 protocol violations (crossover to standard PCI because of low thrombus burden)	Standard PPCI n = 49, all eligible for standard PCI 1 failure to cross the lesion with guidewire, 2 died during PCI. 4 protocol violations (crossover to thrombusaspiration because of high thrombus burden/distalembolism) After crossing of the target lesion with the guidewire, direct stent implantation was	30 day	All-cause mortality Reinfarction, Stroke Target vessel revascularisation	None stated

0.125 microgram/kg/minute for 12 aspirating attempted if hours) was intravenously device: Diver judged administered in all patients CE (Invated, possible by undergoing PPCI, whereas in those Brescia, the with failed fibrinolysis, abciximab Italy). A operator, in use was left to the operator's rapid the discretion. exchange, 6remaining F compatible cases, prethrombusdilation with aspirating an catheter undersized balloon was used before stent implantatio

NB. Patients who died during the procedure and had non-crossable target lesions were considered to have no reperfusion in the analyses. Data of the patients whose treatment crossed over were included in the assigned group and analysed according to ITT principle.

Baseline characteristics

No differences reported between groups for clinical or angiographic characteristics.

	Thrombus aspiration PPCI (n = 50)	No thrombectomy (n = 49)	p value
Age (years) (mean (SD)	61 (13	60 (13	0.76
Gender: males (%) / females (%)	45 (90) / 5 (10)	38 (77.6) / 11 (22.4)	0.09
Risk factors, n (%)			
Smokers	31 (62)	26 (53.1)	0.37
 Hyperchloesterolemia 	27 (54)	17 (34.7)	0.06
Hypertension	31 (62)	28 (57.1)	0.62
 Diabetes mellitus 	11 (22)	9 (18.4)	0.65
 Positive family history 	15 (30)	11 (22.4)	0.96
Previous history of ischaemic heart disease, n (%)	10 (20)	10 (20.4)	0.96
Pre-infarction angina, n (%)	14 (28)	16 (32.7)	0.61
Anterior myocardial infarction, n (%)	20 (40)	25 (51)	0.27
Symptoms to angiography time (min)	274 (137)	300 (202)	0.28

Referred after failure of fibrinolysis	16 (32)	12 (24.5)	0.41
Use of abciximab, n (%)	34 (68)	31 (63.3)	0.53
Renal failure (creatine ≥ 1.2mg/dl)	6 (12)	8 (16.3)	0.53
Killip class III or IV, n (%)	15 (30)	14 (28.6)	0.88
Cardiogenic shock, n (%)	4 (8)	5 (10.2)	0.74
Multivessel disease, n (%)	17 (34)	21 (42.9)	0.36
LAD as culprit vesse, n (%)	20 40)	25 (51)	0.15
Culprit lesion with proximal location, n (%)	26 (52)	21 (42.9)	0.27
Pre-intervention TMI flow grade, n (%)			
• 0	32 (64)	34 (69.4)	0.48
• 1	11 (22)	10 (20.4)	
• 2	2 (4)	3 (6.1)	
• 3	5 (6.3)	2 (4.1)	

Results (at 30 days)		
	Thrombus aspiration PPCI (n = 48)	No thrombectomy (n = 48)
All-cause mortality, n (%)		
In the cath lab	1 (2)	2 (4.1)
After PCI	2 (4)	1 (2.1)
Reinfarction, n (%)	2 (4)	2 (4.1)
Stroke, n (%)	1 (2)	1 (2.1)
Target lesion revascularisation, n (%)	1 (2)	1 (2.1)

Table 64: TAPAS 2008^{109,118}

Reference	Study type	Number of patients	Intervention	Length of follow-up	Outcome measures	Source of funding
Vlaar P, Svilaas T, van der Horst, Diercks G, et al. Cardiac death and reinfarction after 1 year in the thrombus aspiration during Percutaneous coronary intervention in acute myocardial infarction study (TAPAS): a 1-year follow-up study. Lancet. 2008; 371(9628):1915-1920 Svilaas T, Vlaar PJ, van der Horst IC, Diercks GF, de Smet BJ, van den Heuvel AF et al. Thrombus aspiration during primary percutaneous coronary intervention. New England Journal of Medicine. 2008; 358(6):557-567.	RCT Netherlands Single centre Data collected January 2005 to December 2006 Patients randomised BEFORE coronary angiography	Inclusion criteria: symptoms suggesting acute myocardial ischaemia >30 minutes, time from symptoms onset <12 h, ST-segment elevation > 0.1 mV in 2 or more lead on the ECG Exclusion: rescue PCI after fibrinolysis and known existence of concomitant disease with life expectancy less than 6 months	Thrombus aspiration with Export aspiration catheter versus conventional PPCI.	30 days and 1 year	All-cause mortality Myocardial reinfarction Target vessel revascularisation	Medtronic

Baseline characteristics:

	Thrombus aspiration PPCI (n = 535)	Conventional PCI (n = 536)
Age, mean (SD)	63 (13)	63 (13)
Men, n (%)	363 (67.9%)	392 (73.1%)
Diabetes, n/total (%)	56/530 (10.6%)	67/532 (12.6%)
Hypertension, n/total (%)	171/517 (33.1%)	195/526 (37.1%)
Hypercholesteroleamia, n/total (%)	115/485 (23.7%)	130/480 (27.1%)
Previous myocardial infarction, n/total (%)	50/528 (9.5%)	57/533 (10.7%)

Previous PCI, n/total (%)	39/526 (7.4%)	38/531 (7.2%)
Previous CABG, n/total (%)	17/529 (3.2%)	22/533 (4.1%)
Family history, n/total (%)	235/509 (46.2%)	229/514 (44.6%)
Body mass index, mean (SD)	27 (4)	27 (4)
Current smoking, n/total (%)	213/463 (46.0%)	225/469 (48.0%)
Total ischaemic time (min), median (IQR)	190 (110–270)	185 (107–263)
Infarct –related vessel, n/total (%)		
Left anterior descending artery	221/515 (42.9%)	223/517 (43.1%)
Left circumflex artery	93/515 (18.1%)	79/517 (15.3%)
Right coronary artery	180/515 (36.7%)	204/517 (39.5%)
Other	12/515 (2.3%)	11/517 (2.1%)
Initial TIMI flow, n/total (%)		
0/1	288/526 (54.7%)	316/531 (59.5%)
2	102/526 (19.4%)	85/531 (16.0%)
3	136/526 (25.9%)	130/531 (24.5%)

Concomitant therapy: All patients pre-treated with aspirin (500 mg followed by 80–100 mg per day), heparin (5000 IU), and clopidogrel (loading dose 600 mg followed by 75 mg per day). Unless contraindicated, abciximab given during procedure and additional heparin guided by activated clotting time. Standard therapies after PCI included beta-blockers, lipid-lowering agents, and angiotensin-converting-enzyme inhibitors, or angiotensin-II receptor antagonists, according to current guidelines.

Results:

nesures.				
30 day follow-up	Thrombus aspiration, n = 529	No thrombectomy, n = 531	Risk ratio (95% CI)	p value
All-cause mortality, n (%)	11 (2.1)	21 (4.0)	0.52 (0.26-1.07)	0.07
Reinfarction, n (%)	4 (0.8)	10 (1.9)	0.40 (0.13-1.27)	0.11
Target vessel revascularisation, n (%)	24 (4.5)	31 (5.8)	0.77 (0.46-1.30)	0.34
Major bleeding, n (%)	20 (3.8)	18 (3.4)	1.11 (0.60-2.08)	0.11
1 year follow-up	Thrombus aspiration, n = 535	No thrombectomy, n = 536	Hazard ratio (95% CI)	p value
All-cause mortality, n (%)	25 (4.7)	41 (7.6)	1.67 (1.02–2.75)	0.042
Cardiac death, n (%)	19 (3.6)	36 (6.7)	1.93 (1.11–3.37)	0.020
Reinfarction, n (%)	12 (2.2)	23 (4.3)	1.97 (0.98–3.96)	0.05
Target vessel revascularisation, n (%)	60 (12.9)	69 (11.2)	1.19 (0.84–1.68)	0.34
Second PCCI target vessel, n (%)	37 (6.9)	51 (9.5)		

CABG target vessel, n (%)	25 (4.7)	20 (3.7)		
Cardiac death or non-fatal reinfarction, n (%)	30 (5.6)	53 (9.9)	1.81 (1.16–2.84)	0.009
Major adverse cardiac events, n (%)	89 (16.6)	109 (20.3)	1.26 (0.95–1.67)	0.10

Table 65: VAMPIRE 2008⁵⁷

Reference	Study type	Number of patients	Intervention	Length of follow-up	Outcome measures	Source of funding	
lkari Y, Sakurada et al. Upfront thrombus aspiration in primary coronary intervention for patients with ST-segment elevation acute myocardial infarction: report of the VAMPIRE (VAcuuM asPIration thrombus REmoval) trial. IACC: cardiovascular inventions. Vol. 1, No. 4, 2008.	RCT Japan Patients enrolled 2003- 2005 ITT analysis	Inclusion: 21 years with AMI presenting 30 min but 24 h after symptom onset, with 2 mm or more of ST-segment elevation in 2 or more contiguous leads or with a presumably new left bundle-branch block Exclusion: presence of primary fibrinolysis prior to randomisation, cardiogenic shock, history of cardiac arrest, history or coronary bypass surgery, chronic renal failure (Cr >2.0 mg/dl) or haemodialysis, left main disease, or target vessel <2.5 mm or >5 mm in diameter	PPCI with or without thrombus aspiration using Nipros's TransVascular Aspiration Catheter.	In-hospital 8 months	All-cause mortality, MACE (composite death, recurrence of myocardial infarction, and target lesion revascularisation)	Not stated	
Baseline characteristics:							
		Thrombus aspiration PPCI (n = 180)	No thrombecto	omy (n = 175)	p Value		
Age, (years), mean (SD) 63.2		63.2 (10.6)	63.5 (9.9)	63.5 (9.9)			
Male, %		80.6	77.7	77.7			
Body mass index, (kg/m²)		24.1	24.3	24.3		0.53	
Hypertension, %		54.8	59.0		0.45		

Hyperlipidemia, %	50.0	48.5	0.78
Diabetes, %	23.3	29.9	0.16
Insulin use, %	3.6	3.1	
Smoking			
Current smoking, %	56.6	50.9	0.26
Ex-smoker, %	12.1	9.3	
Family history of CAD, %	13.9	14.4	0.89
Previous PCI, %	3.4	5.2	0.42
Killip class > 1, %	11.2	8.4	0.17
Onset to hospital (h), mean (SD)	4.5 (5.0)	5.2 (5.5)	0.13
Stent use, %	94.1	93.4	0.79
IABP or percutaneous cardiopulmonary support, %	9.4	13.4	0.25
Transfemoral approach, %	94.8	92.4	0.34
Glycoprotein IIb/IIIa inhibitors, %	0	0	1
Procedural success, %	98.9	98.3	0.64
Vascular access complication, %	2.9	4.1	0.54

Concomitant therapy: Aspirin and intravenous heparin boluses were administered during he procedure to maintain an activated clotting time ≥300 s. No drug eluting stents were allowed. Use of fibrinolytic agents or GPI IIb/IIIa inhibitors was not allowed.

Results:

	Thrombus aspiration PPCI (n = 180)	No thrombectomy (n = 175)	p value
In-hospital mortality (composite death, recurrence of myocardial infarction, and target lesion revascularisation) , n	1	1	NS
MACE (in-hospital) , n	1 (0.6)	2 (1.2)	NS
Myocardial Infarction (in-hospital), n	0	1 (subacute thrombus)	NS
Target lesion revascularisation (in-hospital) , n	0	1	NS
8-month MACE (composite death, recurrence of myocardial infarction, and target lesion revascularisation) , n	12.9%	21.0%	< 0.05
All-cause mortality (8 months) , n	2	1	NS

Myocardial infarction (8 months) , n	0	1	NS
Target lesion revascularisation (8 months) , n	20	31	0.05

Table 66: X AMINE ST 2005⁷²

Reference	Study	Number of patients and Patient characteristics	Intervention	Comparison	Length of outcome	Outcomes	Source of funding
Lefevre, Garcia, Reimers, Lang et al. X-sizer for thrombectomy in acute myocardial infarction improves ST-segment resolution. Results of the X-sizer in AMI for negligible embolization and optimal ST resolution (X AMINE ST) trial. Journal of the American College of Cardiology. 2005: 46 (2) 246-52	RCT (14 European centres) France, Spain, Italy, Austria, Germany, UK Randomisation 1:1 ratio Allocation concealment Blinding Coronary angiograms were analysed by an independent core laboratory that was blinded to other data. ITT analysis used	Inclusion criteria Patients suffering an AMI who were amenable to PIC were included. AMI < 12 hours (that is, evidence of ischaemic chest pain for > 30 minutes and new ST-segment elevation for ≥ 2mm in 2 or more contiguous electrocardiographic leads, de novo lesion, single vessel treatment in a native vessel ≥ 2.5mm in diameter and occluded, thrombuscontaining, TIMI flow grade 0 to 1 infarct related artery. Exclusion criteria Previous PCI in IRA, rescue PCI, Killip class ≥3, left or right bundle branch block, IRA with excessive proximal tortuosity or severe calcification, left ventricular ejection fraction <30%, contraindication to emergency coronary artery bypass grafting and current participation in another study protocol	n = 100 Mechanical thrombus extraction X-sizer catheter system used. A 2- lumen over the wire system (diameters 1.5 and 2 mm) with a helical shape cutter at its distal tip. 2 or more passages across the lesion from proximal to distal were performed by slowly advancing the activated catheter. Subsequently, additional balloon angioplasty or coronary stenting was performed Before the intervention, all patients received aspirin. Heparin (70 U/kg) was given to maintain an activated	n = 101 PCI (excluding anything but balloon angioplasty and stent)	1 and 6 months ECG recorded 60 min after the procedure	All-cause mortality Stroke MI Target vessel revascularisation	Not stated

	characteristics see below Drug therapy Glycoprotein IIb/IIIa inhibitors were used according to the operator's judgement	250s	
Baseline characteristics			_
	No thrombectomy (n = 101)	Mechanical thrombus extraction (n = 100)	p value
Age (years), mean (SD)	62 (11)	61 (13)	NS
Female, %	27	24	NS
Risk factors, %			
Diabetes	18	25	NS
Hypertension	50	54	
Dyslipidemia	61	58	
Current smoking	51	52	
Previous MI, %	6	10	NS
Baseline angiographic and procedural dat	ta		
Farget vessel, %			
Left anterior descending coronary	48	55	NS
artery	5	7	
Left circumflex coronary artery	43	37	
Right coronary artery			
TIMI flow grade pre-PCI, %	0.18 (0.52	0.15 (0.46	NS
Glycoprotein IIb/IIIa inhibitors, %	65	55	NS
Coronary stenting, %	99	100	NS
Postprocedural angiographic data			
TIMI flow, % grade 3	89	95.9	0.105
Procedural success, %	75	84.5	0.112

Results						
One month	No thrombectomy	Thrombectomy	p value			
All-cause mortality, n	4	4	NS			
Stroke, n	0	2	NS			
Reinfarction	3	1	NS			
Target vessel revascularisation, n	0	2	NS			
6 months						
All-cause mortality, n	4	6	NS			
Stroke, n	0	2	NS			
Reinfarction, n	4	2	NS			
Target vessel revascularisation, n	5	3	NS			

G.5 Culprit versus complete revascularisation **Updated, see the 2020 evidence review**

Table 67: APEX-AMI 113

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Toma M, Buller CE, Westerhout CM, Fu Y, O'Neill WW, Holmes DR, Jr. et al. Non- culprit coronary artery percutaneous coronary intervention during acute ST-segment elevation myocardial infarction: insights from the APEX-AMI trial. European Heart Journal. 2010; 31(14):1701- 1707.	Post hoc analysis of RCT (APEX- AMI), subgroup population assessed as cohort	n = 2201	Inclusion criteria presentation within 6 hours high risk electrographic characteristics; 2 mm ST- elevation in 2 anterior lateral leads or at least 2 mm ST- elevation in 2 inferior leads couples with ST-depression in 2 contiguous anterior leads for a total of 8 mm or more or left bundle branch block Presence of maximum percent stenosis of 70% or greater in more than major epicardial coronary artery or a non-infarct-related vessel requiring intervention by PCI operators Exclusion criteria Rescue PCI Isolated inferior MI Pregnant or breastfeeding Complement deficiency Serious infection Serious medical condition that would likely alter recovery	Culprit only revascularisation (COR); the infarct related artery only was treated and the other arteries were left untreated n = 1984	Multivessel revascularisation(M VR); the infarct related artery and the non-infarct related artery within the same procedure n = 217	90 days	All-cause mortality	Proctor 8 Gamble Pharmace uticals Alexion Pharmace uticals

	Demographics ar characteristics See below Drug therapy After stent impla patients received 75 mg per day fo	ntation all I clopidogrel		
	weeks, other adjutreatments were administered at to of the operator	unctive		
Baseline characteristics				
	COR (n = 1984)	MVD (n = 217)	p value	
Male, %	79.2	79.4	77.4	0.498
Age, median (25 th and 75 th percentile)	64 (55, 73)	64 (55, 73)	64 (53, 74)	0.937
Heart rate, BPM, median (25th and 75th percentile)	75 (65, 87)	75 (64, 87)	77 (67, 88)	0.106
SBP, mmHg, median (25th and 75th percentile)	134 (116, 150)	134 (117, 150)	130 (112, 148)	0.069
Killip class >1, %	11.6	11.4	13.8	0.288
Inferior MI, %	44.7	45.5	37.0	0.017
Hypertension, %	54.8	55.6	47.5	0.022
Prior MI, %	15.5	15.9	12.4	0.185
Prior PCI, %	11.5	11.7	9.7	0.366
Prior CABG, %	3.6	4.0	0.5	0.009
Prior congestive heart failure,%	4.0	3.9	4.6	0.602
Diabetes mellitus, %	19.2	20.0	11.5	0.003
Current smoker, %	39.7	39.9	38.2	0.635
Prior MI, %	15.5	15.9	12.4	0.185

Creatine clearance, ml/min, median (25 th and 75 th percentile)	77.9 (59.8, 100.8)	78.4 (59.9, 100.7)	74.0 (58.6, 101.1)	0.632
IRA, %				
LAD	49.0	48.3	55.8	< 0.001
LCX	12.3	11.4	19.8	
RCA	38.5	40.0	24.4	
Extent of multivessel disease, %				
2-vessel disease	37.7	68.2	63.4	0.534
3-vessel disease	31.2	31.8	25.9	
Time from symptom onset to PCI, hours, median (25 th and 75 th percentile)	3.4 (2.6, 4.7)	3.4 (2.6, 4.7)	3.5 (2.7, 4.9)	0.436
In-hospital GPI use, %	72.6	71.9	78.8	0.030
Stent, %	94.7	94.6	95.9	0.436
Target vessel revascularisation, n (%)	28 (7.9)	1 (3.8)	8 (6.3)	0.66
Length of hospital stay, days, median (25 ^t , 75 th percentile)	6 (4,9)	6 (4,9)	6 (4,9.5)	0.203
Results				
90 day mortality				
	COR (n = 1984)	MVD (n = 214)	p value	
All-cause mortality, n (%)	119 (5.6)	27 (12.5)	< 0.001	
Procedure time, min	42	56	< 0.001	
GUSTO severe bleeding	0.6	1.8	p = 0.065	
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Table 68: Corpus 2004 ²⁹

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Corpus RA, House JA, Marso SP, Grantham JA, Huber KCJ, Laster SB et al. Multivessel xtfpercutaneo us coronary intervention in patients with multivessel disease and acute myocardial infarction. American Heart Journal . 2004; 148(3):493- 500.	Design Retrospect ive cohort study, 1998 to 2002, USA	n = 1982	Inclusion criteria clinical evidence of acute MI, defined as symptoms consistent with ongoing myocardial ischaemia, ECG evidence of acute MI (≥ 1mm ST elevation in ≥ 2 contiguous leads), new left bundle branch block, or true posterior infarction), or both presentation ≤ 12 hours before symptom onset procedure generally begun within 30 to 60 min of patients arrival at hospital Exclusion criteria patients undergoing PPCI of vein graft or left main coronary artery lesions patients undergoing PPCI for acute occlusion after coronary angioplasty or arteriography patients with multivessel disease undergoing stage revascularisation procedures of the non-infarct related artery after discharge from hospital	Culprit only revascularisation (COR); the infarct related artery only was treated and the other arteries were left untreated n = 354	Multivessel revascularisation(M VR); the infarct related artery and the non-infarct related artery within the same procedure n = 26 Staged revascularisation (SR); the infarct related artery was treated immediately and the non-infarct related artery was staged within the indexed hospitalisation n = 126	In-hospital ≤ 30 days 1 year	All-cause mortality Reinfarction Target vessel revascularisation CABG Major bleed	None stated

Baseline characteristics	characteristics see below Drug therapy After stent impla patients received 75 mg per day for weeks, other adj treatments were administered at of the operator	d clopidogrel or at least 4 unctive		
baseline characteristics	COD (** 354)	MANUS and CD associated to		
	COR (n = 354)	MVR and SR combined (n = 152)	p value	
Age, years, mean (SD)	63 (14)	64 (13)	0.27	
Diabetes, n (%)	60 (17)	29 (19)	0.56	
Male, n (%)	245 (69)	109 (72)	0 .57	
Hypertension, n (%)	194 (55)	74 (49)	0.21	
Creatine > 1.5mg /dl, n (%)	12 (3.4)	5 (3.3)	0.87	
Ejection fraction, %	35 (18)	35 (19)	0.96	
Killip class IV on admission, n (%)	12 (3.4)	5 (3.3)	0.95	
Glycoprotein IIb/IIIa inhibitors	139 (39)	44 (29)	0.03	
Stent, n (%)	307 (87)	148 (97)	< 0.001	
	COR (n = 354)	MVR (n = 26)	SR (n = 126)	p value
In-hospital	26 (7.2)	4 (2.0)	C (4.7)	0.54
Major bleeding, n (%)	26 (7.3)	1 (3.8)	6 (4.7)	0.51
All-cause mortality, n (%)	20 (5.6)	5 (19)	3 (2.4)	0.003
≤ 30 days	COR (n = 354)	MVR (n = 26)	SR (n = 126)	p value

Mortality, n (%)	23 (6.5)	5 (19)	10 (7.9)	0.06
Reinfarction, n (%)	2 (0.6)	0 (0)	14 (11)	< 0.001
Target vessel revascularisation, n (%)	28 (7.9)	1 (3.8)	8 (6.3)	0.66
CABG, n (%)	28 (8.0)	1 (3.8)	2 (2.4)	0.07
1 year	COR (n = 354)	MVR (n = 26)	SR (n = 126)	p value
Mortality, n (%)	42 (12)	5 (19)	12 (9.5)	0.36
Reinfarction, n (%)	10 (2.8)	1(3.8)	19 (15)	< 0.001
Target vessel revascularisation, n (%)	53 (15)	3 (12)	35 (28)	0.004
CABG, n (%)	41 (12)	2 (67.7)	8 (6.3)	0.21

Table 69: EUROTRANSFER Reg 2010³⁷

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Dziewierz A, Siudak Z, Rakowski T, Mielecki W, Dubiel JS, Dudek D. Impact of multivessel coronary artery disease and non-infarct related artery revascularisatio	Retrospective cohort study, 15 STEMI hospital networks from 17 European hospitals, Nov 2005 to Jan 2007	n = 777	Inclusion criteria STEMI ≥ 1 significantly stenosed epicardial coronary artery, without previous CABG Exclusion criteria None stated Demographics and	Culprit only revascularisation (COR); the infarct related artery only was treated and the other arteries were left untreated n = 707	Multivessel revascularisation(M VR); the infarct related artery and the non-infarct related artery within the same procedure n = 77	≤ 30 days 12 months	All-cause mortality Reinfarction Repeat revascularisation, Major bleeding,	Eli Lilly and Company, Critical Care Europe, Geneva, Switzerlan
n on outcome of patients with ST- segment elevation myocardial			baseline Not reported for cohort population comparing culprit- only PPCI versus multivessel PCI					

infarction transferred for primary percutaneous coronary intervention. EuroInterventi on. 2010; 20100525(201 00528).		Drug therapy Not reported for cohort population comparing culprit- only PPCI versus multivessel PCI			
Results ≤ 30 days	COR (n = 707)	MVR (n = 70)	p value	Adjusted OR (95% CI)	Adjusted p value
Death	42 (5.9%)	9 (12.9%)	0.039	2.42 (0.96, 6.06)	0.06
Major bleeding requiring transfusion	12 (1.7%)	2 (2.9%)	0.36	1.81 (0.35, 9.27)	0.48
Reinfarction	13 (7.5%)	0 (5.7%)	Not reported	Not reported	Not reported
All bleeding	62 (8.8%)	11 (15.7%)	0.08	1.97 (0.90, 4.29)	0.09
12 months					
Death					

Table 70: HELP-AMI 2004³³

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Outcome measures	Source of funding
Di Mario C, Mara S, Flavio A, Imad S, Antonio M, Anna P et al. Single vs multivessel creatment during primary angioplasty: results of the multicentre randomised HEpacoat for cuLPrit or multivessel stenting for Acute Myocardial Infarction HELP AMI) Study. International cournal of Cardiovascular Interventions. 2004; 6(3- 4):128-133.	Design RCT; multicentre Enrolment Dates not reported Randomisation Not detailed Allocation concealment Not detailed Blinding Open label Sample size calculation For primary outcome of repeat revascularisation; as 75% of the patients in the culprit lesion group were expected to need a second procedure at 1- year follow-up (mainly because of elective treatment	n = 69 Length of follow-up: 12 months	Inclusion criteria presence of ischaemic chest pain started less than 12 hours before hospital admission and / or ST segment elevation of at least 1 mm in 2 or more contiguous electrocardiographic leads (peripheral leads) or 2 mm in the pre-cordial leads multivessel CAD (defined as < 70% diameter stenosis of 2 or more epicardial coronary arteries or their major branches by visual estimation multivessel disease with the technical possibility of coronary and revascularisation with stents of at least 2 lesions (infarct related artery and 1 or more, to a maximum of 3 lesions in a major non-culprit related artery) Exclusion criteria presence of significant lesions in vein grafts or arterial conduits or in segments previously treated with angioplasty or stent implantation. recent fibrinolysis (less than 1 week) cardiogenic shock, defined as hypertension with systolic blood pressure less than 90 mmHg and	Culprit lesion only revascularisation (COR) n = 17 Stent implantation in the culprit artery using 1 or more heparin coated Bx Velocity stents, and subsequent interventions on the non-culprit lesions performed at the investigator's discretion; need and timing of the subsequent interventions were decided according to clinical status (persistent or recurrent angina), evidence of ischaemia in non-invasive tests (perfusion scintigraphy or stress echocardiogram), angiographic severity of non-culprit lesions and clinical relevance of the affected vessels, as well as organisation standards at participating centres	Multivessel revascularisation (MVR) n = 52 Revascularisation of all suitable lesions, using coated Bx Velocity stents in these lesions	Primary; 12 month incidence of repeat revascularisatio n (any revascularisatio n, infarct related artery as well as non- infarct related artery) Secondary; In-hospital repeat revascularisatio n, reinfarction All-cause mortality	None stated

of other lesions), a total of 70 patients were randomised in a 3:1 ratio (53 patients assigned to MVR treatment group and 17 patients assigned to COR group); this was required in order to test for a 55% difference in the incidence of new revascularisation with a power of 0.80 and a 2-sided alpha error equal to 0.05, this unbalanced enrolment strategy was adopted in order to increase the reliability of the main secondary end point (the incidence of adverse in-hospital events in the 2 groups) ITT analysis Not done	tachycardia greater than 100 BPM, not due to hypovolemia or requiring inotropic support or balloon counter pulsation single vessel disease the presence of left main stenosis of 50% or more intention to treat more than 1 totally occluded major epicardial vessel diffuse calcification or severe tortuosity in the culprit and the non-culprit arteries preventing the implantation of the study stents a side branch larger than 2.0 mm which was required to be covered by the stent, unless the operator was willing and technically able to maintain patency of this side branch with either further balloon angioplasty or stent replacement Demographics and baseline characteristics see below Drug therapy All patients received aspirin ≥ 100mg and clopidogrel 75 mg or ticlopidine 500 mg for at least 1 month after the procedure				
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Baseline characteristics			
	COR (n = 17)	MVR (n = 52)	p value
Age (years), n (%)	65.3 (7.4)	63.5 (12.4)	0.575
Male, (%)	84.6	88.2	0.531
Pre-PCI, (%)	2.0	0	0.796
Pre-CABG, (%)	9.6	23.5	0.144
LVEF, (%)	48.9 (8.6)	48.4 (9.9)	0.883
Diabetes, (%)	41.2	11.5	0.012
Hypercholesterolemia, (%)	52.9	41.2	0.285
Smoke, (%)	81.0	66.6	0.514
Hypertension, (%)	58.8	36.5	0.092
Clinical presentation of acute MI			
	COR (n = 17)	MVR (n = 52)	p value
Time onset of symptoms-hospital (min)	167 (180)	122 (97)	0.247
Time hospital to cath lab (min), mean (SD)	69 (54)	88 (90)	0.423
Q wave infarction, (%)	75.0	80.4	0.445
r-tPA (%)	5.9	7.7	0.641
Glycoprotein IIb/IIIa	82.4	75.0	0.397
Anterior infarction (%)	58.8	51.9	0.491
Killip 2 – 3 (%)	18.8	20.0	0.318
Systolic BP (mmHg), mean (SD)	141 (24)	136 (25)	0.474
Diastolic BP (mmHg), mean (SD)	85 (18)	83 (15)	0.528
Heart rate (BPM), mean (SD)	78 (18)	76 (19)	0.651
Two vessel disease, (%)	52.9	69.2	0.432
Three-vessel disease, (%)	47.1	30.8	
Procedural characteristics (all lesions)			
	COR (n = 17)	MVR (n = 52)	p value
Treated lesion/patient, mean (SD)	1.00 (0)	2.36 (0.64)	0.001

Stent/lesion, mean (SD)	1.29 (0.61)	1.12 (0.33)	0.008
Stent/patient, mean (SD)	1.29 (0.61)	2.73 (0.78)	0.001
Mean stent length, (mm), mean (SD)	19.9 (8.4)	16.4 (5.0)	0.088
Maximum balloon pressure, (atm), mean (SD)	13.6 (2.6)	14.1 (2.5)	0.561
Procedure duration (min), mean (SD)	53 (24)	69 (38)	0.032
Contrast used, (ml), mean (SD)	242 (106)	341 (163)	0.025
Results			
In-hospital	COR (n = 17)	MVR (n = 52)	p value
Death, (%)	0	1 (1.9)	0.754
Reinfarction, (%)		0	0
PPCI, (%)	0	1 (1.9)	0.675
CABG, (%)	0	0	-
12 months	COR (n = 17)	MVR (n = 52)	p value
Death (%)	0	1 (1.9)	0.754
Reinfarction, (%)	1 (5.9)	1 (1.9)	0.435
PPCI or CABG, (%)	6 (35.3)	9 (17.3)	0.174

Table 71: KAMIR 2012⁷¹

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Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow- up	Outcome measures	Source of funding
Lee HW, Hong TJ, Yang MJ, An SG, Oh J-H, Choi JH, Lee HC, Cha KS, Hong JY. Comparison of infarct-related artery vs multivessel revascularizatio n in ST-segment elevation myocardial infarction with multivessel disease: Analysis from Korea acute myocardial infarction registry. Cardiology Journal. 2012; 19(3):256-266.	Design Prospective cohort study, Korea Acute Myocardial Infarction Registry carried out in about 50 tertiary hospitals Enrolment 2005-2007; STEMI patients with MVD who had had PPCI Patients divided into 2 groups: patients with IRA (infarct related revascularisation) and patients with multivessel revascularisation. Confounders taken into account in the covariates analysis	n = 1,644	Inclusion criteria patients with STEMI with multivessel disease who had PPCI Exclusion criteria not specifically stated Demographics and baseline characteristics see below Drug therapy see below	IRA revascularisation n = 1106, 67.3%	Multivessel revascularisation n = 538, 32.7%	In hospital One month 12 months	In hospital: Cardiac death Non-cardiac death IABP Cerebrovascular accident Acute renal failure Defib/cardioversio n due to VT or VFib Major bleeding New onset HF Short- and long-term follow-up: Primary: MACE at	2 year research grand of Pusan National University

			short and long- term follow-up All-cause mortality Reinfarction Repeat revascularisation CABG Stent thrombosis
Baseline characteristics			
%	IRA revascularisation (n = 1106, 67.3%)	Multivessel revascularisation (n = 538, 32.7%	p value
Male,	800 (72.3%)	413 (76.8%)	0.055
Age, mean (SD)	63.6 (12.0)	62.1 (11.1)	0.014
Hypertension	604 (55.2%)	256 (47.9%)	0.005
Dyslipidemia	103 (10.8%)	40 (8.1%)	0.106
Diabetes	323 (29.6%)	161 (30.3%)	0.769
Current smoker	484 (44.0%)	263 (49.1%)	0.055
Familial history IHD	81 (8.1%)	54 (10.6%)	0.103
Previous PCI	59 (5.3%)	17 (3.2%)	0.049
Previous CABG	7 (0.6%)	1 (0.2%)	0.286
Previous MI	29 (2.6%)	13 (2.4%)	0.804
Previous cerebrovascular accident	73 (6.6%)	26 (4.8%)	0.157
Discharge medication:			
Aspirin	1081 (97.7%)	526 (97.8%)	0.969
Clopidogrel	1070 (94.7%)	523 (97.2%)	0.608
Cilostazol	308 (27.8%)	285 (53.0%)	< 0.0001

Beta-blocker	818 (74.0%)	417(77.5%)	0.118
ACE-1	749 (67.7%)	381 (70.8%)	0.0204
Statin	838 (75.8%)	423 (87.6%)	0.199
Follow-up duration (days)	370.5 (356.0 – 394.8)	358.5 (329.3 – 374.5)	0.017
N – cumulative incidences	IRA revascularisation	Multivessel revascularisation (n = 538,	p value
	(n = 1106, 67.3%)	32.7%)	
In-hospital outcomes			
Cardiac death	6 (0.5%)	2 (0.4%)	NS
Non-cardiac death	0	0	NS
IABP	51 (4.6%)	27 (5.0%)	0.715
Stroke	7 (0.6%)	0	0.104
Acute renal failure	3 (0.3%)	1 (0.2%)	NS
Defib/cardioversion	50 (4.5%)	13 (2.4%)	0.037
Major bleeding	2 (0.2%)	1 (0.2%)	NS
New onset HF	10 (0.9%)	1 (0.2%)	0.115
Events at 1-month follow-up			
Cardiac Death	7 (0.6%)	5 (0.9%)	0.0.543
Non-cardiac death	0	0	0
Re-MI	4 (0.4%)	3 (0.6%)	0.689
re-PCI	22 (2.0%)	4 (0.7%)	0.057
CABG	0	2 (0.4%)	0.107
MACE (hierarchical)	35(3.2%)	14 (2.6%)	0.529
Events at 12 month follow-up			
Cardiac Death	15 (1.4%)	8 (1.5%)	0.836
Non-cardiac Death	10 (0.9%)	1 (0.2%)	0.115
Re-MI	7 (0.6%)	4 (0.7%)	0.799
re-PCI	129 (11.7%)	66 (12.3%)	0.732
CABG	4 (0.4%)	2 (0.4%)	0.976
MACE (hierarchical)	165 (14.9%)	81 (15.1%)	0.953
Definite / probable stent thrombosis	4 (0.9%)	6 (2.6%)	0.097

Table 72: Meliga 2011⁸²

Reference	Study type	Number of patients	Patient charac	teristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Meliga E, Fiorina C, Valgimigli M, Belli R, Gagnor A, Sheiban I et al. Early angio- guided complete revascularizati on versus culprit vessel pci followed by ischemia- guided staged PCI in STEMI patients with multivessel disease. Journal of Interventional Cardiology. 2011; 24(6):535-541.	Design Retrospective cohort study, multicentre (6 tertiary care centres), consecutive patients treated between 2004 to 2008, Italy	n = 800 Decision to address patients to a multivessel PCI or culprit-only PPCI approach was dependent on the policy of the single institution. Incomplete was the default strategy in 3/6 institutions	Inclusion criter patients with S with multivess disease who had at the centres Exclusion criter patients who do survive to PPCI underwent mu PCI during the procedure patients with e CAD in whom is surgery was like required within days patients in who non-IRA was a total occlusion vessel < 2mm of the procedure patients in who non-IRA was a total occlusion vessel < 2mm of the procedure patients in who non-IRA was a total occlusion vessel < 2mm of the procedure patients in who non-IRA was a total occlusion vessel < 2mm of the procedure patients in who non-IRA was a total occlusion vessel < 2mm of the procedure patients in who non-IRA was a total occlusion vessel < 2mm of the procedure patients in who non-IRA was a total occlusion vessel < 2mm of the procedure patients in who non-IRA was a total occlusion vessel < 2mm of the procedure patients who distributed in the procedure patients with the procedure patients in who non-IRA was a total occlusion vessel < 2mm of the procedure patients with the patients with the procedure patients with the patients w	el ad PPCI ria lid not lor who altivessel index extensive pypass sely to be n ≤ 30 om the chronic or a diameter and	Culprit only revascularisation (COR) n = 383	Staged revascularisation during index stay (SR) n = 417	Mean follow-up 642 (545) days	All-cause mortality Reinfarction Repeat revascularisation	None stated
Baseline charact	eristics								
		COR (n = 383)		MVR (n =	= 417)	p value			
Male, mean (SD)		78.3		74.6		0.63			

Age, mean (SD)	64.5 (11.3)	66.9 (12.4)	0.21
Hypertension	42.8	49.2	0.2
Dyslipidemia	35.5	44.4	0.05
Diabetes	19.3	12.0	0.22
Active smokers	33.4	34.5	0.13
Extracardiac arteriopathy	7.8	1.9	0.08
Familial history	13.8	17.7	0.82
Previous PCI	9.9	10.1	0.94
Previous CABG	1.8	1.9	0.81
Previous acute MI	13.6	8.4	0.31
Number of diseased vessels			
2	61.6	64.8	0.32
3	38.6	35.2	0.11
Culprit lesion site			
LAD	42.2	39.7	0.35
Сх	19.2	23.5	0.29
RCA	38.6	36.8	0.38
Culprit lesion site			
LAD	42.2	39.7	0.01
Cx	19.2	23.5	0.01
RCA	38.6	36.8	0.01
stenosis (nonculprit lesion)	84	81	0.13
RVD	3.22 (0.67)	3.04 (0.34)	0.14
Bare metal stent for culprit lesion	96.2	91.1	0.58
Bare metal stent for nonculprit lesion	65.5	33.9	0.01
Time to PCI on non-IRAs	7 (4.2)	41 (23)	0.01
patients having PCI on non-IRAs	100	68.4	0.01
LVEF	54.8 (13.6)	55.1 (12.9)	0.64

Glycoprotein IIIb/IIa used	89.4	87.5	0.38
Results			
In-hospital events	SR (n = 417)	COR (n = 383)	p value
Death	10	17	0.11
Reinfarction	58	12	0.01
Repeat revascularisation	15	6	0.08
21 months events	SR (n = 417)	COR (n = 383)	p value
Death	19	28	0.13
Reinfarction	20	26	0.22
Revascularisation	72	34	0.012

Table 73: Nat'l CV Data Reg 2009²⁴

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Outcome measures	Source of funding
Cavender MA, Milford- Beland S, Roe MT, Peterson ED, Weintraub WS, Rao SV. Prevalence, predictors, and in- hospital outcomes of non-infarct artery intervention during primary percutaneous coronary intervention for ST-	Design Retrospective cohort study; National Cardiovascular Data registry, data from > 600 sites across USA, April 2004 to March 2007	n = 28936 Length of follow-up: 'In-hospital'	Inclusion criteria patients presenting with STEMI coronary artery disease in >1 major artery STEMI is defined as: Indicate whether the patient was hospitalised for an ST Elevation Myocardial Infarction (STEMI) documented in the medical record. At least one of the following biochemical indicators for detecting myocardial necrosis must be present (see below for a definition of Reference Control Limits): 1) Troponin T or I: a) Maximal concentration of troponin T or I > the MI decision limit on at least 1 occasion during the first 24	Culprit only revascularisation (COR); the infarct related artery only was treated and the other arteries were left untreated n = 25802	Multivessel revascularisatio n (MVR) n = 3134	All-cause mortality Cerebrovascular accident / Stroke Bleeding complications Renal failure	American College of Cardiology , Washingto n DC and Society for Cardiac Angiograp hy and Interventi ons, Washingto n, DC

segment elevation myocardial infarction (from the National Cardiovascular Data Registry). American Journal of Cardiology. 2009; 104(4):507- 513.	hours after the index clinical event. 2) CK-MB: a) Maximal value of CK-MB > 2× the upper limit of normal on 1 occasion during the first hours after the index clinical event; OR b) Maximal value of CK-MB, preferable CK-MB mass, > upper limit of normal on 2 successive samples. 3) Total CK a) In the absence of availability of a troponin or CK-MB assay, total CK > 2× the upper limit of normal, or the B fraction of CK may be employed, but these last 2 biomarkers are considerably less satisfactory than CK-MB; and one of the following ECG changes: 1) ST-segment elevation: New or presumed new ST-segment elevation at the J point in 2 or more contiguous leads with the cut-off points ≥ 0.2 mV in leads V1, V2, or V3, or ≥ 0.1 mV in other leads; OR 2) Development of any Q wave in leads V1 through V3, or the development of a Q-wave ≥ 30 ms (0.03 seconds) in leads I, II, aVL, aVF, V4, V5, or V6. (Q wave changes must be present in any 2 continguous leads, and be ≥ 1 mm in depth.) in patients with multivessel PCI, the infarct-related artery was considered the artery with the greatest preoperative stenosis (in percentages), in the event of a tie, the artery with	

the worst pre-PCI TIMI flow was considered the infarct-related artery patients with cardiogenic shock included; patients with the systolic blood pressure < 80 mmHg, or a cardiac index < 1.8 despite maximal treatment or requiring intravenous ion tropes, or an intra-aortic balloon pump to maintain the systolic blood pressure at > 80 mmHg and or cardiac index > 1.8 litres/minute/m² were classified as being in cardiogenic shock Exclusion criteria Patients receiving PPCI of the left main coronary artery Patients receiving staged PCI (multiple PCI procedures before hospital discharge) Patients receiving fibrinolytics Demographics and baseline characteristics see below Drug therapy Not detailed

Baseline characteristics

	COR (n = 25,802)	MVR (n = 3134)	p value
Age (years) median (IQR)	62 (53–73)	60 (52–72)	< 0.01
Male gender, (%)	72.1	71.5	0.32
Previous MI, (%)	19.3	17.4	< 0.01
Previous PCI, (%)	17.4	15.1	< 0.01

Previous CABG, (%)	9.9	5.1	< 0.01	
Cerebrovascular disease, (%)	7.7	8.2	0.37	
Congestive heart disease, (%)	9.8	13.2	< 0.01	
Diabetes, (%)	23.4	24.7	0.06	
Hypertension, (%)	63.2	60.4	< 0.01	
Hypercholesterolemia, (%)	58.6	56.5	0.05	
Renal failure, (%)			> 0.99	
Interval from symptom onset to	admission			
≤ 6 hours, (%)	78.8	74.4		
> 6 and ≤ 12 hours, (%)	10.1	10.6		
> 12 and ≤ 24 hours (%)	5.3	6.7		
> 24 and ≤ 48 hours, (%)	2.6	3.8		
> 48 hours and ≤ 7 days, (%)	2.6	4.1		
No time / silent MI, (%)	0.5	0.6		
LVEF, (%)	7.0	10.0		
Cardiogenic shock on presentation, (%)	10.3	13.8		
Emergent or salvage PCI status, (%)	91.3	87.3		
Intra-aortic balloon pump use, (%)	11.1	16.0		
Culprit coronary lesion, (%)		< 0.01		
Proximal LAD artery	16.9	19.8		
Proximal right, mid LAD, proximal circumflex artery	35.5	32.2		
Other	47.2	47.6		
Contrast volume (ml)		< 0.01		
Median	200	255		
Quartile 1, quartile 3 150, 262	200, 336			
Fluoroscopy time (min)		< 0.01		

Median 11.5 16.2 Quartile 1, quartile 3 7.6, 17.9 11.3, 24.0 COR (n = 605) MVR (n = 193) p value Results All patients with STEMI n = 25802 n = 3134 p value In-hospital all-cause mortality, n(%) 1321 (5.12) 246 (7.85) < 0.01 Cerebrovascular accident/stroke, n(%) 144 (0.56) 22 (0.72) 0.31 Bleeding complications, n(%) 1368 (5.30) 210 (6.71) < 0.01 Renal failure, n(%) 467 (1.81) 72 (2.31) 0.09 Patients with STEMI without in-hospital all-cause mortality, n(%) n = 2701 0.09 Cerebrovascular accident/stroke, n(%) 88 (3.26) 0.09 Bleeding complications, n(%) 106 (0.46) 12 (0.45) 0.90 Cerebrovascular accident/stroke, n(%) 1049 (4.53) 151 (5.67) 0.02 Renal failure, n(%) 280 (1.21) 23 (1.23) 0.96 Patients with STEMI with STEMI with In-hospital all-cause mortality, n(%) 737 (27.77) 158 (36.49) < 0.01 Petients with STEMI with STEMI with In-hospital all-cause mortality, n(%) 737 (27.77) 158 (36.49) < 0.01 Petients with STEMI with In-hospital all-cause mortality, n(%) 60 (13.81) 0.18 0.044				
Results Name Paulue All patients with STEMI n = 25802 n = 3134 p value In-hospital all-cause mortality, n (%) 1321 (5.12) 246 (7.85) < 0.01	Median	11.5	16.2	
Results All patients with STEMI n = 25802 n = 3134 p value In-hospital all-cause mortality, n (%) 1321 (5.12) 246 (7.85) < 0.01	Quartile 1, quartile 3	7.6, 17.9	11.3, 24.0	
All patients with STEMI n = 25802 n = 3134 p value In-hospital all-cause mortality, n 321 (5.12) 246 (7.85) < 0.01 Cerebrovascular accident/stroke, n (%) 1368 (5.30) 210 (6.71) < 0.01 Renal failure, n (%) 467 (1.81) 72 (2.31) 0.09 Patients with STEMI without n = 23.146 n = 2701 p value In-hospital all-cause mortality, n 585 (2.53) 88 (3.26) 0.09 Cerebrovascular accident/stroke, n (%) 1049 (4.53) 151 (5.67) 0.02 Renal failure, n (%) 280 (1.21) 23 (1.23) 0.96 Patients with STEMI with n = 2654 n = 433 p value In-hospital all-cause mortality, n (%) 280 (1.21) 158 (36.49) ≤ 0.01 Patients with STEMI with n = 2654 n = 433 p value In-hospital all-cause mortality, n (%) 280 (1.21) 28 (3.49) ≤ 0.01 Cerebrovascular accident/stroke, n (%) 11 (2.56) 0.18 Bleeding complications, n (%) 331 (12.48) 60 (13.81) 0.44		COR (n = 605)	MVR (n = 193)	p value
In-hospital all-cause mortality, n (%) 1321 (5.12) 246 (7.85) 22 (0.72) 0.31 Cerebrovascular accident/stroke, n (%) 1368 (5.30) 210 (6.71) < 0.01 Renal failure, n (%) 467 (1.81) 72 (2.31) 0.99 Patients with STEMI without n (%) 106 (0.46) 12 (0.45) 23 (1.23) 0.99 Patients with STEMI without n (%) 1049 (4.53) 151 (5.67) 0.90 Patients with STEMI with n (%) 1049 (4.53) 151 (5.67) 0.96 Patients with STEMI with n (%) 280 (1.21) 23 (1.23) 0.96 Patients with STEMI with n (%) 280 (1.21) 158 (36.49) 50 (0.18 (3.26) 0.18 (3.26) 0.18 (3.26) 0.18 (3.26) 0.18 (3.26) 0.18 (3.26) 0.18 (3.26) 0.18 (3.26) 0.20 (3.26) 0.18 (3.26) 0.18 (3.26) 0.18 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.26) 0.20 (3.	Results			
n (%) Cerebrovascular accident/stroke, n (%) 144 (0.56) 22 (0.72) 0.31 Bleeding complications, n (%) 1368 (5.30) 210 (6.71) < 0.01	All patients with STEMI	n = 25802	n = 3134	p value
accident/stroke, n (%) Bleeding complications, n (%)	•	1321 (5.12)	246 (7.85)	< 0.01
Renal failure, n (%) 467 (1.81) 72 (2.31) 0.09 Patients with STEMI without In-bospital all-cause mortality, n (%) n = 2701 point of the property		144 (0.56)	22 (0.72)	0.31
Patients with STEMI without $n = 23.146$ $n = 2701$ $p \ value$ (8) (8) (8) (9) (9) (9) (9) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10) (10)	Bleeding complications, n (%)	1368 (5.30)	210 (6.71)	< 0.01
In-hospital all-cause mortality, n (%) Cerebrovascular accident/stroke, n (%) Bleeding complications, n (%) 1049 (4.53) 151 (5.67) 0.02 Renal failure, n (%) 280 (1.21) 23 (1.23) 0.96 Patients with STEMI with n = 2654 n = 433 p value $1000000000000000000000000000000000000$	Renal failure, n (%)	467 (1.81)	72 (2.31)	0.09
n (%) Cerebrovascular accident/stroke, n (%) Bleeding complications, n (%) 1049 (4.53) 151 (5.67) 0.02 Renal failure, n (%) 280 (1.21) 23 (1.23) 0.96 Patients with STEMI with n = 2654 n = 433 p value ln-hospital all-cause mortality, 737 (27.77) 158 (36.49) ≤ 0.01 n (%) Cerebrovascular accident/stroke, n (%) Bleeding complications, n (%) 331 (12.48) 60 (13.81) 0.44	Patients with STEMI without	n = 23,146	n = 2701	p value
accident/stroke, n (%) Bleeding complications, n (%) 1049 (4.53) 151 (5.67) 0.02 Renal failure, n (%) 280 (1.21) 23 (1.23) 0.96		585 (2.53)	88 (3.26)	0.09
Renal failure, n (%) 280 (1.21) 23 (1.23) 0.96 Patients with STEMI with $n = 2654$ $n = 433$ $p \times 1000$ value $n = 1000$ in (%) $n = 10$		106 (0.46)	12 (0.45)	0.90
Patients with STEMI with $n = 2654$ $n = 433$ p value In-hospital all-cause mortality, $n = 737 (27.77)$ $n (\%)$ $n $	Bleeding complications, n (%)	1049 (4.53)	151 (5.67)	0.02
In-hospital all-cause mortality, n (%) 737 (27.77) 158 (36.49) ≤ 0.01 Cerebrovascular accident/stroke, n (%) 40 (1.49) 11 (2.56) 0.18 Bleeding complications, n (%) 331 (12.48) 60 (13.81) 0.44	Renal failure, n (%)	280 (1.21)	23 (1.23)	0.96
In-hospital all-cause mortality, n (%) 737 (27.77) 158 (36.49) ≤ 0.01 Cerebrovascular accident/stroke, n (%) 40 (1.49) 11 (2.56) 0.18 Bleeding complications, n (%) 331 (12.48) 60 (13.81) 0.44				
n (%) Cerebrovascular 40 (1.49) 11 (2.56) 0.18 accident/stroke, n (%) Bleeding complications, n (%) 331 (12.48) 60 (13.81) 0.44				
accident/stroke, n (%) Bleeding complications, n (%) 331 (12.48) 60 (13.81) 0.44	n (%)			≤ 0.01
		40 (1.49)	11 (2.56)	0.18
	Bleeding complications, n (%)	331 (12.48)	60 (13.81)	0.44
Renal failure, n (%) 197 (7.41) 42 (9.72) 0.03	Renal failure, n (%)	197 (7.41)	42 (9.72)	0.03

Table 74: NYS Angioplasty Reg 2006⁶⁴

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Kong JA, Chou ET, Minutello RM, Wong SC, Hong MK. Safety of single versus multi-vessel angioplasty for patients with acute myocardial infarction and multi-vessel coronary artery disease: report from the New York State Angioplasty Registry. Coronary Artery Disease. 2006; 17(1):71-75.	Design Retrospective cohort study; New York State Angioplasty Registry database 2000 – 2001	n = 1982	Inclusion criteria acute MI multivessel disease >70% stenosis in at least 2 major coronary arteries PCI within 24 hours of acute MI Exclusion criteria cardiogenic shock haemodynamic instability cardio pulmonary resuscitation left main stenosis >50% Haemodynamic instability (ie, Killip state > 2 or cardiogenic shock) history of prior MI, PCI, or coronary artery bypass surgery Demographics and baseline characteristics see below Drug therapy see below	Culprit only revascularisation (COR); the infarct related artery only was treated and the other arteries were left untreated n = 1350	Multivessel revascularisation (MVR) n = 632	In-hospital	All-cause mortality Confounders taken into account in the propensity analysis (stepwise multiple logistic regression analysis for independent predictors); age, sex, hypertension, diabetes, tobacco use, prior stroke, chronic renal failure, peripheral vascular disease, congestive heart failure, ejection fraction, GP IIb/III a inhibitor therapy, fibrinolytic therapy, PCI total occlusion, proximal LAD lesion present, proximal LAD lesion present, no PCI, stent	None stated

Baseline characteristics				
	COR (n = 1350)	MVR (n = 632)	p value	
Age (years), n (%)	62.0 (13.0)	60.0 (12.3)	0.002	
Female (%)	27.9	22.8	0.016	
Hypertension (%)	61.3	31.9	NS	
Diabetes (%)	20.5	16.8	0.051	
Tobacco use (%)	36.7	37.2	NS	
Prior stroke (%)	3.9	1.3	0.001	
Chronic renal failure (%)	1.0	0.6	NS	
Peripheral vascular disease (%)	8.9	4.7	0.001	
Clinical and procedural character	ristics			
	COR (n = 1350)	MVR (n = 632)	p value	
Congestive heart failure on admission (%)	7.9	6.6	NS	
Ejection fraction mean (SD)	46.5 (11.2)% (n = 1125)	48.3 (10.5) % (n = 582)	0.002	
GP IIb/IIIa inhibitor therapy (%)	20.0	21.2	NS	
PCI total occlusion (%)	52.0	41.8	< 0.001	
Proximal LAD lesion present (%)	26.1	25.6	NS	
Proximal LAD lesion present, no PCI (%)	7.0	0.9	< 0.001	
Stent(%)	91.3	98.9	< 0.001	
Congestive heart failure on admission (%)	7.9	6.6	NS	
Longth of stay (days) maan (CD)	F 4 (6.7)	F 2 (0 1)	NC	
Length of stay (days) mean (SD) Angiographic characteristics	5.4 (6.7)	5.3 (8.1)	NS	
Angiographic characteristics	COP (n = 1350)	M\/D (n = 622)	nyaluo	
Number of locions > 700/	COR (n = 1350)	MVR (n = 632)	p value	
Number of lesions >70%, per patient	3.2	3.5	< 0.001	

Α	12.0	8.4	< 0.001
В	60.6	67.5	< 0.001
С	27.5	24.1	0.004
Lesions treated / lesions present	(%)		
A	209/511 (40.9)	175/186 (94.1)	< 0.001
В	1362/2584 (52.7)	1403/1488 (94.3)	< 0.001
С	757/1172 (64.6)	496/531 (93.4)	< 0.001
Total lesions treated/present (%)	2328/4267 (54.5)	2074/2205 (94.1)	< 0.001
Results			
	COR (n = 1350)	MVR (n = 632)	p value
All-cause mortality (%)	2.3	0.8	0.018
Emergent bypass surgery (%)	0.7	0.5	NS
Acute occlusion or stent thrombosis (%)	0.7	0.9	NS
Stroke (%)	1.0	0.8	NS
Renal failure requiring dialysis (%)	0.2	0.3	NS

Table 75: NYS PCIRS 2010⁵¹

Reference	Study type	Number of patients	Patient charac	teristics	Intervention		Comparison	Outcome measures	Source of funding
Hannan EL, Samadashvili Z, Walford G, Holmes DR, Jr., Jacobs AK, Stamato NJ et al. Culprit vessel percutaneous coronary intervention versus multivessel and staged percutaneous coronary intervention for ST-segment elevation myocardial infarction patients with multivessel disease. JACC Cardiovascular Interventions. 2010; 3(1):22-31.	Design Retrospective cohort study obtained from New York State's Percutaneous Coronary Interventions Reporting System (PCIRS), January 2003 to June 2006, USA	n = 4024, outcome data available for 1006 patients. Length of follow-up: In-hospital 12, 24, and 42 months	Inclusion criter all multivessel patients (New State residents experienced a within 24 hour undergoing PC Exclusion crite patients with rejection fractic left main disea previous opensurgery cardiogenic she fibrinolytic the before PCI Demographics baseline charasee below Drug therapy not detailed	disease York s) who STEMI rs before I ria missing on se -heart ock rapy	Culprit only revasculari (COR); the infarct relationly was treated and the arteries were left untrendered in a state only was treated and the arteries were left untrendered in a state of the state of	ed artery he other eated atched with atched with I re whether ngle vessel e tched on acteristics ed samples cs included the left bronary disease estive u, several	Multivessel revascularisation (MV R); revascularisation of infarct related artery and other diseased vessels n = 259 Staged multivessel revascularisation (Inhospital SR); revascularisation of index infarct related artery and other vessels during index visit n = 259 Staged multivessel revascularisation (within 60 days SR); revascularisation (within 60 days SR); revascularisation of index infarct related artery and other vessels within 60 days n = 538	All-cause mortality	None stated
Baseline characteristics Demographic factors for COR PPCI versus MVR PCI population									
Risk factor		Percent with cul revascularisation PPCI, n = 503	prit vessel		with multivessel arisation at time of 03	Standard di	ifference (%)		

Age				
59 or less	55.47	54.67	1.60	
60 to 69	20.68	21.67	2.43	
70 to 79	15.7	14.12	4.47	
80 or more	8.15	9.54	4.90	
Female	21.27	25.05	8.96	
Cardiac factors				
Number of diseased vessels				
Trainiber of diseased ressels				
2 no proximal LAD	49.30	50.30	1.99	
2 with proximal LAD	24.25	24.06	0.46	
3 no proximal LAD	15.71	17.10	3.76	
Ejection fracture				
19% or less	2.39	3.38	5.94	
20% to 29%	6.16	7.55	5.51	
30% to 39%	16.30	16.30	0.07	
40% to 49%	28.63	28.23	0.88	
50% or more	46.52	44.53	3.99	
Haenodynamic status				
Unstable	2.98	4.77	9.28	
Chronic heart failure history				
Chronic total occlusion	5.57	5.37	0.87	
TIMI flow grade ≤ 2	40.16	43.54	6.86	
Comorbidities				
Cerebrovascular	3.38	3.78	2.14	
Peripheral vascular	3.58	4.17	3.09	
Ventricular arrhythmia	1.79	1.79	0.07	

COPD	4.97	5.37	1.80	
Renal failure	0.80	1.60	7.33	
Type of PCI				
Only drug eluting stent	57.85	57.06	1.61	
Bare metal stent	40.95	41.35	0.81	
No stents	1.19	1.59	3.39	
Baseline characteristics for	COR PPCI and SR PCI populations			
Demographic factors				
Age, years				
Risk factor	Percent with culprit vessel revascularisation at time of PPCI, n = 503	Percent with staged revascularisation at time of PPCI, n = 503	Standard difference (%)	
59 or less	47.88	49.03	2.32	
60 to 69	28.96	28.96	0.00	
70 to 79	16.60	16.99	1.03	
80 or more	6.56	5.02	6.62	
Female	18.90	15.83	8.16	
Cardiac factors				
Number of diseased vessels				
2 no proximal LAD	40.15	42.08	3.92	
2 with proximal LAD	18.15	15.83	6.17	
3 no proximal LAD	26.64	27.80	2.60	
Ejection fracture				
19% or less	1.16	1.16	0.00	
20% to 29%	7.34	5.79	6.24	

30% to 39%	18.53	19.31	1.97				
40% to 49%	31.66	29.34	5.03				
50% or more	41.31	44.40	6.24				
Haemodynamic status							
Unstable	3.47	3.09	2.17				
Chronic heart failure history							
This admission	4.10	2.32	9.08				
Chronic total occlusion	6.56	7.72	4.50				
TIMI flow grade ≤ 2	50.58	53.28	5.41				
Comorbidities							
Cerebrovascular	3.86	4.63	3.83				
Peripheral vascular	2.70	2.32	2.47				
Ventricular arrhythmia	0.39	1.16	8.83				
COPD	6.56	6.95	1.54				
Renal failure	0.39	0.39	0.00				
Type of PCI							
Only drug eluting stent	65.64	66.80	2.45				
Bare metal stent	31.27	30.5	1.67				
No stents	3.09	2.70	2.30	2.30			
Results							
Mortality rates for propensity matched STEMI patients; COR versus in-hospital MVR, n (%)							
	COR	In-hospital SR	percentage difference	p value			
In-hospital	10 (2.0)	17 (3.4)	1.4	0.14			
12 months	28 (5.5)	35 (7.1)	1.6	0.23			
24 months	33 (6.6)	43 (8.6)	2.0	0.17			
42 months	54 (10.8)	60 (11.8)	1.0	0.23			
In-hospital	10 (2.0)	17 (3.4)	1.4	1.4 0.14			
Mortality rates for propensity matched STEMI patients; COR versus in-hospital SR, n (%)							
	COR	In-hospital SR	percentage difference	p value			

In-hospital	5 (1.9)	3 (1.2)	0.7	0.48
12 months	14 (5.5)	10 (3.9)	1.6	0.53
24 months	19 (7.4)	16 (6.3)	1.1	0.71
42 months	22 (8.4)	16 (6.3)	2.1	0.72

Table 76: Politi 2010⁹²

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Outcome measures	Source of funding
Politi L, Sgura F, Rossi R, Monopoli D, Guerri E, Leuzzi C et al. A randomised trial of target-vessel versus multi-vessel revascularisati on in ST-elevation myocardial infarction: major adverse cardiac events during long-term follow-up. Heart (British Cardiac Society). 2010; 96(9):662-667.	Design RCT single centre Italy Enrolment Jan 2003 to Dec 2007 Randomisation Before the angioplasty, dedicated software generated a number (1 for the COR group, 2 for the capital SR group and 3 for the MVR group) Allocation concealment Not detailed Blinding Open label Sample size	Excluded patients: n = 4 patients with unsuccessful procedure included in randomisation but not in final analysis Length of follow-up: Inhospital Mean (SD) 2.5(1.4) years	Inclusion criteria presence of prolonged (more than 30 min) chest pain started less than 12 hours before hospital arrival and ST segment elevation of at least 1 mm in 2 or more contiguous electrocardiographic leads (peripheral leads) or 2 mm in the pre-cordial leads multivessel coronary artery disease (defined as greater than 70% diameter stenosis of 2 or more epicardial coronary arteries, or their major branches by visual estimation Exclusion criteria <18 years Active bleeding or significant increased risk of bleeding (severe hepatic insufficiency, current peptic ulceration, proliferative diabetic retinopathy) uncontrolled hypertension carcinogenic shock	Culprit only revascularisatio n (COR); the infarct related artery only was treated and the other arteries were left untreated n = 84	1. Multivessel revascularisation(MV R); the infarct related artery was opened followed by dilation of other significantly narrowed arteries during the same procedure n = 65 2. Stage revascularisation (SR); the infarct related artery only was treated during the primary intervention while the complete revascularisation was planned in the second procedure n = 65	In-hospital All-cause mortality Long-term All-cause mortality Reinfarction Repeat revascularisatio n Cardiac death	None stated

calculation severe peripheral vascular disease precluding a femoral Sample size was approach calculated on the primary end point previous coronary bypass (major adverse surgery with use of >1 internal cardiac event (MACE) mammary artery defined as cardiac or Previously entered in the study non-cardiac death, Investigational treatment (drug in-hospital death, or drug device) within the reinfarction, reprevious ≤ 30 days hospitalisation for acute coronary Demographics and baseline syndrome and repeat characteristics coronary see below revascularisation). Note MACE is not outcome of interest Drug therapy for clinical question. Before procedure patients In an expected rate were treated with aspirin, of MACE of 17% full unfractionated heparin, and groups undergoing abciximab bolus followed by 12 multivessel hour infusion. In addition, the revascularisation bolus of N- acetylcysteine 1200 (both simultaneous mg and hydration was saline for and staged) versus 12 hours after contrast 50% for the culpritexposure as infusion rate of 1 only group, aiming ml/kg per hour. Iodixamol was for a 0.05 alpha and used as a contrast media in all 0.90 power, the total patients. Post PPCI medical oral of 123 patients treatment included aspirin, needed to be statins and clopidogrel, unless enrolled (41 patients contraindicated, which was per group). recommended for ≤ 30 days in case of bare metal stent ITT analysis implantation and 12 months in case of drug eluting stents. See Yes below for further information.

Baseline characteristics				
	COR (n = 84)	SR (n = 65)	MVR (n = 65)	p value
Age (years)	66.9 (13.2)	64.1 (11.1)	64.5 (11.7)	0.413
Male gender, n (%)	64 (76.2)	52 (80.0)	50 (76.9)	0.849
Diabetes, n (%)	20 (23.8)	12 (18.5)	9 (13.8)	0.305
Hypertension, n (%)	50 (59.5)	42 (64.6)	32 (49.2)	0.192
Systolic blood pressure	136.2 (30.2)	136.2 (31.4)	136.0 (24.3)	0.999
before PPCI, mm Hg, mean (SD)				
Anterior location of STEMI, n (%)	35 (41.7)	28 (43.8)	31 (47.7)	0.761
Killip class that admission, mean (SD)	1.48 (0.61)	1.40 (0.70)	1.24 (0.53)	0.083
Three-vessel disease, n (%)	21 (25.0)	29 (44.6)	19 (29.2)	0.033
TIMI flow grade before PPCI, mean (SD)	0.76 (1.21)	0.89 (1.21)	1.11 (1.32)	0.244
Door to balloon time, min mean (SD)	63.1 (27.2)	65.7 (19.4)	60.1 (22.6)	0.824
Drug eluting stent, n (%)	10 (11.9)	6 (9.2)	5 (7.7)	0.722
Chronic renal failure, n (%)	24 (29.3)	16 (24.6)	17 (26.6)	0.816
Plasma creatine before PPCI mg/dl, mean (SD)	1.14(0.69)	1.13 (0.99)	0.98 (0.27)	0.369
Contrast induced neuropathy n (%)	3 (3.6)	2 (3.1)	1 (1.5)	0.748
Therapy at discharge	n = 77	n = 65	n = 63	
aspirin, n (%)	74 (96.1)	65 (100)	62 (98.4)	0.239
clopidogrel, n (%)	71 (92.2)	65 (100)	61 (96.8)	0.054
beta-blockers, n (%)	62 (80.5)	52 (80.0)	52 (82.5)	0.927
nitrates, n (%)	16 (20.8)	4 (6.2)	4 (6.3)	0.007
ACE inhibitors, n (%)	48 (62.3)	38 (58.5)	35 (55.6)	0.594
Results				

In-hospital outcomes			
	COR (n = 84)	SR (n = 65)	MVR (n = 65)
In-hospital all-cause mortality, n (%)	7 (8.3)	0 (0)	2 (3.1)
Length of hospital stay, days, n (%)	5.3 (2.5)	5.4 (3.1)	4.8 (2.6)
Rehospitalisation, n (%)	30 (35.7)	9 (13.8)	8 (12.3)
Length of hospital stay, days, n (%)	5.3 (2.5)	5.4 (3.1)	4.8 (2.6)
Follow-up (mean(SD)) 2.5(1.4) ye	ears		
	COR (n = 84)	SR (n = 65)	MVR (n = 65)
Repeat revascularisation, n (%)	28 (33.3)	8 (12.3)	6 (9.2)
All-cause mortality, n (%)	13 (15.5)	4 (6.2)	6 (9.2)
Cardiac death, n (%)	10 (11.9)	2 (3.1)	4 (6.3)
Reinfarction, n (%)	7 (8.3)	4 (6.2)	2 (3.1)

G.6 Cardiogenic shock

Table 77: SHOCK

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
30 DAYS and 6 MONTHS	Design: RCT	n = 302	Inclusion criteria ST-segment elevation, Q-wave	(n = 152) Early	(n = 150) Initial medical	30 days, 6 months	1° All- cause	National Heart, Lung
Hochman JS et al. 1999 ⁵³	Enrolment: April 1993 to November 1998.	Drop-outs (lost to follow-up):	infarction, a new left bundle- branch block or a posterior infarction with anterior ST- segment depression,	Revascularisatio n (ERV)	stabilisation (IMS)	1 year 6 years	mortality 2°	and Blood Institute. & American Heart
1 YEAR DATA Hochman JS et al. 2001 SHOCK Investigators 2001 54	Setting 30 sites in USA and Canada and 20% in other countries.	Hospital survivors n = 143	complicated by shock due to left ventricular dysfunction Onset of shock <36 hours of infarction and randomisation no more than 12 hours after diagnosis of shock.	Intra-aortic balloon pump (IABP/Pharmaco logical support Possible prior fibrinolysis	IABP/Pharmacol ogical support Fibrinolysis unless absolute contraindication Delayed	MILQ, NYHA 2 weeks after discharge, 6 and 12	Multidime nsional Index of Life Quality (MILQ) –	foundation.
6- YEAR DATA Hochman JS et al. 2006 SHOCK	Randomisation Patients were enrolled by means	2 patients were lost (n = 1 ERV	ulagilosis of silock.	Emergency early PCI/CABG	revascularisatio n at a minimum of 54 hours	months	instrumen	
Investigators 55	of computerised telephone randomisation,	and n = 1 IMS)	Exclusion criteria Severe systemic illness Mechanical or other cause of	Hospital survivors (n = 77)	Hospital	Notes: Hospital survivors,	t point questions,	
QUALITY OF LIFE and HEART FAILURE	with randomisation at each site performed	1 YEAR n = 1 (IMS patient	shock Sever vulgar disease Dilated cardiomyopathy	Notes: Had to be	survivors (n = 66)	median follow-up was 5.9	Andrew's life'.	
Sleeper LA et al. 2005 SHOCK Investigators	according to a permuted-block design.	was omitted from	Inability to gain catheterisation Unsuitable for	performed of randomisation	Notes: Intra-	(up to 11 years)	New York Heart	
103	Allocation concealment:	survival analysis)	revascularisation	Intra-aortic balloon	aortic balloon counterpulsatio n and	An updated vital status	Associatio congestive	
ADVERSE EVENTS Hochman J.	Yes. Central telephone system.	6 YEARS (April 2005)	Demographics and baseline characteristics see below	counterpulsatio n was recommended.	fibrinolytic therapy recommended.	was obtained for all	failure	

1999 (SHOCK) ⁵² SUBGROUP ANALYSIS	Blinding: Angiograms and echocardiograms were viewed by readers blind to enrolling site and	ERV n = 70 (24 dead, 46 alive)	Definitions Used Index MI- Prolonged (> 30 minutes) chest pain or equivalent symptoms	Delayed revascularisatio n at minimum of 54 hours after randomisation.	patients in 1999–2000 regardless of randomisa tion date.	class – using 4 standard questions	
DIABETES Farkouh ME, et al. 2006 SHOCK Trial Investigators	Sample size calculation: 328	IMS n = 51 (20 dead, 31 alive)	ECG criteria at time of MI diagnosis: ≥ 2 leads with ST elevation or ≥ 1 mm precordial ST depression known to reflect posterior		Follow-up visits were conducted annually from until	Renal failure Major/min or	
Shindler DM et al.	patients would give the study 90 percent power, with an overall type I error rate of 0.05		STEMI OR new left bundle branch black (known not to be old) OR new pathologic Q waves in ≥ 2 related leads or posterior		April 2005.	Intracrania I bleeding	
AGE Dzavik V et al.	absolute difference.		Q waves manifested by R/S ration V1 or V2 > 1 Cardiogenic shock —			IABP – intra-aortic balloon pump	
2005 (SHOCK) ³⁶ Jeger RV et al. 2011	30 DAYS, 6 MONTHS ITT analysis: Yes 1 YEAR		Clinical criteria were hypotension (SBP < 90 mm Hg for at least 30 minutes or supportive measure to				
	ITT analysis: Yes 6 YEARS		maintain SBP ≥ 90 mm Hg) and end-organ hypoperfusion (cool extremities or a urine output of < 30ml per hour and heart rate of ≥ 60 beats per				
	AGE SUBGROUPS: ITT analysis: Yes		minute). Haversusemodynamic criteria were a cardiac index of no				
	DIABETES ITT analysis: Yes		more than 2.2 litres/minute/m²body-surface area and pulmonary-capillary wedge pressure of at least				

QUALITY OF LIFE	15mm Hg.		
ITT analysis:			
No. PPA			

End point definitions:

Acute renal failure – serum creatine level above of 3.0mg per decilitre

NYHA Congestive Heart Failure -

Class I: no limitation is experienced in any activities; there are no symptoms from ordinary activities.

Class II: slight, mild limitation of activity; the patient is comfortable at rest or with mild exertion.

Class III: marked limitation of any activity; the patient is comfortable only at rest.

Class IV: any physical activity brings on discomfort and symptoms occur at rest.

Baseline characteristics. Patients in 2 groups were similar except that more patients in IMS had previously undergone bypass surgery

	Early Revascularisation (n =152)	Initial Medical Stabilisation (n = 150)
Mean age, years (SD)	65.5 ± 10.0	66.2 ± 10.9
Men (%)	63.2	72.7
Hypertension (%)	49.0	43.5
White race, non-Hispanic	72.4	78.7
Previous MI (%)	29.6	35.3
Hypertension (%)	49.0	43.5
Diabetes mellitus (%)	34.2	27.9
Congestive heart failure (%)	4.0	8.2
Renal insufficiency (%)	4.6	6.9
Prior coronary artery by-pass grafting (%)	2.0	10.0
Cigarette smoking (%)	52.6	56.8
Eligible for fibrinolytic therapy (%)	94.1	94.6
Transfer admission (%)	55.3	55.3
Anterior index MI (%)	63.6	57.4
Highest total creatine kinase (IU/L)	3068 (1322–6350)	3464 (1543–5411)
Median time from MI to shock (hr)	5.0 (2.2–12.0)	6.2 (2.4–15.5)

Median time from MI to randomisation (hr)	11.0 (5.9–19.4)	12.0 (6.3–21.8)
<6 hr from MI to randomisation (%)	25.0	23.7
Lowest systolic blood pressure (mm Hg)	66.4 ± 14.3	69.8 ± 11.3
Systolic blood pressure (mm Hg)	89.0 ± 22.8	86.5 ± 17.4
Diastolic blood pressure (mm Hg)	53.9 ± 16.8	55.1 ± 13.6
Heart rate (beats/minute)	103.3 ± 22.0	100.1 ± 22.7
Pulmonary-capillary wedge pressure (mm Hg)	24.2 ± 7.1	24.3 ± 7.7
Cardiac index (litres/minute/m²)	1.8 ± 0.7	1.7 ± 0.5
Left ventricular ejection fraction (%)	29.1 ± 10.6	32.5 ± 13.9
Number of diseased vessels (%)		
≤1	14	11.5
2	21.7	24
3	64.3	64.6
Left main coronary artery disease (%)	23.4	17.5

Treatment (includes treatment recommended or required according to the protocol i.e fibrinolytic therapy & revascularisation)

	Early Revascularisation n = 152 n (%)	Initial Medical Stabilisation n = 150 n (%)
Cardiopulmonary resuscitation, VT, or VF before randomisation N/total (%)	37/113 (32.7%)	27/113 (23.9%)
Fibrinolytic therapy, N (%)	75 (49.3%)	95 (63.3%)
Inotropes or vasopressors, N (%)	151 (99.3%)	148 (98.6%)
Intraaortic balloon counterpulsation, N (%)	131 (86.2%)	129 (86.0%)
Pulmonary-artery catheterisation, N (%)	142 (93.4%)	144 (96.0%)
Left ventricular assist device, N (%)	4/111 (3.6%)	1/110 (0.9%)
Heart transplantation, N (%)	3 (2.0%)	1 (0.7%)
Coronary angiography, N (%)	147 (96.7%)	100 (66.7%)
Angioplasty or coronary-artery bypass grafting, N (%)	132 (86.8%)	38 (25.3%)
Angioplasty, N (%)	83 (54.6%)	21 (14.0%)

• Stent placed		35.	7			52.3	
Platelet glycoprotein IIb/	IIIa receptor antagonist: N/t	otal (%) 25/	(60 (41.7%)			5/20 (25.0%)	
Coronary-artery bypass gra	fting, N (%)	57	57 (37.5%)		17 (11.3%)		
Median time from randomisation to revascularisation (hr) (interquartile rage)			(0.6–2.8)			102.8 (79.0–16	52.0)
Revascularisation more tha	n 6 hours after revascularisa	tion 10	(6.6%)				
VT: sustained ventricular ta	chycardia, VF: sustained ven	tricular fibrillation					
Treatment post randomisa	tion						
		Ear	ly Revascula	risation		Initial Medical	Stabilsation
		(n =	= 152)			(n = 150)	
No Revascularisation attem	pt	20				112	
 Died prior to 		5				?	
IABP		86%	6			86%	
Early Revascularisation atte	empt	125	5			•	ess than 54 hours after (protocol violation))
• PCI		83				1	
• CABG		42				2	
Late Revascularisation atte	mpt	· · · · · · · · · · · · · · · · · · ·	Performed motocol violati	ore than 6 hours after rai on))	ndomisation	a minimum of	or this group: Performed at 54 hours after n (if clinically appropriate))
• PCI		1				20	
• CABG		6				15	
Results: Mortality at 30 day	ys, 6 months, 1 year and 6 y	ears					
	Early Revascularisation (ERV) n = 152	Initial Me Stabilisation (IN		RR 95% CI ERV versus IMS		latio 95% CI rsus IMS	HR 95% CI ERV versus IMS
30-day mortality N/Total (%)			,	LIVY VEISUS IIVIS	LIV	1343 HVI3	LIVA ACIONO HAID
Total ⁵³	71/152 (46.7%)	84/150 (5	56%)	0.83 (0.67 to 1.04)			
• Age < 75 years	53/128 (41.4%)	67/118 (5		0.73 (0.56 to 0.95)			
• ,	•			,			

A 75	10/24/75 00()	47/22/52 42/	4.44 (0.05 : 0.44)	
• Age > 75 years	18/24 (75.0%)	17/32 (53.1%)	1.41 (0.95 to 2.11)	
 Age > 75 years⁵⁸ 			0.93 (0.39 to 2.25)	
 Age ≤ 75 years 			0.83 (0.62 to 1.11)	
• Age 71–75			0.83 (0.48 to 1.43)	
• Age 68–70			0.82 (0.51 to 1.32)	
• Age 56–65			0.81 (0.52 to 1.26)	
• Age ≤ 55			0.86 (0.32 to 2.26)	
• Diabetes ⁴⁵				0.73 (0.42 to 1.27)
Non-diabetes ⁴⁵				0.54 (0.23 to 1.24)
6 month mortality N/Total (%) ⁵³ 2 patients (1 EVR and 1	IMS) were lost to follow-up		
Total	76/151 (50.3%)	94/149 (63.1%)	0.80 (0.65 to 0.98)	
• Age < 75 years	57/127 (44.9%)	76/117(65.0%)	0.70 (0.56 to 0.89)	
• Age > 75 years	19/24 (79.2%)	18/32 (56.3%)	1.41 (0.97 to 2.03)	
• Diabetes, N/total (%) ¹⁰¹	23/51 (45%)	26/41 (63%)	0.71 (0.49 to 1.04)	
• Without diabetes, N/total (%)	49/98 (50%)	66/106 (62%)	0.80 (0.63 to 1.03)	
1 year mortality, N/total (%)				
otal ^{54,55}	83/151 (55%)	109/150 (72.7%)	0.72 (0.54 to 0.95)	
otal ⁵⁸			0.83 (0.70 to 0.96)	
Age > 75 years ⁵⁸			0.93 (0.56 to 1.53)	
Age > 75 years ³⁶	19/24 (79.2%)	21/32 (65.6%)		
Age < 75 years ³⁶	62/127 (48.4%)	78/117 (66.7%)		
• Age ≤ 75 years ⁵⁸			0.79 (0.63 to 0.99)	
• Age 71–75			0.86 (0.60 to 1.23)	
• Age 68–70			0.72 (0.48 to 1.09)	
• Age 56–65			0.78 (0.52 to 1.17)	
• Age≤55			0.82 (0.30 to 2.26)	
6 year mortality, N/total				

(%)				
Total ⁵⁵	107/151 (71%)	119/150 (79.3%)		0.74 (0.57 to 0.97)
Renal Failure 30 days and I	ntracranial bleeding ⁵²			
		Revascularisation	Medical therapy	
Acute renal failure n/Total	(%)	20/152 (13%)	36/150 (24%)	
Intracranial bleeding n/Tot	al (%)	0/152	2/150 (0.7%)	
		Revascularisation Max n = 41	Medical therapy Max n = 2	3
2 weeks after discharge		17.1 ± 5.2	15.9 ± 7.9	
6 months post MI				
1 year post MI		19.9 ± 6.4	17.3 ± 5.7	
•		19.9 ± 6.4 19.1 ± 6.2	17.3 ± 5.7 19.3 ± 4.4	
NYHA Congestive Heart Fai	lure Class of 126 SHOCK tri	19.1 ± 6.2		
	lure Class of 126 SHOCK tri	19.1 ± 6.2		
	lure Class of 126 SHOCK tri	19.1 ± 6.2 al hospital survivors	19.3 ± 4.4	

	Revascularisation	Medical therapy
2 weeks after discharge	n = 58	n = 48
Class I	n = 27 (46.6%)	n = 18 (37.5%)
Class II	n = 17 (29.3%)	n = 12 (25.0%)
Class III	n = 6 (10.3%)	n = 6 (12.5%)
Class IV	n = 8 (13.8%)	n = 12 (25.0%)
6 months post MI	n = 55	n = 37
Class I	n = 30 (54.6%)	n = 20 (54.1%)
Class II	n = 9 (16.4%)	n = 6 (16.2%)
Class III	n = 7 (12.7%)	n = 3 (8.1%)
Class IV	n = 9 (16.4%)	n = 8 (21.6%)
1 year post MI	n = 54	n = 35
Class I	n = 31 (57.4%)	n = 20 (57.1%)

Class II	n = 15 (27.8%)	n = 8 (22.9%)
Class III	n = 1 (1.9%)	n = 1(2.9%)
Class IV	n = 7 (13.0%)	n = 6 (17.1%)

Further analysis:

30 days: Adverse events including stroke and haemorrhage were similar in both groups 52 NB Abstract, no numbers provided.

1 year: No significant difference between treatment effect and gender or diabetes mellitus⁵⁴

6 years: Long-term survival analysis of the entire cohort identified no interactions between treatment assignment and any subgroup factor, including age (≤75 versus ≥75 years), sex, diabetes. ⁵⁵

Figure 1: Study flow

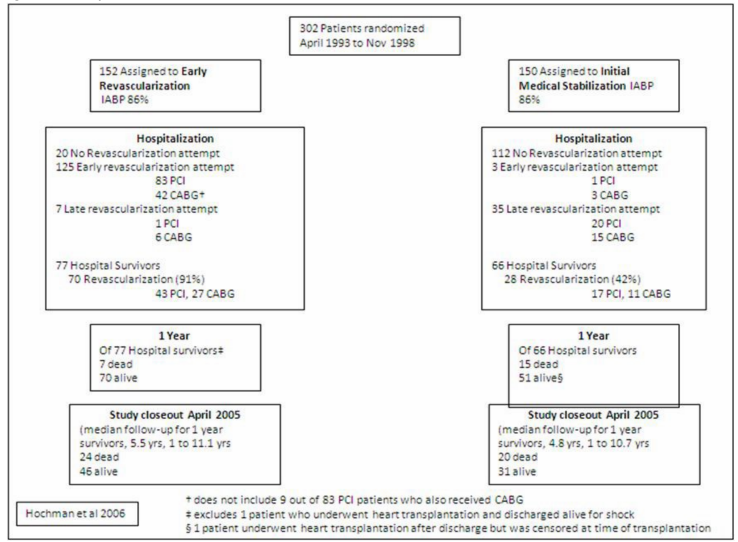


Table 78: SMASH

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow- up	Outcome measures	Source of funding
Urban P et al. 1999 Shock- (S)MASH	RCT (conducted with 2 parallel centres) Enrolment: Jan 1992 to May 1996 Setting 9 settings in 3 European countries Randomisation carried out by contacting a central 24/24 hours randomisation service by telephone. It was stratified for each individual centre. Allocation concealment Yes. Central telephone system Power calculation: Accepting a probability of a type I error of 0.05, and	n = 55	Inclusion criteria cardiogenic shock present for 30 minutes or more, with a systolic blood pressure of 90 mmHg or less for at least 30 minutes despite inotropic and intravenous volume administration as needed acute myocardial infarction onset <48 h prior to randomisation; the diagnosis was made if at least 2 of the following 3 elements were present: chest pain; ST segment elevation of at least 0·1 mV in limb leads or 0·2 mV in precordial leads or left bundle branch block; serum creatine-phosphokinase MB isoenzyme elevation above twice the upper limit of normal. if measured, capillary wedge pressure >15 mmHg and thermodilution cardiac index <2·2 l. min . m; coronary angiography technically feasible (vascular access and catheterisation laboratory available); informed consent given if requested. Exclusion criteria ongoing manual cardiopulmonary resuscitation; prior cardiac arrest with presumed severe cerebral damage; shock not primarily cardiogenic in origin, evidence of ventricular rupture (free wall or septum), acute severe mitral regurgitation, or pericardial tamponade;	Emergency Coronary Angiography n = 32 Notes: Patients were taken to the catheterisation laboratory as soon as possible, coronary angiography was completed, and PCI or CABG were attempted if considered feasible. Revascularisati on procedures were performed with as little delay as possible.	Initial medical management n = 23 Notes: patients did not undergo immediate coronary angiography, but were admitted to the intensive care unit. Late coronary angiography, with or without subsequent revascularisati on, was not considered a protocol violation.	30 days & after 1 year.	mortality from all- causes (cardiac and non- cardiac) 30 days after randomisa tion 2° need for non- emergency PCI or CABG during hospital stay; New York Heart Associatio n (NYHA) heart failure class at discharge from hospital mortality, cardiac	Swiss Cardiolo gy Foundat ion, Bristol- Myers Squibb AG, Switzerl and, Schneid er- Worldwi de, Arterom ed SA, Medtro nic Europe, Cook (Switzerl and) AG

End point definitions

Success of percutaneous revascularisation: the intervention was deemed a technical success by the operator if he/she estimated the residual diameter stenosis as < 50%

and the flow in the culprit vessel after the procedure as TIMI grade 2 or 3.

Reinfarction: No definition provided

Unplanned revascularisation: Counted as patients, post-hospital discharge, who required late percutaneous transluminal coronary angioplasty and late cardiac surgery

Stroke: No definition provided

Baseline characteristics at randomisation.

	Invasive (n = 32)	Conservative (n = 23)
Age (years, mean ± SD)	66 ± 10	64 ± 8
Age >65	19 (59%)	11 (48%)
Male sex	23 (72%)	14 (61%)
Hypertension 1	11 (34%)	6 (26%)
Diabetes	6 (19%)	4 (18%)
Smoking 3	19 (59%)	15 (65%)
Hypercholesterolaemia	6 (19%)	3 (14%)
revious acute myocardial infarction	9 (28%)	10 (43%)
Previous PCI or CABG	3 (9%)	4 (17%)
Prior heart failure (NYHA >I)	9 (28%)	5 (22%)
Prior angina (CCS >I)	12 (38%)	10 (45%)
Median (range) time from acute myocardial infarction to shock (h)	2·5 (0–27)	5 (0–47)
Median (Range) time from shock to randomisation (h)	3 (0–33)	2 (1–7)
leart rate (beats . min ⁻¹)	101 ± 30	105 ± 30
systolic blood pressure	77 ± 10	78 ± 13
Cardiac index (I . min ⁻¹ . m ⁻²)	1·7 ± 0·3 (n = 7)	1·8 (n = 1)
Pulmonary capillary wedge pressure	21 ± 4 (n = 8)	29 ± 6 (n = 4)
Anterior or lateral acute myocardial infarction	15 (47%)	10 (43%)
T segment elevation	26 (81%)	18 (78%)
Right ventricular acute myocardial infarction	5 (16%)	4 (17%)
/F or sustained VT since acute myocardial infarction onset	9 (28%)	9 (39%)
Cardiopulmonary resuscitation required	9 (28%)	8 (35%)
ntravenous inotropes	30 (94%)	23 (100%)
ibrinolysis	11 (34%)	9 (39%)
ABP	3 (9%)	1 (4%)
emporary RV pacing	5 (16%)	4 (17%)
Fracheal intubation	15 (47%)	14 (61%)
Creatine phosphokinase > 2× upper limit of normal	15 (47%)	14 (61%)

Arterial pH	7·3 ± 0·1 (n = 20)	7·3 ± 0·1 (n = 13)
Lactates (mmol . I ⁻¹)	7.5 ± 4.8 (n = 11)	7.8 ± 6.0 (n = 8)

CCS: Canadian Cardiology Society (classification); IABP: intra-aortic balloon pump; VT: sustained ventricular tachycardia; VF: sustained ventricular fibrillation; RV: right ventricle

Outcomes

NYHA

- Class I: no limitation is experienced in any activities; there are no symptoms from ordinary activities.
- Class II: slight, mild limitation of activity; the patient is comfortable at rest or with mild exertion.
- Class III: marked limitation of any activity; the patient is comfortable only at rest.
- Class IV: any physical activity brings on discomfort and symptoms occur at rest.

Treatments

	Invasive	Conservative
30 days	n = 32	n = 23
		Subsequent revascularisation was not considered a protocol violation.
No Revascularisation attempt	4 (2 died before coronary angiography, 1 died after angiography before revascularisation, and 1 other patient was not revascularised because the residual lesion was considered non-significant)	
Early Revascularisation attempt		
• PCI	27	
• CABG	1	
Intra-aortic counterpulsation balloon (IABP) during hospital study, either prior to or following hospitalisation	21 (66%)	7 (30%)
Main outcomes at 30 days and 30 days to 1 year		
	Invasive	Conservative
30 days	n = 32	n = 23
All-cause mortality	22 (69%)	18 (78%)
Reinfarction	1 (3%)	1 (4%)

Stroke	0	2 (13%)
Late PCI	0	1 (4%)
Late cardiac surgery	2 (6%)	0
Median Heart Failure, NYHA	II (I-III)	II (I-IV)
30 days to 1 year	n = 10	n = 5
Mortality	1 (10%)	1 (20%)
Reinfarction	0	0
Late PCI	0	0
Late cardiac surgery	1 (10%)	0

G.7 People who remain unconscious after a cardiac arrest

Table 79: Bulut 2000¹⁸

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow- up	Outcome measures	Effect sizes	Source of funding
Bulut S, Aengevaere n WRM, Luijten HJE, Verheugt FWA. Successful out-of- hosnital cardiopulmo narv resuscitatio n: What is the out-inel in-hospital treatment strategy? Resuscitatio n. 2000; 47(2):155- 161.	Retrospective cohort study, single centre, Netherlands	n = 30	GP 11b/111a inhibitor	PPCI n = 10 Note 6/10 patients conscious prior to PPCI ot reported	Usual care n = 20 Note No data given for number of patients that were conscious after resuscitation	In- hospital	All-cause mortality Successful procedure, not defined (PPCI group only) Neurological recovery, not defined; (PPCI group only)	PPCI; 6/10 (60%) patients (4/6 patients due to neurological cause and 2/6 due to cardiac cause) Usual care; 13/20 (65%) patients (9/13 patients due to neurological cause) PPCI; 8/10 patients PPCI; 3/4 patients	None stated
			• N	or reported					

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow-	Outcome measures	Effect sizes	Source of funding
						up			

Univariate (Pearson's Spearman rho and Kendall's tau model)

Out of 21 possible predictive factors, the following items associated with in-hospital all-cause mortality;

- Glasgow coma scale at admission (GSC), r = -0.77
- Need for ventilation in emergency department, r = -0.61
- Tracheal intubation on admission, r = -0.56
- Need for inotropic drugs at hospital, r = -0.54
- Need for inotropic drugs during transportation to hospital, r = -0.51
- Ventilation at admission, r = -0.51
- Duration of cardiopulmonary resuscitation, r = -0.46
- Cardiac rhythm on admission, r = -0.38
- Initial cardiac rhythm on admission, r = −0.37
- Presence of witness during collapse, r = −0.34
- Duration of stay in intensive care unit, r = −0.24

Multivariable stepwise regression analysis of 11 univariate factors associated with in-hospital all-cause mortality

• GCS most important predictor, r = -0.76 (p < 0.001)

Excluding GCS, following were predictive factors for in-hospital all-cause mortality

• Initial cardiac rhythm on admission, r = -0.65 (p < 0.001)

Ventilation at admission r = -0.60 (p < 0.001)

Table 80: Liu 2012⁷⁶

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Outcome measures	Source of funding
Liu Hw, Pan W, Wang Lf, Sun Ym,	Retrospective cohort study,	n = 81	Inclusion criteria Out-of-hospital coronary arrest	PCI n = 49	Usual care TH, restricted to	All-cause mortality Stroke	None stated
Li Zq, Wang Zh. Impact of emergency percutaneous coronary intervention on outcomes of ST-	China	Length of follow-up: In hospital	 Typical chest pain with ST-segment elevation ≥ 1 mm in at least 2 consecutive precordial or inferior limbs, or chest pain with new onset of complete bundle branch block 		comatose patients n = 32	Acute renal failure Confounders taken into account in the covariates analysis	
segment elevation			Exclusion criteria: Not stated				
myocardial infarction patients			Demographics and baseline characteristics: See below				
complicated by out-of-hospital			Drug therapy: All patients received aspirin, clopidogrel, atorvastatin and				
cardiac arrest. Chinese Medical Journal. 2012;			subcutaneous low molecular weight heparin for 2–3 days. Beta blockers,				
125(8):1405- 1409.			ACEIs or ARBs and aldosterone receptor antagonists were administered early according to heart rate and blood pressure.				
Baseline character	istics of the OHCA pa	tients with and					
			Without PCI (n = 32)	With P	PCI (n = 49)	p value	
Male, n (%)			23 (71.9)	40 (81	.6)	0.4477	
Age (years), mean(SD)		59 (16)	54 (13)		0.1267	
History of MI, n (%)			1 (3.1)	2 (4.1)		0.7048	
Hypertension, n (%)		14 (43.8)	20 (40		0.9751	
Diabetes mellitus, i	n (%)		4 (12.5)	6 (12.2	2)	0.7555	

Renal insufficiency, n (%)	2 (6.3)	2 (4.1)	0.9329
Smokers, n (%)	17 (53.1)	26 (53.1)	0.8242
Onset to admission time (minutes), mean (SD)	138 (42)	122 (38)	0.0794
On admission			
Systolic blood pressure (mmHg), mean (SD)			
Heart rate (beats/minutes), mean (SD)	101 (23)	96 (19)	0.2903
Killip IV, n (%)	15 (46.9)	19 (38.8)	0.6229
GCS ≤ 7, n (%)	10 (31.27)	14 (28.6)	0.9926
Results: During hospitalisation n, (%)			
Stroke			
Acute renal failure	3 (9.3)	1 (2.0)	0.3346
All-cause mortality	27 (84.3)	18 (36.7)	0.0001

Table 81: Pleskot 2008⁹¹

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Effect sizes	Source of funding
Pleskot M,	Retrospective	n = 26	Inclusion criteria	n = 20	Usual care	In-hospital	All-cause	PPCI; 6/19	None
Babu A, Hazukova R,	cohort study, multi-centre		 Out-of-hospital cardiac arrest 	underwent urgent	n = 6	1 year	mortality; in- hospital;	patients	stated
Stritecky J,	April 2002 to		 Δcute STFMI defined as 	coronary	Noto			Usual care; 5/6 patients	
Bis J. Matejka of-hospital cardiac arrests in	Aug 2004, Czech Republic		dynamic ST segment elevations on the ECG along with a typical rise (a minimum 3 times above the upper border	angiography, underwent PPCI	conscious at admission		All-cause mortality; 12 months	PPCI; 0/14 patients	
patients with acute ST			of normal values) and	Note				Usual care; 0/1 patients	

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Effect sizes	Source of funding
elevation myocardial infarctions in the East Bohemian region over the period 2002-2004. Cardiology. 2008; 109(1):41-51.			fall in biochemical markers of myocardial necrosis (serum creatine kinase and its MB fraction) with the consequent development of a pathologic Q wave on the ECG • Due to disputable interpretation of biochemical markers of myocardial necrosis following a cardiopulmonary resuscitation as a diagnostic indicator, the coronary angiogram was taken into consideration prior to direct PCI, myocardial wall kinetics (ultrasound) and in some cases the autopsy report	2/20 patients conscious at admission			Good cerebral performance using Glasgow-Pittsburgh Outcome Categorisation (CPC) of brain injury; discharged from hospital (defined as CPC = 1); in-hospital Good cerebral performance using Glasgow-Pittsburgh Outcome Categorisation (CPC) of brain injury (defined as CPC = 1); 12 months follow-up	PPCI; 11/14 patients Usual care; 0/1 patients PPCI; 13/14 patients Usual care; 0/1 patients	

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Effect sizes	Source of funding
			Exclusion criteria Individuals with cardiac arrest occurring in the presence of emergency medical services Toxic, traumatic, suicidal etiologies for unconsciousness including drowning and terminal illness Demographics and baseline characteristics See below Drug therapy PPCI arm Prior to transport to the cardiac catheterisation laboratory (mostly from intensive care units) patients in this subgroup received 500 mg aspirin and 5,000–10,000 units of unfractionated heparin intravenously Usual care Patients in this subgroup were treated with intravenous unfractionated heparin				Successful procedure; optimal angiographic results after direct P were considered to be achieved when < 30% of residual luminal stenosis remained in the target vessel, that is, fibrinolysis in MI (TIMI) flow grade II and III	PPCI; 17/19 patients	

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Effect sizes	Source of funding
			without fibrinolysis. Stents PPCI group; 17/19 (85%) GP 11b/111a inhibitor Not reported All patients; beta-blockers, nitroglycerine, sedatives, analgesia, angiotensin- converting enzyme inhibitors, statins (similar in both groups) Ticlopidine/clopidogrel given in cases of stent implantation						

Clinical, pre-hospital and hospital characteristics of hospitalised individuals

	With coronary angiography (n = 20)	Without coronary angiography (n = 6)				
Age, years						
Median	55.5	61.5				
Range	35–79	43–69				
Mean (SD)	56.38 (10)	56.78 (8.2)				
> 70 years	2 (10)	0				
Males, n (%)	18 (90)	4 (67)				
Risk factors for coronary artery disease, n (%)						
0	10 (50)	0				
1	7 (35)	3 (50)				

į		Number of					Length of	Outcome		Source of
Reference	Study type	patients	Patient ch	aracteristics	Intervention	Comparison	follow-up	measures	Effect sizes	funding
2				3 (15)			3 (50)			
	diastolic pressur	re, >90 mm H	g), n (%)	6 (30)			2 (33)			
Diabetes mellit				0			2 (33)			
	rolemia (cholesto	erol >5.7 mm	ol/l), n (%)	2 (10)			2 (33)			
Smoking, n (%)				7 (35)			3 (50)			
Previous IHD, n	ı (%)			10 (50)			3 (50)			
Initial cardiac rl	hythm, n (%)									
Ventricular fibr	rillation, n (%)			20 (100)			3 (50)			
Asystole, n (%)				0			1 (17)			
Pulseless electi	rical activity, n (%	%)		0			1 (17)			
Atrioventricula	r block, n (%)			0			1 (17)			
Distance from	the location of c	ardiac arrest	to cardiac c	atheterisation la	boratory, n (%)					
≤ 20 km, n (%)				16 (80)			3 (50)			
> 20 km, n (%)				4 (20)			3 (50)			
Infarct location	n (ECG) on admis	ssion, n (%)								
Anterior				10 (50)			4 (67)			
Inferior				9 (45)			0			
Lateral				1 (5)			0			
Others (combin	nation)			0			2 (33)			
GCS at admissi	on, n (%)									
3–5				17 (85)			6 (100)			
6–10				1 (5)			0			
11–15				2 (10)			0			
Cardiogenic sho	ock, n (%)			8 (40)			4 (67)			
Postanoxic enc	ephalopathy, n ((%)		7 (35)			4 (67)			
Artificial pulmo	nary ventilation	ı, n (%)		17 (85)			6 (100)			
Left ventricular	ejection fractio	n ≤ 35% (Ech	o), n (%)	8 (40)			4 (67)			

G.8 Hospital volumes of PPCI

Table 82: Kumbhani et al. 2009⁶⁶

				1		
Reference	Study type	Number of patients	Intervention	Length of follow- up	Outcome measures	Source of funding
Kumbhani D, Cannon C, Fonarow G, Liang L, Askari A, Peacock W, Peterson E, Bhatt D. Association of hospital primary angioplasty volume in ST-segment elevation myocardial infarction with quality and outcomes. 2009. JAMA. Vol. 302, No. 20: p. 2007 - 2213	Retrospective observational analysis of American Heart Association's get with the guidelines – coronary artery disease national database, July 2001 to December 2007.	n = 29,513 Inclusion: ECG evidence of new ST-segment elevation or new left bundle branch block. Cardiac diagnosis of STE Exclusion: People without STEMI, no I patients receiving fibrinolytic therapy patients with 25% or more missing da patients treated at hospitals that submitted less than 30 PPCI cases over year duration of study.	and high (>70 procedures per year). PPCI, Total angioplasty volume divided into tertiles of low (< 200 procedures per year),		In-hospital mortality	Merck/Schering Plough Parmaceutical.
Baseline characteristics:						
	Hospital P	imary Angioplasty Volume				
Characteristic	Low (n = 3	900) Medium (n = 90	08) High (n = 16,605)		p value	
Age	60.5 (13.0)	60.5 (12.9)	61.1 (13.3)		0.002	
Female sex	1089 (27.9	2480 (27.5)	4875 (29.4)		0.01	
BMI, mean (SD)	28.6 (5.9)	28.8 (6.0)	28.9 (6.0)		0.02	
White	2502 (64.2	6874 (76.3)	13,887 (83.6)			
Black	211 (5.4)	605 (6.7)	793 (4.8)		0.001	
Hispanic	483 (12.4)	651 (7.2)	821 (4.9)			
Diabetes	893 (22.9)	2003 (22.2)	3537 (21.3)		0.01	
Hyperlipidemia	1495 (38.3	4016 (44.6)	7070 (42.6)		0.01	

Peripheral vascular disease	140 (3.6)	346 (3.8)	798 (4.8)	< 0.001
Prior myocardial infarction	503 (12.9)	1155 (12.8)	2506 (15.1)	< 0.001
History of heart failure	172 (4.4)	365 (4.1)	953 (5.7)	< 0.001
Stroke or TIA	149 (3.8)	317 (3.5)	690 (4.2)	0.07
Chronic or recurrent AF	120 (3.1)	259 (2.9)	587 (3.5)	0.02
COPD or asthma	268 (6.9)	664 (7.4)	1500 (9.0)	< 0.001
Smoking history	1671 (42.9)	4015 (44.6)	7099 (42.8)	< 0.001
Chronic renal insufficiency	113 (2.9)	262 (2.9)	533 (3.2)	0.17
Renal dialysis	33 (0.9)	55 (0.6)	112 (0.7)	0.47
Anaemia	12 (0.3)	27 (0.3)	51 (0.3)	0.98
Depression	23 (0.6)	127 (1.4)	143 (0.9)	0.63
Alcohol misuse	200 (5.1)	433 (4.8)	594 (3.6)	< 0.001

Data are shown as no. (%) unit. Low volume indicates less than 36 primary angioplasty procedures performed per year; medium volume, 36 to 70 primary angioplasty procedures per year; and high volume, more than 70 primary angioplasty procedures per year.

Results:

	Odds ratio (95% confidence interval)					
Model	Low versus high volume hospital (< 36 versus 36–70 PPCI/year)	p value	Medium versus high volume hospitals (36–70 versus > 70 PPCI/year)	p value	Volume as a continuous variable b	p value
Unadjusted	1.21 (0.87–1.69)	0.26	1.00 (0.77–1.31)	0.99	1.08 (0.93-1.26)	0.32
Adjusted for demographics and hospital characteristics	1.23 (0.87–1.74)	0.24	1.04 (0.79–1.38)	0.77	1.11 (0.93–1.32)	0.23
Adjusted for demographics, hospital characteristics, and past medical history	1.30 (0.91–1.87)	0.15	1.09 (0.81–1.47)	0.56	1.13 (0.95–1.36)	0.17
Adjusted for demographics,	1.22 (0.78–1.91)	0.38	1.14 (0.78–1.66)	0.49	1.13 (0.93–1.37)	0.23

characteristics, past medical history, and acute use of aspirin	hospital			
acute use of aspirin	characteristics, past			
	medical history, and			
	acute use of aspirin			
and beta blockers	and beta blockers			

^a crude in-hospital mortality rates were 3.9% for low-volume hospitals, 3.2% for medium-volume hospitals, and 3.0% for high-volume hospitals.

Table 83: Magid et al. 2000⁷⁸

Reference	Study type	Number of patients	Intervention	Outcome measures	Source of funding
Magid D, Calonge B, Rumsfeld J, Canto J, Frederick P, Every N, Barron H. relation between hospital primary angioplasty volume and mortality for patients with acute MI treated with primary angioplasty vs thrombolytic therapy. Journal of American Medical Association. 2000. Vol. 284, No. 24, 3131- 3138.	Retrospective cohort National registry of myocardial infarction (NRMI) (voluntary, database). United States. Data collected June 1 1994 to July 31 1999.	Inclusion: arrival at hospital within 12 hours of AMI onset; initial ECG ST-segment elevation or left bundle branch block; absence of cardiogenic shock; no contraindications to fibrinolytic therapy. Exclusions: hospital that did not regularly report data to the NRMI registry and those who had participated <6 months; patients who did not complete their hospital stay at a single hospital.	PPCI	In-hospital mortality	Emergency Medicine Foundation and Genetech

Baseline characteristics:

Characteristic	Low volume (≤ 16 PPCI procedures per year) (n = 1423)	Intermediate volume (≤ 16 PPCI procedures per year) (n = 8817)	High volume (≤ 49 PPCI procedures per year) (n = 11,733)
Time from onset of MI symptoms to hospital arrival, hours			
< 2	59.6	59.7	59.8
2–6	30.4	30.3	31.1
7–12	10.1	10.0	9.0

^b For every decrease in 50 procedures per year

bay (8am-4pm) 59.4 55.5 49.8 Evening (4pm-midnight) 20.6 24.3 27.7 Light (midnight-8am) 20.0 20.2 22.5 Light experientation 92.9 94.6 96.2 Fulse ≥ 100 beats/minute 11.9 11.0 9.6 Post ≥ 100 beats/minute 4.9 4.1 3.8 100-120 23.6 22.9 23.2 1-120 71.6 73.0 73.1 Hillip class 89.0 89.8 (Incheart failure) 89.0 89.8 (Incheart failure) 8.6 8.6 8.3 II (pulmonary edema) 2.4 2.4 1.9 AT-segment elevation 98.6 98.9 98.9 AT-segment of T-wave changes 6.1 4.6 3.2 Valves 14.9 16.1 14.0 BBB 2.6 2.8 2.1 BBB 2.6 2.8 2.1 Values of infarct 40.4 3.2 </th <th>Time of hospital arrival</th> <th></th> <th></th> <th></th>	Time of hospital arrival			
Seening (4pm—midnight) 20.6 24.3 27.7 Sight (midnight—Sam) 20.0 20.2 22.5 Schest pain at presentation 92.9 94.6 96.2 Schest pain at presentation 11.9 11.0 9.6 Systolic blood pressure, mm Hg 19.0 4.1 3.8 Seen Seen Seen Seen 3.8 Seen Seen	·	59.4	55.5	49.8
vilight (midnight—8am) 20.0 20.2 22.5 chest pain at presentation 92.9 94.6 96.2 vistolic blood pressure, mm Hg 11.9 11.0 9.6 vistolic blood pressure, mm Hg 4.9 4.1 3.8 10-120 23.6 22.9 23.2 1-120 71.6 73.0 73.1 villip class 89.0 89.8 (In loe art failure) 8.6 8.6 8.3 Il (pulmonary edema) 2.4 2.4 1.9 vit-segment devation 98.6 98.9 vit-segment depression 38.3 44.4 46.3 vonsp. ST-segment or T-wave changes 6.1 4.6 3.2 Quaves 14.9 16.1 14.0 BBB 2.6 2.8 2.1 Visibility of videout of infarct 40.2 40.4 37.2 videout videout MI 94.6 95.8 96.2 videout videout MI 94.6 95.8 96.2 videout videout Milin 24 h 4.2 4.4 4.3 4.2				
Lhest pain at presentation 92.9 94.6 96.2 bulse ≥ 100 beats/minute 11.9 11.0 9.6 bystolic blood pressure, mm Hg 4.9 4.1 3.8 10-120 23.6 22.9 23.2 120 71.6 73.0 73.1 dillip class (no heart failure) 89.0 89.8 89.8 (la (heart failure) 8.6 8.6 8.3 Il (pulmonary edema) 2.4 2.4 1.9 Ti-segment elevation 98.6 98.9 98.9 Ti-segment depression 38.3 44.4 46.3 Alonsp. ST-segment or T-wave changes 6.1 4.6 3.2 Quaves 14.9 16.1 14.0 BBB 2.6 2.8 2.1 BBB 2.6 2.8 2.1 Waterior location of infarct 40.2 40.4 37.2 viderior rule-out MI 94.6 95.8 96.2 Unstable angina 3.2 2.8 2.5 </td <td></td> <td></td> <td></td> <td></td>				
Systolic blood pressure, mm Hg Systolic blood pressure, mm Hg	Chest pain at presentation			
Systolic blood pressure, mm Hg Systolic blood pressure, mm Hg	Pulse ≥ 100 beats/minute	11.9	11.0	9.6
4.9 4.1 3.8 4.1 4.1 4.1 4.1 4.1 4.1 4.1 4.1 4.1 4.1	Systolic blood pressure, mm Hg			
120 73.0 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 73.1 7	< 90	4.9	4.1	3.8
dillip class (no heart failure) 89.0 89.8 (heart failure) 8.6 8.6 8.3 Il (pulmonary edema) 2.4 1.9 AT-segment elevation 98.6 98.9 AT-segment depression 38.3 44.4 46.3 Nonsp. ST-segment or T-wave changes 6.1 4.6 3.2 Quaves 14.9 16.1 14.0 BBB 2.6 2.8 2.1 BBB 3.6 3.4 3.4 Attention location of infarct 40.2 40.4 37.2 Admission diagnosis 40.4 37.2 Admission diagnosis 95.8 96.2 Unstable angina 3.2 2.8 2.5 Other 2.2 1.4 1.3 Medication within 24 h 49.2 91.7 92.9	90–120	23.6	22.9	23.2
(no heart failure)	> 120	71.6	73.0	73.1
(no heart failure)	Killip class			
In (pulmonary edema) 2.4 2.4 2.4 1.9	I (no heart failure)	89.0	89.0	89.8
ST-segment elevation 98.6 98.9 98.9 98.9 98.9 98.9 98.9 98.9	II (heart failure)	8.6	8.6	8.3
ST-segment depression 38.3 44.4 46.3 Alonsp. ST-segment or T-wave changes 6.1 4.6 3.2 Qwaves 14.9 16.1 14.0 BBB 2.6 2.8 2.1 ABBB 3.6 3.4 3.4 Anterior location of infarct 40.2 40.4 37.2 Admission diagnosis WI or rule-out MI 94.6 95.8 96.2 Unstable angina 3.2 2.8 2.5 Other 2.2 1.4 1.3 Medication within 24 h 48.9 91.7 92.9	III (pulmonary edema)	2.4	2.4	1.9
ST-segment depression 38.3 44.4 46.3 Alonsp. ST-segment or T-wave changes 6.1 4.6 3.2 Qwaves 14.9 16.1 14.0 BBB 2.6 2.8 2.1 ABBB 3.6 3.4 3.4 Anterior location of infarct 40.2 40.4 37.2 Admission diagnosis WI or rule-out MI 94.6 95.8 96.2 Unstable angina 3.2 2.8 2.5 Other 2.2 1.4 1.3 Medication within 24 h 48.9 91.7 92.9	ST-segment elevation	98.6	98.6	98.9
Nonsp. ST-segment or T-wave changes 6.1 4.6 3.2 4.0 4.0 4.0 4.0 4.0 4.0 4.0 4.0 4.0 4.0	_			
Qwaves 14.9 16.1 14.0 BBB 2.6 2.8 2.1 RBBB 3.6 3.4 3.4 Anterior location of infarct 40.2 40.4 37.2 Admission diagnosis WI or rule-out MI 94.6 95.8 96.2 Unstable angina 3.2 2.8 2.5 Other 2.2 1.4 1.3 Medication within 24 h 89.2 91.7 92.9				
RBBB 3.6 3.4 3.4 Anterior location of infarct 40.2 40.4 37.2 Admission diagnosis 40.6 95.8 96.2 Unstable angina 3.2 2.8 2.5 Other 2.2 1.4 1.3 Medication within 24 h 89.2 91.7 92.9	Q waves			
Anterior location of infarct 40.2 40.4 37.2 Admission diagnosis MI or rule-out MI 94.6 95.8 96.2 Unstable angina 3.2 2.8 2.5 Other 2.2 1.4 1.3 Medication within 24 h Aspirin 89.2 91.7 92.9	LBBB	2.6	2.8	2.1
Admission diagnosis MI or rule-out MI 94.6 95.8 96.2 Unstable angina 3.2 2.8 2.5 Other 2.2 1.4 1.3 Medication within 24 h Aspirin 89.2 91.7 92.9	RBBB	3.6	3.4	3.4
WI or rule-out MI 94.6 95.8 96.2 Unstable angina 3.2 2.8 2.5 Other 2.2 1.4 1.3 Medication within 24 h 89.2 91.7 92.9	Anterior location of infarct	40.2	40.4	37.2
WI or rule-out MI 94.6 95.8 96.2 Unstable angina 3.2 2.8 2.5 Other 2.2 1.4 1.3 Medication within 24 h 89.2 91.7 92.9	Admission diagnosis			
Other 2.2 1.4 1.3 Medication within 24 h 89.2 91.7 92.9	MI or rule-out MI	94.6	95.8	96.2
Medication within 24 h Aspirin 89.2 91.7 92.9	Unstable angina	3.2	2.8	2.5
Aspirin 89.2 91.7 92.9	Other	2.2	1.4	1.3
	Medication within 24 h			
Market 40.0	Aspirin	89.2	91.7	92.9
-blocker 49.5 52.6	b-blocker	49.6	49.3	52.6
ACE inhibitor 16.4 15.3 16.6	ACE inhibitor	16.4	15.3	16.6
Values are percentages unless otherwise indicated.	• Values are percentages unless otherwis	e indicated.		

• MI – myocardial infarction; ECG – electrocardiogram; LBBB – left bundle branch block; RBBB – right bundle branch block; ACE – angiotensin-converting enzyme.

Results:

	Low volume (≤16 PPCI procedures per year)	Intermediate volume (≤16 PPCI procedures per year)	High volume (≤49 PPCI procedures per year)
In-hospital death (%)	6.2	4.5	3.4

Table 84: Srinivas et al. 2009¹⁰⁴

Reference	Study type	Number of patients	Intervention Comparison	Outcome measures	Source of funding
Srinivas VS, Hailpern S, Koss E, Monrad ES, Alderman M. Effect of	Retrospective cohort	n = 7321	PPCI for acute myocardial infarction	In-hospital mortality	
physicians volume on the relationship between hospital	Data collected 2000	Inclusion: AMI presenting within 12h of			
volume and mortality during primary angioplasty. Journal of the American College of Cardiology. Vol. 53. No. 7. 2009	State PCI reporting	fibrinolytics.			
Baseline characteristics:	Hospital volume (PPCI	/year) ≤ 50 (n = 18)	Hospital volume (PPC	l/year) ≥ 50 (n =	18)
Patients	1148		6173		
Age, mean (SD)	61.1 (13.0)		61.2 (13.0)		
Female	30.3		28.3		
Hispanic ethnicity	13.0		6.7‡		
Black race	9.4		5.2		
Hypertension	62		60		
Diabetes	22.4		18.2¥		
Current smoker	29.4		32		
Previous MI	15.2		16.5		
Previous PCI	16.7		17.2		
Previous cardiac surgery	5.8		7.7†		

Renal failure	1.1	1.2
Need for dialysis	1.3	0.65
Current heart failure	8.9	9.2
Previous heart failure	2.8	2.3
Chronic lung disease	5.3	5.7
Vascular disease	4.2	6.6
Prior stroke	4.3	3.9
Malignant arrhythmia	4.3	3.4
Haemodynamic unstable	7.4	5.8†
Cardiogenic shock	2.5	2.1
Cardiopulmonary resuscitation	0.7	1.5
Balloon pump	6.3	6.3
Stent thrombosis	2.2	3.0
Myocardial infarction ≤ 6 hours	72	75.4†
Ejection fraction		
< 20%	1.9	1.2
20% to 29%	7.6	6.8
Single vessel disease	53.3	56.4
Left main disease	1.2	1.9
Stent used	91.1	90.9
PCI risk score, mean (SD)	10.4 (4.1)	10.2 (3.8)
Hospital mortality	5.4	3.4¥
†p < 0.05; ‡p < 0.0001; ¥p < 0.001		

Risk adjusted in-hospital mortality	у		
Annual hospital volume threshold for PPCI	N	Mortality (%)	OR (95% CI)
≤ 25 years	410	5.37	Reference

> 25 years	6911	3.62	0.61 (0.34–1.10)
450	4440	F 40	Deference
≤ 50 years	1148	5.40	Reference
> 50 years	6173	3.40	0.58 (0.38–0.88)
. 55 / 545	02.0	3	3.33 (3.33 3.33)
≤ 75 years	3159	4.24	Reference
> 75 years	4162	3.32	0.82 (0.57–1.17)
> 75 years	4102	5.52	0.02 (0.37-1.17)
^Risk adjusted mortality was calc	ulated as the rate of observed morta	lity to predicted mortality multiplied	by the statewide mortality rate of 3.75%

Table 85: Canto et al. 2000²⁰

Reference	Study type	Number of patients	Intervention	0	utcome measures	Source of funding
Canto J, Every N, Magid D, Rogers W, Malmgren J, Frederick P, French W, Tiefenbrunn A, Misra V, Kiefe C, Barron H. The volume of primary	Retrospective cohort Data collected June 1994 and March 1998	n = 36,535	PPCI patients divided into 4 of according to volume of PPCI procedures the hospital perfoyear.		-hospital mortality	Genetech and the Agency for Health Care policy and research
and survival after acute	Myocardial Infarction					
New England Journal of	patients hospitalised with					
21: 1573 – 1580.						
Baseline characteristics:	Quartile 1 (5–11 PPCI/year)	Quartile 2 (12–20 PPCI/year)	Quartile 3 (21–33 PPCI/year)	Quartile 4 (> 3 PPCI/year)	p value	
Number of patients	2825	5245	9303	19162		
Age (years) mean	61.7	61.6	61.7	62.0		
Female	30.5	31.0	30.4	30.4		
White race	82.5	84.1	87.3	90.5	≤ 0.01	
Diabetes mellitus	20.7	20.2	20.3	18.5	≤ 0.01	
Hypertension	45.4	48.4	46.6	45.4	≤ 0.01	
Current smoking	34.0	34.9	33.9	35.4		
Family history of	31.5	33.4	33.3	32.6		

Reference	Study type	Number of patients	Intervention	Out	come measures	Source of
						funding
coronary disease						
High serum cholesterol	27.2	29.3	30.9	31.6	≤ 0.01	
Angina	16.9	14.3	14.6	13.8	≤ 0.01	
Myocardial infarction	19.7	19.9	19.3	18.4		
Heart failure	4.0	4.4	4.3	3.7		
PCI	13.3	13.4	13.7	13.5	≤ 0.01	
CABG	7.4	7.3	8.6	8.3	≤ 0.01	
Stroke	5.4	5.7	5.8	4.8	≤ 0.01	
Median interval between onset of symptoms and arrival at hospital (minutes)	109.8	111.0	108.0	105.0		
Mean systolic blood pressure	136.8	137.6	138.8	139.0	≤ 0.01	
Mean pulse (beats/minute)	78.2	78.1	78.3	77.9		
Killip class (% of patients)						
I	84.9	83.8	84.9	85.7	≤ 0.01	
II	9.0	9.0	8.6	8.8		
III	2.1	3.2	2.9	2.4	≤ 0.01	
IV	4.0	4.0	3.5	3.0	≤ 0.01	
Type of myocardial infarction (% of patients)						
Q wave	72.1	75.8	75.3	76.2	≤ 0.01	
Anterior	36.5	39.1	36.5	36.1	≤ 0.01	
Length of stay	6.1	6.0	5.9	5.8		
Mean	7.5	7.6	7.5	7.3		
Crude death rate (%)	7.7	7.5	7.0	5.7	≤ 0.01	

Reference	Study type	Number of patients	Interventi	on		Outcome mea	asures	Source of funding
Adjunctive pharmacologic	al treatments and arrival to an	gioplasty:						
	Quartile 1 (n = 2825)	Quartile 2 (n = 5245)	Quartile	3 (n = 9303)	Quartile 4	(n = 19,162)	p value	
Aspirin or an antiplatelet agent (%)	88.1	38.6	89.4		90.9		< 0.01	
Beta-blocker	43.8	16.9	44.5		46.4		< 0.01	
Heparin	91.8	93.1	92.7		94.7		< 0.01	
Arrival to angioplasty, mean (minutes)	198.6	196.8	189.6		170.4			
Arrival to angioplasty, median (minutes)	129.0	135.0	129.0		118.8			
	relative risk of death among pa	tients who underwent P	PCI					
	relative risk of death among pa	tients who underwent P Quartile 2 (12–20 PP		Quartile 3 (21–	33 PPCI/year	·) Quart	ile 4 (> 33 P	PCI/year)
Unadjusted and adjusted i	relative risk of death among pa			Quartile 3 (21–3	· •		ile 4 (> 33 P 0.62–0.84)	PCI/year)
Unadjusted and adjusted i	-	Quartile 2 (12–20 PP		-)	0.72 (0	•	PCI/year)
Unadjusted and adjusted in Model* Unadjusted Adjusted for demographic	-	Quartile 2 (12–20 PP 0.96 (0.81–1.14)		0.89 (0.76–1.04)	0.72 (0 0.71 (0	0.62–0.84)	
Unadjusted and adjusted of Model* Unadjusted Adjusted for demographic Adjusted for demographic history	characteristics characteristics and medical characteristics, medical history,	Quartile 2 (12–20 PP 0.96 (0.81–1.14) 0.92 (0.76–1.10) 0.92 (0.77–1.10)		0.89 (0.76–1.04 0.84 (0.71–1.00)))	0.72 (0 0.71 (0 0.72 (0	0.62–0.84) 0.61–0.83)	
Unadjusted and adjusted and Model* Unadjusted Adjusted for demographic history Adjusted for demographic clinical presentation, and many adjusted for demographic clinical presentation.	characteristics characteristics and medical characteristics, medical history, nedication within 24 hours characteristics, medical history, cations within 24 hours, year,	Quartile 2 (12–20 PP 0.96 (0.81–1.14) 0.92 (0.76–1.10) 0.92 (0.77–1.10) 0.87 (0.71–1.07)		0.89 (0.76–1.04 0.84 (0.71–1.00 0.84 (0.71–1.00)))	0.72 (0 0.71 (0 0.72 (0	0.62–0.84) 0.61–0.83) 0.61 – 0.84)	

Table 86: Every et al. 2000⁴⁴

Reference	Study type	Number of patients	Intervention	Outcome me	asures Source of funding
Every N, Maynard C, Schulman K, Ritchie J. The	Retrospective cohort	n = 2623 (STEMI subgroup) n = 6124 (total AMI population)	PPCI in patients wi	th STEMI 30-day and 1 mortality	year None stated.
association between		(Volume quartiles b	ased on	
angioplasty procedure	Cardiovascular	and July 1995.	angioplasty proced	ures	
elderly Americans. Journal	database –		observation period	. Total	
2000. Vol. 12, no. 6: 303 -	improvement	years) with confirmed AMI who	Medicare patients	and PCI all	
	United States	within 12 hours of hospital admi	ssion.		
		Exclusion: Patients admitted wit cardiogenic shock or who receive fibrinolytic therapy prior to the performance of angioplasty; pat without data for the first 12 hou treatment.	ed ients rs of		
Baseline characteristics:	25 th quartile (~4.4 PPCI/year – medicare patients) (~10.2 PPCI/year – all patients)	patients) (~31.4	75 th quartile (~24.3 PPCI/year - medicare patients) (~56.5 PPCI/year – all patients)	100 th quartile (~47.2 PPCI/year - medicare patients) (~109.8 PPCI/year – all patients)	p value
Age	71.2 ± 8.2	71.5 ± 8.1	71.2 ± 8.2	71.6 ± 7.6	0.46
Male (%)	61.2	57.9	60.7	59.7	0.26
White race (%)	89.6	92.3	93.3	94.7	< 0.001
Diabetes	26.3	22.8	22.6	23.2	0.069
Myocardial infarction	25.2	22.7	23.8	24.2	0.451
	6.6	7.5	7.6	6.2	0.34
Heart failure	0.0	7.5	7.0	·	0.54
Apache score (mean ±)	7.8 ± 3.6	7.6 ± 3.5	7.5 ± 3.4	7.4 ± 3.4	0.008

Reference	Study type	Number of patients		Intervention		Outcome measures	Source of funding
±)							
Heart failure during hospitalisation (%)	27.6	27.5	27.8		25.3	0.163	
Recurrent myocardial infarction (%)	4.9	5.0	3.0		3.9	0.038	
Bypass surgery during hospitalisation (%)	9.7	9.8	8.3		8.5	0.328	

Results (STEMI subgroup):

Time to treatment in lowest volume centre 2.7 \pm 1.9h versus highest volume centre 2.4 \pm 1.8h; p < 0.0001

Multivariate adjustment for baseline differences by hospital 30-day mortality, OR = 0.95 per quartile, 95% CI 0.91 – 1.00, NS (30 day and 1-year mortality % presented by quartiles graphically in paper. None of the results were statistically significant but patients admitted to higher volume primary angioplasty trended toward lower 30-day and 1-year mortality).

G.9 Pre-hospital versus in-hospital fibrinolysis

Table 87: BARBASH1990A⁶; ROTH1990A⁹⁴

Reference	Study type	Number of patients	Patient Characteristics	Intervention	Comparison	Outcome	Funding source
Barbash , G., Roth, A., Hod, H., Miller, H., et al., Improved survival but not left ventricular function with early and pre-hospital treatment with tissue plasminogen activator in acute myocardial infarction. The Journal of Cardiology. 1990. 66: 261-266 Roth, A., Barbash, G., Hanoch, H., Miller, H., et al. Should thombolytic therapy be administered in the mobile intensive care unit in patients with evolving myocardial infarction? JACC, Vol 15, No. 5 April 1990:932-6	Enrolment: via the emergency ward and the mobile intensive care units which were staffed by a physician and paramedic. October 1986 to December 1987. Randomisation: 'by month' into pre-hospital or in-hospital treatment Allocation concealment: no Blinding: no ITT analysis	n = 191 Male: n = 161; 56 ± 9.4 years old Female: n = 29; 62 ± 7.1 years old; p < 0.001 Except for age there was no significant difference between men and women in either baseline characteristic or clinical outcome. 165 (87%) with first MI: 76 (40%) had anterior wall and 89 (47%) inferior wall infarctions. An additional 25 patients (13%) had a previous MI. 8 patients in whom initial clinical diagnosis of AMI was not confirmed.	Inclusion: < 72 years old, had severe chest pain for > 30 minutes, but not > 4 hours; had ST-segment elevation of 0.1 mV in at least 2 contiguous ECG leads Exclusion: LBBB on ECG; history of congestive heart failure, terminal illness or bleeding predisposition; prior cardiac surgery, SBP > 120 mmHg. Diagnosis: the team radioed senior physician on call to describe patient's history and ECG.	Rt-PA infusion initiated at home.	Same drug therapy as intervention but patients treated as soon as possible after admission with recombinant tissue plasminogen activator (rt-PA) infusion initiated in emergency room or on admission to intensive coronary care unit.	Mortality, bleeding, reinfarction, length of stay, heart failure Length of follow-up 24 months	Not stated

			Patient				Funding
Reference	Study type	Number of patients	Characteristics	Intervention	Comparison	Outcome	source

End point definitions:

Infarction – confirmed by occurrence of ischaemic chest pain lasting > 30 minutes and accompanied by ST-segment elevation of at least 0.1 mV in 2 contiguous ECG leads, followed by either an increase of creatine kinas-MB fraction to values >5% of total creatine kinase, or appearance of a Q wave not present on entry ECG. Patency – TIMI grade 2 or 3 was defined as patent. Occluded infarct-artery was accordingly defined as TIMI grade 0 or 1.

Treatment:

- 120 mg recombinant tissue plasminogen activator (rt-PA) during 6-hour infusion starting with 10mg bolus followed by continuous infusion of 50 mg during first hour, 20 mg in second hour and 10 mg during following 4 hours.
- Heparin bolus 5000IU continuing with 25000 IU/24 hours. 1.5 to 2 times each patient's baseline value. Continued 5 days. 250 mg aspirin daily.
- Antianginal and anticongestive therapy administered as needed. 2g/day intravenous lidocaine for first 24 hours.
- All patients underwent coronary angiography 72 hours after treatment unless their clinical situation necessitated earlier catheterisation. Decision to proceed to angioplasty was based on coronary anatomy.

Baseline characteristics:

	Enrolment by mobile coronary unit			Enrolment by time to treatment		
	Pre-hospital (n = 43)	In-hospital (n = 44)	Emergency ward (n = 103)	< 120 minutes (n = 96)	≥ 120 minutes (n = 94)	
Age (yrs)	59 ± 7	58 ± 8	56 ± 10.0	57 ± 8	57 ± 10	
Sex (%male)	79.1	81.8	88.3	81.0	88.0	
Clinical prognostic groups						
Anterior AMI (%)	37.2	31.8	44.6	54.1	39.4*	
Inferior AMI (%)	46.5	47.7	46.7	32.4	47.8	
Previous AMI (%)	16.3	20.5	8.7	13.5	12.8	
Angina pectoria (% patients)						
Absent	46.5	52.3	54.3	49.0	55.0	
< 6 months	27.9	36.4	34.9	32.3	35.1	
> 6 months	25.6	11.4	10.6	18.7	9.5	
Functional classification (% patients)						

Reference	Study type	Number of patients	Patient Characteristics	Intervention	Comparison	Outcome	Funding source
0 to 1	81.0	84.1	87.3	81.3		89.3	
2 to 3	19.0	15.9	12.6	18.7		10.8	
Rates over 1/3 lung fields (% patients)	19.0	9.3	10.6	15.6		8.5^	
% patients with 2- and 3- vessel CAD	48.8	60.5	56.4	53.1		57.7	
Time to treatment (minutes)	1.6 ± 0.6 ^{&}	2.2 ± 0.7	2.0 ± 0.8	1.3 ± 0.	.4	2.7 ± 0.5	

^{*}statistical difference between the prevalence of anterior wall infarction in the early (< 120 minutes) versus late (≥ 120 minutes) treatment groups, p = 0.06

Results:

Mortality by enrolment groups and time to treatment

	Mobile coronary unit		Emergency		
Time to treatment	Pre-hospital	Hospital	Ward	Total	
< 120 minutes	n = 33	n = 14	n = 51	n = 96	
≥ 120 minutes	n = 10	n = 10	n = 52	n = 94	
All patients	n = 43	n = 44	n = 103	n = 190	p value*
Mortality at 60 days					
< 120 minutes	0	0	0	0	
≥ 120 minutes	1	3	2	6	0.01
Mortality at 24 months					
(all)					
< 120 minutes	0	0	1	1	
≥ 120 minutes	3	3	3	9	0.01
Mortality at 24 months					
(cardiac only)					
< 120 minutes	0	0	1	1	
≥ 120 minutes	1	3	3	7	0.03

[^] by Canadian classification

[&]p < 0.0001

Reference Stud	y type Ni	ımber of patients	Patient Characteristics	Intervention	Comparison	Outcome	Funding source
*statistically significant different		•			•		Jource
statistically significant unferent	in mortality between the	total early and late tre	atment groups. Examin	ed by Fisher's exa	ict test for 2x2 tab	nes.	
In-hospital outcomes	Group A (MICU	, n =72)	Group B (CCU, n =	:44)	p value		
Bleeding (%)	14		9		NS		
PCI (%)	42 (65*)		56 (43*)		NS		
CABG (%)	7		5		NS		
Reinfarction (%)	13.9		13.6		NS		
Mortality (%)	5.5^		6.8 ^{\$}		NS		
Length of stay (days)	11		14		NS		
Congestive heart failure (at disch %)	arge 5 (7)		7 (16)		NS		
*No. of patients catheterised. CA	BG=coronary artery bypas	ss graft					
^ 2 x electromechanical dissocia	cion, 1 x cardiogenic shock	, 1 x rupture of left ve	ntricular free wall, 1 x af	ter coronary bypa	ass operation.		
\$ 2 x cardiogenic shock, 1 x rupto	ire of left ventricular wall.						

Table 88: Castaigne et al. 1989²³

Reference	Study type	Number of patients	Patient Characteristics	Intervention	Comparison	Length of follow- up	Outcome	Funding source
Castaigne, A., Herve, C., et al. Prehospital use of APSAC: results of a placebo- controlled study. The American Journal of Cardiology. Vol.	Design: RCT Enrolment: Val de Marne district of Paris. Randomisation: yes Allocation concealment: coded injections.	n = 100 (91 male; 9 female) Mean age 55 years for men [range 35 to 75] and 63 years for women [range 45 to 75]	Inclusion: < 75 years who had typical ischaemic chest pain for > 30 minutes and < 3 hours that did not respond to nitrates. ST-segment elevation ≥ 0.2mV in at least 2 standard leads (posterior infarction) or 3 precordial leads	n = 57^ 30U of APSAC injected > 4 minutes before arriving at the hospital.	n = 36* On arrival at hospital the code was broken and if the patient received placebo at home the physician decided	In- hospital	All-cause mortality	Not stated

Reference	Study type	Number of patients	Patient Characteristics	Intervention	Comparison	Length of follow- up	Outcome	Funding source
62: 30A-33A	Blinding: Mobile care unit physician was blinded to the treatment given but could break the code if he or she thought it necessary, or if a cardiologist was present when the ambulance reached the patient's home.	Loss to follow-up:	(anterior infarction). Exclusion: history of severe hypertension or any other 'classic contraindication' to fibrinolytic therapy.		whether fibrinolytic treatment was appropriate.			
	Sample size calculation: not stated		Diagnosis: Diagnosis made by anesthesiologist in mobile care unit.					

Coronary angiography was confirmed as soon as possible after admission. Angioplasty was indicated if the infarct-related artery was patent and if there were no other severe coronary stenoses. Coronary artery bypass graft was indicated if the infarct-related artery was patent and if there was 1 or more other coronary stenoses.

30 patients underwent angioplasty 1 to 10 days after MI. 18 patients underwent coronary artery bypass grafting 1 to 15 days after MI.

Results:	Pre-hospital (n = 57)	In-hospital (n = 36)
All-cause mortality; pre-hospital	1	1
All-cause mortality; in-hospital	2	1

^{^7} patients transferred from the placebo to the active treatment group.

^{*7} patients did not receive fibrinolytic treatment in the home or hospital.

Table 89: McAleer et al. 2006⁸⁰

Reference	Study type	Number of patients	Patient Characteristics	Intervention	Comparison	Length of follow- up	Outcome measures	Source of funding
McAleer, B., Feasibility and long term outcome of home vs hospital initiated thrombolysis. Irish journal of medical science, 2006, Volume 175, number 4.	Design: RCT Enrolment: Northern Ireland district hospital, 1988-1992 Randomisation: on call rota (SHO) Allocation concealment: no Blinding: no Sample size calculation: no No ITT analysis	n = 248	Inclusion criteria: Patients whose major symptoms were less than 6 hours; ECG evidence of MI: ST-segment elevation of at least 1 mm in 2 or more standard leads or at least 2 mm in 2 or more praecordial leads. Exclusion criteria: bleeding disorder, concurrent anticoagulant therapy, active peptic ulceration, recent stroke (< 3 months), recent surgery (< 4 weeks), severe HT (BP > 220/120 mmHg) and any patient whose life expectancy is less than 2 years.	All patients assessed by MCCU staff (physician at SHO level) then randomly allocated to receive fibrinolysis at home.	All patients assessed by MCCU staff then randomly allocated to wait to receive fibrinolysis at the hospital coronary care unit.	5 years	Mortality, time to treatment, adverse events.	Not reported

Treatment:

All patients received 1.5 million units of streptokinase in 100ml of N saline over 30 minute period. Immediately followed by IV infusion of heparin in dosage of 1000–1500U/hour such that PTTK ration was maintained at 1.5–3 times the baseline value. After 5 days of heparin therapy oral anticoagulation was substituted and continued for at least 3 months in the vast majority of cases. Ancilliary therapy (such as aspirin, beta-blockers, ACE inhibitors) was prescribed along standard post-infarct guidelines.

End point definitions:

Reperfusion:

- Rapid relief of chest pain and improvement in haemodynamic parameters
- Regression of ST-segment elevation (50% reduction in ST elevation)
- Early CK and CK-MB peak
- $\bullet \ \ Occurrence \ of specific \ reperfusion \ arrhythmias: idioventricular \ rhythm, \ ventricular \ fibrillation, \ ventricular \ ectopic \ beats$

Reference	Study type	Number of patients	Patient Characteristics	Intervention	Comparison	Length of follow- up	Outcome measures	Source of funding
Baseline charac		patients	MCCU		spital	ир	incusures	Turrumg
Male (%)			76.8	76.				
Age (mean, yea	rs)		61.0	60.	5			
Age (range, yea	rs)		37–78	37-	-95			
Previous infarct	ion		17.1	22.	3			
Anterior infarct	(%)		40.2	41.	6			
Angina (%)			43.9	41.	0			
VF prior to treat	tment (%)		6.1	7.2				
Results:								
Mortality*			MCCU	Hospital		p va	alue	
• 30 days			4/82 (4.9%)	26/166 (15.7%	5)	0.0	14	
• 1 year			8/82 (9.8%)	39/166 (23.5%	5)	0.00	09	
• 2 years			9/82 (11.0%)	46/166 (27.7%	5)	0.00	03	
• 5 years			14/81 (17.3%)	58/165 (35.2%	5)	0.00	05	
Minor bleeding	(bleeding at venepuncture, I	naematuria)	16%	15%				
Major bleeding	events		0%	0%				
• 1 patient lost	to follow-up in each group							

Table 90: Schofer et al. 98

Reference	Study type	Number of patients	Patient Characteristics	Intervention	Comparison	Length of follow- up	Outcome measures	Source of funding
Schofer, J., Butter, J., Geng., Gutschmidt. K., Herden, H., Mathey, D.,	Design: RCT Enrolment: Setting: Randomisation: assi	n = 78 (66 male, 12 female, mean age ± standard	Inclusion criteria: severe chest pain typical for myocardial ischaemia lasting > 30 minutes, arrival of	n = 40 Mobile care unit staffed by a physician and 2	n = 38 After hospital admission a 12 lead ECG performed and	In- hospital	Mortality in-hospital, bleeding, myocardial reinfarctio	Not stated.

Reference	Study type	Number of patients	Patient Characteristics	Intervention	Comparison	Length of follow- up	Outcome measures	Source of funding
Moecke, H., Polster, P., Raftopoulo, A., Sheehan, F., Voelz, P. Prehospital thrombolysis in acute myocardial infarction. The American Journal of Cardiology. 1990	to the next in the series of ampule pairs of either urokinase in ampule A and placebo in ampule B, or vice versa. Allocation concealment: Blinding: double-blind Sample size calculation: ITT analysis? No (the diagnosis of an AMI could not be confirmed at hospital admission — ne nad pulmonary embolism)	deviation 55 ± 8 years)	ambulance doctor within 4 hours of onset of symptoms, ≥ 2 mm ST-elevation in ≥ 2 ECG leads for inferior AMI and ≥ 3 mm ST elevation in ≥ 2 precordial leads for anterior AMI, age ≤ 70 years, no prior AMI, and no contrindications against fibrinolysis. Exclusion criteria: prior AMI, contraindications to fibrinolysis; age > 70 years.	emergency technicians.1 st injection given at home before hospital admission.	if patient still meets inclusion criteria and had no contraindications to fibrinolytic therapy received an IV injection of ampule B followed by heparin (1000U/hour). Patient then transferred to ICU.		n	

Definitions:

Reinfarction – new onset of symptoms combined with typical ECG changes or a secondary peak of creatine kinase to more than twice the upper limit of normal, or both.

Coronary patency – complete filling of the suspected infarct artery with a delayed or normal runoff of the contrast medium (TIMI perfusion grade 2 or 3).

Treatment:

All patients who were given fibrinolytic therapy received urokinase (2 million units IV). Administration of ampule B was followed by heparin (1000 U/hour). Coronary angiography was performed before hospital discharge unless clinical instability necessitated earlier catheterisation.

Angiography was performed before hospital discharge in 31 of 40 patients in group A and in 30 of 38 patients in group B. Angiography was not performed in 17 patients because of death before discharge (n = 3), refusal by the patient (n = 4), a wrong diagnosis (n = 1), and logistic difficulties in studying patients from other cities (n = 9).

Baseline characteristics:	Pre-hospital (Group A)	In-hospital (Group B)
Number of patients	40	38

Reference	Study type	Number of patients	Patient Characteristics	Intervention	Comparison	Length of follow- up	Outcome measures	Source of funding
Age, years (mea		57 ±		intervention	55 ± 8	чρ	incusures	Turiumg
Anterior AMI (%		47	1		32			
					32			
	owed coronary arteries (⁵⁰) ·	1\^		38 (30)			
1 2		40 (3			41 (30)			
3		7 (31			21 (30)			
Initial clinical sta	atus	, (31	ı		21 (30)			
Blood pressure								
Systolic (mean ±	· = ·	141 -	± 23 (39)		137 ± 32			
Diastolic (mean			14 (39)		82 ± 17			
Cardiogenic sho		6	()		6			
Pulmonary cong		17			14			
Resuscitation (%		5			3			
*Disease ≥ 50%	diameter stenosis							
^Where data ar	e incomplete, the numbe	er of observations is in	dicated in parentheses					
Results:		Pre-l	nospital (Group A)		In-hospital (Group I	3)		
Pre-hospital								
Bleeding		0			0			
Wrong diagnosi	S	1			1			
Death		0			0			
In-hospital								
Bleeding compli	ications							
		1			1			
• Puncture site		0			0			
		0 0			0			

Reference	Study type	Number of patients	Patient Characteristics	Intervention	Comparison	Length of follow- up	Outcome measures	Source of funding
Reinfarction		4			5			
Death		1			2			
PCI / CABG		4/1			1/0			
CABG = coronary ar	tery bypass surgery							

Table 91: Weaver et. Al. & Brouwer et al. 16,120

Reference	Study type	Number of patients	Patient Characteristics	Intervention	Comparison	Length of follow- up	Outcome measures	Source of funding
Weaver, W., Cerqeuira, M., Hallstrom, A., Litwin, P., et al. Pre-hospital initiated vs hospital initiated thrombolytic therapy, JAMA, 1993, Volume 270, No. 10. 120 Brouwer, M., Martin, J., Maynard, C., et al. Influence of early pre- hospital thrombolysis on mortality and event free	Enrolment: Paramedics obtained directed history and physical examination, then transmit an ECG to a base station emergency physician for all potentially suitable candidates for fibrinolytic therapy. Findings discussed by phone or radio with this physician before decision is made to proceed with therapy. Setting: 19 hospitals and 5 paramedic systems in the city of Seattle and	Total n = 360 (from n = 1973 who met history criteria and had ECG done) 353 (98%) of the 360 patient enrolled had subsequent evidence of acute myocardial infarction, as determined by serial enzyme tests and ECGs. 5 had no enzyme elevation and diagnosed as UA/NSTEMI; most had residual ST elevation from a prior infarction. 1 had pericarditis; 1 had peptic ulcer disease. ECG evidence of anterior injury in 39% of patients, inferior wall injury in 58%,	Inclusion criteria: <75 years who were alert, orientated and had on going chest discomfort for ≥ 15 minutes and 6 hours. SBP > 80 and < 180 mmHg, DBP < 120 mmHg and systolic BP difference between arms < 20 mmHg. 2 or more contiguous leads in a group location displayed ST-segment elevation ≥ 1 mm. Exclusion criteria: any known	Active kit containing aspirin (325 mg) and alteplase (100 mg) were administered immediately, the latter infused in open-label manner over 3 hours using special infusion pump)	No placebo was given pre-hospital, but the active treatment kit (identical to pre-hospital) was available in the emergency department.	5 years	Mortality, heart failure	Not stated

survival (The myocardial infarction triage and intervention [MITI] randomised trial. American inurnal of cardiology 1996: 78: 497-502 ¹⁶	Randomisation: kits were identical for weight and halance Allocation concealment for patients allocated to treatment in-hospital, no treatment was giver in the field. Sample size calculation 360 patients would provide 90% power to detect a 5% difference in infarct size in the composite end point with an α level of 0.05.	Reasons given were death before treatment initiated (n = 4), direct coronary angioplasty after initial physician assessment inhospital (n = 6), not treated due to spontaneous improvement in ECG abnormality by time of admission (n = 4), or because of admitting	bleeding condition, a history of stroke, seizures or transient cerebral ischaemic attacks, major surgery in preceding 2 months gastrointestinal hleeding in nast year cancer or other terminal illness known liver disease or jaundice, renal insufficiency or insulindependent diabetes, a history of active colitis, recent trauma or central line placement and warfarin therapy
Treatment:			

All patients received basic medical care (oxygen, intravenous cannulation, and rhythm monitoring) during pre-hospital period. Morphine sulphate was used for pain relief, lidocaine and atropine for arrhythmias and vasopressors and diuretics for treatment of hypotension or pulmonary oedema if prescribed by remote physicians. All patients received sodium heparin at time of hospital arrival and continued for at least 48 hours.

Baseline characteristics:

	Pre-hospital treatment group (n = 175)	Hospital treatment group (n = 185)	p value
Age, years ± SD	57 ± 10	59 ± 10	0.04
Male (%)	82	82	0.99
Prior cardiac histories			
• angina (%)	33	28	0.30
 myocardial infarction (%) 	21	20	0.69

congestive failure (%)	2	4	0.60
coronary bypass surgery (%)	4	9	0.11
ECG findings			
anterior ST elevation* (%)	39	39	0.99
inferior ST elevation (%)	58	58	0.99
• no ST elevation (%)	2	2	0.99
Haemodynamic findings			
 heart rate, beats/minute ± SD 	76 ± 17	73 ± 18	0.11
 systolic pressure, mmHg ± SD 	133 ± 23	131 ± 23	0.29
 heart rate >100 beats/min, no. 	6	4	0.72
 systolic pressure <100mmHg (%) 	8	9	0.83
Onset of symptoms to calling 911, minutes, median time (25 th percentile– 75 th percentile)	27 (30–60)	28 (11–58)	0.31
Calling 911 to randomisation, minutes, mean \pm SD	30 ± 9	28 ± 10	0.007
Randomisation to treatment, minutes, mean $\pm\text{SD}$	15 ± 11	53 ± 21	< 0.001
Randomisation to hospital arrival, mean ± SD	38 ± 12	23 ± 9	< 0.001

^{*}atypical (V_{4i} V_{5i} V₆) and lateral (I, aVL) changes were grouped with anterior (V_{1i} V_{2i} V₃) or inferior (II, II, aVF) if both abnormalities were present.

Results:

Up to discharge	Pre-hospital treatment group	Hospital treatment group	p value
All patients, mortality^, %	5.7 (2.3–9.1)	8.1 (4.2–12.0)	0.49
Anterior infarct, mortality, %	5.9	12.3	0.30
Inferior infarct, mortality, %	5.9	5.6	0.99
First infarction, mortality, %	4.4	7.5	0.40
First anterior infarction, mortality %	5.3	11.9	0.35

'Serious' bleeding, %	6	6	
Stroke	4	2	95% CI: 0.3% to 3.0%

^{*}first infarction excludes patients with known prior infarction

^{^6} due to stroke; 1 to arrhythmia; 11 to myocardial failure; 2 to cardiac rupture; 1 to pulmonary embolism; 2 to cardiac surgery; 2 associated with multisystem disease from complications following acute myocardial infarction.

	Number of events		Number of events/100 patients/year	•	
	< 70 minutes (n = 82)	≥ 70 minutes (n = 254)	< 70 minutes (n = 82)	≥ 70 minutes (n = 254)	
Angiography	27	89	11.0	13.1	0.45
Angioplasty	22	46	9.0	6.8	0.35
Bypass surgery	9	29	3.7	4.3	0.69
Congestive heart failure	4	19	1.6	2.8	0.38
Myocardial infarction	8	35	3.3	5.1	0.18
Recurrent ischaemia	11	49	4.5	7.2	0.18

Table 92: Kuhn et al. 13,65

Reference	Study type	Number of patients	Patient Characteristics	Intervention	Comparison	Outcome measures	Source of funding
Kuhn et al. Pre- hospital thrombolytic	Design:	n = 5469; n = 2750 pre-hospital;	Inclusion criteria: pain characteristic of MI and	30 units of anistreplase	Placebo when first	Mortality at 30 days	Smith Kline
therapy in patients	Enrolment: October 1988 to	n = 2719 in-hospital	≥ 30 minutes duration, or	when first	seen outside		Beecha
with suspected acute	lanuary 1992 by 163 centres in		pain < 30 minutes but not	seen by	hospital,		m
myocardial infarction	15 Furonean countries and	Dron outs: n = 73	responsive to nitrates and	emergency medical	then anistreplase		
(The European	Canada	didn't receive anv	who underwent 12-lead	personnel	after		
myocardial infarction project group). The		study treatments (39 pre-hospital, 34	ECG.	(physician)	hospital		
New England Journal	Randomisation: stratified on	in-hospital – main	Exclusion criteria: those	outside the	admission.		
of Medicine. Vol 329.	whether ST-segment elevation	III-1105pitai – IIIaiii	Exclusion criteria. those	hospital,			
N	was present in ECG: and	reasons broken	receiving oral	placebo after			
		problems with	(but aspirin, dipyridamole,	hospital			
Boissel, J. P. The		computer).	or any other antiplatelet	admission.			
European Myocardial	Allocation concealment:	343 received only	was allowed, known to				
Infarction Project: an		first injection, but	have haemorrhagic				
assessment of pre-	Double-blinded.	89 died before the	diathesis or recently active				
hospital		second injection	peptic ulcer, stroke,				
thrombolysis.	Sample size calculation: sample	could be	surgery or major trauma in				
of Cardiology 49	reduction in mortality, with a		>200mmHg or DBP >120				
Suppl, S29-S37. 1995.	power of 90 percent and an	Length of follow	mmHg; known or				
	ITT analysis	30 days	or PCI in previous 2 weeks.				
Treatment: Other trea	tments were given at the discretion	of the attending physic	cian.				
Baseline characteristic	s:						
		Pre-hospital group (n =	2750)	Hospital grou	ıp (n = 2719)		
Male sex (%)		76.8		77.0			
Age (years)		61.1 ± 12.2		61.2 ± 12.1			
Previous myocardial in	farction (%)	19.0		19.2			
•	` '						

Previous angina pectoris (%)	44.0	45.3
Previous atherosclerotic diseases (%)	15.3	16.3
Ventricular fibrillation (%)	1.5	1.3
Shock (%)	7.3	8.1
Systolic blood pressure (mmHg)	131 ± 29	131 ± 28
Diastolic blood pressure (mmHg)	79 ± 21	79 ± 15
Heart rate (beats/minute)	77 ± 19	76 ± 20
ST-segment elevation (%)	87.0	87.3
Final diagnosis		
Myocardial infarction	2408 (87.6)	2396 (88.1)
Probable myocardial infarction*	46 (1.7)	42 (1.5)
Acute coronary syndrome	206 (7.5)	184 (6.8)
Pericarditis	9 (0.3)	12 (0.4)
Aortic dissection	6 (0.2)	3 (0.1)
Other cardiac disease	25 (0.9)	32 (1.2)
Noncardiac disease	50 (1.8)	50 (1.8)
Compliance		
Both injections received	2499 (90.9)	2462 (90.5)
Pre-hospital injection only received	164 (6.0)	179 (6.6)
Hospital injection only received	0	1 (< 0.1)
No study injection received	39 (1.4)	34 (1.3)
Incomplete information	48 (1.7)	43 (1.6)
Died before hospital injection	49 (1.8)	40 (1.5)
± values are means ± SD		
* these patients died before a diagnosis could be made		
Results:		
	Pre-hospital group (n = 2750)	Hospital group (n = 2719)
Pre-hospital mortality	36	24
Total mortality at 30 days	266	303

G.10 Use of antithrombin as an adjunct to fibrinolysis

None.

G.11 Rescue PCI

Table 93: REACT 2005^{22,49}

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow- up	Protocol outcome measures	Source of funding
Gershlick AH, et al. Rescue angioplasty after failed thrombolyti c therapy for acute myocardial infarction. N Engl J Med. 2005; 353(26):275 8-68. Carver A, et al. Longer- term follow- up of patients recruited to the REACT	Design: RCT (Multicentre – 35 sites in UK) Enrolment Dec 1999 – Mar 2004. Terminated prematurely due to declining recruitment and funding issues Randomisation/allocation concealment: 24-hour computer generated allocation system Blinding: Open-label. Investigators blinded	n = 427 Drop-outs: 0 Crossover 18 (see below)	Inclusion criteria Acute myocardial infarction with ST- segment elevation of > 0.1 mV in at least 2 contiguous leads, excluding V1 Aspirin and fibrinolysis administered within 6 hours of onset of symptoms Age 21 to 85 years Ability to give informed consent At 90 minutes (± 15 minutes) after the beginning of initial fibrinolytic therapy, electrocardiogram shows failed fibrinolytic therapy – less than 50% resolution of the ST segment in the lead showing the greatest ST- segment elevation measured from the baseline (isoelectric line) to 80ms beyond the J point, with or without chest pain Rescue angioplasty, if assigned, can be performed within 12 hours of the onset of pain	Rescue PCI (n = 144) Coronary angiography, followed by angioplasty if required (< TIMI grade 3 flow and >50% stenosis in the infarct-related artery). Adjunctive strategies (stenting or GPIs) were used at the discretion of the interventionist.	Repeated fibrinolysis (n = 142) Alteplase or reteplase and IV heparin - LMWH was not used in the first 24 hours Conservative therapy (n = 141) Standard medical therapy for MI without	30 days; 6, 12 months; and mortality at a median of 4.4 years	Composite of death, recurrent MI, cerebrovas cular event, and severe heart failure at 6 months Compone n ts of the primary end point Bleeding Revascular isation	British Heart Foundation; Roche Pharmaceut icals provided reteplase for repeated fibrinolysis (its use was optional)

(Rescue Angioplasty Versus Conservativ e Treatment or Repeat Thrombolysi s) trial. J Am Coll Cardiol. 2009; 54(2):118- 26.	to outcomes. Sample size calculation: During study design (1998), it was estimated that the primary end point rate in the conservative group would approach 20%. To detect a 40% relative reduction in the rescue-PCI group 1200 patients would be required (80% power, $\alpha = 0.05$). In December 2001, on the basis of new published evidence, it was determined that a sample size of 156 patients in each group would provide 80% power ($\alpha = 0.05$) to detect the same 40% relative reduction. ITT analysis: Yes. At 6 months all components of the primary end point were recorded for 406/427 subjects.	Exclusion criteria Probable inability to gain femoral access for intervention (for example, severe peripheral vascular disease) Left bundle-branch block Life expectancy <6 months owing to noncardiac cause Previous inclusion in this trial at any time, or in any other clinical trial during the previous month Contraindication to fibrinolysis (for example, cardiopulmonary resuscitation after first fibrinolytic treatment) Haemoglobin greater than 1.5 g/dl below normal range within previous 6 hours Platelet count below normal range within previous 6 hours For patients ≥75 years: SBP >200 mm Hg, DBP >100 mm Hg, or both at any time during the current episode of pain, even if successfully reduced by therapy For patients <75 years of age: after prescription of first fibrinolytic therapy, SBP >200 mm Hg, DBP >100 mm Hg, or both on more than 1 occasion Estimated body weight <65 kg Cardiogenic shock, either in the opinion of the investigator or defined as persistent (lasting >30 minutes) systolic hypotension (<90 mm Hg) with oliguria and autonomic	Crossover between treatment groups was discouraged but allowed if a patient had on going or further chest pain associated with ST-segment reelevation or new elevation in at least 2 contiguous leads or had cardiogenic shock.	fibrinolysis or PCI. To ensure a standardised group, conservative therapy included intravenous heparin for 24 hours, irrespective of the first fibrinolytic agent. Notes: Heparin administrati on in the repeated-fibrinolysis and conservative -therapy groups was titrated to an activated partial-thrombopla s tin time ratio of 1.5 to 2.5.		(See below for definitions)	
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Mortality status was confirmed for the

	remainder (R-PC RT = 6; CT = 6). At 12 months complete clinical follow-up was available for 388 patients. At a median of 4 years, mortality obtained for 416 (missing: R-PCI = RT = 3; CT = 7). Data on missing subjects was censored at times	/427 .4 was 6/427 1;	and consider ventricular d to any other Administration weight hepat 12 hours	on of low-molecu rin within the pre cs and baseline	than					
C	last follow-up									
Crossover		Poscuo D	CI → conservative	Rescue PCI →		Penested	fibrinolysis →		Repeated fibrino	lveis 🛆
		therapy	Ci -7 conservative	repeated fibrin	olysis	-	ive therapy		rescue PCI	119313 -7
Patients who on their randomly treatment		14		2		1		:	1	
Demographics	and baseline cha									
			Repeated fibrinolysis (n	= 142)	Conservative th	nerapy (n = 1	41)	Rescue P (n = 144)		
Mean age±SD	(range) – years	(61.3±10.3 (40-85)		61.0±10.7 (37–8	85)		61.1±11.	9 (34–85)	
Male sex – no. Medical histor			114 (80.3)	3) 111		111 (78.7)		113 (78.5	5)	
Angina			32 (22.5)		29 (20.6)			32 (22.2)		

activation, with or without pulmonary oedema despite

Acute MI	23 (16.2)	17 (12.1)	14 (9.8)*
PCI	6 (4.2)	4 (2.8)	6 (4.2)
CABG	7 (4.9)	4 (2.8)	7 (4.9)
Diabetes	23 (16.2)	16 (11.3)	21 (14.6)
Hypertension	60 (42.3)	53 (37.6)	47 (32.6)
Smoking history			
Currently smoked	70 (49.6)*	65 (46.1)	68 (47.2)
Formerly smoked	41 (29.1)*	42 (29.8)	40 (27.8)
Never smoked	30 (21.3)*	34 (24.1)	36 (25.0)
Anterior infarct – no. (%)	54 (38.0)	66 (46.8)	61 (42.7)*
First fibrinolytic therapy – no. (%)			
Reteplase	43 (30.3)	28 (19.9)	42 (29.2)
Streptokinase	82 (57.7)	88 (62.4)	84 (58.3)
Tenecteplase	2 (1.4)	5 (3.5)	3 (2.1)
Tissue plasminogen activator	15 (10.6)	20 (14.2)	15 (10.4)
Time to first fibrinolytic therapy (min)			
Median	135	150	140
Interquartile range	94–217	100–210	95–240
* Data were missing for 1 patient			

Definitions of end points

Reinfarction

During index admission: further chest pain lasting more than 30 minutes and accompanied by new electrocardiographic changes (new Q waves above 0.04 second or ST-segment elevation above 0.1 mV in 2 leads for more than 30 minutes), further enzyme rise, or both

Late chest pain lasting more than 30 minutes and accompanied by new electrocardiographic changes, enzyme rise, or both

Cerebrovascular event

A new focal neurologic deficit of presumed vascular cause persisting for more than 24 hours and without evidence of a nonvascular cause according to a neurologic imaging study

Severe heart failure

Early heart failure: any new-onset cardiogenic shock or heart failure with pulmonary oedema that is resistant to medical therapy and that occurs during the index

admission and after randomisation

Late heart failure: admission to hospital for treatment of heart failure (New York Heart Association class III or IV)

Bleeding

Major bleeding: decrease in haemoglobin of at least 5 g/dl during index admission, severe bleeding event (for example, intracranial haemorrhage, haemopericardium, or haemodynamic compromise, with or without transfusion), or both

Minor bleeding: observed bleeding during index admission, with or without a decrease in haemoglobin of at least 5 g/dl, with or without transfusion

Effect Size

Rescue PCI

Of the 144 patients assigned to rescue PCI, 88 (61.1%) were recruited from hospitals with interventional capabilities

The median transfer time for patients from hospitals without interventional capabilities was 85 minutes (IQR 55 to 120). At 6 months, among patients assigned to rescue PCI, there was no significant difference in event rates between those who were transferred for intervention (16.4 %) and those who were recruited in hospitals with on-site facilities for intervention (14.6%, p = 0.80)

16 patients in this group crossed from their assigned therapy, and 128 proceeded to angiography, 13 of whom did not require angioplasty because of patent vessels Of the remaining 115 patients, only 9 were deemed to have had an unsuccessful rescue-PCI procedure; in 6 of these patients the artery was deemed not amenable to PCI, in 1 instance affecting 1 patient there was a technical failure of x-ray equipment, and in 2 patients the attempts to open the artery were unsuccessful

Rescue PCI was commenced (the wire crossed the lesion) a median of 414 minutes after the onset of pain (IQR 350 to 505). At 6 months, logistic-regression analysis indicated that the time to repeated PCI (up to 12 hours) had no significant effect on outcome

Stents were deployed in 68.5% of patients, and a glycoprotein IIb/IIIa receptor inhibitor (abciximab) was administered in 43.4%

For patients assigned to rescue PCI rather than repeated fibrinolysis, the median additional delay in the time to the assigned treatment was 84 minutes (4.6 hours for rescue PCI versus 3.2 hours for repeated fibrinolysis)

Outcomes

0 – 30 days§			0 – 6 months	0 – 6 months				0 – 12 months		
	RT (n = 142)	CT (n = 141)	R-PCI (n = 144)	RT (n = 142)	CT (n = 141)	R-PCI (n = 144)	Overall p value	RT (n = 142)	CT (n = 141)	R-PCI (n = 144)
Primary end point	Not reported									
Death from any cause (%)		15 (11)	7 (4.9)	18 (12.7)	18 (12.8)	9 (6.2)	0.12	20 (14.1)	21 (14.9)	11 (7.6)
Death from cardiac		Not reported		15 (10.6)	14 (9.9)	8 (5.6)	0.26	Not reported		

causes (%)									
Recurrent acute MI (%)	9 (6.4)	1 (0.7)	15 (10.6)	12 (8.5)	3 (2.1)	<0.01			
Cerebrovasc ular event (%)	1 (0.7)	2 (1.4)	1 (0.7)	1 (0.7)	3 (2.1)	0.63	1 (0.7)	1 (0.7)	3 (2.1)
Severe heart failure (%)	10 (7.1)	6 (4.2)	10 (7.0)	11 (7.8)	7 (4.9)	0.58	13 (9.2)	13 (9.2)	8 (5.6)
Secondary end point									
Major bleeding (no. deaths)	Not reported		7 (5)	5 (3)	4 (0)	0.65	Not reported		
Minor bleeding (no. sheath- related)			10 (3)	8 (0)	33 (28)	<0.001			
Revascularis ation – PCI or CABG (%)			33 (23.2)	29 (20.6)	19 (13.2)	0.08†	41 (28.9)	40 (28.4)	25 (17.4)
& Data reported in Kunadian I	Potal Am Hoa	rt 2007 · 152/	51.762 71 ⁶⁷						

§ Data reported in Kunadian B, et al. Am Heart J. 2007; 153(5):763-71.

Mortality at median 4.4 years

	Repeat fibrinolysis (n = 139)	Conservative therapy (n = 134)	Rescue PCI (n = 143)
All-cause mortality	31	30	16
Cardiovascular deaths*	28	23	13

^{*}Defined as cardiac or cerebrovascular death

Hazard ratios

	At 6 months			At 12 months			Median 4.4 years		
	HR	95% CI	p value	HR	95% CI	p value	HR	95% CI	p value
All-cause mortality*									

 $[\]dagger$ p = 0.05 by the log-rank test.

Rescue PCI versus repeat fibrinolysis	0.42	0.19-0.94	<0.04	Not reported			0.41	0.22-0.75	0.004
Rescue PCI versus conservative therapy	0.42	0.19–0.94	<0.04				0.43	0.23-0.79	0.006
Repeat fibrinolysis versus conservative therapy	Not repo	rted					1.04	0.63–1.72	0.89
Cardiovascular mortality									
Rescue PCI versus repeat fibrinolysis	Not repor	rted					0.43	0.22-0.83	0.0116
Rescue PCI versus conservative therapy							0.52	0.27–1.04	0.06280
Repeat fibrinolysis versus conservative therapy							0.82	0.47–1.42	0.4746
Recurrent MI									
Rescue PCI versus repeat fibrinolysis	0.23	0.09-0.62	0.004	Not reported			Not reported		
Rescue PCI versus conservative therapy	0.33	0.12-0.93	0.04						
Repeat fibrinolysis versus conservative therapy	Not repo	rted							
Revascularisation									
Rescue PCI versus repeat fibrinolysis	0.5	0.29-0.88	<0.02	0.53 [§]	0.32-0.86	0.011	Not reported		
Rescue PCI versus conservative therapy	0.58	0.33-1.04	<0.07	0.50 [§]	0.30-0.83	0.007			

Repeat fibrinolysis	1.17	0.71-1.92	0.56	1.05 [§]	0.68-1.62	0.84
versus conservative						
therapy						

^{*} Adjusted for age and diabetes at 6 months and age, previous history of angina, and diabetes at 4.4 years – the only baseline characteristics that were identified as predictors of mortality by multivariate analysis

Table 94: MERLIN 2004^{67,107,108}

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow- up	Protocol outcome measures	Source of funding
Sutton AG, et al. A randomized trial of rescue angioplasty versus a conservative approach for failed fibrinolysis in ST-segment elevation myocardial infarction: the Middlesbrou gh Early Revasculariz ation to Limit Infarction (MERLIN) trial. J Am	Design: RCT (Multicentre – 3 sites in UK) Enrolment Feb 1999 – June 2002 Randomisation: Standard random number charts and was a block randomisation process (block size 4) Allocation concealment: Telephone/sealed envelope Blinding: Open-label	n = 307 Drop-outs: 100% follow-up at 3 years Crossover: Not stated	Patients with STEMI and evidence of failure to respond to the administration of fibrinolytic therapy Presentation to the hospital within 10 h of the onset of major symptoms was required Myocardial infarction was defined by the presence of ischaemic chest pain lasting more than 30 min, unrelieved by sublingual nitrate and associated with typical ST segment elevation on the 12-lead electrocardiogram (ECG; at least 2 mm of ST-segment elevation in 2 or more contiguous chest leads and at least 1 mm in 2 or more contiguous limb leads) Failure to respond to fibrinolytic therapy was defined by a second 12-lead ECG obtained 60 minutes after the onset of fibrinolytic therapy, showing failure of the ST-segment elevation in the worst lead (the lead with maximal ST-segment elevation) to have resolved by 50%, as compared with the pre-treatment ECG (ST-segment measured 80 ms after the J	Rescue PCI (n = 153) Coronary followed by required. Unfractionated administered who had a activated clotting time of 300 s at the time of intervention. Stents, IABP, other mechanical devices, and	Conservative therapy (n = 154) Standard medical treatment after the administration of fibrinolytic therapy. Note: The use of repeat fibrinolytic therapy was discouraged but allowed during the trial if this was standard local policy. The use of heparin and other	30 days, 6 months, 1 and 3 years	1° All-cause mortality at 30 days 2° Composite of death, reinfarctio n, stroke, heart failure, and clinically driven subsequen t revasculari sation within 30 days Length of stay	Guidant, Boston- Scientifi c, Jomed, and Datasco pe

[§]Adjusted for first fibrinolytic treatment and previous PCI

Coll Cardiol. 2004; 44(2):287-96. Sutton AG, et al. One year results of the Middlesbrou gh early revascularis ation to limit infarction (MERLIN) trial. Heart. 2005; 91(10):1330-7. Kunadian B, et al. Early invasive versus conservative treatment in patients with failed fibrinolysis-no late survival benefit: the final analysis of the Middlesbrou gh Early Revascularis	Sample size calculation: Based on available data, 150 patients would be required in each arm to detect a mortality difference between 18% in the conservative group and 6% in the salvage group, with 90% power and a p value of 0.05. ITT analysis: Yes (100% follow-up at 3 years)	point), as well as the absence of an accelerated idioventricular rhythm at the time of the 60-minute ECG EXCLUSION CRITERIA Cardiogenic shock (defined by SBP ≤90 mm Hg, oliguria, and poor peripheral perfusion with or without clinical or radiologic evidence of pulmonary edema) Confounding features on the prefibrinolytic ECG, preventing ST-segment reduction analysis Reinfarction in the same ECG territory within 2 months of an original infarction Absent femoral pulses Pregnancy Presence of significant coexisting pathology likely to affect the prognosis during the follow-up period Demographics and baseline characteristics see below	adjunctive pharmacologic therapy were used at the discretion of the attending cardiologist Note: In both arms electrocardiogr aphy was performed every 3 h after the initiation of therapy for 24 hours and all patients were treated initially with 300 mg aspirin and thereafter ≥75 mg/day aspirin; use of mechanical devices and additional pharmacothera py was discretionary (see below for in-hospital treatment) If ≥1 stents were used, ticlopidine 250	treatments was at the discretion of the attending physician. Early crossover to the rescue angioplasty arm was not allowed, except for cardiogenic shock Angiography and revascularisati on were permitted for reinfarction (defined subsequently) or for recurrent ischaemia or a positive exercise test during the hospital admission.	Minor bleeding (See below for definitions)
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ation to Limit Infarctio (MERLIN) randomized trial. Am Heart J. 2007; 153(5):763- 71. Demographics and baseline characteristics		mg twice daily following an initial immediate dose of 500 mg, or clopidogrel 75 mg/day following an initial dose of 300 mg, was administered.	
	Conservative arm (n = 154)		Rescue angioplasty arm (n = 153)
Mala (0/)			
Male (%)	114 (74)		108 (71)
Age (yrs)	62.7 ± 10.9		63.0 ± 11.2
History of hypertension (%)	47 (30.5)		62 (40.5)
History of diabetes (%)	23 (14.9)		18 (11.8)
Insulin therapy (%)	7 (4.5)		4 (2.6)
Known hyperlipidemia (%)	24 (15.6)		29 (19.0)
Total cholesterol on admission (mmol/l)	5.54 ± 1.17		5.82 ± 1.32
Blood glucose on admission (mmol/l)	8.9 ± 3.0		9.0 ± 3.3
Current smoker (%)	57 (37.0)		64 (41.8)
Ex-smoker (%)	51 (33.1)		45 (29.4)
Previous MI (%)	20 (13.0)		17 (11.1)
Anterior MI (%)	62 (40.3)		74 (48.4)
Fibrinolytic therapy			
Streptokinase (%)	149 (96.8)		147 (96.1)
rt-PA (%)	5 (3.2)		6 (3.9)
Pain to lysis time (min)	170 ± 96		180 ± 120
Lysis to laboratory time (min)			146 ± 37
,			

Pain to laboratory time (min)	-	327 ± 121
ECG to laboratory time (min)	-	85 ± 36

In-hospital treatment

	Conservative arm (n = 154)	Rescue angioplasty arm (n = 153)
Immediate coronary angiography (%)	0	149 (97.4)
Immediate coronary angioplasty (%)	0	100 (65.4)
Stent(s) deployment (%)	0	77 (50.3)
Glycoprotein IIb/IIIa inhibitor (%)	0	5 (3.3)
IABP (%)	0	19 (12.4)
Additional fibrinolytic therapy (%)	18 (11.7)	0
Transfusion (%)	2 (1.3)	17 (11.1)
Discharge medication*		
Aspirin (%)	135 (97.1)	136 (97.8)
Thienopyridine (%)	21 (15.1)	84 (60.4)
Warfarin (%)	4 (2.9)	6 (4.3)
Beta-blocker (%)	114 (82.0)	110 (79.1)
Lipid-lowering medication (%)	101 (72.7)	98 (70.5)
ACE inhibitor (%)	97 (69.8)	109 (78.4)

^{*}In each arm, 139 patients were discharged from the hospital. Medication use was comparable at 3 years

Definitions of end points

Reinfarction

Repeat episode of ischaemic chest pain after recovery from the initial event, associated with typical ST-segment re-elevation on the ECG and lasting for >30 minutes despite opiate and nitrate therapy.

Stroke

Any new neurologic deficit lasting >24 hours; computed axial tomography was performed when possible

Heart failure

Requirement for diuretic treatment in the presence of typical chest X-ray characteristics, or auscultatory crackles extending at least one-third of the way up the lung fields without a previous history of chronic pulmonary disease, or

A third heart sound with persistent tachycardia

Subsequent revascularisation

Any catheter-based or surgical intervention in the conservative group and any additional revascularisation procedure in the rescue group that was not planned after the initial coronary angiogram

Minor bleeding

Transfusion was reserved for those with a fall in haemoglobin of ≥ 2 g/dl, and only if this took the total haemoglobin to <10 g/dl.

Effect Size*

	at 30 days			at 6 months	*	at 1 year			at 3 year		
	CT (n = 154)	R-PCI (n = 153)	p value	CT (n = 154)	R-PCI (n = 153)	CT (n = 154)	R-PCI (n = 153)	p value	CT (n = 154)	R-PCI (n = 153)	p value
All-cause death (%)	17 (11.0)	15 (9.8)	0.7	19 (12.3)	17 (11.1)	20 (13.0)	22 (14.4)	0.7	26 (16.9)	27 (17.6)	0.9
CAD death (%)	17 (11.0)	13 (8.5)	0.4	Not reported	d	Not reported	d		Not reported	d	
Unplanned revasculari sation (%)	31 (20.1)¥	10 (6.5)€	<0.01	40 (25.8)	19 (12.4)	46 (29.9)†	19 (12.4)§	<0.001	52 (33.8)	22 (14.4)	<0.01
Stroke (%)	1 (0.6)	7 (4.6)	0.03	2 (1.3)	7 (4.6)	2 (1.3)	8 (5.2)	0.06	4 (2.6)	10 (6.5)	0.1
Reinfarctio n (%)	16 (10.4)	11 (7.2)	0.3	20 (12.6)	12 (7.9)	22 (14.3)	16 (10.5)	0.3	23 (15)	17 (11.1)	0.3
Heart failure (%)	46 (29.9)	37 (24.2)	0.3	48 (31.3)	39 (25.3)	48 (31.2)	40 (26.1)	0.3	50 (32.5)	44 (28.8)	0.5
Minor bleeding	2 (1.3)	17 (11.1)	<0.001	NA							
Median hospital stay, days (range)	7 (2–23)	7 (2–46)	0.95	NA							

^{*}p value not reported at 6 months

¥16 patients underwent emergency revascularisation (PCI 15, 1 CABG); 1 urgent CABG for unstable angina and severe triple vessel disease; 14 unscheduled PCI for post infarction angina with ECG changes (8) or positive exercise treadmill test (6)

€7 patients underwent emergency revascularisation (5 PCI, 2 CABG); 1 urgent CABG after readmission for unstable angina; 2 patients who did not undergo rescue PCI

underwent unplanned PCI for post infarction angina (1) or positive exercise treadmill test (1)

†In addition to 30 days: 6 CABG; 9 PCI § In addition to 30 days: 5 CABG; 4 PCI

HR for all-cause mortality was calculated at 3 years from Kaplan-Meier curve 0.97 [0.57–1.65]

Table 95: RESCUE I 1994³⁹

. , , , , , , , , , , , , , , , , , , ,	igioplasty t	Conservative therapy (n = 73)	30 days	Death	None
Randomized comparison of rescue angioplasty with conservative management of patients with early failure of thrombolysi s for acute anterior myocardial infarction. Circulation. 1994; 90(5):2280-4 Randomized Comparison USA, Europe, Japan) Drop-outs: 0 precordial leads with cardiac catheterisation within 6 hours of chest pain onset; with severe ongoing chest pain, the time window could be extended to within 8 hours at the discretion of the investigator. Treatment with any acceptable intravenous fibrinolytic regimen (including but not limited to streptokinase [1.5 million U], tissue-type plasminogen activator [TPA; 100 to 125 mg], and urokinase [3 million U]) Aged 21–79 years TIMI flow grade 0–1 in the left anterior descending coronary artery (LAD) after intracoronary nitrate administration and at least 90 minutes after initiation of fibrinolytic therapy Ability to give informed consent Drop-outs: O Crossover: conservative the discretion of the investigator. Treatment with any acceptable intravenous fibrinolytic regimen (including but not limited to streptokinase [1.5 million U], tissue-type plasminogen activator [TPA; 100 to 125 mg], and urokinase [3 million U]) Aged 21–79 years TIMI flow grade 0–1 in the left anterior descending coronary artery (LAD) after intracoronary nitrate administration and at least 90 minutes after initiation of fibrinolytic therapy Ability to give informed consent Enrolment Angional served to within 8 hours at the discretion of the investigator. Treatment with any acceptable intravenous fibrinolytic regimen (including but not limited to streptokinase [1.5 million U]) Aged 21–79 years TiMI flow grade 0–1 in the left anterior descending coronary artery (LAD) after intracoro	atheterisation A atients b ceived s pirin (325 mg p	Note: Angioplasty or bypass surgery was proscribed for 72 hours.		Severe heart failure (New York Heart Associatio n functional class III or IV)	stated

Sa ca pa tc di ej (1 2- ar	emained blinded o study outcome t all times.	bradycardia <60 beats per minute) Prior myocardial infarction Left main stenosis ≥50% in diameter Demographics and baseline characteristics see below	urokinase (1 million U) was given to patients who had received fibrin-specific agents Use of IABP was discretionary Notes: Patients in both arms received aspirin (80 to 325 mg/d), IV nitrates for ≥24 hours, and IV or high-dose (>10 000 U BID) sc heparin for ≥3 days, as tolerated.			
Demographics an	nd baseline characteristics					
		Angioplasty (n = 78)		Conservative (n = 73)		
Age,y		59±11		59±11		
Sex, % male		79		85		
Diabetes, %		16		11		
Smoking, %		44		56		
Time from MI, h		4.5±1.9		4.5±1.9		
Systolic BP, mm F	Hg	126±23		135±26		

Heart rate, bpm	84±15	83±17
Killip class ≥2, %	21	26
Multivessel disease, %	34	40
Ongoing angina at the time of catheterisation, %	81	67
Proximal occlusion site, %	46	51
TIMI 1 flow, %	36	46
Angiographic collaterals, %	32	37

Effect Size

Procedures and outcomes

Angioplasty (n = 78)	Conservative (n = 73)	p value
72 (92.3)	1* (50.0)	-
7 (9.0)	4 (5.6)	0.42
	(n = 78) 72 (92.3)	(n = 78) (n = 73) 72 (92.3) 1* (50.0)

[†]Final TIMI flow ≥ 2 and stenosis ≤ 50%

^{*2} patients 'crossed over' to urgent PCI

Table 96: RESCUE II 2000⁴⁰

Reference S	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Protocol outcome measures	Source of funding
al. Review of dimmediate angioplast y after fibrinolytic therapy for acute myocardial infarction: insights from the RESCUE I, RESCUE II, and other contempor ary clinical experience s. Am si Heart J. 2000; 139(6):104 6-53. Acc	Design: RCT (no further details) Enrolment Sep 1995 – Jan L998. Terminated brematurely due to funding issues and concern that fandomising batients with TIMI If flow to CT was mappropriate Randomisation: Permuted block design stratified by site and time from infarct onset (<5 mours, >5 hours) Allocation concealment: Closed envelope Blinding: Dpen-label Gample size calculation:	n = 29 Drop-outs: Not stated Crossover: Not stated	INCLUSION CRITERIA Acute MI with ST-segment elevation ≥2.0 mV in ≥ 4 leads of a standard 12-lead electrocardiogram Aged ≥ 21 years Receipt of any accepted IV fibrinolytic regimen Cardiac catheterisation within 12 hours of infarct onset (usually precipitated by suspected incomplete fibrinolysis) TIMI 2 flow and diameter stenosis ≥ 60% after intracoronary nitroglycerin ≥ 90 minutes after onset of fibrinolytic therapy Suitability for PCI by local standards Capacity to sign informed consent EXCLUSION CRITERIA None stated Demographics and baseline characteristics see below	Rescue PCI (n = 14) Note: Stents were suggested when PCI failed to achieve a <40% stenosis or TIMI 3 flow All patients received aspirin and heparin (titrated to achieved clotting time ≥350 seconds during coronary intervention and to an activated partial thromboplast in time 60–80 seconds for ≥ 48 hours thereafter Use of β-blockers, ACEi	Conservative therapy (n = 15) No further details	30 days, 1 year	All-cause mortality Reinterven tion during index admission Reinfarction Length of stay	No details

No details ITT analysis: Yes		and IABP was discretionary	
Demographics and baseline charac	taristics		
Demographics and baseline charac	R-PCI (n = 14)	CT (n = 15)	
Age (y)	66 ± 10	59 ± 9	
Men (%)	93	93	
Hours to			
Fibrinolytic therapy	3.5 ± 2.6	2.9 ± 2.1	
Catheterisation	4.9 ± 4.2	5.4 ± 2.5	
Anterior MI	79†	40	
Cardiogenic shock (%)	0	0	
Infarct artery			
Left anterior descending (%)	79†	40	
Left circumflex (%)	0†	40	
Right coronary (%)	21	20	
LVEF (%)	44 ± 10	45 ± 9	
Treatment			
Stent (%)	29	0	
Abciximab (%)	7	0	
Technical success (%)*	100 (21% TIMI 2)	-	
†p ≤ 0.05 versus CT			
*Defined as diameter stenosis <509	% and TIMI flow ≥2		
Effect Size			
	R-PCI (n = 14)	CT (n = 15)	
In hospital			
Reintervention	0	5*	

Reinfarction	0	0
Length of stay (d)	7.6 ± 3.0	6.4 ± 1.6
30 days		
All-cause mortality	1	0
1 year		
All-cause mortality	1	1
Reintervention	4 (3 PCI, 1 CABG)	7 (all PCI)

Table 97: Belenkie 1992⁷

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Protocol outcome measures	Source of funding
Belenkie I, et al. Rescue angioplast y during myocardial infarction has a beneficial effect on mortality: a tenable hypothesis . Can J Cardiol. 1992; 8(4):357-62	Design: RCT (4 sites, Canada) Enrolment: Aug 1986 – Oct 1988 Randomisation: Method not stated Allocation concealment: Not stated Blinding: Open label Sample size	n = 28 Drop-outs: 0 Crossove r: Not stated	INCLUSION CRITERIA Aged <76 Chest pain characteristic of myocardial ischaemia present for >30 minutes unrelieved by sublingual nitroglycerin ST segment elevation >0.1 mV in ≥2 limb leads or >0.2 mV in ≥2 precordial leads Treatment with a fibrinolytic agent feasible within 3 h of symptom onset Persistently occluded IRA >3 hours after the onset of infarction EXCLUSION CRITERIA Fibrinolytic therapy contraindicated Demographics and baseline characteristics see below	Rescue PCI (n = 16) Note: All patients received fibrinolytic therapy (streptokinas e or rt-PA), heparin and indefinite aspirin. Cardiac catheterisati performed immediately after initiation of fibrinolytic therapy	Conservative therapy (n = 12)	In-hospital	Mortality Bleeding	Foothills Hospital Research and Develop ment the Heart Stroke on of Calgary

calculation: Not stated ITT analysis: Yes					
Demographics and baseline characteristics					
	Rescue PCI (n = 16)	Conservative (n = 12)			
Age (years)	58 ± 8	61 ± 13			
Male (%)	7 (44)	6 (50)			
Study myocardial infarction:					
Anterior (%)	9 (56)	6 (50)			
Inferior (%)	7 (44)	6 (50)			
1 vessel disease (%)	7 (44)	5 (42)			
2 vessel disease (%)	6 (37)	4 (33)			
3 vessel disease (%)	3 (19)	3 (25)			
Previous myocardial infarction	2 (12)	1 (8)			
Time to PCI (minutes)	257 ± 57	-			
Mortality (total)	1/16 (6.3)	4/12 (33.3)*			
Mortality (successful PCI)	0 (13)	-			
*p = 0.13 versus patients randomised to rescu	ie PCI				
Effect Size					
Outcomes – In-hospital					
	Rescue PCI (n = 16)	Conservative (n = 12)			
All-cause mortality	1	4			
Complications	1: gastrointestinal bleeding1: groin hematoma requiring transfusion	1: severe groin hematoma			
PCI was successful in 13 of 16 patients					

Table 98: Mounsey 1995⁸⁴

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow- up	Protocol outcome measures	Source of funding
Mounsey JP, et al. Rescue thrombolysi s: alteplase as adjuvant treatment after streptokinas e in acute myocardial infarction. Br Heart J. 1995; 74(4):348-53.	Design: RCT (UK) Enrolment: Not stated Randomisation: : Minimisation programme Allocation concealment: Not stated Blinding: Double-blind Sample size calculation: >32 randomly allocated patients — calculated to detect an improvement in left ventricular ejection fraction of 10%, assuming $\sigma = 10\%$, $\alpha = 0.05$ and $\beta = 0.2$	n = 37 Drop-outs: 0 Cross-over: Not stated	INCLUSION CRITERIA Presented within 6 hours of the onset of a first acute myocardial infarction, defined as chest pain of more than 30 minutes' duration unresponsive to glyceryl trinitrate; electrocardiographic ST segment elevation, either ≥2 mm in ≥2 contiguous leads V1-V6 or ≥1 mm in ≥2 contiguous limb leads; or ST depression of ≥2 mm with tall R waves in leads V1-V3 suggesting true posterior infarction Evidence of failed reperfusion (<25% reduction of ST elevation in the electrocardiographic lead with maximum ST shift on a pre-treatment electrocardiogram) 30 minutes after 1.5 MU iv streptokinase over 60 minutes EXCLUSION CRITERIA Previous Q wave myocardial infarction Previous coronary surgery Pre-existing right or left bundle branch block or fascicular block Left ventricular hypertrophy Any of the general contraindications to fibrinolysis There was no upper age limit Demographics and baseline characteristics see below	Repeat fibrinolysis (n = 19) Alteplase (rtPA) 100 mg over 3 hours Notes: All patients received heparin following initial fibrinolysis for ≥ 24 hours, and aspirin; all other treatments were discretionary	Placebo (n = 18)	6 weeks	Mortality Bleeding	Boehrin ger Ingelhei m provide d alteplas e and identical placebo

Demographics and baseline characteristics				
	Streptokinase + rtPA (n = 19)	Streptokinase + placebo (n = 18)	p value*	
Age (years)	63 (10)	63 (10)	0.9	
No (%) of men	11 (58)	13 (72)	0.4	
No (%) with infarct site:				
Anterior	10 (53)	10 (56)		
Inferior	8 (42)	6 (33)		
other	1 (5)	2 (11)	0.7	
Time to streptokinase (h)	3.4 (1.9)	4.3 (2.1)	0.2	
Time to rt-PA (h)	5.6 (1.9)	6.5 (2.2)	0.3	

^{*}For difference between rt-PA and placebo assessed by t or $\chi 2$ test.

Effect Size

Outcomes		
	Streptokinase + rtPA (n = 19)	Streptokinase + placebo (n = 18)
Mortality	1	1

There were 5 episodes of bleeding. Only 1, a gastrointestinal haemorrhage in a patient randomly allocated placebo, required transfusion. Minor bleeding included 2 haemoptyses, 1 epistaxis, and 1 rectal bleed; it was not specified which arm these patients were allocated to. There were no cerebral haemorrhages

Table 99: Sarullo 2000⁹⁷

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow- up	Protocol outcome measures	Source of funding
Sarullo FM, et al. Efficacy of rescue thrombolysi s in patients with acute myocardial infarction: preliminary findings. Cardiovasc Drugs Ther. 2000; 14(1):83-9	Design: RCT (3 sites – Italy) Enrolment: Jan 1995–Dec 1997 Randomisation: Sequentially numbered boxes Allocation concealment: Not stated Blinding: Double-blind Sample size calculation: Not stated ITT analysis: Yes	n = 90 Drop-outs: 0 Cross-over: Not stated	First episode of AMI Aged <70 years Admitted to hospital and given fibrinolysis within 4 hours of onset of pain Killip class I-II plus acceptable echocardiographic window to allow electrocardiographic images of adequate technical quality ST elevation of >1 mm in the peripheral leads or 2 mm in the pericardial leads, involving >4 leads, with concomitant alterations of the segmentary kinetics in the ECG performed at entry Basal creatine kinase within normal range Pain and ST segment elevation showed lack of response 120 minutes after starting treatment (<50% ST segment resolution) EXCLUSION CRITERIA Patients unsuitable for fibrinolysis or who had LBBB on the admission ECG History of cardiomyopathy or heart failure Patients receiving β-blockers Patients who showed no enzymatic alterations after fibrinolysis (classified as having unstable angina)	Repeat fibrinolysis (n = 45) rTPA 50 mg (10 mg bolus, 40 mg in 60 minutes) Note: All patients received nitrates, aspirin, metoprolol where possible and heparin. A continuously adjusted maintenance infusion of heparin was administered to keep the aPTT at 1.5–2.5 times laboratory control values (45s). aPTT was determined within 4 hours after heparinisation on the first day and every 12 hours the next 4 days	Placebo (n = 45)	In- hospital	Reinfarction Major bleeding Minor bleeding Urgent revasculars ation	None stated

	Demographics and baseline characteristics see below		
Demographics and baseline characteristics	see below		
	Repeat fibrinolysis (n = 45)	Placebo (n = 45)	
Sex F/M	10/35	11/34	
Age, years	56±9	57±8	
Onset of symptoms (min)	107±53	116±54	
Anterior AMI	26	24	
Lateral + inferior AMI	10	11	
Anterior + inferior AMI	9	10	
Beta-blockers	18	20	
Hypertension	30	23	
Diabetes	15	13	
Hypercholesterol	18	13	
Smokers	30	24	
Effect Size			
Outcomes – In-hospital	Repeat fibrinolysis (n = 45)	Placebo (n = 45)	
All-cause mortality (%)	3 (6.6)	13 (28.8)	
Non-fatal reinfarction (%)	7* (15.5)	0	
Major bleeding (%)	1 (2.2)	0	
Minor bleeding (%)	20 (44.4)	7 (15.5)	
Urgent revascularisation (PCI or CABG) (%)	14 (31)	1 (2.2)	
	•	tion 10–50 minutes after the start of additional rTPA administration minutes) after standard fibrinolysis	

G.12 Routine early angiography following fibrinolysis

Table 100: GRACIA-1⁴⁶

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Fernandez-Avilés F, et al. Routine invasive strategy within 24 hours of thrombolysi s versus ischaemiaguided conservative approach for acute myocardial infarction with ST-segment elevation (GRACIA-1): a randomised controlled trial. Lancet. 2004; 364(9439):1 045-53.	Design: RCT (22 sites – Spain, Portugal) Enrolment: Mar 2000 – Nov 2001 Setting 15/22 sites had onsite interventional facilities. When required patients were transferred to an interventional centre Randomisation Computerised block (6, 8 or 10 patients) randomisation Allocation concealmisedent: Central telephone system Blinding: Open label. Outcome	n = 500 Drop-outs: 10 (invasive strategy: 4 lost to 12 month follow-up; conservat ive: 1 withdrew consent; 5 lost to 12 month follow-up)	Inclusion criteria Patients aged >18 years Chest pain lasting 30 minutes — 12 hours unresponsive to nitroglycerin with ST-segment elevation ≥ 1 mm in ≥ 2 contiguous leads or a non- diagnostic ECG due to LBBB or paced rhythm Received fibrinolytic treatment with accelerated dose of alteplase within 12 hours of pain onset Exclusion criteria Cardiogenic shock (sustained SBP < 90 mm Hg, with no response to fluids, or SBP > 100 mm Hg with vasopressors (in absence of bradycardia) Suspicion or evidence of mechanical complication Non- cardiac condition with expected survival < 1 year Women with positive pregnancy test	Routine invasive (n = 248) Routine angiography and revascularisation if indicated within 6–24 hours of fibrinolysis (direct stenting of culprit artery attempted if morphologically suitable; non-culprit lesions only stented if a large amount of myocardium was threatened by stenosis. Surgery was done if stenting not feasible or when: continued ischaemia secondary to failure of culprit artery stenting; any other lesion with continuing ischaemia of a functionally important territory unsuitable for stenting; ≥ 70% left main artery stenosis Notes:	Ischaemia-guided (n = 252) Predischarge angiography and revascularis ation were only done in cases with spontaneous recurrent ischaemia with ECG changes or a non-invasive stress test, under β-blockade which identified ischaemia with a heartrate of < 100 bpm or functional capacity < 5 METS, hypotension	In-hospital, 30 days, 12 months	1° Combined death, non-fatal reinfarctio n or ischaemi-induced revasculari sation at 1 year 2° Mortality Reinfarcti on Ischaemia-induced revasculari sation Major bleeding Intracrania I bleeding	Spanish Ministry of Health, Spanish Networ for Cardiov scular Research , Spanis Society of Cardiology, Guidant , Lilly

reviewers (all events) blinded to treatment assignment Sample size calculation: 250 patients in each group based on ≥ 11% difference in primary end point (80% power; β error = 0.2) ITT analysis: Yes, 1 patient from ischaemia-guided strategy who withdrew consent excluded from analysis	Current use of warfarin or other anticoagulant drug Active bleeding or major surgery within past 2 weeks prohibiting use of heparin or antiplatelet therapy Aspirin, ticlopidine, clopidogrel or heparin contraindication Known renal failure (creatine > 221 micromol/L Any kind of stroke in the past year or haemorrhagic stroke ever Inclusion in other clinical trial Known multivessel coronary artery disease not suitable for revascularisation Major surgery pending in coming year Peripheral vascular disease prohibiting catheterisation Demographics and baseline characteristics see below	Given oral ticlopidine (500 mg) or clopidogrel (300 mg). Abciximab was strongly recommended in patients with clear evidence of thrombus. Heparin was interrupted at time of stent implantation.	on effort or ventricular tachycardia on effort Patients in both arms were given chewable aspirin (200–500 mg), IV fibrinolysis, β-blockers and ACEI. Secondary prevention measures were explained and strongly encouraged Heparinisati on maintained ≤ 48 hours after fibrinolysis in stable patients assigned to conservative treatment	Length of hospital stay Recurrent ischaemia (see below for definitions)
	Invasive (n = 248)		Conservative (n = 251)	
Mean age, years (SD)	60 (12)		61 (12)	

Man (9/)	215 (87)	214/05\
Men (%)	215 (87)	214 (85)
Hypertension (%)	82 (33)	86 (34)
Cholesterol (>5.5 mmol/L) (%)	93 (38)	105 (42)
Diabetes mellitus (%)	31 (13)	34 (14)
Family history (%)	48 (19)	53 (21)
Current smoker (%)	141 (57)	141 (56)
Previous MI (%)	17 (7)	22 (9)
Previous angina (%)	40 (16)	35 (14)
Previous treatment (%)		
Antianginal medication (%)	32 (13)	30 (12)
Aspirin (%)	32 (13)	28 (11)
Statin (%)	57 (23)	48 (19)
Anterior MI (%)	107 (43)	93 (37)
Time from onset to fibrinolysis (h)	3.04 (1.88)	3.12 (2.01)
Time from onset to randomisation (h)	14.8 (4.4)	14.2 (4.6)
Time from onset to angiography (h)*	19.6 (5.5)	NA
Medication at discharge		
Aspirin (%)	233 (94)	233 (93)
β blockade (%)	205 (83)	206 (82)
Statins (%)	178 (72)	163 (65)
ACEI (%)	119 (48)	136 (54)
Calcium antagonist (%)	15 (6)	22 (9)
Medication at 1 year	n = 244	n = 246
Aspirin (%)	228 (92)	226 (90)
β blockade (%)	201 (81)	203 (81)
ACEI (%)	126 (51)	128 (51)
Calcium antagonist (%)	32 (13)	35 (14)
*The time between fibrinolysis and angiography was (17.7 [SD 7.5] hours versus 14.6 [8.9] hours, p = 0.01)	onger for the 61 patients who were transferred from a non	-interventional centre to an interventional hospital

National Clinical Guideline Centre, 2013.

Definitions of end points

Reinfarction

Typical chest pain lasting > 30 minutes with a new increment of creatine kinase MB isoenzyme with or without new ECG abnormalities

Myocardial ischaemia

Spontaneous (at rest) or stress-induced recurrence of typical angina pectoris (or anginal equivalent) that had to coincide with new ECG abnormalities, or abnormal stress test

Ischaemia-driven revascularisation

- Any revascularisation procedure (percutaneous or surgical) involving any diseased coronary artery after identification of severe myocardial ischaemia that had to meet at least 1 of the following criteria: (a) spontaneous typical angina (at rest) with ECG changes; (b) grade III or IV effort angina (Canadian classification); and (c) stress test under β-blockade showing unequivocal ECG changes, perfusion defects, or regional contractility abnormalities along with 1 feature of poor prognosis (appearance of ischaemia before 100 bpm or 5 METS are reached); functional capacity under 5 METS; hypotension on effort; or ventricular tachycardia on effort
- Revascularisation of patients assigned to the conservative group after spontaneous ischaemia in hospital, or after detecting high-risk ischaemia in a predischarge non-invasive test, was regarded as part of this strategy. Consequently, predischarge revascularisation in the conservative group was analysed as a secondary end point, but not included as part of the primary end point. Instead, post-discharge revascularisation was regarded as part of the primary end point.

Major bleeding or vascular complication

Any complication causing death, need for surgery or transfusion, or extended time in hospital

Angiographic profile and results	
	Invasive (n = 248)
One-vessel disease (%)	146 (59)
Multivessel disease (%)	86 (35)
Angiographically non-significant CAD or normal coronary arteries (%)	16 (6)
Angioplasty with stenting of the culprit artery	199
Angioplasty of an additional artery	51
Pre/post PCI TIMI flow 0 (%)	16 (6)/ 0 (0)
Pre/post PCI TIMI flow 1 (%)	9 (4)/ 0 (0)
Pre/post PCI TIMI flow 2 (%)	22 (9)/ 4 (2)

Pre/post PCI TIMI flow 3 (%)		201 (81)/ 164 (98)	
Abciximab		64	
Not eligible for culprit artery stenting		49	
Medical treatment		40	
Non-culprit stenting		3	
CABG		2	
		C	
Condition and the standard by the form disable and 100		Conservative (n = 251)	
Cardiac catheterisation before discharge (%)	/	52 (21)	
Predischarge ischaemia-driven revascularisation	(stenting in all cases)	51*	
Unsuitable for any type of revascularisation (%)		1	
*successfully achieved in all cases			
Outcomes			
	Invasive (n = 248)		Conservative (n = 251)
Deaths, n (%)			
Index hospital stay	5 (2)		6 (2)
By 30 days	6 (2)		6 (2)
By 1 year	9 (4)		16 (6)
Non-fatal reinfarction, n (%)			
Index hospital stay	3 (1)		2 (1)
By 30 days	3 (1)		4 (2)
By 1 year	9 (4)		15 (6)
Revascularisation, n (%)			
Index hospital stay			
Induced by spontaneous ischaemia	6 (2)		30 (12)
Induced by noninvasive stress tests	0		21 (8)
By 1 year after discharge	9* (4)		30† (12)
Readmission due to ischaemia by 1 year	37 (15)		62 (25)
Major bleeding, n (%)	4 (1.6)		4 (1.6)

Intracranial haemorrhage, n (%)	0 (0)	1 (0.4)
Mean index hospital duration (days)	7.1 ± 5.6	10.5 ± 5.7

*PCI in 8 and CABG in 1. †PCI in 25 and CABG in 5

Transfer

Patients who were transported and those who were not, did not differ with respect to the frequency of minor or major cardiac events between fibrinolysis and angiography, and no patient died during this period. Only 1 event (angina) took place during transportation (1.6%); it was not reported what group this patient was allocated to.

Table 101: SIAM III²⁸

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures
Scheller B, et al. Beneficial effects of immediate stenting after thrombolysis in acute myocardial infarction. J Am Coll Cardiol. 2003; 42(4):634-41.	Design: RCT (5 sites – Germany) Enrolment: July 1998 – April 2001 Setting Patients were recruited from community hospitals without cathlab facilities located	n = 163 Drop-outs: 0	No details on whether high or low risk patients INCLUSION CRITERIA Age >18 yrs Patients presenting with symptoms of MI present for <12 hours and having, on the basis of 12-lead electrocardiography, ST-segment elevation of ≥1 mm in ≥2 limb leads, ST-segment elevation of ≥2 mm in the precordial leads, or new LBBB Patients eligible for fibrinolysis	Immediate stenting (n = 82) Transferred within 6 hours after fibrinolysis for coronary angiography, including stenting of the infarct-related artery (IRA) Note:	Elective stenting (n = 81) Elective coronary angiography 2 weeks after fibrinolysis with stenting of the IRA or earlier in case of ongoing ischaemia	In- hospital, 30 days, 6 months	Death, reinfarction, ischaemic events, and target lesion revascularisation at 6 months
Clever Y, et al. Long-term follow-up of early versus delayed invasive approach after fibrinolysis in acute	within 35 km of an interventional centre Randomisation: Computer algorithm Allocation concealment: Not stated		No secondary or iatrogenic infarction No chronic renal insufficiency requiring dialysis Secondary angiographic inclusion criteria Indication for angioplasty independent of the study Infarct-related lesion in a native coronary artery >2.5 mm	Reteplase was administered in 2 boluses of 10 MU 30 minutes apart. Patients received 250 mg of aspirin intravenously and a bolus of 5,000 IU heparin. GPIs use was discretionary.			component of primary end point Major bleeding Cerebral bleeding

myocardial infarction. Circ. Cardiovascula r interventions. 2011; 4:342-348.	Blinding: Open-label. Unclear whether outcome assessors were blinded to treatment allocation Sample size calculation: 163 patients, with a 2- sided type I error rate of 0.05, yielded 90% power to detect a decrease in the incidence of the primary end point from 50% – 25% by immediate stenting.	Diameter stenosis of ≥70% or TIMI flow <grade 3="" 6="" anatomy="" and="" angiographic="" anticipated="" area="" baseline="" below<="" characteristics="" clearly="" coronary="" criteria="" defined="" demographics="" exclusion="" for="" in="" indication="" infarct-related="" lesion="" mi="" months="" not="" of="" placement="" previous="" revascularisation="" secondary="" see="" stent="" surgical="" td="" the="" unsuitable="" vessel="" within=""><td>Heparin was continued by an infusion of 1,000 IU/h. The initial rate of heparin infusion was reduced to 800 weighing <80 kg and was adjusted to maintain an activated partial thromboplastin time of 50–70 s in all patients. Additional heparin was given in the cathlab (target activating clotting time was 250 seconds). Aspirin and clopidogrel were continued for 4</td><td>(see below for definitions)</td></grade>	Heparin was continued by an infusion of 1,000 IU/h. The initial rate of heparin infusion was reduced to 800 weighing <80 kg and was adjusted to maintain an activated partial thromboplastin time of 50–70 s in all patients. Additional heparin was given in the cathlab (target activating clotting time was 250 seconds). Aspirin and clopidogrel were continued for 4	(see below for definitions)
	Yes		weeks.	
Demographics	and baseline characteristics			
		Immediate stenting (n = 82)	Delayed stenting (n = 81)	
Male gender		76.8%	80.2%	
Age		62.4 ± 11.2 years	63.4 ± 9.9 years	
Time from sym	ptom onset to fibrinolysis	3.2 ± 2.2 hours	3.6 ± 2.6 hours	
CK		945 ± 874 IU/I	969 ± 684 IU/I	
CK-MB		125 ± 101 IU/I	116 ± 95 IU/I	
Cardiogenic sho	ock at randomisation	3.7%	1.2%	
Anterior wall in	farction	43.9%	40.7%	
Diabetes mellit	us	29.3%	34.6%	
Smoker		42.7%	38.3%	

Hypertension	62.2%	63.0%
Hyperlipidemia	51.2%	60.5%
In-hospital medical treatment		
β-blocker	77.3%	76.2%
Nitrates	57.6%	77.8%
ACE inhibitors	93.9%	93.7%
AT antagonists	1.5%	1.6%
Statins	62.1%	57.1%
Time from symptom onset to first angiography	6.7 ± 2.9 h	11.7 ± 6.8 days
Time from fibrinolysis to angiography	3.5 ± 2.3 h	11.7 ± 6.8 days
Unplanned premature angiography	0	23.5%
Abciximab	9.8%	16.0%
Stents	100%*	100%*

^{*}Based on methods section (no further details were reported in the results section)

Angiographic profile and results	Immediate stenting (n = 82)	Delayed stenting (n = 81)
Three-vessel disease (%)	28	25.9
TIMI flow before intervention (%)		
0/ 1	21 (5.8 with collaterals)	24.3 (11.1 with collaterals)
2	18.5	16.2
3	60.5	59.5
TIMI 3 flow post intervention	97.5	91.9
TIMI 3 flow after 6 months	95.5	91.2

Definitions of end points

Reinfarction:

Two or more of the following: (1) chest pain lasting for more than 30 minutes; (2) a new significant ST-elevation; (3) rise in the serum creatine kinase level to >3X ULN

Target lesion revascularisation:

Any reintervention or CABG involving the infarct-related vessel $\,$

Ischaemic events

Unplanned hospitalisation or unplanned angiography due to postinfarction angina, recurrent angina pectoris lasting >15 m despite the administration of nitrates or being accompanied by ECG changes, pulmonary oedema, or hypotension

Major bleeding

Defined as need for transfusion, bleeding requiring surgical intervention with a timely connection with the coronary intervention, bleeding documented by computed tomography or ultrasound, intracerebral as well as ocular, retroperitoneal, abdominal, intestinal, or urogenital, or a decrease in haemoglobin >4g% within 72 hours with a timely connection with the coronary intervention

Effect Size

Outcomes			
Acute phase	Early PCI (n = 82)	Delayed PCI (n = 81)	p value
Cardiogenic shock at randomisation	3 (3.7%)	1 (1.2%)	0.315
Cardiogenic shock at follow-up	5 (6.1%) (new 2)	7 (8.6%) (new 6)	0.374
Major bleeding	8 (9.8%)	6 (7.4%)	0.400
Cerebral bleeding	1 (1.2%)	2 (2.5%)	
Stroke after fibrinolysis	2 (2.4%)	2 (2.5%)	0.685
30 days			
CABG	0	0	1.000
Target lesion reintervention	2 (2.4%)	2 (2.5%)	0.685
Ischaemic events	3 (3.7%)	20 (24.7%)	0.001
Reinfarction	2 (2.4%)	2 (2.5%)	0.685
Death	4 (4.9%)	8 (9.9%)	0.179
6-months follow-up			
Follow-up	272.4 ± 191.0 days	302.0 ± 255.8 days	0.404
CABG	6 (7.3%)	6 (7.4%)	0.609
Target lesion reintervention	16 (19.5%)	19 (23.5%)	0.336
Ischaemic events	4 (4.9%)	23 (28.4%)	0.001
Reinfarction	2 (2.4%)	2 (2.5%)	0.685
Death	4 (4.9%)	9 (11.1%)	0.119
Transfer No details			

Longer-term follow-up	Early PCI (n = 82)	Delayed PCI (n = 81)	Hazard ratio (95% CI)	p
Follow-up time, years (SD)	8.1 (3.1)	7.9 (3.6)		0.603
Death	16	26	0.59 (0.32–1.11)	0.101
Ischaemic events	4	23	0.18 (0.06–0.51)	0.001
Target lesion revascularisation	23	24	0.87 (0.49–1.55)	0.646
Stroke	5	9	0.55 (0.19–1.65)	0.287
Reinfarction	6	6	0.96 (0.31–2.98)	0.944

Table 102: CAPITAL-AMI⁶⁸

Reference Study type	patients	Patient characteristics	Intervention	Comparison	Outcome measures	Source of funding
et al. RCT (4 sites – Combined Canada) angioplasty and Enrolment: gical 2004 versus s alone in When required acute patients were myocardial transferred to	n = 170 Drop-outs: 2 from TNK alone group. 1 lost to follow-up at 30 days; 1 lost to follow-up at 6 months	INCLUSION CRITERIA Patients presenting ≤6 hours of the onset of chest discomfort of ≥ 30 minutes duration and having ≥1 mm ST-segment elevation in 2 or more contiguous leads or left bundle branch block on a 12-lead electrocardiogram were eligible if they had 1 of the following highrisk criteria: 1) anterior infarction with ST-segment elevation ≥2 mm in each of 2 contiguous precordial leads; 2) extensive non-anterior infarction: 8 or more leads with ≥ 1 mm ST-segment elevation or depression or both, or the sum of ST-segment elevation >20 mm; 3) Killip class 3; or 4) systolic blood pressure <100 mm Hg EXCLUSION CRITERIA	Tenecteplase-facilitated angioplasty (n = 86) Fibrinolysis followed by immediate transfer to interventional facility. Heparin was stopped upon arrival at the cath lab and coronary angiography performed as soon as possible. Angioplasty was performed unless angiography identified diffuse disease not amenable to revascularisation or the IRA had TIMI flow grade 3 and <70% stenosis at the culprit site. Coronary stenting was performed unless the IRA was small	TNK alone (n = 84) Indications for acute angiography were persistent chest pain and ST-segment elevation ≥90 minutes after initiation of fibrinolysis or deteriorating haemodynamic status.	Composite of death, reinfarction, recurrent unstable ischaemia, or stroke at 6 months 2° Bleeding Heart failure Length of stay (index hospitalisation) Unplanned revascularisation	Canadian Institutes of Health Research (CIHR) and a CIHR Industry- Partnered with La-Roche Limited, Canada, and Corporatio

Demographics and baseline characteri	stics*		
	Tenecteplase-alone (n = 84)	Tenecteplase-facilitated angioplasty (n = 86)	p value
Age, yrs	58 (51, 66)	57 (50, 67)	0.82
Age ≥75 yrs	10.7	18.6	0.19
Male gender	76.1	75.6	1.00
Hypertension	44.0	44.2	1.00
Diabetes	11.9	20.9	0.15
Current smoking	63.1	54.7	0.28
Previous angina	16.7	23.2	0.34
Previous myocardial infarction	10.7	16.3	0.37
Previous angioplasty	3.6	5.8	0.72
Anterior index myocardial infarction	47.6	52.3	0.65
Critical time intervals, minutes			
 Onset of symptoms to hospital arrival 	64 (48, 141)	68 (45, 115)	0.68
Hospital arrival to randomisation	37 (25, 54)	34 (22, 50)	0.48
 Randomisation to tenecteplase bolus 	5 (5, 6)	5 (5, 10)	0.12
 Hospital arrival to tenecteplase bolus 	45 (34, 61)	43 (32, 55)	0.40
 Onset of symptoms to tenecteplase 	120 (90, 208)	120 (90, 153)	0.61
 Randomisation to first balloon inflation 		95 (73, 106)	
 Randomisation to first balloon inflation in 		104 (95, 111)	
transferred patients, n = 39			
 Randomisation to first balloon inflation in 		88 (63, 102)	
non-transferred patients, n = 46			

 Hospital arrival to first balloon inflation 		132 (113, 150)	
 Onset of symptoms to first balloon inflation 		204 (172, 250)	
Cardiac medication at discharge (at	6 months)		
• clopidogrel	57%	91%	<0.001
• Aspirin	96% (96%)		
Beta-blockers	93% (84%)		
• ACEI	90% (84%)		
• Lipid-lowering drugs	92% (91%)		

^{*}Values are given as percentages or medians (25th, 75th percentiles)

Definitions of end points

Reinfarction

Recurrent ischaemic symptoms at rest lasting \geq 30 minutes and accompanied by: 1) new or recurrent ST-segment elevation of \geq 1 mm in any contiguous leads; 2) new left bundle branch block; or 3) re-elevation in serum creatine kinase level to greater than twice the upper limit of normal and \geq 50% above the lowest level measured after infarction. If reinfarction occurred within 18 h, enzyme criteria were not used.

Recurrent unstable ischaemia

Recurrent symptoms of ischaemia at rest associated with new ST-segment or T-wave changes, hypotension, or pulmonary oedema.

Stroke

Focal neurological deficit, compatible with damage in the territory of a major cerebral artery with signs or symptoms persisting for >24 hours and was classified as haemorrhagic or non-haemorrhagic according to computerised tomography.

Congestive heart failure

When any 2 of the following were present: 1) dyspnea; 2) pulmonary venous congestion with interstitial or alveolar oedema on chest radiograph; 3) crackles greater than or equal to one-third of the way up the lung fields; and 4) third heart sound associated with tachycardia.

Bleeding: Classified as minor or major according to the TIMI criteria

Effect Size

Angioplasty

- Among the 86 patients assigned to tenecteplase-facilitated PCI, 40 required ambulance transfer, and 85 (99%) had coronary angiography within 3 hours from the time of randomisation. Of the latter, 79 patients (91%) underwent PCI. Stents were implanted in 77 patients (89%), and PCI with balloon alone was performed in 2 patients (2.3%)
- Platelet glycoprotein IIb/IIIa inhibitors were prescribed in 12 patients (14%) and used only when the angiographic result was suboptimal
- Angiographic success (stenosis of<50% and TIMI flow grade 3) was observed in 92% of the 79 patients who underwent facilitated PCI.

Outcomes†

	In-hospital		At 30 days		At 6 months	
	Tenecteplase alone (n = 84)	Angioplasty (n = 86)	Tenecteplas e alone (n = 84)	Angioplasty (n = 86)	Tenecteplase alone (n = 84)	Angioplasty (n = 86)
Death	3 (3.6)	2 (2.3)	3 (3.6)	Death	3 (3.6)	2 (2.3)
Reinfarction	11 (13.1)	3 (3.5)	11 (13.3)	Reinfarction	11 (13.1)	3 (3.5)
Recurrent unstable ischaemia*	15 (17.9)	5 (5.8)	15 (18.1)	Recurrent unstable ischaemia*	15 (17.9)	5 (5.8)
Stroke (due to intracranial haemorrhage)	1 (1.2)	1 (1.2)	1 (1.2)	Stroke (due to intracranial haemorrhage)	1 (1.2)	1 (1.2)
Congestive heart failure**			10 (12.1)	Congestive heart failure**		
Major bleeding	6 (7.1)	7 (8.1)		Major bleeding	6 (7.1)	7 (8.1)
Minor bleeding	11 (13.1)	20 (23.3)		Minor bleeding	11 (13.1)	20 (23.3)
Length of stay	6.0 (5.5, 8.0)	5.0 (4.0, 7.0)		Length of stay	6.0 (5.5, 8.0)	5.0 (4.0, 7.0)

†Values are given as number (percentages) or medians (25th, 75th percentiles). *Includes patients with reinfarction. **Includes patients with cardiogenic shock

Unscheduled cardiac procedures†

	In-hospital		At 30 days		At 6 months	
	Tenecteplase alone (n = 84)	Angioplasty (n = 86)	Tenecteplas e alone (n = 84)	Angioplasty (n = 86)	Tenecteplase alone (n = 84)	Angioplasty (n = 86)
Coronary angiogram	56 (66.6)	12 (14.0)	56 (67.5)	Coronary angiogram	56 (66.6)	12 (14.0)
 Time to angiography (days) 	2.5 (0, 6.0)	3.5 (1.5, 6.0)		Time to angiography (days)	2.5 (0, 6.0)	3.5 (1.5, 6.0)

 Indication 				Indication		
 Recurrent ischaemia 	33 (39.3)	5 (5.8)		Recurrent ischaemia	33 (39.3)	5 (5.8)
 Failed fibrinolysis 	8 (9.5)	-		Failed fibrinolysis	8 (9.5)	-
• Other	15 (17.8)	7 (8.1)		Other	15 (17.8)	7 (8.1)
PCI	42 (50.0)	12 (14.0)*	42 (50.6)	PCI	42 (50.0)	12 (14.0)*
CABG	2 (2.4)	0 (0.0)	2 (2.4)	CABG	2 (2.4)	0 (0.0)
†Values are given as nu	umber (percentages)					
*In 7 patients this invo	lved a coronary artery ot	her than the IRA				
Transfer No details						

Table 103: WEST⁴

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Armstrong PW, et al. A comparison of pharmacologic therapy with/without timely coronary intervention vs. primary percutaneous intervention early after ST-elevation myocardial infarction: the WEST (Which Early ST-elevation myocardial	Design: RCT (4 sites – Canada) Enrolment: Not stated Setting The protocol emphasised pre-hospital randomisation and treatment where possible Randomisation:	n = 204 Drop-outs: not stated	INCLUSION CRITERIA Eligible patients were male or non- pregnant females (≥18 years) with symptoms presumed secondary to STEMI lasting at least 20 minutes accompanied by ECG evidence of high risk. These included: ≥2 mm of ST-elevation in 2 or more contiguous precordial leads or limb leads; or ≥1 mm ST-elevation in 2 or more limb leads coupled with ≥1 mm ST-depression in 2 or more contiguous precordial leads (total ST-deviation ≥4 mm) or presumed new left bundle branch block Reperfusion therapy (PPCI, fibrinolysis or transfer for rescue PCI) was feasible within 3 hours of randomisation	Invasive strategy (n = 104) Weight-adjusted tenecteplase and mandatory invasive management within 24 hours of enrolment including protocol-specified rescue PCI, if the admission ST-	TNK only (n = 100) Weight- adjusted TNK followed by the usual standard of care	In- hospital, 30 days	1° Composite of 30-day death, reinfarctio n, refractory ischaemia, congestive heart failure, cardiogeni c shock, and major ventricular arrhythmia	Hoffma nn- LaRoch e Limited, Aventis Pharma, and Eli- Lilly.

Eur Heart I. 2006;	infarction	No details.		elevation failed	Individual
2006; 27(13):1530-8 PPLY Was deemed to be available within 1 hour of diagnosis in-hospital randomisation and time from symptom onset to randomisation (5.2 hours versus > 2 hours) Allocation concealment: No details Blinding: Open-label. Outcome assessors blinded to treatment allocation for reinfarction, recurrent ischaemia, and the indications for rescue PCI Sample size calculation: PPLY Was deemed to be available within 1 hour of diagnosis in hour of diagnosis in minutes after tenecteplase tenerapy or if haemodynamic or electrical instability on coccurred ge Note: Note: Note: Note: Note: Note: All patients received aspirin (160-325 mg) and sc enoxaparin (1 mg/kg) at randomisation with (see below for other particular parti	Therapy) study.			to decrease by	· ·
and the deprehospital randomisation and time from symptom onset to candomisation (≤ 2 hours) Allocation concealment: No details Blinding: Open-label. Outcome assessors blinded to treatment allocation for reinfarction, recurrent ischaemia, and the indications for rescue PCI Sample size calculation: Sample size calculation: Sample size calculation: Contraindications to fibrinolysis therapy or if haemodynamic or electrical intracrania or electrical or electrical intracrania or					
Inhospital inhospital inhospital randomisation and time from symptom onset to personal content of the first o	27(13):1530-8	included pre-	_	· ·	
randomisation and time from symptom onset to candomisation (≤ 2 hours versus > 2 hours) Allocation concealment: No details Blinding: Open-label. Outcome assessors blinded to treatment allocation for reinfarction, recurrent ischaemia, and the indications for rescue PCI Sample size calculation: Taybor of adys Allocation symptom onset to person and baseline characteristics see below Allocation concealment: No details Outcome assessors Blinding: Open-label. Outcome assessors Disabling stroke Note: All patients received aspirin (160-325 mg) and sc enoxaparin (1 mg/kg) at randomisation with subsequent use definitions for a 27 hours; additional iv enoxaparin (0.3-0.5 mg/kg) was permitted during PCI, post-PCI use was discretionary. Sample size calculation:		The state of the s			
symptom onset to randomisation (≤ 2 hours versus > 2 hours) Allocation concealment: No details Blinding: Open-label. Outcome assessors blinded to treatment allocation for reinfarction, recurrent ischaemia, and the indications for rescue PCI Sample size calculation: Sample size calculation: Demographics and baseline characteristics see below Note: All patients received aspirin (160- 325 mg) and sc enoxaparin (1 mg/kg) at pleeding randomisation with (see below subsequent use definitions recommended every 12 hours additional iv enoxaparin (0.3-0.5 mg/kg) was premitted during PCI, post-PCI use was discretionary. Abciximab was recommended		The state of the s	Glycoprotein IIb/IIIa antagonist use within		Intracrania
onset to randomisation (≤ 2 hours versus > 2 hours) Allocation concealment: No details Blinding: Open-label. Outcome assessors blinded to treatment allocation for reinfarction, recurrent ischaemia, and the indications for rescue PCI Sample size calculation: See below Demographics and baseline characteristics see below Note: All patients received aspirin (150-325 mg) and sc enoxaparin (1 mg/kg) at pleeding systemic bleeding randomisation with subsequent use definitions procommended every 12 hours for ≥ 72 hours; additional iv enoxaparin (0.3-0.5 mg/kg) was permitted during PCI, post-PCI use was discretionary. Sample size calculation:			7 days		haemorrha
randomisation (s ≤ 2 hours versus > 2 hours) All patients received aspirin (160— 325 mg) and sc enoxaparin (1 mg/kg) at no details Blinding: Open-label. Outcome assessors blinded to treatment allocation for reinfarction, recurrent ischaemia, and the indications for rescue PCI Sample size calculation: see below Note: All patients All patients received aspirin (160— 325 mg) and sc enoxaparin (1 mg/kg) at support (100— systemic bleeding randomisation with (see below for subsequent use definitions (see below for erommended every 12 hours additional iv enoxaparin (0.3–0.5 mg/kg) was permitted during PCI, post-PCI use was discretionary. Abciximab was recommended			Domographics and has alima share starieties	occurred	ge
(s 2 hours versus > 2 hours) Note a spirin (160—				Note:	
hours) Allocation Allocation concealment: No details Blinding: Open-label. Outcome assessors blinded to treatment allocation for reinfarction, recurrent ischaemia, and the indications for rescue PCI Sample size calculation: Allocation Amajor aspirin (160— aspirin			See Below		_
Allocation concealment: No details Blinding: Open-label. Outcome assessors blinded to treatment allocation for recurrent ischaemia, and the indications for rescue PCI Sample size Calculation: 325 mg) and sc enoxaparin (1 mg/kg) at mg/kg) at mg/kg) at penoxaparin (2 subsequent use definitions vith subsequent use every 12 hours additional iv enoxaparin (0.3-0.5 mg/kg) was permitted during PCI, post-PCI use was discretionary. Abciximab was recommended					Stroke
Allocation concealment: No details Blinding: Open-label. Outcome assessors blinded to treatment allocation for reinfarction, recurrent ischaemia, and the indications for rescue PCI Sample size calculation: Allocation mg/kg) at mg/kg) at plant bleeding mg/kg) at plant blee					The state of the s
No details Sample size Calculation: Calcula				The state of the s	
Blinding: Open-label. Outcome assessors blinded to treatment allocation for recurrent ischaemia, and the indications for rescue PCI Sample size calculation: with subsequent use definitions for definitions for every 12 hours additional iv every 12 hours; additional iv enoxaparin (0.3–0.5 mg/kg) was permitted during PCI, post-PCI use was discretionary. Sample size calculation:				= =:	bleeding
Blinding: Open-label. Outcome assessors blinded to treatment allocation for recurrent ischaemia, and the indications for rescue PCI Sample size calculation: use recommended every 12 hours for ≥ 72 hours; additional iv enoxaparin (0.3–0.5 mg/kg) was permitted during PCI, post-PCI use was discretionary. Abciximab was recommended		No details			(see below
Open-label. Outcome assessors for ≥ 72 hours; blinded to treatment allocation for reinfarction, recurrent ischaemia, and the indications for rescue PCI Sample size calculation: use recommended every 12 hours additional enoxaparin (0.3-0.5 mg/kg) was permitted during PCI, post-PCI use was discretionary. Abciximab was recommended		Blinding:			
assessors blinded to treatment allocation for reinfarction, recurrent ischaemia, and the indications for rescue PCI Sample size calculation: for ≥ 72 hours; additional iv enoxaparin (0.3–0.5 mg/kg) was permitted during PCI, post-PCI use was discretionary. Abciximab was recommended		_			definitions)
blinded to treatment allocation for (0.3–0.5 reinfarction, recurrent ischaemia, and tinizeral permitted during PCI, the indications for rescue PCI Sample size calculation: blinded to additional iv enoxaparin (0.3–0.5 mg/kg) was permitted during PCI, the indications post-PCI use was discretionary. Abciximab was recommended					
treatment allocation for ceinfarction, recurrent ischaemia, and the indications for rescue PCI Sample size calculation: enoxaparin (0.3–0.5 mg/kg) was permitted during PCI, post-PCI use was discretionary. Abciximab was recommended				·	
reinfarction, recurrent ischaemia, and the indications for rescue PCI Sample size calculation: (0.3-0.3 mg/kg) was permitted during PCI, post-PCI use was discretionary. Abciximab was recommended					
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for rescue PCI was discretionary. Sample size calculation: recommended		-		during PCI,	
Sample size Calculation: discretionary. Abciximab was recommended					
calculation: recommended					
		Feasibility		for all PCI	

nc in fib ar dis he dis th er ev th in ph th	on- nferiority of brinolytic rms (as iscussed ere) with PCI arm (not iscussed	procedures unless performed within 3 hours of fibrinolytic therapy. Clopidogrel use was according to ACC/AHA PCI guidelines.
	es	

Demographics and baseline characteristics		
	Tenecteplase only (n = 100)	Invasive strategy (n = 104)
Age, median (IQR) (years)	58 (51–69)	57 (50–67)
Gender (female)	25 (25.0)	19 (18.3)
Hypertension (yes)	37 (37.0)	56 (53.8)
Diabetes mellitus, yes (total)	18 (18.0)	8 (7.7)
Family history of early CAD (yes)	39 (39.0)	44 (42.3)
History of angina (yes)	25 (25.0)	31 (29.8)
Previous MI (yes)	14 (14.0)	12 (11.5)
Previous PCI (yes)	9 (9.0)	8 (7.7)
Smoking status (current smoker)	45 (45.0)	53 (51.0)
MI location on Q-ECG (anterior)	42 (42.0)	37 (35.6)

Killip class		
•	93 (93.9)	99 (97.1)
•	6 (6.1)	3 (2.9)

Definitions of end points

Refractory ischaemia

Symptoms of ischaemia with ST-deviation or definite T-wave inversion persisting for at least 10 minutes despite medical management while in hospital.

Recurrent MI:

- In the first 18 hours after randomisation: (a) Recurrent signs and symptoms of ischaemia at rest accompanied by new or recurrent ST-segment elevations of ≥0.1 mV in at least 2 contiguous leads lasting ≥ 30 minutes.
- After 18 h: (a) New Q-waves (by Minnesota Code Criteria) in 2 or more leads or enzyme evidence of reinfarction: re-evaluation of CK-MB or troponin to above the upper limit of normal and increased by >50% over the previous value. (b) The total CK must either be re-elevated to 2 times or more the upper limit of normal and increased by >25% or be re-elevated to >200 U/mL over the previous value. (1) If re-evaluated to less than 2 times the upper limit of normal, the total CK must exceed the upper limit of normal by >50% and exceed the previous value by 2-fold or be re-elevated to > 200 U/mL.
- Reinfarction after PCI (+/-stenting): (a) CK greater than 3 times the upper limit of normal and 50% greater than the previous value, or new Q-waves (Minnesota Code) in 2 or more contiguous leads.
- Reinfarction after CABG surgery: (a) CK greater than 5 times the upper limit of normal and ≥ 50% greater than the previous value, or new Q-waves (Minnesota Code) in 2 or more contiguous leads.

Congestive heart failure:

(i) Physician's decision to treat congestive heart failure with a diuretic, intravenous inotropic agent or intravenous vasodilator and either (a) the presence of pulmonary oedema or pulmonary vascular congestion on chest X-ray believed to be of cardiac cause or (b) at least 2 of the following: (1) rales greater than one-third up the lung fields believed to be due to congestive heart failure. (2) PCWP >18 mm Hg (3) Dyspnoea, with documented pO2 less than 80 mm Hg on room air or O2 saturation <90% on room air, without significant lung disease.

Major bleeding:

Bleeding that causes haemodynamic compromise requiring blood or fluid replacement, inotropic support, ventricular assist devices, surgical intervention, or cardiopulmonary resuscitation to maintain a sufficient cardiac output.

Median (IQR) minutes from symptom onset to treatment

	Tenecteplase only (n = 100)	Invasive strategy (n = 104)
Symptom onset to randomisation	105 (63–158)	114 (67–172)

Symptom onset to Tenecteplase		113 (74–1	79)		130 (75–185)	
Symptom onset to PCI		395 (294–	3711), n = 58		425 (288–1331), n = 8	1†
Symptom onset to rescue PCI		299 (270–	325), n = 14		277 (213–381), n = 29	
Symptom onset to non-rescue PC	1	1498 (341	–5465), n = 44		926 (398–1454), n = 5	2
First medical contact* to Tenecte	plase	51 (37–75			54 (38–77)	
First medical contact* to PCI		350 (245–	3561)		324 (218–1216)	
${\ ^\dagger Protocol\text{-}mandated procedure}.$						
*First medical contact refers to an	mbulance arrival or hos	pital arrival				
Effect Size						
Outcomes						
At 30 days		Tenectepl	ase only (n = 100)		Invasive strategy (n =	104)
• Death		4 (4.0)			1 (1.0)	
• Re-MI		9 (9.0)			6 (5.8)	
Heart failure		15 (15.0)			15 (14.4)	
Refractory ischaemia		0			3 (2.9)	
In-hospital						
Intracranial haemorrhage		0			0	
Non-haemorrhagic stroke		0			1 (1.0)	
Major systemic bleeding		1 (1.0)			2 (1.9)	
Cardiac procedures	In-hospital			At 30 days	;	
	Tenecteplase only (n = 100)		Invasive strategy (n = 104)			Tenecteplase only (n = 100)
Cardiac catheterisation	Not reported		102 (within 24 hours)	Cardiac ca	theterisation	Not reported
Revascularisation	60		89*	Revascula	risation	60
Rescue PCI	14, at a median of 197 minutes after randomisation (IQR 1 minutes).	72–280	38 (29 protocol-mandated; 6 ECG criteria)	Rescue PC	1	14, at a median of 197 minutes after randomisation (IQR 172–280 minutes).
* Of these, 8 patients received CA	BG					

- 48% of the 91 patients undergoing PCI within 24 hours of randomisation in the invasive strategy arm received abciximab; 81% of these patients had TIMI 3 and 12% had TIMI 2 flow at the end of the procedure
- Coronary stents were used in over 97% of all patients undergoing PCI

Transfer

There were no significant differences (in the primary end point) across the treatment groups in either setting (pre-hospital or in-hospital)

Table 104: TRANSFER-AMI²¹

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Cantor WJ, et al. Routine early angioplasty after fibrinolysis for acute myocardial infarction. N Engl J Med. 2009; 360(26):270 5-18.	Design: RCT (52 sites – Canada) Enrolment: July 2004 – Dec 2007. Terminated prematurely due to slowing enrolment, lack of additional funding, and loss to follow- up that was lower than anticipated Setting Enrolment was at centres that did not have the capability of performing PCI Randomisation: Performed by the	n = 1059 Drop-outs: At 30 days: 1 patient in the early-PCI group At 6 months: 9 in the early PCI group, 11 in the standard treatment group	INCLUSION CRITERIA Patients with myocardial infarction with ST-segment elevation who presented within 12 hours of symptom onset and were treated with tenecteplase ST-segment elevation of 2 mm or more in 2 anterior leads or if they had ST-segment elevation of 1 mm or more in 2 inferior leads and at least 1 of the following high- risk characteristics: systolic blood pressure of less than 100 mm Hg, heart rate of > 100 bpm, Killip class II or III, ST-segment depression of 2 mm or more in the anterior leads, or ST-segment elevation of 1 mm or more in right-sided lead V4 (V4R), which is indicative of right ventricular involvement. EXCLUSION CRITERIA Cardiogenic shock before randomisation PCI within the previous month Previous coronary-artery bypass surgery Availability of PPCI with an anticipated	Routine early PCI (n = 537) Patients were transferred to a PCI centre for angiography and PCI 6 hours after fibrinolysis. PCI was performed when persistent occlusion or substantial stenosis of IRA (stenosis of ≥70% or stenosis of 50 to 70% with thrombus, ulceration, or spontaneous	Standard treatment (n = 522) 12-lead electrocardiog raphy was repeated 60 to 90 minutes after randomisation; patients with persistent ST-segment elevation (a decrease in ST-segment elevation of less than 50%) and chest pain or with haemodynami c instability were transferred to	In- hospital, 30 days, 6 months	Combined incidence of death, reinfarctio n, recurrent ischaemia, new or worsening heart failure, or cardiogeni c shock at 30 days. 2° Death or reinfarctio n at 6 months Bleeding complicati ons	Canadia n Institute s of Health Researc h, Roche Canada, and Abbott Vascular Canada

coordinating	door-to-balloon time of less than 60	dissection) was	another		
centre or	minutes	present.	hospital for	Ison	below
physician		Stents were	rescue PCI. All		Delow
delegated by		used whenever	other patients	for	nitions
centre and was	Demographics and baseline characteristics		remained at	detii	nitions
stratified by site	see below	technically	their)	
and patient's age		possible. Use	presenting		
(> 75 versus ≤ 75		of GPIs during	hospital for at		
years)		PCI and for 12	least 24 hours.		
yearsy		hours (for			
		abciximab) or	Patients		
Allocation		18 hours (for	underwent		
concealment:		eptifibatide)	cardiac		
No details		after PCI was	catheterisatio		
		discretionary	n within 2		
Blinding:		Note:	weeks after		
Open label. All		All patients	randomisation		
		received	•		
non-fatal		tenecteplase,			
components of		aspirin, and			
the primary end		unfractionated			
point were		heparin or			
adjudicated by		enoxaparin in			
assessors blinded		the emergency			
to treatment		department.			
allocation		After April			
		2005			
Sample size		concomitant			
calculation:		treatment with			
A sample size of		clopidogrel at			
1200 patients		the time of			
(80% power to		fibrinolysis (at			
show a relative		an initial dose			
risk reduction of		of either 300			
30% with the		mg for			
early-PCI strategy,		participants			
at an alpha level of		aged ≤75 years			
0.05) assuming an		or 75 mg for			

event rate for the primary end point of 21% with the	participants aged >75 years) was
standard-	recommended.
treatment strategy and a 5% loss to	Patients aged >75 years did
follow-up.	not receive
ITT analysis:	enoxaparin.
Yes	

Demographics and baseline characteristics

	Standard treatment (n = 522)	Routine early PCI (n = 537)	p value
Age — yr			
Median	56	57	0.45
 Interquartile range 	50–66	51–66	
Age >75 yr — no. (%)	46 (8.8)	52 (9.7)	0.62
Female sex — no. (%)	105 (20.1)	111 (20.7)	0.82
Prior congestive heart failure — no. (%)	11 (2.1)	3 (0.6)	0.03
Prior myocardial infarction $-$ no. (%)	51 (9.8)	59 (11.0)	0.52
Prior PCI — no. (%)	22 (4.2)	34 (6.3)	0.12
Prior stroke or transient ischaemic attack — no. (%)	5 (1.0)	16 (3.0)	0.02
History of smoking — no./total no. (%)	316/519 (60.9)	332/533 (62.3)	0.64
Hypertension — no. (%)	178 (34.1)	173 (32.2)	0.52
Dyslipidemia — no. (%)	149 (28.5)	147 (27.4)	0.67
Diabetes — no. (%)	80 (15.3)	79 (14.7)	0.78
Killip class — no./total no. (%)			0.39
• 1	480/522 (92.0)	488/535 (91.2)	
• II	37/522 (7.1)	36/535 (6.7)	
• III	4/522 (0.8)	6/535 (1.1)	
• IV	1/522 (0.2)	5/535 (0.9)	

ST-segment elevation — no. (%)

Anterior	271 (51.9)	302 (56.2)	0.16
• Inferior¶	250 (47.9)	236 (43.9)	0.20
• With systolic BP <100 mm Hg	48 (9.2)	52 (9.7)	0.79
 With heart rate >100 bpm 	29 (5.6)	28 (5.2)	0.81
With Killip class II or III	17 (3.3)	13 (2.4)	0.41
 With ≥2-mm ST depression in the anterior leads 	162 (31.0)	157 (29.2)	0.52
 With ≥1-mm ST elevation in lead V₄R 	101 (19.3)	96 (17.9)	0.54
Time from symptom onset to administration of tenecteplase — min			
• Median	115	113	0.72
 Interquartile range 	75–191	74–182	
Time from hospital presentation to administration of tenecteplase — min‡			
• Median	25	27	0.07
• Interquartile range	16–41	17–44	
, , , , ,	patients presenting with inferior ST-segmeneristics: systolic blood pressure less than 100	•	
	depression of 2 mm or more in the anterior	<u>=</u>	
of 1 mm or more in lead V4R.			
Data were calculated on the basis of 52 ‡ Data were calculated on the basis of 52	22 patients in the standard-treatment group 22 patients in the standard-treatment group	and 535 in the early-PCI group and 536 in the early-PCI group.	
Interventions used in the 2 study groups	;		
	Standard treatment (n = 522)	Routine early PCI (n = 537)	p value
Drug therapy			
Before admission or within the first 6 hours – no. (%)			

AspirinClopidogrel

509 (97.5)

359 (68.8)

525 (97.8)

475 (88.5)

0.79

< 0.001

• With fibrinolysis – no. (%)			
Unfractionated heparin	239 (45.8)	266 (49.5)	0.22
• Enoxaparin	282 (54.0)	269 (50.1)	0.20
• At discharge – no./total no. (%)†			
Aspirin	468/506 (92.5)	487/513 (94.9)	0.11
Clopidogrel	412/506 (81.4)	463/513 (90.3)	<0.001
Beta-blocker	432/506 (85.4)	462/513 (90.1)	0.02
 ACE inhibitor or angiotensin- receptor blocker 	409/506 (80.8)	425/513 (82.8)	0.40
• Statin	453/506 (89.5)	463/513 (90.3)	0.70
Cardiac catheterisation			
• Total – no. (%)	463 (88.7)	529 (98.5)	<0.001
• Time from randomisation to insertion of arterial sheath – hours‡			
Median	32.5	2.8	<0.001
Interquartile range	4.0–69.1	2.2–3.8	
• Single-vessel coronary artery disease – no./total no. (%)	216/463 (46.7)	227/529 (42.9)	0.24
• Baseline TIMI flow – no./total no. (%)			
• 0	83/405 (20.5)	88/511 (17.2)	
• 1	37/405 (9.1)	68/511 (13.3)	
• 2	56/405 (13.8)	89/511 (17.4)	
• 3	229/405 (56.5)	266/511 (52.1)	
PCI			
• Total – no. (%)	352 (67.4)	456 (84.9)	<0.001
 Time from randomisation to first balloon inflation – hours§ 			
Median	21.9	3.2	<0.001
 Interquartile range 	3.9–73.8	2.5–4.2	

 Time from tenecteplase administration to first balloon inflation – hours¶

Median	22.7	3.9	<0.001
 Interquartile range 	4.5–74.3	3.1-4.9	
 Access – no./total no. (%) 			
• Femoral	285/349 (81.7)	382/455 (84.0)	0.39
 Thrombectomy – no./total no. (%) 	11/352 (3.1)	19/456 (4.2)	0.44
 Stent implanted – no./total no. (%) 	349/352 (99.1)	445/456 (97.6)	0.09
 Glycoprotein IIb/IIIa inhibitor use – no. (%) 	286/352 (81.2)	381/456 (83.6)	0.39
Coronary-artery bypass grafting – no. (%)	45 (8.6)	38 (7.1)	0.35

[†] The total number represents the number of patients who were discharged alive.

Definitions of end points

Reinfarction

- During the first 18 hours after enrolment, reinfarction was diagnosed on the basis of recurrent ST-segment elevation and recurrent chest pain lasting at least 30 minutes
- After 18 hours, the diagnosis of reinfarction required that there be an elevation in the MB fraction of creatine kinase to higher than the upper limit of the normal range (more than 3 times the upper limit of normal after PCI and more than 5 times the upper limit of normal after coronary-artery bypass surgery) or new Q waves

Recurrent ischaemia

• Chest pain lasting 5 minutes or longer associated with ST-segment or T-wave changes.

New or worsening heart failure

• Heart failure that required treatment 6 hours or more after enrolment and either pulmonary oedema on a chest radiograph, rales, or a pulmonary-capillary wedge pressure greater than 18 mm Hg

[‡] Data were from 449 patients in the standard-treatment group and 528 in the early-PCI group.

[§] Data were from 348 patients in the standard-treatment group and 455 in the early-PCI group.

[¶] Data were from 348 patients in the standard-treatment group and 454 in the early-PCI group.

Bleeding complications

• Classified according to TIMI and GUSTO severity scales

Effect Size

Outcomes

	Standard treatment (n = 522)	Routine early PCI (n = 536)†	p value
Efficacy end points at 30 days – no. (%)			
• Death	18 (3.4)	24 (4.5)	0.39
Reinfarction	30 (5.7)	18 (3.4)	0.06
Recurrent ischaemia	11 (2.1)	1 (0.2)	0.003
 New or worsening congestive heart failure 	29 (5.6)	16 (3.0)	0.04
Efficacy end points at 6 mo – no./total no. (%)			
• Death	23/511 (4.5)	30/528 (5.7)	0.39
Reinfarction	33/511 (6.5)	21/528 (4.0)	0.07
Safety end points during index hospitalisation – no. (%)			
 Intracranial haemorrhage 	6 (1.1)	3 (0.6)	0.34
TIMI bleeding			
• Minor	17 (3.3)	26 (4.8)	0.19
• Major	47 (9.0)	40 (7.4)	0.36
 Major, non–CABG-related 	25 (4.8)	18 (3.4)	0.24
 GUSTO bleeding 			
• Mild	47 (9.0)	70 (13.0)	0.04
Moderate	29 (5.6)	34 (6.3)	0.59
• Severe	8 (1.5)	6 (1.1)	0.55
Severe, non–CABG-related	7 (1.3)	5 (0.9)	0.53
Final TIMI flow – no./total no. (%)			0.58
• 0	3/339 (0.9)	2/451 (0.4)	

• 1	3/339 (0.9)	4/451 (0.9)	
• 2	12/339 (3.5)	24/451 (5.3)	
• 3	321/339 (94.7)	421/451 (93.3)	

†In this group, the efficacy end points were calculated on the basis of 536 patients because 1 patient was lost to follow-up at 30 days. The safety end points were calculated on the basis of all 537 patients in the group.

Transfer

Complications requiring treatment developed during transfer in 3.0% of the patients in the standard treatment group who were transferred to a PCI facility and in 2.4% of the patients in the early-PCI group (all of whom were transferred to a PCI facility). The most common complication was hypotension. The only death that occurred during transfer was of a patient in the standard treatment group who was being transferred for rescue PCI

Table 105: NORDISTEMI^{11,12}

Reference	Study type	Number of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Bøhmer E, et al. Efficacy and safety of immediate angioplasty versus ischaemiaguided managemen t after thrombolysi s in acute myocardial infarction in areas with very long transfer distances results of the	Design: RCT (5 sites – Norway) Enrolment: Feb 2005 – April 2008 Settings: Community hospitals in rural area with long transfer distances (100–400 km) to PCI and well established systems for pre-hospital fibrinolysis Randomisation: Permuted block	n = 266 Drop-outs: 0 for clinical outcomes 7 patients were unwilling to register the 15D questionna ire and were excluded from QoL analysis; 2 patients surviving	INCLUSION CRITERIA Age 18–75 yrs Symptoms of myocardial infarction present for <6 h ECG indicative of an acute STEMI: ≥2 mm ST-segment elevation in 2 contiguous precordial leads or ≥1 mm ST-segment elevation in 2 contiguous extremity leads or new left bundle branch block Expected time delay from first medical contact to PCI >90 min Receiving fibrinolytic treatment with tenecteplase	Early invasive group (n = 134) Patients were transferred to a PCI center as soon as possible after fibrinolysis, for immediate angiography and angioplasty of the IRA if indicated (≥50% diameter stenosis). Choice of stent type and use of GPIs were discretionary. As was referral for surgery in case of left main coronary artery disease or serious 3-	Conservative group (n = 132) Patients were admitted to (in case of pre-hospital fibrinolysis) or kept in community hospitals for continued care, with referral for urgent angiography if persistent chest pain and <50% reduction of ST- segment elevation 60 minutes after initiation of fibrinolysis (rescue indication) or haemodynamic	In- hospital, 30 days, 12 months	1° Composite of death, reinfarctio n, stroke, or new myocardial ischaemia at 12 months 2° Composite stroke, or n at 12 months	Scientific Board of the Eastern Norway Regional Health Authority, Hamar, Norway; Ada and Hagbarth Waage's Humanitære og Veldedige Stiftelse, Oslo,

NORDISTEM I (Norwegian study on District treatment of ST-elevation myocardial infarction). J Am Coll Cardiol. 2010; 55(2):102-10. Bøhmer E, et al. Health and cost consequenc es of early versus late invasive strategy after thrombolysi s for acute myocardial infarction. Eur J Cardiovasc Prev Rehabil. 2011; 18(5):717-23	randomisation stratified by site. The random allocation sequence was generated at Oslo University Hospital Allocation concealment: Sealed envelopes Blinding: Open-label. A blinded, independent committee adju- dicated all possible events related to the primary outcome. Sample size calculation: Total of 266 patients required (2-sided alpha of 5% and a power of 80%) based on a 30% and 15% occurrence of the primary end point in the conservative group and early invasive group respectively. ITT analysis: Yes	cerebral haemorrha ge were unable to report QoL for the first 3 months. Here, missing data were replaced by age- and sex- adjusted data from an unpublishe d Finnish study of stroke patients	EXCLUSION CRITERIA Standard exclusion criteria for tenecteplase Cardiogenic shock or serious arrhythmias at randomisation Pregnancy Known serious renal failure (serum creatine > 250 mmol/l) Other diseases with life expectancy <12 months Psychiatric disease, mental retardation, dementia, drug abuse, alcoholism, or conditions that can severely reduce compliance Demographics and baseline characteristics see below	vessel disease. Note: All patients received standard weight-adjusted dose tenecteplase, aspirin 300 mg orally, and iv enoxaparin 30 mg followed by a sc dose of 1 mg/kg repeated every 12 hours up to hospital discharge or revascularisation for a maximum of 7 days. All patients also received clopidogrel 300 mg on the first day. Beta-blockers and statins were given unless contraindicated, ACE inhibitors when indicated. Patients receiving stents were recommended clopidogrel 75 mg daily for 9 months. Other patients were recommended clopidogrel until angiography or for 9 months	instability. Referral for early angiography was recommended if the patient had spontaneous recurrent ischaemia with or without ECG changes, or if signs of ischaemia (chest pain, significant ST-segment changes, hypotension, or ventricular tachycardia) occurred in the exercise ECG recommended before hospital discharge. Angiography was encouraged in all other patients as a routine assessment after successful fibrinolysis within 2 to 4 weeks after discharge.		health quality of life (HRQoL) Bleeding at 30 days Hospital stay (see below for definitions)	Norway; and the Innland et Hospital Trust, Hamar, Norway
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Demographics and baseline character	istics		
	Early invasive group (n = 134)	Conservative group (n = 132)	p value
Age (yrs)	60 (55, 67)	61 (53, 68)	0.98
Men	107 (80%)	94 (71%)	0.13
Treated hypertension	33 (25%)	50 (38%)	0.03
Diabetes mellitus	8 (6%)	10 (8%)	0.78
Current or previous smoker	106 (79%)	104 (79%)	0.93
Family history of coronary heart disease	77 (58%)	78 (59%)	0.88
Previous angina	17 (13%)	17 (13%)	0.89
Previous myocardial infarction	15 (11%)	14 (11%)	0.97
Previous coronary bypass graft	4 (3%)	3 (2%)	1.0
Previous angioplasty	9 (7%)	12 (9%)	0.62
Infarct location on ECG			
• Anterior	59 (44%)	51 (39%)	0.44
New left bundle branch block	4 (3%)	0 (0%)	0.12
Nonanterior	71 (53%)	81 (61%)	0.21
Time intervals (minutes)			
 Symptom onset to first medical contact 	67 (34, 122)	65 (40, 135)	0.68
• Symptom onset to tenecteplase	117 (80, 195)	126 (80, 195)	0.72
Pre-hospital tenecteplase	80 (60%)	71 (54%)	0.40

- At discharge, 98% of patients were prescribed aspirin, 92% clopidogrel, 90% beta-blockers, and 99% statins, with no difference between groups
- At 1 year, medication was unchanged except for clopidogrel, which was used in only 21% of patients (no difference between groups)

Invasive procedures

	Early invasive group (n = 134)	Conservative group (n = 132)	p value
Coronary angiography performed	133 (99%)	125 (95%)	0.04
Time from tenecteplase to arrival at	130 (105, 155) min	5.5 (0, 17.5) days	< 0.001
catheterisation laboratory			

PCI performed	119 (89%)	94 (71%)	0.001
Time from tenecteplase to first balloon	163 (137, 191) min	3.0 (0, 13) days	<0.001
Radial access	111 (83%)	118 (89%)	0.17
Stents implanted	115 (86%)	90 (68%)	0.001
Abciximab	16 (14%)	8 (6%)	0.14
Thrombectomy	0 (0%)	1 (0.8%)	0.5
Intra-aortic balloon pump	2 (1.5%)	2 (1.5%)	1.0
CABG performed	9 (7%)	16 (12%)	0.19
Repeat angiography during follow-up	23 (17%)	17 (13%)	0.42
Repeat PCI during follow-up	15 (11%)	9 (7%)	0.30

Data are n (%) or median (25th, 75th percentiles).

• In the conservative group, 36 patients (27%) were transferred for rescue coronary intervention, and 32 patients underwent rescue PCI

Angiographic characteristics of the early invasive group and the rescue population of the conservative group

	Early invasive group (n = 134)	Conservative/ rescue group (n = 36)	p value
Time from tenecteplase to arrival at catheterisation laboratory (minutes)	130 (105, 155)	187 (149, 240)	<0.001
Immediate/rescue PCI performed	118 (88%)	32 (89%)	1.0
Time from tenecteplase to first balloon (minutes)	162 (137, 189)	20 (193, 247)	<0.001
Symptom onset to first balloon (minutes)	302 (236, 380)	340 (279, 400)	0.04
TIMI flow grade pre-PCI	n = 133	n = 36	
• 0	17 (13%)	9 (25%)	0.12
• 1	10 (8%)	2 (6%)	1.0
• 2	39 (29%)	8 (22%)	0.54
• 3	67 (50%)	17 (47%)	0.91
TIMI flow grade post-PCI	n = 118	n = 32	
• 0	2 (2%)	1 (3%)	0.52
• 1	2 (2%)	0 (0%)	1.0

• 2	11 (9%)	3 (9%)	1.0
• 3	103 (87%)	28 (88%)	1.0

Data are n (%) or median (25th, 75th percentiles).

Definitions of end points

Reinfarction

- In the first 18 hours was defined as recurrent symptoms of ischaemia at rest accompanied by new ST-segment elevation of ≥0.1 mV in at least 2 contiguous leads, lasting ≥30 min.
- After 18 h, the definition was: new Q waves in 2 or more leads, or new increase in concentrations of creatine kinase-MB or troponins above the upper limit of normal (>3X upper limit of normal after PCI and >5X upper limit of normal after coronary artery bypass graft), and >50% higher than the previous value.

Stroke

• A new focal, neurological deficit of vascular origin lasting more than 24 h.

New myocardial ischaemia

• Unstable angina (chest pain at rest suspicious for coronary disease with or without ECG changes), recurrent angina grade II to IV (Canadian Cardiovascular Society classification) or serious arrhythmias (ventricular tachycardia/ventricular fibrillation) that appeared more than 12 hours after randomisation.

HRQoL

Health-related quality of life (HRQoL) was assessed using the 15D instrument. This is a generic, multidimensional, standardised, self-administered evaluative tool with 15 dimensions and 5 levels for each dimension (no problems to severe problems). The 15D scores were translated into a single index score with values from zero (dead) to 1.0 (perfect health) using a simple algorithm.

Bleeding

Classified according to the GUSTO severity scale

Effect Size

Clinical outcomes

	Early invasive group (n = 134)	Conservative group (n = 132)
Hospital stay (index admission)	5 days (IQR 4–6)	5 days (IQR 4–7)
At 30 days		
• Death	3 (2.2%)	3 (2.3%)
• Reinfarction	2 (1.5%)	7 (5.3%)

• Stroke	3 (2.2%)	5 (3.8%)
Recurrent ischaemia	8 (6.0%)	16 (12.1%)
 Severe bleeding, including intracranial haemorrhage (All severe bleedings were caused by intracranial haemorrhage) 	2	3
Moderate bleeding	0	3
Minor bleeding	14	13
At 12 months		
• Death	3 (2.2%)	4 (3.0%)
Reinfarction	4 (3.0%)	12 (9.1%)
• Stroke	3 (2.2%)	7 (5.3%)
Recurrent ischaemia	20 (15.0%)	20 (15.2%)
Hardah malatad marktur of life (AFD account)		

Health-related quality of life (15D scores)

р*

Values are mean±SD

Transfer

One patient (0.7%) died during transfer, and 4 (3%) were successfully defibrillated in the early invasive group. In the conservative group, 2 patients had ventricular tachycardia treated with intravenous drugs

^{†4} days before STEMI

^{*7} patients were unwilling to register the 15D questionnaire and were excluded from the analysis. The study did not report which group these patients were originally allocated to.

Table 106: AGATI et al¹

gotil et al Designa	patients	Patient characteristics	Intervention	Comparison	follow- up	Outcome measures	of funding
ngati L, et al. noes noronary ngioplasty fter timely hrombolysi improve nicrovascul r perfusion nd left entricular unction fter acute nyocardial nfarction? Im Heart J. 1007; 54(1):151- Sopen label. No details on whether outcome assessors were blinded ITT analysis: Yes RCT (Multicentre — 3 sites in Italy) Enrolment: Dec 2004 — Nov 2005 Allocation: No details Allocation concealment: No details Allocation concealment: No details ITT analysis: Yes	N = 96 in 3 arms. For fibrinolysis alone versus fibrinolysis within 24 hours followed by PCI, N = 60 Drop-outs: None stated	No details on whether high or low risk patients INCLUSION CRITERIA Patients with STEMI who presented within 3 hours of symptom onset EXCLUSION CRITERIA Patients who underwent revascularisation procedure as a result of failed fibrinolysis, early reinfarction, or ischaemia after the initial treatment (lysis or PCI) Patients aged ≥80 years History of previous MI, cardiomyopathy or CABG surgery Contraindications to glycoprotein IIb/IIIa receptor antagonists, PPCI, or lysis Patients in cardiogenic shock Demographics and baseline characteristics see below	Fibrinolysis alone (n = 30) Enoxaparin was repeated every 12 hours up to 7 days after lysis and 100 mg aspirin once a day was given indefinitely Note: All patients received a full dose of tenecteplase, aspirin and sc enoxaparin (1 mg/kg) at randomisation. All patients were maintained on ACE inhibitors and betablockers throughout hospital stay and at discharge.	Fibrinolysis followed within 24 hours by PCI (n = 30) Patients received abciximab immediately before the interventional procedure (0.25 mg/kg body weight) followed by a 12- hour infusion (0.125 microgram/kgminut es) and loading dose of clopidogrel 300 mg. After the procedure, 75 mg clopidogrel was prescribed for 6 months and 100 mg aspirin once a day indefinitely. Patients received IRA stenting using a standard procedure	In- hospital	Major bleeding Minor bleeding	No details

Mean age (y)	59±7	57±7
Male	25 (83)	26 (86)
Hypertension	26 (86)	24 (80)
Current smoker	17 (56)	19 (63)
Diabetes	6 (20)	5 (16)
Time to fibrinolysis (minutes)	118 (72–170)	129 (74–167)
PCI time after fibrinolysis	20±2 hours	-
Killip class >1	11 (36)	11 (36)
Multivessel disease	11 (36)	10 (33)
Anterior MI	24 (80)	22 (73)
TIMI 3 flow before PCI	20 (66)	-
TIMI 3 flow after PCI	26 (86)	-
Data presented are mean value ±SD or number (%)		
Effect Size		
	Fibrinolysis plus PCI (n = 30)	Fibrinolysis alone (n = 30)
Major bleeding	0	0
Minor bleeding (at access site)	4	0

Appendix H: Economic evidence tables

H.1 Time to reperfusion

Table 107: BRAVO VERGEL 2007¹⁵

Bravo Vergel Y, Palmer S analysis. Heart 93(10): 12	_	E, de Belder M, Abrams K et al.	Is primary angioplasty cost effo	ective in the UK? Results of a comprehensive decision
Study details	Population & interventions	Costs	Health outcomes	Cost effectiveness
Economic analysis: CUA (health outcome = QALYs)	Population: Acute STEMI patients	Total costs (mean per patient): Average (PPCI-related time delay 54 minutes)	QALYs (mean per patient): Average (PPCI-related time delay 54 minutes) Intervention 1: 6.83	ICER (Intervention 2 versus Intervention 1): Average (PPCI-related time delay 54 minutes) £9241 per QALY gained (pa) CI: NR
Study design: Probabilistic decision analytic model.	Cohort settings: Start age = 61 Male = NR	Intervention 1: £10,080 Intervention 2: £12,760 Incremental (2–1): £2680	Intervention 2: 7.12 Incremental (2–1): 0.29 (CI NR; p = NR)	Probability cost effective (£20K/30K threshold): 90%/95% Sensitivity analyses by PPCI-related time delay:
Approach to analysis: Short-term decision tree that captures differential outcomes up to 6 months (IHD death, new non-fatal MI; non-fatal stroke or no further event [alive with IHD]). Followed by Markov model to extrapolate to lifetime costs and QALYs. Markov model with 1 year cycles and 8 states (no further event [IHD] year 1, no further event [IHD]	Intervention 1: Fibrinolysis Intervention 2: PPCI with PPCI- related time delay of 54 minutes (Average delay from trials) Sensitivity analyses by time delay: • 30 minutes • 60 minutes • 90 minutes	(CI NR; p = NR) Sensitivity analyses by PPCI-related time delay: 30 minutes Intervention 1: £10,080 Intervention 2: £12,820 Incremental (2-1): £2740 (CI NR; p = NR) 60 minutes Intervention 1: £10,080 Intervention 2: £12,750 Incremental (2-1): £2670 (CI NR; p = NR) 90 minutes Intervention 1: £10,080	Sensitivity analyses by PPCI-related time delay: 30 minutes Intervention 1: 6.83 Intervention 2: 7.23 Incremental (2–1): 0.40 (CI NR; p = NR) 60 minutes Intervention 1: 6.83 Intervention 2: 7.09 Incremental (2–1): 0.26 (CI NR; p = NR) 90 minutes Intervention 1: 6.83 Intervention 1: 6.83 Intervention 1: 6.83	30 minutes £6850 per QALY gained (pa) CI: NR Probability cost effective (£20K/30K threshold): 98%/99% 60 minutes £10,269 per QALY gained (pa) CI: NR Probability cost effective (£20K/30K threshold): 83%/91% 90 minutes £64,750 per QALY gained (pa) CI: NR Probability cost effective (£20K/30K threshold): 36%/45%

year 2+, new non-fatal MI, post-MI, new non- fatal stroke, post- stroke, IHD death, other death. The	Intervention 2: £12,670 Incremental (2–1): £2590 (CI NR; p = NR)	Incremental (2–1): 0.04 (CI NR; p = NR)	Analysis of uncertainty: Probabilistic analysis results and time delay SA results are reported in main table. In a threshold analysis the time-delay up to which PPCI
possibility of needing further revascularisation was also modelled in short-	Currency & cost year: 2003/4 UK pounds Cost components		remained cost effective (at a £20K CE threshold) was 79 minutes (84.5 minutes at a £30K threshold). With a reduced length of stay with PPCI compared to fibrinolysis (assumed the same in base case) ICERs were reduced to £5448, £4087, £6038 and £37,250
Perspective: UK NHS and PSS Time horizon: lifetime	incorporated: Short-term model: initial interventions (drug acquisition costs, procedure costs and associated		respectively. The probability PPCI cost effective also increased. Only using fibrin-specific trials to estimate effectiveness reduced the cost effectiveness of PPCI. ICERs were £9833, £7284 and £11,500 for average, 30-minute and
Treatment effect duration: 6 months Discounting: Costs = 3.5%; Outcomes = 3.5%	hospital length of stay), further revascularisations, repeat MIs and stroke. Long- term model: management of non-fatal MI and stroke year1 and year 2+.		60-minute delays. And PPCI was dominated by fibrinolysis with a 90-minute delay. With both a reduced length of stay with PPCI and only using t-Pa trials ICERS were £5788, £4324 and £6739 for average, 30-minute and 60-minute delays. And PPCI was dominated by fibrinolysis with a 90-minute delay.

Data sources

Health outcomes: Initial 6-month decision tree: Baseline risks with fibrinolysis and relative risk with PPCI for clinical outcomes (non-fatal MI, non-fatal stroke, death) at 6 months were based on a systematic review and Bayesian meta-regression of 22 RCTs with the impact of PPCI-related time delay analysed using the mean additional time delay compared to fibrinolytic administration as a covariate in a random effects model – parallel clinical paper included in clinical review (Asseburg et al. 2007⁵). Post-6-month Markov model: Non-IHD death by 10 year age bands: Office of National Statistics data (does not appear to take account of gender split). Probabilities of transitioning between other health states based on Nottingham Heart Attack Registry (NHAR) data analysis. Quality-of-life weights: published EQ-5D estimates for health states, tariff not stated, who completed questionnaire not stated. Resource use and cost sources: Initial intervention: Assumed all PPCI patients required angiogram, GPIs and stent and all fibrinolysis patients did not require angiogram and received alteplase (most expensive agent). Initial length of hospital stay: assumed equal in base case based on average from Hospital Episode Statistics; in SA 5.76 days for PPCI and 12.12 days for fibrinolysis based on personal communication of estimates from a sample of 80 patients from a London hospital. Further revascularisation rates: based on analysis of same studies as health outcomes. Long-term costs associated with Markov model health states: MI resource use was based on analysis of NHAR hospitalisation data combined with national estimates of length of stay and published data to take account outpatient/day case care of costs outside the hospital setting. Stroke resource use was based on probability of being disabled, severity and discharge location from published literature. IHD death based on probability of dying in hospital from published literature. Unit costs: UK national sources (BNF, NHS reference costs) or published sources, inflate

was dominated by fibrinolysis with a 90-minute delay.

Comments

Source of funding: Study funded by unrestricted educational grant to University of York from Cordis Ltd (manufacturer of medical devices used in PCI). Some authors

also declared having received previous research funding or consultancy fees from various manufacturers of medical devices such as stents. **Limitations:** More recent estimates of initial resource use that better reflect current healthcare system are used in Wailoo 2010 update to analysis. Some uncertainly about measurement and valuation methods of health-related quality of life due to unclear reporting but considered minor limitation (EQ-5D used). Relative effectiveness of PPCI compared with fibrinolysis and the impact of time from study-level meta-regression based on systematic review of literature (Asseburg 2007⁵). This found no mortality benefit at 90 minutes with PPCI; however, an IPD analysis (Boersma 2006¹⁰) found benefit for 1-month mortality in patents 79–120 minutes and so there is uncertainty that relative treatment effects are from best available source. Three new RCTs that meet inclusion criteria have been published since Asseburg but considered likely to have small impact on effect estimates as low patient numbers relative to meta-analysis total. Ambulance costs not incorporated – may be higher with PPCI due to more transfers or longer journeys, although other PPCI cost assumptions generally conservative. **Other:** None.

Overall applicability*: minor limitations Overall quality**: partially applicable

Abbreviations: CI = confidence interval; CUA = cost—utility analysis; EQ-5D = EuroQol 5 dimension utility instrument, scale 0.0 (death) to 1.0 (full health), negative scores indicate a state worse than death; ICER = incremental cost-effectiveness ratio; IHD = ischaemic heart disease; MI = myocardial infarction; NR = not reported; pa = probabilistic analysis; QALYs = quality-adjusted life years

Table 108: WAILOO 2010, 119 GOODACRE 2008 50

Wailoo A, Goodacre S, Sampson F, Alava MH, Asseburg C, Palmer S et al. Primary angioplasty versus thrombolysis for acute ST-elevation myocardial infarction: An economic analysis of the National Infarct Angioplasty project. Heart 96(9): 668-672, 2010.

Goodacre S, Sampson F, Carter A, Wailoo A, O'Cathain A, Wood S et al. Evaluation of the National Infarct Angioplasty Project. Report for the National Co-ordinating Centre for NHS Service Delivery and Organisation (NCCSDO). National Coordinating Centre for the Service Delivery and Organisation (NCCSDO).

CLIA (I. III.	opulation:	-		
QALYs) Col Study design: Probabilistic decision analytic model incorporating cost analysis of UK observational data. Approach to analysis: Development of Bravo Vergel model: 1) updated with 'real-life' Col Ma	cute STEMI patients chort settings: tart age = 64.3 years fale = NR chervention 1: chrinolysis-based service Fibrinolysis: 72.6% (median CTN 67 min) PPCI: 15.8%; with PPCI- related time delay 52 minutes (median CTB 119 minutes)	Total costs (mean per patient): Average (time delay 64 minutes) Intervention 1: £10,700 Intervention 2: £11,600 Incremental (2-1): £829 (CI £130, £1440; p = NR) Sensitivity analyses by time delay: Transferred patients (delay 100 minutes) Intervention 1: £10,700 Intervention 2: £11,400 Incremental (2-1): £664 (CI -£324, £1390; p = NR) Non-transferred patients (delay	QALYs (mean per patient): Average (time delay 64 minutes) Intervention 1: 6.40 Intervention 2: 6.58 Incremental (2-1): 0.183 (CI -0.0764, 0.415; p = NR) Sensitivity analyses by time delay: Transferred patients (delay 100 minutes) Intervention 1: 6.40 Intervention 2: 6.32 Incremental (2-1): -0.0848	Primary ICER (Intervention 2 versus Intervention 1): Average (time delay 64 minutes) £4520 per QALY gained (pa) CI: NR Probability cost effective (£20K/30K threshold): 90%/95% Sensitivity analyses by time delay: Transferred patients (delay 100 minutes) Intervention 1 dominates CI: NR Probability cost effective (£20K/30K threshold): 38%/NR Non-transferred patients (delay 53 minutes)

^{*} Directly applicable / Partially applicable / Not applicable; ** Minor limitations / Potentially serious Limitations / Very serious limitations

updated with 'real-life' treatment NIAP delay estimates; 3) took system level perspective where some patients receive fibrinolysis in PPCI service, and vice versa; 4) adjusted baseline mortality rates with fibrinolysis to reflect the effectiveness of fibrinolysis as a function of the observed patient presentation delay in these patients.

Perspective: UK NHS and PSS Time horizon: lifetime Treatment effect

duration:6 months
Discounting: Costs =

3.5%; Outcomes =3.5%

Note that untreated patients were excluded from model.

Intervention 2:

PPCI-based service

- Fibrinolysis: 4.3% (median CTB 67 minutes)
- PPCI: 67.1%; with PPCIrelated time delay 64 minutes (median CTB 131 minutes) (Average delay from NIAP)

Sensitivity analyses by time delay:

- Transferred patients: delay 100 minutes (median CTB 167 minutes)
- Non-transferred patients: delay 53 minutes (median CTB 120)
- Direct to cath lab: delay 56 minutes (median CTB 123 minutes)
- Not direct to cath lab: delay 73 minutes (median CTB 140 minutes)

53 minutes)

Intervention 1: £10,700 Intervention 2: £11,600 Incremental (2–1): ~£900

(CI NR; p = NR)

<u>Direct to cath lab (delay 56</u> minutes)

Intervention 1: £10,700 Intervention 2: £11,600

Incremental (2-1): ~£900

(CI NR; p = NR)

Not direct to cath lab (delay 73 minutes)

Intervention 1: £10,700 Intervention 2: £11,500

Incremental (2–1): $^{£800}$

(CI NR; p = NR)

Currency & cost year:

2006/7 UK pounds

Cost components incorporated:

Short-term model: initial interventions (drug acquisition costs, procedure costs and associated hospital length of stay), further revascularisations, repeat MIs and stroke. Long-term model: management of non-fatal MI and stroke year 1 and year 2+.

Real-world initial costs did not account for potential additional ambulance time for PPCI centres operating a 'bypass' system.

(CI -0.831, 0.343; p = NR) Non-transferred patients

(delay 53 minutes)

Intervention 1: 6.40

Intervention 2: 6.64 Incremental (2–1):0.24

(CI NR; p = NR)

<u>Direct to cath lab (delay 56</u> minutes)

Intervention 1: 6.40

Intervention 2: 6.63

Incremental (2-1): 0.23

(CI 0.135, 0.597; p = NR)

Arrive via emergency department or CCU (delay

73 minutes)

Intervention 1: 6.40 Intervention 2: 6.53

Incremental (2-1): 0.13

(CI NR; p = NR)

£3635 per QALY gained (pa)

CI:NR

Probability cost effective (£20K/30K

threshold): 95%/NR

Direct to cath lab (delay 56 minutes)

£3817 per QALY gained (pa)

CI: NR

Probability cost effective (£20K/30K

threshold): 95%/NR

Arrive via emergency department or CCU

(delay 73 minutes)

£6112 per QALY gained (pa)

CI: NR

Probability cost effective (£20K/30K

threshold): 75%/NR

Analysis of uncertainty:

Probabilistic analysis results and time delay SA results are above.

Replacing the post-acute revascularisation rates from the evidence synthesis in the York model with those observed in NIAP increased the ICER to £7070 due to reduced benefit of PPCI in terms of repeat PCI. Probability cost effective (£20K threshold) was 85%.

Data sources

Health outcomes: The distribution of patient presentation delays was used to estimate baseline mortality for fibrinolysis-treated patients. Baseline event rates were modified by presentation time (call to arrival in hospital) and treatment times (arrival in hospital to reperfusion treatment) based on analysis of 10 NIAP sites implementing PPCI (n = 2083) and 5 other sites routinely providing fibrinolysis (n = 919). Relative effectiveness was not updated from Bravo-Vergel model (see evidence table). Quality-of-life weights: Not updated from Bravo Vergel model –published EQ-5D estimates for health states, tariff not stated, who completed questionnaire not stated. Resource use and cost sources: Initial intervention: Costs calculated using patient-level NIAP resource use and length of stay data for control site fibrinolysis (£3509), NIAP site fibrinolysis (£4361) and NIAP site PPCI (£5176). Control site PPCI had insufficient information and was assumed to be the same as in NIAP sites.

Resource use was based on analysis of 10 NIAP sites implementing PPCI (n = 2083) and 5 other sites routinely providing fibrinolysis (n = 919) supplemented by data from questionnaire (n = 50) where necessary. Further revascularisation rates: not updated from Bravo Vergel model in base case – based on analysis of same studies as health outcomes. Long-term costs associated with Markov model health states: not updated from Bravo Vergel model – MI resource use was based on analysis of NHAR hospitalisation data combined with national estimates of length of stay and published data to take account outpatient/day case care of costs outside the hospital setting. Stroke resource use was based on probability of being disabled, severity and discharge location from published literature. IHD death based on probability of dying in hospital from published literature. Unit costs: UK national sources (BNF, NHS reference costs, PSSRU) and NIAP site costs. Other: None.

Comments

Source of funding: National Co-ordinating Centre for NHS Service Delivery and Organisation R&D. **Limitations:** Some uncertainly about measurement and valuation methods of health-related quality of life due to unclear reporting but considered minor limitation (EQ-5D used). Relative effectiveness of PPCI compared with fibrinolysis and the impact of time from study-level meta-regression based on systematic review of literature (Asseburg 2007⁵). This found no mortality benefit at 90 minutes with PPCI; however, an IPD analysis (Boersma 2006¹⁰) found benefit for 1-month mortality in patients at 79–120 minutes and so there is uncertainty that relative treatment effects are from the best available source. Three new RCTs that meet inclusion criteria have been published since Asseburg but considered likely to have small impact on effect estimates as low patient numbers relative to meta-analysis total. Ambulance costs to first hospital not incorporated – may be higher with PPCI due to longer journeys. **Other:** 'Real-world' data collection (NIAP) took place 2005–06. Data analyses were adjusted for case-mix.

Overall applicability*: minor limitations **Overall quality**:** directly applicable

Abbreviations: CI = confidence interval; CUA = cost—utility analysis; EQ-5D = EuroQol 5 dimension utility instrument, scale 0.0 (death) to 1.0 (full health), negative scores indicate a state worse than death; ICER = incremental cost-effectiveness ratio; IHD = ischaemic heart disease; MI = myocardial infarction; NR = not reported; pa = probabilistic analysis; QALYs = quality-adjusted life years

†Total costs were reported in paper rounded to nearest hundred whereas incremental costs were reported to nearest whole pound therefore resulting in some discrepancies between total and incremental costs reported here; where incremental cost was not reported an approximate incremental cost is calculated from the rounded total costs. Total QALYs were reported in paper rounded to 2 decimal places and incremental QALYs rounded to 3 significant figures resulting in some discrepancies between total and incremental QALYs reported here; where incremental QALYs were not reported they were calculated from the totals and so are reported to 2 decimal places here. Cost effectiveness ratios are as reported in the paper.

* Directly applicable / Partially applicable / Not applicable; ** Minor limitations / Potentially serious Limitations / Very serious limitations

H.2 Facilitated PPCI

None.

H.3 Radial versus femoral arterial access for PPCI

None.

H.4 Thrombus extraction during PPCI

None.

H.5 Culprit versus complete revascularisation **Updated, see the 2020 evidence review**

Table 109: HELP-AMI³³

Di Mario C, Mara S, Flavio A, Imad S, Antonio M, Anna P et al. Single vs multivessel treatment during primary angioplasty: results of the multicentre randomised HEpacoat for cuLPrit or multivessel stenting for Acute Myocardial Infarction (HELP AMI) Study. International Journal of Cardiovascular Interventions 6(3-4): 128-133, 2004.

Economic analysis: CCA P	N - 4!			
(various health outcomes) P W Study design: Within-trial analysis (RCT: HELP-AMI study -	Population: Patients admitted to hospital with ischaemic chest pain or STEMI with arteriography showing lesions in multiple coronary arteries	Total costs (mean per patient): Intervention 1: £14,771 Intervention 2: £16,183 Incremental (2-1): £1,412 (CI = NR; p = 0.323)	12 month outcomes: From clinical review - same paper (2 versus 1) • Mortality: RR 0.98 (CI: 0.04, 23.03); ARD 19 fewer per 1000	Not applicable Analysis of uncertainty: No sensitivity analysis performed
health outcomes. Initial procedure costs: exact costing methodology unclear, probably individual-level costs, unit cost source unclear. Downstream costs: trial event rates with standard unit costs used. Ir Perspective: Italy health	Patient characteristics: n = 69 Mean age = 63.9 Male = 87.3 (see Tables 1–3 in chapter 9 for further details) Intervention 1: PCI using heparin-coated stents to recanalise all suitable lesions.	Cost breakdown: Initial procedure costs: Intervention 1: £9,659 Intervention 2: £9,141 Incremental (2-1): -£518 (CI = NR; p = 0.263) Downstream costs: Intervention 1: £5,112 Intervention 2: £7,042 Incremental (2-1): £1,930	 Reinfarction: RR 3.06 (CI: 0.20, 46.30); ARD 40 more per 1000 Repeat revascularisation: RR 2.04 (CI: 0.85, 4.90); ARD 180 more per 1000 	

Treatment effect duration: 12 months	Diabetes = 11.5%	(CI = NR; p = 0.185)	
Discounting: Costs = n/a; Outcomes = n/a	Intervention 2: PPCI using heparin-coated stents to recanalise culprit lesion only n = 17	Currency & cost year: 2004†† Euros (presented here as 2004 UK pounds‡)	
	Diabetes = 41.2%	Cost components incorporated: Initial procedure: all materials, stay in hospital including intensive care and cardiology wards. Downstream costs: additional revascularisation procedures (PCI or CABG)	

Data sources

Health outcomes: Within trial analysis. **Quality-of-life weights:** N/a. **Cost sources:** Time in hospital and materials used for initial procedures from within RCT patient-level analysis, source of unit costs not reported. Downstream event numbers for later revascularisation procedures from within RCT, costs based on Disease Related Group price (primary/complex angioplasty) for Lombardy region of Italy.

Comments

Source of funding: NR; 1 author was from Cordis Italia who manufacture the heparin-coated stents used in the trial. Limitations: Intervention used heparin-coated stents which are not routinely used in current practice. Contradictory definitions of study population - not certain whether all patients had STEMI. Study arms had unbalanced proportions of patients with diabetes. Some uncertainty about the applicability of unit costs and resource use from Italy pre-2004 (exact year not stated). Quality of life difference was not measured; QALYs not used. One-year time horizon may not fully capture differences in costs and health outcomes. Quality of life difference was not measured. Within-trial analysis, therefore by definition does not reflect all evidence available (see clinical review for comparison with other studies). Unclear if all relevant costs are included, and some unit cost sources are unclear. The costs of the initial procedures do not appear to account for the additional stents used in multivessel procedures – this would make the multivessel strategy appear cheaper and therefore more likely to be cost effective than it would otherwise have been. No sensitivity analysis undertaken. Funding not reported but 1 author worked for Cordis. Other: None.

Overall applicability*: Partially applicable **Overall quality**:** Very serious limitations

Abbreviations: ARD = absolute risk difference; CABG = coronary artery bypass graft; CCA = Cost-consequence analysis; CI = confidence interval; ICER = incremental cost-effectiveness ratio; MI = myocardial infarction; n/a = not applicable; NR = not reported; PCI = percutaneous coronary intervention; RCT = randomised controlled trial; RR = relative risk †Perspective not reported, but assumed to be Italy as 1 Italian cost is cited. Health service assumed as only direct medical costs included.

- *††Cost year not reported, assumed to be the same as publication year.*
- ‡ Converted using 2004 purchasing power parities 89
- * Directly applicable / Partially applicable / Not applicable; ** Minor limitations / Potentially serious limitations / Very serious limitations

H.6 Cardiogenic shock

None.

H.7 People who remain unconscious after a cardiac arrest

None.

H.8 Hospital volumes of PPCI

None.

H.9 Pre-hospital versus in-hospital fibrinolysis

None.

H.10 Use of antithrombin as an adjunct to fibrinolysis

None.

H.11 Rescue PCI

None.

H.12 Routine early angiography following fibrinolysis

Table 110: NORDISTEMI 2011¹²

Study details	Population & interventions	Costs§	Health outcomes	Cost effectiveness§
Economic analysis: CUA (health outcome = QALYs) Study design: within- trial analysis (RCT – NORDISTEMI) with	See clinical review for more details – NORDISTEMI ¹² Population: ST-elevation MI < 6 hour duration and > 90 minutes expected delay to PPCI, received full dose	Total costs (mean per patient): Intervention 1: £8888 Intervention 2: £9768 Incremental (2-1): £868 (CI -£777, £2545; p = NR) Total cost difference breakdown (2-1): • In-hospital care: £272 (CI -£1142, £1725)	Key outcome measure: QALYs – unadjusted / adjusted (mean per patient) Intervention 1: 0.885 / NR Intervention 2: 0.870 / NR Incremental (2–1): 0.016 (CI –0.023, 0.055; p = NR) / NR	ICER – total costs excluding unrelated (Intervention 2 versus Intervention 1): £62,648 per QALY gained (pa) CI: NR
probabilistic analysis Approach to analysis:	tenecteplase (57% pre- hospital)	 Out-patient care: £65 (CI -£379, £473) Transportation: £575 (CI £381, £769) 		at a threshold of £41,061 intervention 2 was cost effective in 49% of bootstrap iterations.§§
QALYs were calculated based on patient-level utility data recorded	Patient characteristics: • n = 259 (n = 266 in main	• Pharmaceuticals: -£27 (CI -£69, £14)		
over 12 months (see clinical review) and	study; 7 people who were unwilling to register the 15D were excluded from the	Total costs excluding those assumed to be unrelated (mean per patient) (used in base case ICER calculation):	0.008 (CI -0.027, 0.043; p = NR) / NR	Subgroup analyses: none
adjusted for baseline utility. Missing data was imputed. Bootstrapping was used to test for	 cost-effectiveness analysis) Mean age (total population) = 61 (SD: 10) Male (total population) = 	Intervention 1: £8756 Intervention 2: £9257 Incremental (2–1): £501 (CI NR; p = NR) Total cost difference breakdown (2–1):	Adjusted QALYs used to calculate cost effectiveness.	Analysis of uncertainty: When unrelated hospitalisation costs were included the ICER rose to
differences. Perspective: Norway	76% Intervention 1:	 As above except in-hospital care: -£182 (CI NR; p = NR) 		£108,463. When intra-cardiac defibrillator costs were excluded as well as unrelated hospitalisation costs the ICER reduced to £20,077.
healthcare system† Time horizon: 1 year Treatment effect	Deferred/selective angiography – admission to community hospital, with referral for urgent	Hospitalisations assumed to be unrelated were mostly orthopaedic, cancer-related and hernia surgery).		
duration: 1 year Discounting: Costs =	angiography (27%) if rescue indication or haemodynamic	Currency & cost year: 2008 Norwegian Kroner		

n/a; Outcomes = n/a

instability; early angiography was recommended if recurrent ischaemia occurred spontaneously or in a predischarge exercise test.

Otherwise angiography was recommended within 2 weeks of discharge. (86% had angiography with 30 days)

Intervention 2:

Routine early angiography – immediate transfer for angiography with PCI if indicated.

(presented here as 2008 UK pounds‡)

Cost components incorporated: In-hospital care (length of stay of index hospitalisation including the number of hours in the coronary care unit and days in the cardiology ward, number of subsequent hospital admissions, investigations or interventions including coronary angiography, PCI, CABG and implantation of intra-cardiac defibrillator), outpatient care (including GP visits, emergency department, clinic visits, home visits, household task assistance, hotel and rehabilitation) transportation (including ground ambulance, escorting nurse, helicopter ambulance, taxi, train, bus, private car) and pharmaceuticals. Sick leave was costed but is not presented in the results reported here.

Data sources

Health outcomes: within-RCT analysis. Quality-of-life weights: within-RCT analysis (measured at baseline (4 days before ST-elevation MI) and at 1, 3, 7 and 12 months) – 15D instrument, Finnish visual analogue scale valuation set. Study reports an imbalance in 15D at baseline (mean difference 0.011) and therefore adjusts for baseline score in QALY calculations. Cost sources: resource use collected prospectively in RCT from hospital databases, records and patients' self-reporting; for the index hospitalisation and later visits at the invasive centre direct costing using hospital accounts was used, while diagnosis-related group costs were used for other hospital admissions. Healthcare resources were costed using national fees, municipality and institutional accounts; transportation was costed according to type and distance.

Comments

Source of funding: Eastern Norway Regional Health Authority, A and H Waage's Veldedige stiftelse, Oslo Norway and Innlandet Hospital Trust, Brumunddal, Norway. Limitations: Some uncertainty about the applicability of Norway resource use and unit costs. Utility instrument used in QALY estimation does not meet NICE reference case (15D instrument with Finnish VAS-based valuation set). One-year time horizon may not fully capture differences in costs and health outcomes. Within-trial analysis therefore by definition does not reflect all evidence available (see clinical review tables for comparison with other studies) – judged to be one of the more relevant clinical trials, although patients were considered to be fairly low risk. Limited sensitivity analysis. Other: None.

Overall applicability*: partially applicable **Overall quality**:** potentially serious limitations

 $Abbreviations: CI = confidence\ interval;\ CUA = cost-utility\ analysis;\ ICER = incremental\ cost-effectiveness\ ratio;\ n/a = not\ applicable;\ NR = not\ reported;\ pa = probabilistic\ analysis;$

- † Study used societal perspective but results are presented disaggregated and so results have been recalculated to only include healthcare system costs in line with NICE reference case
- ‡ Converted from Euros to Norwegian Kroner using exchange rate reported in paper of €1.00 = NOK8.22 and from NOK to UK pounds using 2008 Purchasing Power Parities⁸⁹
- \S Results have been recalculated to exclude sick leave costs in line with the NICE reference case
- §§ It was not possible to recalculate the probability cost effective, and so this result includes sick leave costs. Since sick leave costs were higher for early than deferred strategy, if this had been excluded it is likely that the probability of the early strategy being cost effective would have been higher.
- * Directly applicable / Partially applicable / Not applicable; ** Minor limitations / Potentially serious Limitations / Very serious limitations