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## 7 **STABLE ANGINA**

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	<b>METHODS, EVIDENCE &amp; GUIDANCE</b>
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16 Produced by the National Clinical Guidelines Centre

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33 Appendices A to K are in separate volumes.

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4           David Wonderling

5

## Glossary and abbreviations

### GLOSSARY

Term	Description
Acute coronary syndrome	A condition in which there is an event in a coronary artery with plaque rupture or erosion, or coronary dissection, with the formation of intra-coronary thrombus. A single term which includes both unstable angina and myocardial infarction.
Acute myocardial infarction	<p>When there is evidence of myocardial necrosis in a clinical setting consistent with myocardial ischaemia, any one of the following criteria meets the diagnosis for myocardial infarction in people presenting with acute chest pain or discomfort:</p> <p>Detection of rise and/or fall of cardiac biomarkers (preferably troponin) with at least one value above the 99th percentile of the upper reference limit (URL) together with evidence of myocardial ischaemia with at least one of the following:</p> <ul style="list-style-type: none"> <li>• Symptoms of ischaemia</li> <li>• ECG changes indicative of new ischaemia (new ST-T changes or new left bundle branch block (LBBB))</li> <li>• Development of pathological Q waves in the ECG</li> <li>• Imaging evidence of new loss of viable myocardium or new regional wall motion abnormality.</li> </ul> <p>The guideline accepts the definition used in the studies included in the evidence review.</p>
Annual risk reduction	The difference between the percentage annual incidence of an adverse outcome in a treatment group compared with that in a control group.
Beta blockers (BBs)	A class of drugs that block beta-adrenergic substances such as adrenaline (epinephrine) in the "sympathetic"

	portion of the autonomic (involuntary) nervous system.
Biomarker	An objective measure of an indicator of a normal biologic process, a pathogenic process, or pharmacologic response to a therapeutic intervention.
Calcium channel blockers (CCBs)	Calcium channel blockers are medicines that slow the movement of calcium into the cells of the heart and blood vessels. This, in turn, relaxes blood vessels, increases the supply of oxygen-rich blood to the heart, and reduces the heart's workload.
Canadian Cardiovascular Society (CSS) Functional Classification of Angina	<p>Class I - Ordinary activity (e.g. walking, climbing stairs at own pace) does not bring on angina. Angina occurs only with strenuous, rapid, or prolonged exertion at work or during recreation.</p> <p>Class II - Slight limitation of ordinary activity. Symptoms occur when walking or climbing stairs rapidly, walking up a hill, walking up stairs after a meal, in cold weather, in wind, or when under emotional stress, or only a few hours after waking, and climbing more than one flight of ordinary stairs at a normal pace and in normal conditions.</p> <p>Class III - Marked limitation of ordinary activity. Symptoms occur after walking 50-100 yards on the level, or climbing more than one flight of ordinary stairs in normal conditions.</p> <p>Class IV - Inability to carry on any physical activity without discomfort. Angina may be present at rest.</p>
Cardiac syndrome X	Presence of exertional angina and angiographically normal epicardial arteries/coronary arteries.
Cardiovascular event	An acute coronary, cerebrovascular or peripheral arterial event.
Cardiovascular risk	The risk of a cardiovascular event occurring.
Clinical classification	A method of allocating patients into different groups based on clinical characteristics.
Clinical risk stratification	A method of allocating patients to different levels of risk of them suffering an adverse event, based on their clinical characteristics.
Coronary angiography	An invasive diagnostic test which provides anatomical information about the degree of stenosis (narrowing) in a coronary artery. It involves manipulation of cardiac catheters from an artery in the arm or top of the leg. A contrast medium is injected into the coronary arteries, and the flow of contrast in the artery is monitored by taking a rapid series of X-rays. It is considered the 'gold standard' for providing anatomical information

	and defining the site and severity of coronary artery lesions (narrowings).
Coronary artery	An artery which supplies the myocardium.
Coronary artery bypass surgery (CABG)	Open-heart surgery in which the rib cage is opened and a section of a blood vessel is grafted to the coronary artery to bypass the blocked section of the coronary artery and improve the blood supply to the heart
Coronary artery disease	Coronary artery disease is a condition in which atheromatous plaque builds up inside the coronary artery. This leads to narrowing of the arteries which may be sufficient to restrict blood flow and cause myocardial ischaemia.
Cost-consequences analysis	A type of economic evaluation where various health outcomes are reported in addition to the costs for each intervention under consideration. There is however no formal synthesis of the costs and health effects.
Cost-effectiveness acceptability curve (CEAC)	A CEAC plots the probability of an intervention being cost-effective compared with alternative intervention(s), for a range of maximum monetary values, that decision-makers might be willing to pay, for a particular unit change in outcome.
Cost-effectiveness analysis	An economic study design in which consequences of different interventions are measured using a single outcome, usually in 'natural' units (for example, life-years gained, deaths avoided, heart attacks avoided, cases detected). Alternative interventions are then compared in terms of incremental costs per unit of effectiveness.
Cost-utility analysis	A form of cost-effectiveness analysis in which the units of effectiveness are quality-adjusted life-years (QALYs).
Discounting	<p>Discounting is the process by which economists make allowances for society's time preference for costs and benefits. All else being equal, society places a higher value on the same unit of cost and benefit today than it does for the same unit in the future. For example, society prefers to receive £100 today as opposed to £100 in <math>n</math> years time. The differential is expressed in terms of the discount factor DF, where</p> $DF = 1 / (1 + r)^n$ <p>and where</p> <p><math>r</math> is the discount rate, and</p> <p><math>n</math> is the number of years forward from the current</p>

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	year.
Dominance	A health intervention is said to be dominant if it is both more effective and less costly than an alternative intervention.
Economic evaluation	Comparative analysis of alternative health strategies (interventions or programmes) in terms of both their costs and consequences.
Emergency	Immediate request leading to an immediate response from the ambulance service with a 'blue light' ambulance.
Equivocal	Where a diagnostic test result is indeterminate because it can be interpreted in one of 2 or more ways.
Evidence statements	A summary of the evidence distilled from a review of the available clinical literature.
Exercise ECG (sometimes known as an exercise test or stress ECG)	An investigation which measures the electrical activity from the heart during exercise, usually used to look for signs of myocardial ischaemia.
Health economic model	An explicit mathematical framework, which is used to represent clinical decision problems and incorporates evidence from a variety of sources in order to estimate costs and health outcomes.
Health economics	The branch of economics concerned with the allocation of society's scarce health resources, between alternative healthcare treatments/programmes, in an attempt to improve the health of the population.
Health related quality of life	An attempt to summarise an individual's or the population's quality of life resulting from the combined effect of their physical, mental, and social well-being.
Incremental cost-effectiveness ratio (ICER)	<p>The difference in the costs of two alternative treatment strategies/programmes, divided by the difference in the effectiveness outcomes of the treatment strategies/programmes for a defined population of interest. That is:</p> $\frac{\text{Cost treatment B} - \text{Cost treatment A}}{\text{Effectiveness treatment B} - \text{Effectiveness treatment A}}$
Intention-to-treat (ITT)	A strategy for analysing data from a randomised controlled trial. All participants are included in the arm to which they were allocated, whether or not they received (or completed) the intervention given to that arm.

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IPD meta analysis	IPD meta analysis involve the central collection, validation and re-analysis of “raw” data, from all clinical trials, world-wide, that have addressed a common research question; obtained from those responsible for the original trials.
Life years	The number of years lived by an individual or a population. For example, if a population of 50 patients live for an average addition 2 years each as the result of receiving a healthcare intervention, then the intervention has provided 100 life years gained.
Minimal important difference (MID)	The MIDs are the threshold for appreciable benefits and harms.
Myocardial infarction	See Acute Myocardial Infarction.
Myocardial perfusion scintigraphy with SPECT (MPS)	MPS involves injecting small amounts of radioactive tracer to evaluate perfusion of the myocardium via the coronary arteries at stress and at rest. The distribution of the radioactive tracer is imaged using a gamma camera. In SPECT the camera rotates round the patient and the raw data processed to obtain tomographic images of the myocardium. Cardiovascular stress may be induced by either pharmacological agents or exercise.
Opioid	An opioid is a chemical that works by binding to opioid receptors, and has pain killing properties. The term opiate is sometimes used as synonym, but this is natural opium alkaloids occurring in the resin of the opium poppy and the semi-synthetic opioids derived from them, and should be restricted to this.
Opportunity cost	The cost in terms of health benefits foregone by allocating resources to one intervention over an alternative intervention. The definition implicitly acknowledges the concept of scarcity of healthcare resources.
Other anti anginal drugs	Nicorandil, ivabradine and ranolazine are the other anti-anginal drugs that are licensed for use in the treatment of stable angina. They are distinguished in this way in the BNF from BBs, CCBs and nitrates.
Percutaneous coronary intervention (PCI).	The management of coronary artery occlusion by any of various catheter-based techniques, such as percutaneous transluminal coronary angioplasty, atherectomy, angioplasty using the excimer laser, and implantation of coronary stents and related devices
Probabilistic sensitivity analysis (PSA)	The process of measuring the degree of uncertainty around outcomes in an economic evaluation by assigning probability distributions to all of the key parameters in the evaluation, and then simultaneously generating values from each of these distributions

	using techniques of random number generation such as Monte Carlo methods.
Quality-adjusted life-year (QALY)	An index of survival weighted to account for quality of life. The year of life is weighted by a utility value U (where $0 \leq U \leq 1$ ). U reflects the health related quality of life, such that a U of zero represents the worst possible quality of life (equivalent to being dead), and a U of 1 represents perfect health. For example, 1 QALY is achieved if one patient lives in perfect health for one year, or alternatively if 2 people live in perfect health for 6 months each. Alternatively, a person living with a quality of life represented by a U value of 0.5 for 2 years is also representative of 1 QALY value. QALYs have the advantage of incorporating changes in both quantity (longevity/survival) and quality of life (morbidity as represented by psychological, physical and social functioning for example). QALYs are core to cost-utility analysis where the QALY is used as the measure of effectiveness in the economic evaluation.
Refractory angina	The European Cardiology Society definition of refractory angina is angina that cannot be controlled with optimal medical therapy and where revascularisation is unfeasible.
Rehabilitation	Cardiac rehabilitation is the process by which people with cardiac disease, in partnership with a multidisciplinary team of health professionals, are encouraged and supported to achieve and maintain optimal physical and psychosocial health.
Relative risk reduction	The ratio of the probability of an event occurring in the treatment group compared to the control group.
Sensitivity	<p>Sensitivity is the proportion of people with the disease who have a positive test. Sensitivity reflects how good the test is at identifying people with the disease. A measure of the diagnostic accuracy in including individuals with the condition.</p> <p>Number of True Positives divided by (Number of True Positives + Number of False Negatives)</p> <p>True positive: People correctly diagnosed with the condition</p> <p>False positive: Healthy people wrongly diagnosed with the condition</p> <p>True negative: Healthy people correctly identified as healthy</p> <p>False negative: People wrongly identified as healthy</p>

Sensitivity analysis	A means of exploring the uncertainty in the results of an economic evaluation/model by varying the parameter values of the included variables one at a time (univariate sensitivity analysis) or simultaneously (multi-variate sensitivity analysis).
Specialist	A healthcare professional that has expert knowledge of and skills in a particular clinical area, especially one who is certified by a higher medical educational organization.
Specificity	<p>Specificity is the proportion of people free of disease who have a negative test. Specificity reflects how good the test is at identifying people without the disease. A measure of the diagnostic accuracy in excluding individuals without the condition.</p> <p>Number of True Negatives divided by (Number of True Negatives + Number of False Positives)</p> <p>True positive: People correctly diagnosed with the condition</p> <p>False positive: Healthy people wrongly diagnosed with the condition</p> <p>True negative: Healthy people correctly identified as healthy</p> <p>False negative: People wrongly identified as healthy</p>
Stable angina	Angina is a symptom of myocardial ischaemia that is recognized clinically by its character, its location and its relation to provocative stimuli. Angina is stable when it is not a new symptom and when there is no deterioration in frequency, severity or duration of episodes.
Stress ECG	See exercise ECG above.
Stress echocardiograph	Echocardiography is an ultrasound examination of the heart. Exercise or pharmacological stress may be used to look for reversible systolic regional wall motion abnormalities consistent with the development of myocardial ischaemia.
Stress magnetic resonance imaging (stress MRI)	MRI is a diagnostic procedure that uses radio waves in a strong magnetic field. The pattern of electromagnetic energy released is detected and analysed by a computer to generate detailed images of the heart. Stress MRI is a specific application in which a contrast agent is used to detect myocardial blood flow at stress and at rest. Pharmacological stress is used to induce cardiovascular stress.

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Syndrome X	See cardiac syndrome X
Technology appraisal	Formal ascertainment and review of the evidence surrounding a health technology, which in this publication refers to technology appraisals undertaken by NICE only.
Technology appraisal guidance (TAG)	Technology Appraisal Guidance (see Technology Appraisal)
Unstable angina	New (within 24 hours) onset angina or abrupt deterioration in previously stable angina, often with prolonged episodes of rest pain.
Utility	A variable usually taking a value between zero (death) and unity (perfect health) which reflects health related quality of life, and which is used in the calculation of QALYs.
Willingness to pay (WTP)	The amount of money that an individual or society is willing to pay in order to achieve a specified level of health benefit. For example, it is generally recognised that the current willingness to pay for an incremental QALY gain in the NHS is somewhere between £20,000 and £30,000.

**Abbreviations**

<b>Abbreviation</b>	<b>Description</b>
2VD	two-vessel disease
3VD	three-vessel disease
AC	attenuation-corrected
ACE inhibitors	angiotensin-converting enzyme inhibitors
ACER	average cost-effectiveness ratio
AMI	acute myocardial infarction
ARB	angiotensin II receptor blocker
BB	beta blocker
BMJ	British Medical Journal
BNF	British National Formulary
CA	coronary angiography
CABG	coronary artery bypass graft
CAD	coronary artery disease
CAD	coronary artery disease
CCB	calcium channel blocker
CCS	Canadian Cardiovascular Society (CSS) Functional Classification of Angina
CFR	coronary flow reserve
CHD	coronary heart disease
CI	confidence interval
CRD	Centre for Reviews and Dissemination
CVD	cardiovascular disease
DTM	decision tree model
EBCT	electron beam computed tomography
EKG	Electrocardiography

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ECHO	Echocardiography
FN	false negative
FP	false positive
GDG	Guideline development group
GTN	glyceryl trinitrate
HR	hazard ratio
ICER	incremental cost-effectiveness ratio
ISMN	Isosorbide mononitrate
ITT	Intention-to-treat
LAD	left anterior descending
LBBS	left bundle branch block
LDL	low-density lipoprotein
LMS	left main stem
LR	likelihood ratio
MBF	myocardial blood flow
MD	Mean difference
MI	myocardial infarction
MID	Minimal Important difference
MPI	myocardial perfusion imaging
MPI	myocardial perfusion imaging
MPS	myocardial perfusion scintigraphy
MRI	magnetic resonance imaging
MVD	multi-vessel disease
NHS	National Health Service
NICE	National Institute for Health and Clinical Excellence
NIDDM	non-insulin dependent diabetes mellitus
NSF	National Service Framework

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OR	odds ratio
PCI	percutaneous coronary intervention
PCT	Primary care trust
PET	positron-emission tomography
PET	positron emission tomography
PTCA	percutaneous transluminal coronary angioplasty
QALY	quality-adjusted life-year
QoL	quality of life
QUADAS	quality assessment of diagnostic accuracy studies
RCT	randomised controlled trial
ROC	receiver operating characteristic
RR	relative risk
SA	sensitivity analysis
SD	Standard deviation
SPECT	single photon emission computed tomography
SRS	summed rest score
SVD	single-vessel disease
TN	true negative
TP	true positive

# 1 Introduction

2 Angina is constricting pain or discomfort that typically occurs in the chest (but may  
3 radiate to the neck, shoulders, jaw or arms) and is brought on by physical exertion or  
4 emotional stress. It is the main symptomatic manifestation of myocardial ischaemia  
5 and is usually caused by obstructive coronary artery disease restricting oxygen  
6 delivery to the cardiac myocytes. Other factors may exacerbate angina either by  
7 further restricting oxygen delivery (for example severe anaemia) or by increasing  
8 oxygen demand (for example left ventricular hypertrophy). Angina symptoms are  
9 associated with other cardiac disease such as aortic stenosis but the management of  
10 angina associated with non-coronary artery disease is outside the scope of this  
11 guideline.

12 *Epidemiology:* Unlike other manifestations of coronary artery disease, angina does not  
13 appear to be declining in incidence[2]. The Health Survey for England (2006)[3]  
14 found that about 8% of men and 3% of women aged between 55 and 64 years  
15 have, or have had angina. For people aged between 65 and 74 years the figures  
16 are about 14% of men and 8% of women. It is estimated that almost 2 million people  
17 in England have or have had angina. Prevalence is higher in men than in women, and  
18 increases sharply with age. Being diagnosed with angina can have a significant  
19 impact on a person's quality of life, which deteriorates progressively in proportion to  
20 the severity of symptoms[4].

21 *Current practice:* Stable angina is a chronic medical condition. The aim of management  
22 is to abolish or minimise symptoms, and to improve quality of life and long-term  
23 morbidity and mortality. Medical management includes pharmacological strategies or  
24 a combination of pharmacological and revascularisation strategies and lifestyle  
25 interventions. Revascularisation may be performed using percutaneous techniques or  
26 by surgery.

27 *Variation in practice.* Completed in 2003, the Euro Heart Survey on Stable Angina  
28 Pectoris included 3,779 ambulatory patients from 36 countries, presenting to a  
29 cardiologist as an outpatient, with new-onset stable angina[5]. The survey revealed  
30 considerable variation between participating countries in the use of non-invasive and  
31 invasive investigations, the prescription of anti-anginal drugs and rates of  
32 revascularisation. Guideline compliant therapy was associated with reduced rates of  
33 myocardial infarction and death.

34 *Current controversy.* The variation in practice documented within the Euro Heart Survey  
35 likely reflects continuing uncertainty about appropriate management strategies in key  
36 clinical areas where the evidence base is incomplete or contradictory. This applies

1 particularly to the role of revascularization for which symptomatic but not prognostic  
2 benefit has emerged as the predominant finding in contemporary clinical trials. This  
3 was highlighted by the COURAGE investigators who were unable to show prognostic  
4 benefit for revascularization in patients already receiving optimal medical treatment.  
5 The failure of revascularization to deliver prognostic benefit for people with angina  
6 has since been confirmed in two other landmark trials, BARI 2D and MASS II, and has  
7 stimulated considerable debate about the role of percutaneous and surgical  
8 management strategies in these patients. While some consensus has emerged around  
9 symptomatic indications, prognostic indications, if any, remain uncertain. Indeed, the  
10 only trials to report prognostic benefit for revascularization were randomized  
11 comparisons of bypass surgery and medical treatment that are now more than 25  
12 years old. It is noteworthy that these trials antedated introduction of statins and other  
13 secondary prevention treatments and the relevance of their findings to contemporary  
14 practice is doubtful.

15 Uncertainty about the effectiveness of revascularization for delivering prognostic  
16 benefit in people with coronary artery disease is heightened by some recent analyses  
17 that have reported excessive incremental cost-effectiveness ratios for percutaneous  
18 revascularization strategies compared with medical therapy. These areas of  
19 uncertainty surrounding the relative roles of medical therapy and revascularization in  
20 managing people with stable angina have received special attention from the  
21 guideline group in making its recommendations.

22 *Relationship between this guideline and NICE Clinical Guideline CG95 'Chest pain of*  
23 *recent onset'.*

24 NICE clinical guideline CG95 makes recommendations on the diagnosis of Stable  
25 Angina. That guideline covers the history, physical examination and investigations  
26 required to make a diagnosis of stable angina. This guideline presumes that a  
27 diagnosis of stable angina has already been made in accordance with NICE Clinical  
28 Guideline CG95 which recommends that angina can be diagnosed on the basis of  
29 history alone or on the basis of history and the results of functional or anatomical  
30 tests.

31 Typical angina is 3 out of 3 of the following: (a) constricting discomfort in anterior  
32 chest, neck, shoulder, jaw or arms; (b) precipitated by physical exertion or  
33 psychological stress and (c) relieved by rest or nitroglycerin within minutes. The  
34 requirement for functional or anatomical tests is dependent on the likelihood of  
35 coronary artery disease. That likelihood is dependent on how typical the history of  
36 angina is, the patient's age and gender and the presence of risk factors.

37

38

## 2 Development of the guideline

### 2.1 What is a guideline?

Our clinical guidelines are recommendations for the care of individuals in specific clinical conditions or circumstances within the NHS – from prevention and self-care through primary and secondary care to more specialised services. We base our clinical guidelines on the best available research evidence, with the aim of improving the quality of health care. We use predetermined and systematic methods to identify and evaluate the evidence relating to specific review questions.

Clinical guidelines can:

- provide recommendations for the treatment and care of people by health professionals
- be used to develop standards to assess the clinical practice of individual health professionals
- be used in the education and training of health professionals
- help patients to make informed decisions
- improve communication between patient and health professional

While guidelines assist the practice of healthcare professionals, they do not replace their knowledge and skills.

19

We produce our guidelines using the following steps:

- guideline topic is referred to NICE from the Department of Health
- stakeholders register an interest in the guideline and are consulted throughout the development process.
- the scope is prepared by the National Clinical Guidelines Centre (NCGC)
- the NCGC establishes a guideline development group
- a draft guideline is produced after the group assesses the available evidence and makes recommendations

- 1 • there is a consultation on the draft guideline
- 2 • the final guideline is produced

3

4 The NCGC and NICE produce a number of versions of this guideline:

- 5 • the **full guideline** contains all the recommendations, plus details of the  
6 methods used and the underpinning evidence
- 7 • the **NICE guideline** presents the recommendations from the full version in a  
8 format suited to implementation by health professionals and NHS bodies
- 9 • the **quick reference guide** presents recommendations in a suitable format for  
10 health professionals
- 11 • information for the public ('**understanding NICE guidance**') is written using  
12 suitable language for people without specialist medical knowledge.

13 This version is the full version. The other versions can be downloaded from NICE  
14 [www.NICE.org.uk](http://www.NICE.org.uk).

15

## 16 **2.2 Remit**

17 On 19 October 2007 the Department of Health formally requested the National  
18 Institute for Health and Clinical Excellence to prepare a clinical guideline as described  
19 in the box below (17<sup>th</sup> Wave Work Programme).

Remit: To prepare a clinical guideline on the management of stable angina.
--

20 NICE commissioned the National Collaborating Centre for Primary Care to develop  
21 this guideline. The National Collaborating Centre for Primary Care merged in 2009  
22 with the National Collaborating Centre for Chronic Condition, the National  
23 Collaborating Centre for Nursing and Supportive Care and the National  
24 Collaborating Centre for acute Care to form the National Clinical Guideline Centre  
25 (NCGC).

26

## 27 **2.3 Who developed this guideline?**

28 A multidisciplinary Guideline Development Group (GDG) comprising professional  
29 group members and consumer representatives of the main stakeholders developed  
30 this guideline (see section on Guideline Development Group Membership and  
31 acknowledgements).

1 The GDG was convened by the NCCPC/NCGC and chaired by Professor Adam  
2 Timmis in accordance with guidance from the National Institute for Health and Clinical  
3 Excellence (NICE).

4 The group met approximately every 6 weeks during the development of the  
5 guideline. At the start of the guideline development process all GDG members  
6 declared interests including consultancies, fee-paid work, share-holdings, fellowships  
7 and support from the healthcare industry. At all subsequent GDG meetings, members  
8 declared arising conflicts of interest, which were also recorded.

9 Members were either required to withdraw completely or for part of the discussion if  
10 their declared interest made it appropriate. The details of declared interests and the  
11 actions taken are shown in Appendix J

12 Staff from the NCGC provided methodological support and guidance for the  
13 development process. They undertook systematic searches, retrieval and appraisal of  
14 the evidence and drafted the guideline.

15

## 16 **2.4 What the guideline covers**

### 17 **2.4.1 Key clinical issues that are covered**

- 18 a) Non-invasive and invasive assessments to assess functional status, underlying  
19 disease, prognosis and plan management
- 20 b) Education programmes for people with angina (and carers and families as  
21 appropriate) that aim to help patients understand and manage their condition.  
22 They include self care, symptom management, medication management and  
23 lifestyle interventions
- 24 c) Psychological interventions for symptom relief and to improve long-term  
25 outcomes.
- 26 d) Pharmacological interventions for symptom relief and to improve long-term  
27 outcomes.
- 28 e) Revascularisation strategies for symptom relief and to improve long-term  
29 outcomes.
- 30 f) Specialised interventions for symptom relief, for example transcutaneous  
31 electrical nerve stimulation (TENS), temporary or destructive sympathectomy,  
32 and enhanced external counter pulsation (EECP).
- 33 g) Rehabilitation programmes.
- 34 h) Cardiac syndrome X

1     **2.4.2     Economic aspects**

2             Developers took into account both clinical and cost effectiveness when making  
3             recommendations involving a choice between alternative interventions. A  
4             review of the economic evidence was conducted and analyses were carried  
5             out as appropriate. The unit of effectiveness was the quality-adjusted life  
6             year (QALY), and the costs considered were from an NHS and personal social  
7             services (PSS) perspective. Further detail on the methods can be found in 'The  
8             guidelines manual' (see 'Further information').

9     **2.4.3     Groups that are covered**

- 10            a) Adults (18 years and older) who have been diagnosed with stable angina due  
11            to atherosclerotic disease
- 12            b) The following subgroups, were included:
- 13                    • people of south Asian origin
  - 14                    • people older than 65 years
  - 15                    • people with chronic refractory angina
  - 16                    • people with diabetes
  - 17                    • people with normal or minimally diseased coronary arteries
  - 18                    • women

19                    For further details please refer to the scope in Appendix [X].

20     **2.4.4     Healthcare settings that are covered**

- 21            a) All NHS primary, secondary and tertiary healthcare settings managing people  
22            with stable angina.

23

24     **2.5    What the guideline does not cover**

- 25            a) People with recent-onset chest pain or discomfort of suspected cardiac origin.
- 26            b) People with acute coronary syndrome.
- 27            c) People with chest pain or discomfort of unknown cause.
- 28            d) People with angina-type pain that is likely to be due to non-cardiac disease,  
29            such as anaemia.
- 30            e) People with angina-type pain associated with other types of heart disease,  
31            such as valvular heart disease (for example, aortic stenosis) or  
32            cardiomyopathy (for example, hypertrophic cardiomyopathy).

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## **2.6 Relationships between the guideline and other national guidance**

### **2.6.1 NICE guidance partly updated as a result of this clinical guideline**

- 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/TA73](http://www.nice.org.uk/TA73)

### **2.6.2 Other related NICE guidance**

- Chronic heart failure (partial update). NICE clinical guideline 108 (2010). Available from [www.nice.org.uk/guidance/CG108](http://www.nice.org.uk/guidance/CG108)
- Chest pain of recent onset. NICE clinical guideline 95 (2010). Available from [www.nice.org.uk/guidance/CG95](http://www.nice.org.uk/guidance/CG95)
- Unstable angina and NSTEMI. NICE clinical guideline 94 (2010). Available from [www.nice.org.uk/guidance/CG94](http://www.nice.org.uk/guidance/CG94)
- Endoscopic saphenous vein harvest for coronary artery bypass grafting. NICE interventional procedure guidance 348 (2010). Available from [www.nice.org.uk/guidance/IPG348](http://www.nice.org.uk/guidance/IPG348)
- Depression in chronic health problems. NICE clinical guideline 91 (2009). Available from [www.nice.org.uk/guidance/CG91](http://www.nice.org.uk/guidance/CG91)
- Medicines adherence. NICE clinical guideline 76 (2009). Available from [www.nice.org.uk/guidance/CG76](http://www.nice.org.uk/guidance/CG76)
- Percutaneous laser revascularisation for refractory angina pectoris. NICE interventional procedures guidance 302 (2009). Available from [www.nice.org.uk/guidance/IPG302](http://www.nice.org.uk/guidance/IPG302)
- Transmyocardial laser revascularisation for refractory angina pectoris. NICE interventional procedures guidance 301 (2009). Available from [www.nice.org.uk/guidance/IPG301](http://www.nice.org.uk/guidance/IPG301)
- Spinal cord stimulation for chronic pain of neuropathic or ischaemic origin. NICE technology appraisal guidance 159 (2008). Available from [www.nice.org.uk/guidance/TA159](http://www.nice.org.uk/guidance/TA159)
- Drug-eluting stents for the treatment of coronary artery disease (part review of NICE technology appraisal guidance 71). NICE technology appraisal guidance 152 (2008). Available from [www.nice.org.uk/guidance/TA152](http://www.nice.org.uk/guidance/TA152)

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- 1           • Lipid modification. NICE clinical guideline 67 (2008). Available from  
2           [www.nice.org.uk/guidance/CG67](http://www.nice.org.uk/guidance/CG67)
- 3           • Smoking cessation services (2008). NICE public health guidance 10. Available  
4           from [www.nice.org.uk/guidance/PH10](http://www.nice.org.uk/guidance/PH10)
- 5           • Ezetimibe for the treatment of primary (heterozygous-familial and non-  
6           familial) hypercholesterolaemia. NICE technology appraisal guidance 132  
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- 8           • Myocardial infarction: secondary prevention. NICE clinical guideline 48  
9           (2007). Available from [www.nice.org.uk/guidance/CG48](http://www.nice.org.uk/guidance/CG48)
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11          (2007). Available from [www.nice.org.uk/guidance/TA123](http://www.nice.org.uk/guidance/TA123)
- 12          • Hypertension. NICE clinical guideline 34 (2006). Available from  
13          [www.nice.org.uk/guidance/CG34](http://www.nice.org.uk/guidance/CG34)
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15          guidance 94 (2006). Available from [www.nice.org.uk/guidance/TA94](http://www.nice.org.uk/guidance/TA94)
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18          [www.nice.org.uk/guidance/IPG98](http://www.nice.org.uk/guidance/IPG98)
- 19          • Off-pump coronary artery bypass grafting. NICE interventional procedure  
20          guidance 35 (2004). Available from [www.nice.org.uk/guidance/IPG35](http://www.nice.org.uk/guidance/IPG35)  
21          (currently being updated with an expected publication in January 2011)
- 22          • Guidance on the use of coronary artery stents. NICE technology appraisal  
23          guidance 71 (2003). Available from [www.nice.org.uk/guidance/TA71](http://www.nice.org.uk/guidance/TA71)
- 24

## 1 **3 Methods**

2 This guidance was developed in accordance with the methods outlined in the NICE  
3 Guidelines Manual[6].

4

### 5 **3.1 Developing the review questions and outcomes**

6 Review questions were developed based on the scope (Appendix A). They were  
7 drafted by the review team and refined and validated by the GDG. Review  
8 questions were developed in a PICO framework (patient, intervention, comparison  
9 and outcome) for intervention reviews, risk scores and prognostic reviews. This was to  
10 guide the literature searching process and to facilitate the development of  
11 recommendations by the GDG.

12

### 13 **3.2 Searching for evidence**

#### 14 **3.2.1 Clinical literature search**

15 Systematic literature searches were undertaken to identify evidence within published  
16 literature in order to answer the review questions as per The NICE Guidelines  
17 Manual[6]. Clinical databases were searched using relevant medical subject headings,  
18 free-text terms and study type filters where appropriate. Non-English studies were  
19 not reviewed and were therefore excluded from searches. All searches were  
20 conducted on core databases, Medline, Embase, Cinahl and The Cochrane Library.  
21 Additional subject specific databases were used for some questions. All searches were  
22 updated on the 22nd of October 2010. No papers after this date were considered.

23 Search strategies were checked by looking at reference lists of relevant key papers,  
24 checking search strategies in other systematic reviews and asking the GDG for known  
25 studies. The questions, the study types applied, the databases searched and the years  
26 covered can be found in Appendix D.

27 During the scoping stage, a search was conducted for guidelines and reports on the  
28 websites listed below and on organisations relevant to the topic. Searching for grey  
29 literature or unpublished literature was not systematically performed. All references  
30 sent by stakeholders were considered.

- 1 • Constituent websites of the Guidelines International Network ([www.g-i-n.net](http://www.g-i-n.net))
- 2 • National Guideline Clearing House ([www.guideline.gov/](http://www.guideline.gov/))
- 3 • National Institute for Health and Clinical Excellence (NICE) ([www.nice.org.uk](http://www.nice.org.uk))
- 4 • National Institutes of Health Consensus Development Program
- 5 ([consensus.nih.gov/](http://consensus.nih.gov/))
- 6 • National Library for Health ([www.library.nhs.uk/](http://www.library.nhs.uk/))

7

### 8 **3.2.2 Health economic literature search**

9 Systematic literature searches were also undertaken to identify health economic  
10 evidence within published literature relevant to the review questions. The evidence  
11 was identified by conducting a broad search relating to the stable angina population  
12 in the NHS economic evaluation database (NHS EED), the Health Economic Evaluations  
13 Database (HEED) and health technology assessment (HTA) databases with no date  
14 restrictions up to 13/9/10. Additionally, the search was run on Medline (years 1950 -  
15 2007) and Embase (1996-2007), with a specific economic filter, to ensure recent  
16 publications that had not yet been indexed by these databases were identified. This  
17 was supplemented by additional searches from (1990-13/9/10) that looked for  
18 economic papers specifically relating to revascularisation, rehabilitation, nicorandil,  
19 long acting nitrates on Medline, Embase, Cochrane (TA's and EE's, as it became  
20 apparent that some papers in this area were not being identified through the first  
21 search.

22 The search strategies for health economics are included in Appendix D. All searches  
23 were updated on the 13<sup>th</sup> Sept 2010. No papers after this date were considered.

24

### 25 **3.3 Reviewing the evidence**

26 The Research Fellow and Health Economist:

- 27 • Identified potentially relevant studies for each review question from the relevant  
28 search results by reviewing titles and abstracts – full papers were then obtained.
- 29 • Reviewed full papers against pre-specified inclusion / exclusion criteria to identify  
30 studies that addressed the review question in the appropriate population and  
31 reported on outcomes of interest (research protocols are included in Appendix C)
- 32 • Critically appraised relevant studies using the appropriate checklist as specified in  
33 The Guidelines Manual[6].
- 34 • Extracted key information about the study's methods and results into evidence tables  
35 (evidence tables are included in Appendix E2)

- 1 • Generated summaries of the evidence by outcome (included in the relevant chapter  
2 write-ups):
  - 3 ○ Randomised studies: meta analysed, where appropriate and reported in  
4 GRADE profiles (for clinical studies) – see below for details
  - 5 ○ Observational studies: each study summarised in a table and narrative  
6 developed
  - 7 ○ Qualitative studies: each study summarised in a table and narrative  
8 developed
  - 9 ○ Economic studies: summarised in NICE economic evidence profiles – see  
10 below for details

### 12 3.3.1 Inclusion/exclusion

13 See the review protocols in Appendix C for full details.

#### 14 **Population**

15 The remit of the guideline was to make recommendations for people with stable  
16 angina. Studies were required to have at least 60% of people with stable angina to  
17 be included. The interventions (e.g. drugs and revascularisation procedures) used in  
18 stable angina are also used commonly in people who are found to have coronary  
19 artery disease or who present with other coronary artery diseases such as unstable  
20 angina or MI. Hence many of the trials for these interventions include a mixed group  
21 of patients including stable angina, unstable angina and/or MI. For this reason the  
22 GDG decided to consider studies with at least 60% stable angina population as this  
23 would be more relevant to the population specified in this guideline.

24 In this guideline we have also looked separately at people with cardiac syndrome X.

#### 25 **Intervention**

26 The following classes of drugs have been considered in this guideline:

- 27 • Short acting nitrates
- 28 • BBs
- 29 • CCBs
- 30 • Long acting nitrates
- 31 • Nicorandil
- 32 • Ivabradine

- 1       • Ranolazine
- 2       • ACE inhibitors
- 3       • ARBs
- 4       • Aspirin
- 5       • Statins

6  
7

The following prognostic tests have been considered in this guideline:

- 8       • Exercise ECG / exercise tolerance test / exercise stress test / stress ECG.
- 9       • Stress echocardiography/exercise, dobutamine, dipyridamole, adenosine- stress  
10       echocardiography.
- 11       • Stress myocardial perfusion imaging/ MPS/ myocardial perfusion scintigraphy /  
12       exercise thallium MPS/ MPS using single photon emission CT (SPECT).
- 13       • Stress magnetic resonance imaging / stress CMR / adenosine, dipyridamole -stress  
14       perfusion imaging / dobutamine -stress induced motion wall abnormalities.
- 15       • Computed tomography CT / CT coronary angiography / multi slice CT,  
16       multidetector CT / CT coronary angiography / CAT
- 17       • Ca scoring , coronary calcium scoring
- 18       • Electron beam CT (EBCT).
- 19       • Coronary Angiography

20  
21

The following revascularisation procedures have been considered in this guideline:

- 22       • Percutaneous coronary intervention (PCI) (includes coronary balloon angioplasty  
23       and coronary stent implantation),
- 24       • Coronary artery bypass surgery (CABG)

25  
26

The details of the interventions can be found in the relevant review sections.

## 27 **Outcomes**

28  
29

The following outcomes are reported in this guideline

*Outcomes in intervention studies*

- 1 • Exercise tolerance
- 2 • Nitroglycerin consumption
- 3 • Angina frequency/severity
- 4 • MI/Non fatal MI
- 5 • Revascularisation
- 6 • Hospitalisation
- 7 • Stroke/cerebrovascular accident
- 8 • Death
- 9 • Cardiac/cardiovascular death
- 10 • Quality of Life
- 11 • Adverse events

12

13 *Outcomes in Prognostic studies*

14 The main outcomes considered in prognostic studies were:

- 15 • Death
- 16 • Cardiac death/cardiovascular death
- 17 • MI/Nonfatal MI
- 18 • Revascularisation

19

20 **3.3.2 Health economic inclusion/exclusion criteria**

21 Full economic evaluations (cost-effectiveness, cost-utility, cost-benefit and cost-  
22 consequence analyses) and comparative costing studies that addressed the review  
23 question in the relevant population were considered to have the potential for inclusion  
24 as economic evidence.

25 Studies that only reported cost per hospital (not per patient), or only reported  
26 average cost effectiveness without disaggregated costs and effects, were excluded.  
27 Abstracts, posters, reviews, letters/editorials, foreign language publications and  
28 unpublished studies were excluded. Studies judged to have an applicability rating of  
29 'not applicable' were excluded (this included studies that took the perspective of a  
30 non-OECD country).

1 Remaining studies were prioritised for inclusion based on their relative applicability to  
2 the development of this guideline and the study limitations. For example, if a high  
3 quality, directly applicable UK analysis was available, other less relevant studies  
4 were not included.

5 For more details about the assessment of applicability and methodological quality  
6 see the economic evaluation checklist (The Guidelines Manual[6], Appendix H and the  
7 health economics research protocol in Appendix C .

8 When no relevant economic analysis was found from the economic literature review,  
9 relevant UK NHS unit costs related to the compared interventions were presented to  
10 the GDG to inform the possible economic implication of the recommendation to make.

### 11 **Quality assessment for inclusion of studies**

12 All studies are quality assessed before being included as part of the systematic  
13 review. The criteria for assessment for different types of studies are listed below.

14 For systematic reviews and meta-analysis, the main criteria considered were:

- 15 • An appropriate and clearly focused question was addressed
- 16 • Methodology was well described
- 17 • The literature search was sufficiently robust to identify all the relevant studies
- 18 • The individual study quality included in the review was assessed and taken into  
19 account
- 20 • The studies were sufficiently similar to make combining them reasonable

21

### 22 ***Intervention studies***

23 The quality assessment criteria as listed in the NICE Guidelines Manual 2009 were  
24 used to assess systematic reviews, meta-analysis, and randomised controlled trials.

25 For randomised controlled trials, the main criteria considered were:

- 26 • An appropriate and clearly focused question was addressed
- 27 • Appropriate randomisation allocation and concealment methods were used
- 28 • Subjects, investigators and outcomes assessors were masked about treatment  
29 allocation
- 30 • The intervention and control groups are similar at baseline
- 31 • The only difference between group is the type of intervention received

- 1 • All outcomes are measured in a standard and reliable method
- 2 • Drop out rates reported and are acceptable, and all participants are analysed in
- 3 the groups to which they were randomly allocated the treatment
- 4 • For multi-centred trials, results are comparable between sites
- 5 Only studies which fulfilled some to all of the criteria included were included in the
- 6 evidence review.

7

### 8 **Prognostic studies**

9 Prospective cohort studies were included for the prognostic questions. The prospective  
10 cohort studies' quality was assessed using the quality checklist in the NICE Guidelines  
11 Manual April 2009. The main criteria considered in assessing study quality were:

- 12 • An appropriate and clearly focused question was addressed
- 13 • The cohort(s) being studied are selected from source populations that are
- 14 comparable in all respects other than the factor under investigation
- 15 • The inclusion or participation rate was reported
- 16 • The likelihood that some eligible subjects might have the outcome at the time of
- 17 enrolment assessed had been taken into account in the analysis
- 18 • The drop out rate was reported and acceptable
- 19 • Comparison by the prognostic status is made between participants who completed
- 20 the study and those lost to follow up
- 21 • The outcomes were clearly defined
- 22 • The assessment of outcome was blind to exposure status or acknowledged where
- 23 this was not possible
- 24 • The methods of assessment used for the prognostic factor and the outcomes were
- 25 valid and reliable
- 26 • The main potential confounders are identified and taken into account adequately
- 27 in the design and analysis
- 28 • Confidence intervals or standard deviation were provided

29

### 30 **3.3.3 Methods of combining clinical studies**

31 *Data synthesis for intervention reviews*

1 Where possible, meta-analyses were conducted to combine the results of studies for  
 2 each review question using Cochrane Review Manager (RevMan5) software. Fixed-  
 3 effects (Mantel-Haenszel) techniques were used to calculate risk ratios (relative risk)  
 4 for the binary outcomes: [death, cardiac death, MI/non fatal MI, revascularisation,  
 5 stroke, number patients free of angina, adverse events]. The continuous outcome(s)  
 6 [exercise tolerance, angina frequency, nitroglycerin consumption] was(were)  
 7 analysed using an inverse variance method for pooling weighted mean differences  
 8 and where the studies had different scales, standardised mean differences were used.  
 9 Statistical heterogeneity was assessed by considering the chi-squared test for  
 10 significance at  $p < 0.05$  or an I-squared inconsistency statistic of  $> 50\%$  to indicate  
 11 significant heterogeneity. When there were a high number of studies, a p-value of  
 12 0.1 was taken as a threshold for heterogeneity. We carried out predefined subgroup  
 13 analyses as defined in the protocol for each question (see Appendix D).

14 The standard deviations of continuous outcomes were required for imputation for  
 15 meta-analysis. However, in cases where this was not reported, calculation based on  
 16 methods outlined in section 7.7.3 of the Cochrane Handbook[7]: 'Data extraction for  
 17 continuous outcomes' were applied to estimate the standard deviations if p values of  
 18 the difference between two means, 95% confidence intervals or standard error of the  
 19 mean (SEM) had been reported'. Where p values were reported as "less than", a  
 20 conservative approach was undertaken. For example, if p value was reported as " $p$   
 21  $\leq 0.001$ ", the calculations for standard deviations will be based on a p value of  
 22 0.001. If these statistical measures were not available then the methods described in  
 23 section 16.1.3 of the Cochrane Handbook (February 2008) 'Missing standard  
 24 deviations' were applied as the last resort.

25 For binary outcomes, absolute event rates were also calculated using the GRADEpro  
 26 software using event rate in the control arm of the pooled results.

27 In the evidence reviews in this guideline we have presented additional data from  
 28 studies along with the GRADE tables. These have been referred to as 'Additional  
 29 data' and refer to data which was not analysed due to lack of sufficient reported  
 30 information and/or outcomes.

31 *Data synthesis for prognostic review*

32 Odds ratio, relative or hazard risks, with their 95% confidence intervals, from  
 33 multivariate analyses were extracted from the papers. Studies were not combined in  
 34 a meta-analysis for observational studies.

35

### 36 **3.4 GRADE (Grading of Recommendations Assessment, Development and** 37 **Evaluation)**

38 The evidence for outcomes from studies which passed the quality assessment were  
 39 evaluated and presented using an adaptation of the 'Grading of Recommendations  
 40 Assessment, Development and Evaluation (GRADE) toolbox' developed by the  
 41 international GRADE working group (<http://www.gradeworkinggroup.org/>). The  
 42 software (GRADEpro) developed by the GRADE working group was used to assess

1 pooled outcome data using individual study quality assessments and results from  
2 meta-analysis.

3 The summary of findings was presented as two separate tables in this guideline. The  
4 “Clinical Study Characteristics” table includes details of the quality assessment while  
5 the “Clinical Summary of Findings” table includes pooled outcome data, where  
6 appropriate, an absolute measure of intervention effect calculated and the summary  
7 of quality of evidence for that outcome. In this table, the columns for intervention and  
8 control indicate pooled sample size for continuous outcomes. For binary outcomes such  
9 as number of patients with an adverse event, the event rates (n/N) are shown with  
10 percentages. Reporting or publication bias was only taken into consideration in the  
11 quality assessment and included in the Clinical Study Characteristics table if it was  
12 apparent.

13 Each outcome was examined separately for the quality elements listed and defined in  
14 Table 3.1 and each graded using the quality levels listed in Table 3.2. The main  
15 criteria considered in the rating of these elements are discussed in the literature  
16 reviewing process (see section 3.4.1 Grading of Evidence). Footnotes were used to  
17 describe reasons for grading a quality element as having serious or very serious  
18 problems. Then, an overall quality of evidence for each outcome was applied by  
19 selecting from the options listed in Table 3.3. The GRADE toolbox is currently  
20 designed only for randomised controlled trials and observational studies but we  
21 adapted the quality assessment elements and outcome presentation for diagnostic  
22 accuracy studies.

23 **Table 3.1: Descriptions of quality elements in GRADE for intervention studies**

<b>Quality element</b>	<b>Description</b>
<b>Limitations</b>	Limitations in the study design and implementation may bias the estimates of the treatment effect. Major limitations in studies decrease the confidence in the estimate of the effect.
<b>Inconsistency</b>	Inconsistency refers to an unexplained heterogeneity of results.
<b>Indirectness</b>	Indirectness refers to differences in study population, intervention, comparator and outcomes between the available evidence and the review question, or recommendation made.
<b>Imprecision</b>	Results are imprecise when studies include relatively few patients and few events and thus have wide confidence intervals around the estimate of the effect relative to the minimal important difference.
<b>Publication bias</b>	Publication bias is a systematic underestimate or an overestimate of the underlying beneficial or harmful effect due to the selective publication of studies.

24  
25 **Table 3.2: Levels for quality elements in GRADE**

<b>Level</b>	<b>Description</b>
<b>None</b>	There are no serious issues with the evidence
<b>Serious</b>	The issues are serious enough to downgrade the outcome evidence by one level
<b>Very serious</b>	The issues are serious enough to downgrade the outcome evidence by two levels

1 **Table 3.3: Overall quality of outcome evidence in GRADE**

<b>Level</b>	<b>Description</b>
<b>High</b>	Further research is <i>very unlikely</i> to change our confidence in the <i>estimate of effect</i>
<b>Moderate</b>	Further research is <i>likely</i> to have an important impact on our confidence in the <i>estimate of effect</i> and may change the estimate
<b>Low</b>	Further research is <i>very likely</i> to have an important impact on our confidence in the estimate of effect and is likely to change the estimate
<b>Very low</b>	<i>Any estimate of effect is very uncertain</i>

2

3 **3.4.1 Grading the quality of clinical evidence**

4 After results were pooled, the overall quality of evidence for each outcome was  
5 considered. The following procedure was adopted when using GRADE:

- 6 1. A quality rating was assigned, based on the study design. RCTs start HIGH and  
7 observations studies as LOW.
- 8 2. The rating was then downgraded for the specified criteria: Study limitations,  
9 inconsistency, indirectness, imprecision and reporting bias. These criteria are  
10 detailed below. Observation studies were upgraded if there was: a large  
11 magnitude of effect, dose-response gradient, and if all plausible confounding  
12 would reduce a demonstrated effect or suggest a spurious effect when results  
13 showed no effect. Each quality element considered as having “serious” or “very  
14 serious” risk of bias was rated down 1 or 2 points respectively.
- 15 3. The downgraded/upgraded marks were then summed and the overall quality  
16 rating was revised. For example, all RCTs started as HIGH and the overall  
17 quality became MODERATE, LOW or VERY LOW if 1, 2 or 3 points were  
18 deducted respectively.
- 19 4. The reasons or criteria used for downgrading were specified in the footnotes.

20 The details of criteria used for each of the main quality element are discussed further  
21 in the following sections x.

22

23 **3.4.2 Study limitations**

24 The main limitations for randomised controlled trials are listed in Table 3.4.

25

1 **Table 3.4: Study limitations of randomised controlled trials**

<i>Limitation</i>	<i>Explanation</i>
<b>Allocation concealment</b>	Those enrolling patients are aware of the group to which the next enrolled patient will be allocated (major problem in “pseudo” or “quasi” randomised trials with allocation by day of week, birth date, chart number etc.).
<b>Lack of blinding</b>	Patient, caregivers, those recording outcomes, those adjudicating outcomes, or data analysts are aware of the arm to which patients are allocated
<b>Incomplete accounting of patients and outcome events</b>	Loss to follow-up not accounted and failure to adhere to the intention to treat principle when indicated
<b>Selective outcome reporting</b>	Reporting of some outcomes and not others on the basis of the results
<b>Other limitations</b>	For example: <ul style="list-style-type: none"> <li>• stopping early for benefit observed in randomised trials, in particular in the absence of adequate stopping rules</li> <li>• use of unvalidated patient-reported outcomes</li> <li>• carry-over effects in cross-over trials</li> <li>• recruitment bias in cluster-randomised trials</li> </ul>

2

3 **3.4.3 Inconsistency**

4 Inconsistency refers to an unexplained heterogeneity of results. When estimates of the  
5 treatment effect across studies differ widely (i.e. heterogeneity or variability in  
6 results), this suggests true differences in underlying treatment effect. When  
7 heterogeneity exists (Chi square  $p < 0.05$  [ $p < 0.1$  for high number of studies] or I-  
8 squared inconsistency statistic of  $> 50\%$ ), but no plausible explanation can be found,  
9 the quality of evidence was downgraded by one or two levels, depending on the  
10 extent of uncertainty to the results contributed by the inconsistency in the results. On  
11 top of the I- square and Chi square values, the decision for downgrading was also  
12 dependent on factors such as whether the intervention is associated with benefit in all  
13 other outcomes or whether the uncertainty about the magnitude of benefit (or harm)  
14 of the outcome showing heterogeneity would influence the overall judgment about net  
15 benefit or harm (across all outcomes).

16 If inconsistency could be explained based on subgroup analysis, the GDG took this  
17 into account and considered whether to make separate recommendations based on  
18 the identified explanatory factors, i.e. population and intervention. In this situation, the  
19 quality of evidence would not be downgraded.

20 **3.4.4 Indirectness**

21 Directness refers to the extent to which the populations, intervention, comparisons and  
22 outcome measures are similar to those defined in the inclusion criteria for the reviews.  
23 Indirectness is important when these differences are expected to contribute to a  
24 difference in effect size, or may affect the balance of harms and benefits considered  
25 for an intervention.

1 **3.4.5 Imprecision**

2 The sample size, event rates and the resulting width of confidence intervals were the  
3 main criteria considered. Where the minimal important difference (MID) of an  
4 outcome is known, the optimal information size (OIS), i.e. the sample size required to  
5 detect the difference with 80% power and  $p \leq 0.05$  was calculated and used as the  
6 criteria. The criteria applied for imprecision are based on the confidence intervals for  
7 pooled or the best estimate of effect as illustrated in Figure 3.1 and outlined in Table  
8 3.5.

9 **Table 3.5: Criteria applied to determine precision - Criteria for downgrading an outcome for**  
10 **imprecision**

***Dichotomous and continuous outcomes***

1. 95% confidence interval (or alternative estimate of precision) around the pooled or best estimate of effect:

a) does not cross the threshold for appreciable benefit or harm defined as precise

Rating for precision: 'no serious imprecision'

2. 95% confidence interval (or alternative estimate of precision) around the pooled or best estimate of effect:

a) If the 95% confidence interval crosses either minimal important difference (MID) threshold, defined as imprecise

Rating for precision: 'serious'

3. 95% confidence interval (or alternative estimate of precision) around the pooled or best estimate of effect:

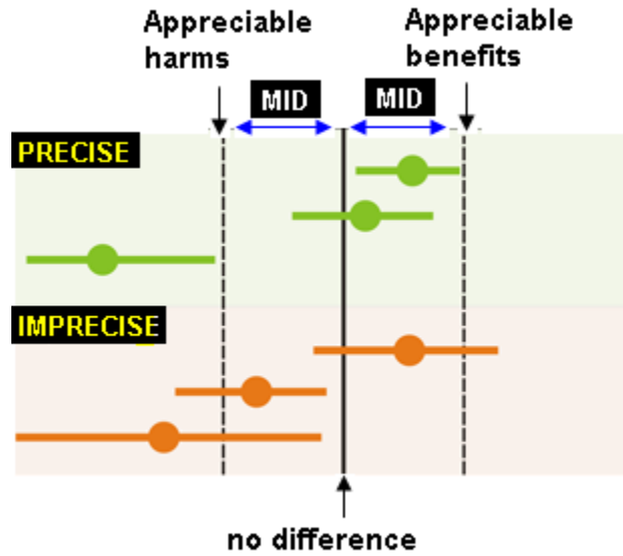
a) crosses both the line of appreciable benefit and harm, defined as imprecise

Rating for precision: 'very serious'

11

12

1 **Figure 3.1: Illustration of precise and imprecise outcomes based on the confidence interval of**  
 2 **outcomes in a forest plot**



3 *MID = minimal important difference determined for each outcome. The MIDs are the threshold for appreciable*  
 4 *benefits and harms. The confidence intervals of the top three points of the diagram were considered precise*  
 5 *because the upper and lower limits did not cross the MID. Conversely, the bottom three points of the diagram*  
 6 *were considered imprecise because all of them crossed the MID and reduced our certainty of the results. Figure*  
 7 *adapted from GRADEPro software.*

9  
 10 The following are the MID for the outcomes and the methods used to calculate the OIS  
 11 in this guideline.

- 12 For continuous outcomes:
- 13 • Anginal attacks per week: -3 to +3 attacks/week
  - 14 • Exercise time (min): +30 to 30 sec (-0.50 to +50 min)

- 15 For all dichotomous outcomes
- 16 • The default confidence intervals in GRADE of 0.75 and 1.25.

17 The MID's for the outcomes were based on the advice from the clinical advisor and  
 18 chair for the guideline.

20 **3.5 NICE economic evidence profiles**

21 The NICE economic profile has been used to summarise cost and cost-effectiveness  
 22 estimates from published studies and analyses conducted for the guideline. The  
 23 economic evidence profile shows, for each economic study, an assessment of  
 24 applicability and methodological quality, with footnotes indicating the reasons for

1 the assessment. These assessments were made by the health economist using the  
 2 economic evaluation checklist from The NICE Guidelines Manual, Appendix H  
 3 (2009). It also shows incremental costs, incremental outcomes (e.g. QALYs) and the  
 4 incremental cost-effectiveness ratio from the primary analysis, as well as information  
 5 about the assessment of uncertainty in the analysis. See Table 3.6 for more details.

6 If a non-UK study was included in the profile, the results were converted into pounds  
 7 sterling using the appropriate purchasing power parity[8].

8

9 **Table 3.6: Content of NICE economic profile**

<i>Item</i>	<i>Description</i>
<b>Study</b>	First author name, reference, date of study publication and country perspective.
<b>Limitations</b>	An assessment of methodological quality of the study*: <ul style="list-style-type: none"> <li>• <b>Minor limitations</b> – the study meets all quality criteria, or the study fails to meet one or more quality criteria, but this is unlikely to change the conclusions about cost effectiveness.</li> <li>• <b>Potentially serious limitations</b> – the study fails to meet one or more quality criteria, and this could change the conclusion about cost effectiveness</li> <li>• <b>Very serious limitations</b> – the study fails to meet one or more quality criteria and this is very likely to change the conclusions about cost effectiveness. Studies with very serious limitations would usually be excluded from the economic profile table.</li> </ul>
<b>Applicability</b>	An assessment of applicability of the study to the clinical guideline, the current NHS situation and NICE decision-making*: <ul style="list-style-type: none"> <li>• <b>Directly applicable</b> – the applicability criteria are met, or one or more criteria are not met but this is not likely to change the conclusions about cost effectiveness.</li> <li>• <b>Partially applicable</b> – one or more of the applicability criteria are not met, and this might possibly change the conclusions about cost effectiveness.</li> <li>• <b>Not applicable</b> – one or more of the applicability criteria are not met, and this is likely to change the conclusions about cost effectiveness.</li> </ul>
<b>Other comments</b>	Particular issues that should be considered when interpreting the study.
<b>Incremental cost</b>	The mean cost associated with one strategy minus the mean cost of a comparator strategy.
<b>Incremental effects</b>	The mean QALYs (or other selected measure of health outcome) associated with one strategy minus the mean QALYs of a comparator strategy.
<b>ICER</b>	Incremental cost-effectiveness ratio: the incremental cost divided by the respective QALYs gained
<b>Uncertainty</b>	A summary of the extent of uncertainty about the ICER reflecting the results of deterministic or probabilistic sensitivity analyses, or stochastic analyses of trial data, as appropriate.

10 \*Limitations and applicability were assessed using the economic evaluation checklist from The Guidelines  
 11 Manual[6], Appendix H

12 Where economic studies compare multiple strategies, results are presented in the economic  
 13 evidence profiles for the pair-wise comparison specified in the review question, irrespective  
 14 of whether or not that comparison was ‘appropriate’ within the analysis being reviewed. A  
 15 comparison is ‘appropriate’ where an intervention is compared with the next most expensive  
 16 non-dominated option – a clinical strategy is said to ‘dominate’ the alternatives when it is

1 both more effective and less costly. Footnotes indicate if a comparison was ‘inappropriate’  
2 in the analysis.

### 3 **3.5.1 Cost-effectiveness criteria**

4 The NICE Guidelines Manual[6] sets out the principles that GDGs should consider  
5 when judging whether an intervention offers good value for money. In general, an  
6 intervention was considered to be cost effective if either of the following criteria  
7 applied (given that the estimate was considered plausible):

8 a) The intervention dominated other relevant strategies (that is, it was both less  
9 costly in terms of resource use and more clinically effective compared with all  
10 the other relevant alternative strategies), **or**

11 b) The intervention cost less than £20,000 per quality-adjusted life-year (QALY)  
12 gained compared with the next best strategy.

### 13 **3.6 Undertaking new health economic analysis**

14 As well as reviewing the published economic literature for each review question, as  
15 described above, new economic analysis was undertaken by the Health Economist in  
16 priority areas. Priority areas for new health economic analysis were agreed by the  
17 GDG after formation of the review questions and consideration of the available  
18 health economic evidence.

19 Additional data for the analysis was identified as required through additional  
20 literature searches undertaken by the Health Economist, and discussion with the  
21 GDG. Model structure, inputs and assumptions were explained to and agreed by  
22 the GDG members during meetings, and they commented on subsequent revisions.

23 See Appendix H for details of the health economic analysis undertaken for the  
24 guideline.

25

### 26 **3.7 Developing recommendations**

27 Over the course of the guideline development process, the GDG was presented with:

28 • Evidence tables of the clinical and economic evidence reviewed from the  
29 literature. All evidence tables are in Appendix E2

30 • Summary of clinical and economic evidence and quality (as presented in  
31 chapters 5-19

32 • Forest plots (Appendix F)

33 • A description of the methods and results of the cost-effectiveness analysis  
34 undertaken for the guideline (Appendices G and H)

35 Recommendations were drafted on the basis of this evidence whenever it was  
36 available.

1 When clinical and economic evidence was absent, of poor quality or conflicting, the  
2 GDG drafted recommendations based on their expert opinion. This was done through  
3 discussions in the GDG. The considerations for making these consensus based  
4 recommendations included the balance between potential harms and benefits,  
5 economic or implications compared to the benefits, current practices, recommendations  
6 made in other relevant guidelines, patient preferences and equality issues. The GDG  
7 also considered whether the uncertainty was sufficient to justify delaying making a  
8 recommendation to await further research, taking into account the potential harm of  
9 failing to make a clear recommendation.

10 The main considerations specific to each recommendation are outlined in the Evidence  
11 to Recommendation Section preceding the recommendation section.

12

### 13 **3.7.1 Research recommendations**

14 When areas were identified for which good evidence was lacking, the guideline  
15 development group considered making recommendations for future research. Decisions  
16 about inclusion were based on factors such as:

- 17 • the importance to patients or the population
- 18 • national priorities
- 19 • potential impact on the NHS and future NICE guidance
- 20 • ethical and technical feasibility

21

### 22 **3.8 Validation process**

23 The guidance is subject to an eight week public consultation and feedback is used to  
24 quality assure the document. All comments received from registered stakeholders are  
25 responded to in turn and posted on the NICE website when the pre-publication check of  
26 the full guideline occurs.

27

### 28 **3.9 Updating the guideline**

29 Following publication, and in accordance with the NICE technical manual, NICE will ask a  
30 National Collaborating Centre or the National Clinical Guidelines Centre to advise  
31 NICE's Guidance executive whether the evidence base has progressed significantly to  
32 alter the guideline recommendations and warrant an update.

33

1 **3.10 Disclaimer**

2 Health care providers need to use clinical judgement, knowledge and expertise when  
3 deciding whether it is appropriate to apply guidelines. The recommendations cited here  
4 are a guide and may not be appropriate for use in all situations. The decision to adopt  
5 any of the recommendations cited here must be made by the practitioners in light of  
6 individual patient circumstances, the wishes of the patient, clinical expertise and  
7 resources.

8 The National Clinical Guideline Centre disclaims any responsibility for damages arising  
9 out of the use or non-use of these guidelines and the literature used in support of these  
10 guidelines.

11

12 **3.11 Funding**

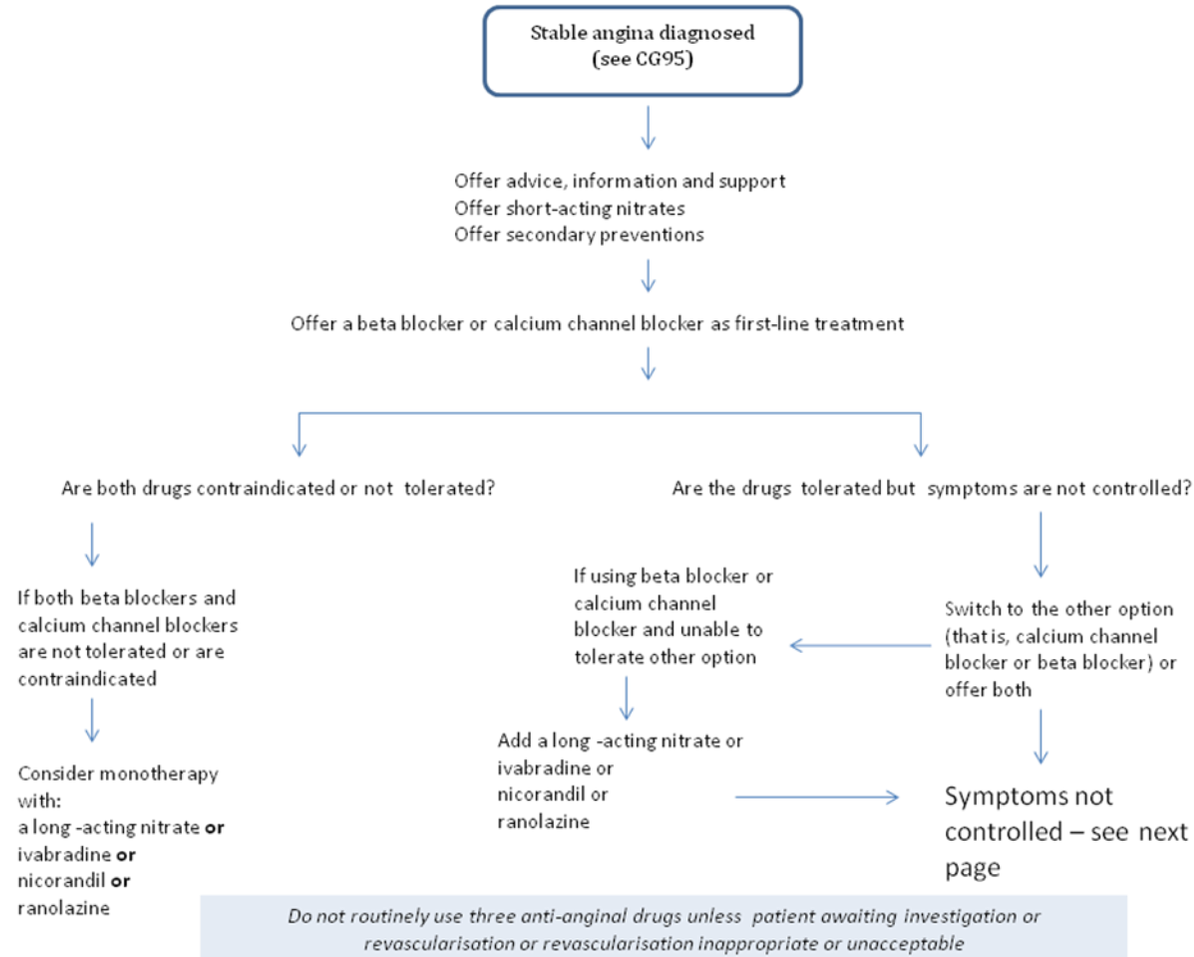
13 The National Clinical Guideline Centre was commissioned by the National Institute for  
14 Health and Clinical Excellence to undertake the work on this guideline.

15

1 **4 Guideline summary**

2 **4.1 Algorithms**

1



2

3

Symptoms not controlled on anti-anginal medication



Consider revascularisation

*If revascularisation is being considered:  
review any functional/anatomical tests performed at diagnosis,  
further non-invasive or invasive functional tests may be needed,  
consider risks and benefits of continuing drug treatment and revascularisation,  
Consider discussing with MDT*

Consider PCI for :  
People with single or multivessel disease (including left main stem disease) if coronary anatomy is suitable.

Consider CABG for:  
People with single or multivessel disease (including left main stem disease) if coronary anatomy is unsuitable for PCI. [People over 65 with multivessel disease and/or with diabetes]

If stable angina doesn't respond to drug treatment or revascularisation,  
offer comprehensive re-evaluation and advice which may include:

*Exploring the person's understanding of their condition / the impact of symptoms on the person's quality of life / reviewing the diagnosis and considering non-ischaemic causes of pain / reviewing drug treatment and considering future drug treatment and revascularisation options / explaining how the person can manage the pain themselves / acknowledging the limitations of future treatment / specific attention to role of psychological factors in pain / development of skills to modify cognitions and behaviours associated with pain.*

1

## 1 **4.2 Key priorities for implementation**

2 From the full set of recommendations, the GDG selected 10 key priorities for  
3 implementation. The criteria used for selecting these recommendations are listed in  
4 detail in The Guidelines Manual[6]. The reasons that each of these recommendations  
5 was chosen are shown in the table linking the evidence to the recommendation in  
6 Appendix I.

7 • Address personal issues including:

8 – self management skills such as pacing activities and goal setting

9 – dealing with stress or depression

10 – advice about physical exertion including sexual activity. (1.1.7)

11 • Do not routinely perform functional tests for myocardial ischaemia or anatomical tests for  
12 obstructive coronary artery disease to stratify risk. (1.2.3)

13 • Do not routinely offer percutaneous coronary intervention (PCI) or coronary artery  
14 bypass grafting (CABG) to people whose symptoms are controlled with drug treatment.  
15 (1.2.4)

16 • Offer people optimal drug treatment for the initial management of stable angina.  
17 Optimal drug treatment consists of one or two anti-anginal drugs as necessary plus drugs  
18 for secondary prevention of cardiovascular disease. (1.3.1)

19 • Consider whether the decision to continue drug treatment or perform revascularisation  
20 (PCI or CABG) needs to be discussed by a multidisciplinary team. The team should  
21 include an interventional cardiologist and a cardiac surgeon. (1.4.6)

22 • Consider the relative risks and benefits of PCI and CABG using a systematic approach  
23 to assess the severity and complexity of the person's coronary disease, in addition to  
24 other relevant clinical factors and comorbidities. (1.4.7)

25 • Consider PCI in preference to CABG for people who have single-vessel disease or  
26 multi-vessel disease, including left main stem disease, and who have continuing  
27 symptoms despite optimal medical treatment and the anatomy is suitable for PCI.  
28 (1.4.8)

29 • Consider CABG for people with single-vessel disease or multi-vessel disease, including  
30 left main stem disease, and continuing symptoms despite optimal medical treatment if  
31 the anatomy is unsuitable for PCI. (1.4.9)

32 • Consider CABG in preference to PCI for people with multi-vessel disease who have  
33 continuing symptoms despite optimal medical treatment and who:

34 – are over 65 years and/or

35 – have diabetes. (1.4.10)

- 1     • Ensure people with stable angina receive balanced information and have the  
2     opportunity to discuss the benefits, limitations and risks of continuing drug treatment, PCI  
3     and CABG to help them make an informed decision about their treatment. (1.4.11)

#### 4     **4.3 Full list of recommendations**

##### 5     ***1.1.Information and support for people with stable angina***

6           1.1.1. Clearly explain stable angina, including factors that can provoke it (for  
7           example, exertion, emotional stress, exposure to cold, a heavy meal) and its  
8           long-term course and management.

9           1.1.2. Encourage the person to ask questions about their angina and its treatment.  
10          Provide opportunities for them to voice their concerns and fears.

11          1.1.3. Discuss the person's, and if appropriate, their family or carer's ideas,  
12          concerns and expectations about their condition, prognosis and treatment.  
13          Explore and address any misconceptions about stable angina and its  
14          implications for daily activities, heart attack risk and life expectancy.

15          1.1.4. Clearly explain to the person when they should seek emergency or  
16          professional help.

17          1.1.5. Discuss with the person the purpose and any risks and benefits of their  
18          treatment.

19          1.1.6. Assess the person's need for lifestyle advice (for example about exercise,  
20          stopping smoking, diet and weight control) and psychological support, and  
21          offer interventions as necessary.

22          1.1.7. Address personal issues including:

- 23                 • self-management skills such as pacing activities and goal setting  
24                 • dealing with stress or depression  
25                 • advice about physical exertion including sexual activity.

##### 26       ***1.2.General principles for treating people with stable angina***

27           1.2.1. Do not exclude people with stable angina from treatment based on their  
28           age alone.

29           1.2.2. Do not investigate or treat symptoms of stable angina differently in men and  
30           women or in different ethnic groups.

31           1.2.3. Do not routinely perform functional tests for myocardial ischaemia or  
32           anatomical tests for obstructive coronary artery disease to stratify risk. [This  
33           recommendation partially updates recommendation 1.2 of 'Myocardial  
34           perfusion scintigraphy for the diagnosis and management of angina and  
35           myocardial infarction' (NICE technology appraisal guidance 73)]

1 1.2.4. Do not routinely offer percutaneous coronary intervention (PCI) or coronary  
2 artery bypass grafting (CABG) to people whose symptoms are controlled with  
3 drug treatment.

#### 4 **Treating episodes of angina**

5 1.2.5. Offer a short-acting nitrate for preventing and treating episodes of angina.  
6 Advise people:

- 7 • how to administer the short-acting nitrate
- 8 • to use it immediately before any planned exercise or exertion
- 9 • that side effects such as flushing, headache and light-headedness may occur
- 10 • to sit down or find something to hold on to if feeling light-headed
- 11 • when treating episodes of angina, to repeat the dose after 5 minutes if the  
12 pain has not gone
- 13 • to call an emergency ambulance if the pain has not gone 5 minutes after  
14 taking a second dose of short-acting nitrate.

#### 15 **Drugs for secondary prevention of cardiovascular disease**

16 1.2.6. Consider aspirin 75 mg daily for people with stable angina, taking into  
17 account the risk of bleeding and comorbidities.

18 1.2.7. Do not offer angiotensin-converting enzyme (ACE) inhibitors to manage  
19 stable angina. Offer ACE inhibitors to treat other conditions, as appropriate.

20 1.2.8. Offer statin treatment in line with 'Lipid modification' (NICE clinical guideline  
21 67).

22 1.2.9. Offer treatment for high blood pressure in line with 'Hypertension' (NICE  
23 clinical guideline 34, currently being updated).

#### 24 **Dietary supplements**

25 1.2.10. Do not offer fish oil or vitamin supplements to treat stable angina. Inform  
26 people that there is no evidence that they help stable angina.

#### 27 **1.3. Anti-anginal drug treatment**

##### 28 **General recommendations**

29 1.3.1. Offer people optimal drug treatment for the initial management of stable  
30 angina. Optimal drug treatment consists of one or two anti-anginal drugs as  
31 necessary plus drugs for secondary prevention of cardiovascular disease.

- 1 1.3.2. Advise people that the aim of anti-anginal drug treatment is to prevent  
2 episodes of angina and the aim of secondary prevention treatment is to  
3 prevent cardiovascular events such as heart attack and stroke.
- 4 1.3.3. Discuss how side effects of drug treatment might affect the person's daily  
5 activities and explain why it is important to take drug treatment regularly.
- 6 1.3.4. Review the person's response to treatment, including any side effects, 2–4  
7 weeks after starting or changing drug treatment.
- 8 1.3.5. Titrate the drug dosage against symptoms up to the maximum tolerable  
9 dosage.
- 10 **Drugs for treating stable angina**
- 11 1.3.6. Offer either a beta blocker or a calcium channel blocker as first-line  
12 treatment for stable angina. Decide which drug to use based on comorbidities,  
13 contraindications and the person's preference.
- 14 1.3.7. If the person cannot tolerate the beta blocker or calcium channel blocker or  
15 if it is contraindicated, switch to the other option (calcium channel blocker or  
16 beta blocker).
- 17 1.3.8. If the person's symptoms are not controlled, consider either switching to the  
18 other option (calcium channel blocker or beta blocker) or using a combination  
19 of the two<sup>1</sup>.
- 20 1.3.9. Do not routinely offer anti-anginal drugs other than beta blockers or calcium  
21 channel blockers as first-line treatment for stable angina.
- 22 1.3.10. If the person cannot tolerate beta blockers and calcium channel blockers or  
23 they are contraindicated, consider monotherapy with one of the following  
24 drugs:
- 25 • a long-acting nitrate
- 26 • ivabradine
- 27 • nicorandil<sup>2</sup> or
- 28 • ranolazine.

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<sup>1</sup> When combining a calcium channel blocker with a beta blocker, a dihydropyridine calcium channel blocker should be used.

<sup>2</sup> At the time of consultation (December 2010), nicorandil did not have UK marketing authorisation for use in this indication. Informed consent should be obtained and documented.

<sup>3</sup> Ivabradine should only be combined with a dihydropyridine calcium channel blocker.

1 Decide which drug to use based on comorbidities, contraindications, the person's  
2 preference and costs.

3 1.3.11. For people on beta blocker or calcium channel blocker monotherapy whose  
4 symptoms are not controlled and the other option (calcium channel blocker or  
5 beta blocker) is contraindicated or not tolerated, consider one of the following  
6 as an additional drug:

7 • a long-acting nitrate

8 • ivabradine<sup>3</sup>

9 • nicorandil<sup>2</sup> or

10 • ranolazine.

11 Decide which drug to use based on comorbidities, contraindications, the person's  
12 preference and costs.

13 1.3.12. Do not offer a third anti-anginal drug to people whose stable angina is  
14 controlled with two anti-anginal drugs.

15 1.3.13. Consider adding a third anti-anginal drug when:

16 • the person's symptoms are not controlled with two anti-anginal drugs and

17 • the person is waiting for revascularisation or it is not considered appropriate  
18 or acceptable.

19 Decide which drug to use based on comorbidities, contraindications, the person's  
20 preference and costs.

#### 21 **1.4. People whose symptoms are not controlled by optimal drug treatment**

22 1.4.1. Consider revascularisation (PCI or CABG) for people whose symptoms are  
23 not controlled with drug treatment.

24 1.4.2. Review the results of any functional and/or anatomical tests performed at  
25 diagnosis when revascularisation is being considered (see 'Chest pain of recent  
26 onset', NICE clinical guideline 95).

27 1.4.3. Offer coronary angiography to guide the revascularisation strategy if not  
28 recently completed during diagnosis. Additional non-invasive or invasive  
29 functional testing may be required. [This recommendation partially updates  
30 recommendation 1.2 of 'Myocardial perfusion scintigraphy for the diagnosis  
31 and management of angina and myocardial infarction' (NICE technology  
32 appraisal guidance 73)].

33 1.4.4. Consider further investigation to confirm the diagnosis of stable angina if the  
34 lack of response to drug treatment raises uncertainty about the diagnosis (see  
35 'Chest pain of recent onset', NICE clinical guideline 95).

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**Revascularisation strategy**

1.4.5. Consider the risks and benefits of continuing drug treatment or performing revascularisation (PCI or CABG) after coronary angiography.

1.4.6. Consider whether the decision to continue drug treatment or perform revascularisation (PCI or CABG) needs to be discussed by a multidisciplinary team. The team should include an interventional cardiologist and a cardiac surgeon.

1.4.7. Consider the relative risks and benefits of PCI and CABG using a systematic approach to assess the severity and complexity of the person’s coronary disease, in addition to other relevant clinical factors and comorbidities.

1.4.8. Consider PCI in preference to CABG for people who have single-vessel disease or multi-vessel disease, including left main stem disease, and who have continuing symptoms despite optimal medical treatment and the anatomy is suitable for PCI.

1.4.9. Consider CABG for people with single-vessel disease or multi-vessel disease, including left main stem disease, and continuing symptoms despite optimal medical treatment if the anatomy is unsuitable for PCI.

1.4.10. Consider CABG in preference to PCI for people with multi-vessel disease who have continuing symptoms despite optimal medical treatment and who:

- are over 65 years and/or
- have diabetes.

1.4.11. Ensure people with stable angina receive balanced information and have the opportunity to discuss the benefits, limitations and risks of continuing drug treatment, PCI and CABG to help them make an informed decision about their treatment.

1.4.12. Explain to the person that:

- The purpose of revascularisation is to improve the symptoms of stable angina.
- PCI and CABG are effective in relieving symptoms.
- CABG is slightly more effective than PCI in relieving symptoms of stable angina in the longer term.
- Repeat revascularisation may be necessary after either PCI or CABG and the rate is higher after PCI than CABG.

- 1                   • Stroke is uncommon after either PCI or CABG, and the incidence is similar  
2                   between the two procedures.

3                   1.4.13. Inform the person about the practical aspects of PCI and CABG. Include  
4                   information about:

- 5                   • vein and/or artery harvesting  
6                   • likely length of hospital stay  
7                   • recovery time  
8                   • drug treatment after the procedure.

9                   **1.5.Pain interventions**

10                  1.5.1. Do not offer the following interventions to manage stable angina:

- 11                  • transcutaneous electrical nerve stimulation (TENS)  
12                  • enhanced external counterpulsation (EECP)  
13                  • acupuncture.

14                  **1.6.Stable angina that has not responded to treatment**

15                  1.6.1. Offer people whose stable angina has not responded to drug treatment  
16                  and/or revascularisation comprehensive re-evaluation and advice, which may  
17                  include:

- 18                  • exploring the person's understanding of their condition  
19                  • exploring the impact of symptoms on the person's quality of life  
20                  • reviewing the diagnosis and considering non-ischaemic causes of pain  
21                  • reviewing drug treatment and considering future drug treatment and  
22                  revascularisation options  
23                  • explaining how the person can manage the pain themselves  
24                  • acknowledging the limitations of future treatment  
25                  • specific attention to the role of psychological factors in pain  
26                  • development of skills to modify cognitions and behaviours associated with  
27                  pain.

28

1        **1.7. Cardiac syndrome X**

2            1.7.1. In people with angiographically normal coronary arteries and continuing  
3            anginal symptoms, consider a diagnosis of cardiac syndrome X.

4            1.7.2. Continue drug treatment for stable angina only if it improves the symptoms  
5            of the person with suspected cardiac syndrome X.

6            1.7.3. Do not routinely offer drugs for the secondary prevention of cardiovascular  
7            disease to people with suspected cardiac syndrome X.

8        **4.4 Key research recommendations**

9            ***Addition of the newer anti-anginal drugs to CCB***

10            What is the clinical and cost effectiveness of adding a newer anti-anginal drug  
11            (nicorandil, ivabradine or ranolazine) to a calcium channel blocker for treating stable  
12            angina?

13            ***Interventional management vs. continued drug treatment in people with stable angina and***  
14            ***evidence of ischaemia on non-invasive functional testing***

15            Do people with stable angina and evidence of reversible ischaemia on non-invasive  
16            functional testing who are on optimal drug treatment benefit from routine coronary  
17            angiography with a view to revascularisation?

18            ***Coronary anatomy investigations***

19            In people with stable angina and multi-vessel disease (including left main stem [LMS]  
20            disease) whose symptoms are controlled on optimal drug treatment, would an initial  
21            treatment strategy of revascularisation be clinically and cost effective compared with  
22            continued drug treatment?

23            ***Cardiac Rehabilitation***

24            Is an 8-week, comprehensive, multidisciplinary, cardiac rehabilitation service more  
25            clinically and cost effective for managing stable angina than current clinical practice?

26            ***Patient Self-Management Plans***

27            What is the clinical and cost effectiveness of a self-management plan for people with  
28            stable angina?

29

30

1

## 2 **5 Patient Information**

### 3 **5.1 Introduction**

4 Stable angina is a chronic condition which people may live with for many years.  
5 People require information to ensure they understand their condition and the  
6 available treatments. Episodes of angina are potentially frightening and it is  
7 important that people are guided as to how to adapt their lifestyle if they have  
8 continuing symptoms. It is equally important however to ensure that people do not  
9 unnecessarily limit their lifestyle because of fear about precipitating angina or  
10 myocardial infarction. The GDG were interested in studies of people with angina  
11 where patients reported their information needs both at the time of diagnosis and  
12 later in the course of the condition. The question for the evidence review was:

13 “What are the information needs of people with stable angina regarding their  
14 condition and its management?”

### 15 **5.2 Information needs of people with stable angina**

#### 16 **5.2.1 Clinical Evidence**

17 The “Review Protocol” for this topic can be found in Appendix C, the “Search  
18 Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
19 E1, and the “Clinical Evidence Tables” in Appendix E2.

20 The studies included in this review were qualitative studies or questionnaires which  
21 reported direct patient experience. Four papers were included in this review; there  
22 were 3 qualitative studies[9-11] and 1 cross-sectional questionnaire study (analysed  
23 quantitatively) (Karlik 1990)[12]. Qualitative studies were critically appraised using  
24 the NICE qualitative methodology checklist. A summary of the quality of studies is  
25 included in Table 5.1. The studies and results are described in narrative format.

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**Table 5.1: Quality of included studies in evidence review for “Patient information”**

Study	Population	Methods	Analysis	Relevance to guideline population
Pier 2008[9]	Well reported	Well reported	Well reported and credible.	Australia.  Patients with MI, CABG, angioplasty or angina from GP practices.
Weetch 2003[10]	Poorly reported	Poorly reported	Poorly reported	UK. People suffering from angina who had been hospitalised in the coronary care ward.
McGillion 2004[11]	Well reported	Well reported	Well reported and credible	Canada.  People with chronic stable angina living at home.
Karlik 1990[12]	Well reported	Well reported	Quantitative analysis.	USA.  In-patients experiencing angina admitted to acute –care hospital for cardiac catheterisation.

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**Narrative report of results**

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**Pier 2008[9]** conducted a qualitative study in Melbourne, using thematic analysis of semi-structured interviews on the types of health information that people with CHD considered useful to assist with the management of their illness. Structured clinical interviews were used to assess current and prior depressive episodes in these patients. The study had 14 patients (12 men and 2 women) with a mean age of 67 years recruited from general practices. The patients had a history of MI, CABG, angioplasty or angina. Eight of these participants had a history of depression.

12

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Five themes relating to information on how patients could manage their cardiovascular health and improve their psychosocial wellbeing were recognised: psychosocial issues; anger management; physical activity; medical information; and information for family.

1 The most important information needs recognised by the patients were: the need for  
2 information on how to establish social networks and access appropriate social and  
3 support groups so as to gain support and to understand their medical condition  
4 particularly from other people with CHD; information regarding how to identify  
5 precipitating symptoms of anger and anger management; information on physical  
6 activity and amount of physical activity that could be done following an event;  
7 information regarding identification and management of risk-related physical  
8 symptoms; and information for family members and spouses, such as how the patient  
9 may react to an adverse cardiac event or medical procedure.

10

11 **Weetch 2003[10]** conducted a qualitative study to determine the level of satisfaction  
12 with the amount and quality of information received by patients suffering from angina  
13 who had been hospitalised in the coronary care ward. The patient survey was done  
14 by using questionnaires. All patients discharged from the ward with a diagnosis of  
15 angina during the study were asked to participate. Thirty patients were identified as  
16 having discharged with a diagnosis of angina during a 3 month period and were  
17 issued with a questionnaire of which 16 were returned. Seven of these 16 patients  
18 had previously been hospitalised with an MI; and 8 had angina but no previous MI.  
19 The average age of the respondents was 59.7 years (range 40 to 78 years), 60% of  
20 the respondents were male and 40% were female.

21 The results showed a very high satisfaction with the overall standard of care.  
22 However, the results showed that 73% of the patients were dissatisfied with the  
23 amount of information that they were given. They wanted to know more about the  
24 causes of angina, its treatment, their medication, and in particular the effect it will  
25 have on their daily activities. Although the patients agreed that nurses gave them the  
26 opportunity to ask questions, many wanted more written and verbal information.  
27 Another significant finding was the lack of satisfaction with the information that  
28 patients had received from health care professionals working in primary care settings.

29 **McGillion 2004[11]** conducted a qualitative study to determine the learning needs of  
30 people with chronic stable angina living at home, in order to inform content of a  
31 chronic stable angina self management programme. Eight (n=8) chronic stable angina  
32 patients were eligible and included in the study. Eligible patients had angina  
33 symptoms for at least 6 months, were experiencing either class I,II or III angina and  
34 had a medical diagnosis of CAD confirmed by nuclear imaging or angiography. The  
35 age of the eight patients ranged from 44 to 70 years, and one had post-secondary  
36 education. There were two women and 6 men in the study and the participants had  
37 angina from 6 months to 10 years.

38 Four focus groups were organised: two with chronic stable angina patients (n=5, n=3)  
39 and two with clinicians. Since the views of clinicians are not relevant to the question  
40 the results for these focus groups are not reported in the review. Each audio taped  
41 session consisted of a semi-structured interview lasting approximately 1.5 hours.

42 The results were organised according to the antecedent constructs of Braden's Self  
43 Help model: Perceived severity of illness; Uncertainty; and Limitation.

44 The patients identified that education on interpreting angina symptoms was a high  
45 priority and felt that they had great difficulty knowing when they were experiencing

1 angina versus some other type of pain. The patients felt that they had difficulty  
2 deciding to seek professional/emergency help because they doubted their own  
3 judgment, the ER was seen as a burden and also because there had some confusion  
4 about how ambulance services and tertiary care centres were organised. Patients  
5 stated that they were concerned about medication schedules, dose, side effects; and  
6 exercise frequency and acceptable duration. The patients felt that for patients  
7 dealing with angina related symptoms needed a forum in which to discuss the  
8 difficulties of identifying safe activity limits;. Patients expressed a need for help in  
9 dealing with their anxiety and also suggested that education on stress management  
10 would be helpful. Patients also gave several suggestions on how to deal with  
11 emotional responses and triggers; the most popular were teaching guided imagery  
12 and progressive muscle relaxation as means to alleviate anxiety, stress and general  
13 tension. Also, a majority of the patients expressed a need for a programme wherein  
14 they could learn to develop their chronic stable angina self-management skills.

15 **Karlik 1990[12]** conducted a questionnaire study to compare the learning needs of  
16 angina patients rated by patients themselves and the nurses who care for them. Since  
17 the review includes only information needs of patients, the results of learning needs  
18 identified by nurses are not reported.

19 The study included 15 patients (11 men, 4 women) aged 26-70 years. The sample  
20 consisted of patients experiencing angina who were selected from inpatients admitted  
21 to an acute care hospital for cardiac catheterisation. The Cardiac Patient Learning  
22 Need Inventory (CPLNI), a 43 item instrument originally designed to measure learning  
23 needs of post MI patients, and the Educator Preference Tool were used to assess the  
24 learning needs and educator preference of the patients.

25 The following 8 informational categories assessed: introduction to hospital unit;  
26 anatomy and physiology; psychologic; risk factors; medications; diet; activity; and  
27 miscellaneous.

28 In the CPLNI assessment, when the information categories were ranked by inpatient  
29 ratings, the categories of risk factors and medications emerged as the most important  
30 to learn and the categories of introduction to the hospital unit and diet emerged as  
31 the least important to learn. The category of risk factors emerged the most important  
32 to learn and the category of medications emerged as the second most important to  
33 learn, and the psychologic category emerged as the least important to learn when  
34 ranked by the post discharge patients.

35 For the Educator Preference Tool, a greater percentage of patients expressed a  
36 preference for physicians alone, rather than for nurses alone, to teach them all 8  
37 informational categories. Nurses received the highest percentage by patients in the  
38 category of introduction to the hospital unit and the lowest percentage in the  
39 categories of risk factors and activity. No patients believed the nurse alone could  
40 teach them dietary information. Physicians received the highest percentage by  
41 patients in the category of activity and the lowest percentage in the category of diet.  
42 Combining the percentages of nurses alone and nurses with others, patients still  
43 preferred physicians to teach them all informational categories except introduction to  
44 hospital unit.

45

1 **5.2.2 Economic evidence**

2 No economic studies were found on this question.

3

4 **5.2.3 Evidence statements**

The following themes have been identified on requirements for information:

- on causes of angina
- treatment of angina
- Purpose of each medication
- Medication schedules distinguishing angina from other types of pain
- prognosis and survival rates
- identification and management of risk factors
- organisation of medical services
- re-introduction of physical activity and exercise options after cardiac event
- Information for family members

Patients requested help with coping with anxiety, depression and stress management and a need for forum to discuss their condition. Patients expressed a need for learning how to manage their condition.

**Economic**

No economic evidence was found on this question.

1 5.2.4 Recommendations and link to evidence

<p><b>Recommendation</b></p>	<p><b>Clearly explain stable angina, including factors that can provoke it (for example, exertion, emotional stress, exposure to cold, a heavy meal) and its long-term course and management.</b></p> <p><b>Encourage the person to ask questions about their angina and its treatment. Provide opportunities for them to voice their concerns and fears.</b></p> <p><b>Discuss the person's, and if appropriate, their family or carer's ideas, concerns and expectations about their condition, prognosis and treatment. Explore and address any misconceptions about stable angina and its implications for daily activities, heart attack risk and life expectancy.</b></p> <p><b>Clearly explain to the person when they should seek emergency or professional help.</b></p> <p><b>Discuss with the person the purpose and any risks and benefits of their treatment.</b></p> <p><b>Advise people that the aim of anti-anginal drug treatment is to prevent episodes of angina and the aim of secondary prevention treatment is to prevent cardiovascular events such as heart attack and stroke.</b></p>
<p><b>Relative values of different outcomes</b></p>	<p>The outcomes considered as important during the development of the review protocol for patient information included information on: the condition, the symptoms, prognosis, treatment (choice of treatment and side effects), need and type of rehabilitation, prevention, activities for daily living, QoL.</p> <p>Evidence based on qualitative studies confirmed the following information themes being considered as important by stable angina patients: causes of angina and management, identification and management of risk factors, organisation of medical services, physical activity, information to family members, education on stress management, forum/groups for discussion of the condition, self-management programmes, management of anger and depression, preference for Educator for delivery of information</p>

<b>Trade off between clinical benefits and harms</b>	The studies reviewed do not provide a report on harms arising from patient information. The GDG considered that patients had a right to information about their condition and did not believe there were harms that would outweigh benefits.
<b>Economic considerations</b>	No economic evidence was found. There is a negligible cost of staff time associated with providing information to the patient. However the benefits are likely to offset the minimal costs.
<b>Quality of evidence</b>	Evidence from 4 moderate quality studies. One UK study.
<b>Other considerations</b>	<p>The GDG used the evidence from the studies, and their own experiences as professionals and patients to develop the recommendations about information required for patients. The GDG considered that information should be individualised to each patient and that exploring the patient's own concerns and ideas about their condition and its treatment was pivotal in addressing the needs of individual patients. They considered that information and advice on management of stress, anxiety, and depression was not necessarily required by all patients but healthcare professionals need to address these and other areas of importance to patients when appropriate.</p> <p>The GDG considered it particularly important that patients be advised about appropriate physical activity including sexual activity. The GDG considered it important that patients were given information about risks and benefits of treatments.</p> <p>The GDG considered it important that patients were informed what different drugs and revascularisation strategies would achieve e.g. improve symptoms and this recommendation was informed by the evidence reviews of interventions.</p>

1 **5.2.5 Research recommendation**

2 The GDG recommended the following research question:

3 ➤ **Research question:** What is the clinical and cost effectiveness of a self-management  
4 plan for people with stable angina?

5 ➤ **Why this is important:** Stable angina is a chronic condition. Evidence suggests that  
6 addressing people's beliefs and behaviours in relation to angina may improve  
7 quality of life, and reduce morbidity and use of resources. Self-management plans  
8 could include: educating people with stable angina about the role of psychological  
9 factors in pain and pain control; and teaching people self-management skills to  
10 modify cognitions, behaviours and affective responses in order to control chest pain.  
11 These skills may include pacing of physical activities, modifying stress using cognitive  
12 reframing and problem-solving techniques, and relaxation training or mindfulness  
13 techniques. The proposed study is a randomised controlled trial in primary care that

1 would assess the clinical and cost effectiveness of self-management plans. This  
2 research would inform future updates of key recommendations in the guideline.  
3 Furthermore the research would be relevant to a national priority area (National  
4 service framework for coronary heart disease [NSF CHD] chapter 4: stable angina  
5 and chapter 7: cardiac rehabilitation) as well as the Coalition White Paper 2010  
6 (Equity and excellence: liberating the NHS) that emphasize the importance of  
7 increasing people's choice and control in managing their condition.

## 1 **6 Treatment & prevention of episodes of** 2 **angina**

### 3 **6.1 Introduction**

4 In people with stable angina short-acting drugs may be used to relieve episodes of  
5 angina and can be taken prophylactically before activities that are likely to bring on  
6 an episode. Short-acting drugs include organic nitrates (e.g. glyceryl trinitrate) and  
7 nifedipine administered via the buccal mucosa. Glyceryl trinitrate (GTN) is available  
8 as a tablet or as a metered dose aerosol spray and has a rapid onset of effect.  
9 Glyceryl trinitrate tablets deteriorate when exposed to air and should be discarded  
10 after eight weeks in use (BNF). Modified release buccal glyceryl trinitrate tablets can  
11 be used for rapid relief of an episode of angina but have a slower onset and longer  
12 duration of effect (BNF). Nifedipine capsules can be used for rapid relief of an  
13 episode of angina by releasing the fluid within the capsule into the oral cavity.

14 Organic nitrates act mainly by venodilatation, but coronary vasodilatation may  
15 contribute to the therapeutic effect. Nitrates may cause headache and flushing, and  
16 repeated use may cause hypotension. Short-acting formulations of nifedipine may  
17 cause reflex tachycardia and hypotension.

18 The GDG were interested in whether there was evidence to support use of nifedipine  
19 and evidence about mode of delivery of GTN.

### 20 **6.2 Short acting nitrates**

#### 21 **6.2.1 Clinical question**

22 What is the clinical /cost effectiveness of short acting drugs for the management of  
23 anginal symptoms?

#### 24 **6.2.2 Clinical evidence**

25 The “Review Protocol” for this topic can be found in Appendix C, the “Search  
26 Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
27 E1, the “Clinical Evidence Tables” in Appendix E2, and the “Forest Plots” in Appendix  
28 F.

29

1 **Table 6.1: Sublingual nifedipine versus placebo**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Sublingual nifedipine	Placebo	Relative (95% CI)	Absolute	
<b>Mean total work time for stepped increase in load (mins) (follow-up mean 1 hour (a); measured with: minutes; better indicated by higher values)</b>											
Atterhog 1975[13]	randomised trials	serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	None	10	10 (c)	-	MD 5.2 higher (0.81 to 9.59 higher)	⊕⊕⊕O LOW
<b>Estimated workload at breakpoint for stepped increase in load (kpm/min) (follow-up mean 1 hour; measured with: kpm/min; better indicated by higher values)</b>											
Atterhog 1975[13]	randomised trials	serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	None	10 (c)	10 (c)	-	MD 146 higher (257.72 to 34.28 higher)	⊕⊕⊕O LOW
<b>Total work for stepped increase in load (kpm) (follow-up mean 1 hour (a); measured with: kpm; better indicated by higher values)</b>											
Atterhog 1975[13]	randomised trials	serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	None	10 (c)	10 (c)	-	MD 3685 higher (6489.71 to 880.29 higher)	⊕⊕⊕O LOW
<b>Mean total work time for continuous increase in load (mins) (follow-up mean 1 hour; measured with: minutes; better indicated by higher values)</b>											
Atterhog 1975[13]	randomised trials	serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	None	10 (c)	10 (c)	-	MD 1.1 higher (2.2 to 0 higher)	⊕⊕⊕O LOW
<b>Estimated workload at breakpoint for continuous increase in load (kpm/min) (follow-up mean 1 hour (a); measured with: kpm/min; better indicated by higher values)</b>											
Atterhog 1975[13]	randomised trials	serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	None	10 (c)	10 (c)	-	MD 112 higher (223.91 to 0.09 higher)	⊕⊕⊕O LOW
<b>Total work for continuous increase in load (kpm) (follow-up mean 1 hour (a); measured with: kpm; better indicated by higher values)</b>											
Atterhog 1975[13]	randomised trials	serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	None	10 (c)	10 (c)	-	MD 1146 higher (1888.83 to 403.17 higher)	⊕⊕⊕O LOW
<b>Mean work capacity at angina threshold (minutes of exercise) (measured with: minutes; better indicated by higher values)</b>											
Atterhog 1975[13]	randomised trials	serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	None	10	10	-	MD 2.1 higher (3.35 to 0.85 higher)	⊕⊕⊕O LOW
<b>Maximal work capacity at maximal exercise level (minutes of exercise) (measured with: minutes; better indicated by higher values)</b>											
Atterhog 1975[13]	randomised trials	serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	None	10 (c)	10 (c)	-	MD 2.3 higher (3.67 to 0.93 higher)	⊕⊕⊕O LOW

(a) There were 4 tests (approx 1hr) in 2 wks after entering the study. Each test was administered within 30 mins of treatment

(b) The randomisation process is not reported and double blinding of all results was not achieved due to side effects which may have jeopardised allocation concealment.

(c) This was a crossover trial

**Additional data:**

2  
3  
4  
5

1 Adverse events: No safety issues are reported in the trial. Patients spontaneously reported a feeling of "heat in the face" at an average  
 2 14 minutes after 11 of 20 administrations of nifedipine.  
 3  
 4

**Table 6.2: Sublingual nifedipine versus no treatment**

Quality assessment							Summary of findings				Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		
							Sublingual nifedipine	no treatment	Relative (95% CI)	Absolute	
<b>Mean exercise time to 1mm ST segment depression (secs) (measured with: seconds; better indicated by higher values)</b>											
Pupita 1993[14]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	10 (b)	10 (b)	-	MD 146 higher (257.13 to 34.87 higher)	⊕⊕⊕⊕ LOW

(a) Randomisation details are not reported. This comparison was not blinded.

(b) This was a crossover trial

**Table 6.3: Sublingual GTN versus sublingual nifedipine**

Quality assessment							Summary of findings				Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		
							Sublingual GTN	sublingual nifedipine	Relative (95% CI)	Absolute	
<b>Mean exercise time to 1mm ST segment depression (secs) (follow-up 4-6 mins (a); better indicated by higher values)</b>											
Pupita 1993[14]	randomised trials	serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	none	10 (c)	10 (c)	-	MD 90 higher (14.07 lower to 194.07 higher)	⊕⊕⊕⊕ LOW
<b>Mean pain severity at 2 minutes post treatment (f) (follow-up 4-6 minutes post drug administration (a); better indicated by lower values)</b>											
Mooss 1989[15]	randomised trials	serious (d)	no serious inconsistency	no serious indirectness	no serious imprecision	none	7	6	-	MD 6.3 lower (8.4 to 4.2 lower)	⊕⊕⊕⊕ LOW
<b>Mean pain severity at 4 minutes post treatment (better indicated by lower values)</b>											
Mooss 1989[15]	randomised trials	serious (d)	no serious inconsistency	no serious indirectness	no serious imprecision	none	7	6	-	MD 5.6 lower (7.08 to 4.12 lower)	⊕⊕⊕⊕ LOW
<b>No. of participants with complete pain resolution at 2 minutes post treatment (follow-up 4 to 6 minutes post drug administration (e))</b>											
Mooss 1989[15]	randomised trials	serious (d)	no serious inconsistency	no serious indirectness	Serious (g)	none	5/7 (71.4%)	0/6 (0%)	RR 9.63 (0.64 to 144.88)	710 more per 1000 (from 340 more to 1090 more)	⊕⊕⊕⊕ LOW

No. of participants with complete pain resolution at 4 minutes post treatment (patient pain intensity scoring)											
Mooss 1989[15]	randomised trials	serious (d)	no serious inconsistency	no serious indirectness	Serious (g)	none	5/7 (71.4%)	0/6 (0%)	RR 9.63 (0.64 to 144.88)	710 more per 1000 (from 340 more to 1090 more)	⊕⊕○○ LOW

- 1 (a) Patients were involved in the study for a duration of approximately 24 days. Assessments from exercise tests were made at the start and end of this period ("off therapy")
- 2 and three times directly following administration of drugs
- 3 (b) Randomisation details are not reported. It is unclear to what extent this comparison was blinded.
- 4 (c) This was a crossover trial
- 5 (d) Randomisation details are not reported. It is unclear to what extent this comparison was blinded. The trial is small - with very few participants in each arm of the parallel
- 6 phase of the trial and only 4 in one arm of the crossover phase.
- 7 (e) Patients were followed for four minutes after receiving their randomised drug. Those who had <50% reduction in pain intensity were crossed over to the alternate therapy
- 8 and followed for another 2 minutes.
- 9 (f) Patients were asked to rate the intensity of their chest pain using a 10 cm visual pain intensity rating scale (0= no pain, 10=most sever pain).
- 10 (g) Adverse events - Mooss 1989[15]: Adverse reactions attributable to nifedipine and nitroglycerin were negligible. No patients complained of side effects following nifedipine
- 11 alone. Two of the nifedipine patients complained of flushing following GTN administration and one of these patients developed a headache. One of the seven patients who
- 12 received GTN alone complained of headache.
- 13

1 **Additional data from two studies:**

2  
3 **A. Sublingual GTN versus Buccal GTN: Ryden 1987[16]**

4 N=126 [n=113 completed the study]. Open RCT with cross over design

5 **Population:** All patients had at least a 6 month history of stable angina with a  
6 minimum of 5 attacks/week

7 Mean age 61+/-8 years (range 38-82)

8 **Intervention:** 2.5mg or 5mg buccal GTN tablet for the treatment or prophylaxis of  
9 angina (tablet held in the cheek for 15 minutes 1) after the relief of angina, 2) after  
10 stopping an activity inducing pain or 3) following cessation of activity, when taken  
11 prophylactically prior to activity starting)

12 **Comparison:** Sublingual GTN

13 **Results:** During the study background medications were kept constant. Outcomes  
14 recorded in patient diaries and from 2 questionnaires administered at weeks 4 and 6.

- 15 • Treatment of anginal attacks: The total number of treated anginal attacks was  
16 31% less during the buccal (n=1381) compared to the sublingual nitroglycerin  
17 (n=1978) period (p<0.001).
- 18 • Prophylactic use: Prophylactic nitroglycerin was altogether utilised on 806  
19 occasions during the sublingual period and on 929 occasions during the buccal  
20 period respectively (p<0.05). The expected attack of angina pectoris was  
21 prevented in 66% of the attempts with sublingual and 74% of the attempts  
22 with buccal nitroglycerin (p<0.05). When angina pectoris developed despite  
23 prophylactic nitroglycerin, the distribution of mild, moderate and severe  
24 attacks did not differ significantly between the two formulations.
- 25 • Adverse events: Four patients withdrew from a cross over RCT due to side  
26 effects of buccal GTN (headache 3 patients, flushing 1 patient). Significantly  
27 more patients receiving buccal GTN reported a smarting sensation in mouth  
28 than those receiving sublingual GTN (p <0.05). There were no significant  
29 differences between patients receiving buccal and sublingual GTN for  
30 occurrence of headache, dizziness or flushing, as reported following active  
31 enquiry.
- 32 • General preference for drug: Given the opportunity to select only one of the  
33 two nitroglycerin formulations for future use 65% (p<0.05) would have  
34 preferred the buccal and 19% the sublingual, while 16% did not have any  
35 particular preference. When patients were asked to give their preference for  
36 one of the two formulations considering solely the prophylactic use, 81%  
37 preferred buccal and 4% sublingual nitroglycerin, while 15% did not express  
38 any preference (p<0.05).

1        **B. Sublingual GTN versus Spray GTN: Sandler 1967[17]**

2  
3        Quasi RCT with crossover design (n=23)  
4

5        **Population:** People with stable angina of duration range 3-72 months with attacks  
6        occurring 3 to 40 times weekly.

7        Previous MI = 4/23 participants

8        Age range 39-69 years

9        Male = 20/23 participants

10       **Intervention:** Glyceryl trinitrate aerosol delivering 0.13 mg of the drug per inhalation

11       **Comparisons:** 1) Placebo aerosol 2) Standard tablets of 0.5 mg of glyceryl trinitrate

12       **Results:** SD (standard deviation) not reported for results. Results reported as  
13       narrative.

14       Exercise tests (using a modification of the Master two-step test) were carried out at  
15       the same time each day, in the same environment, and with the same technical staff.

16       No information about concurrent therapy is reported.

17

18       **Mean change in exercise undertaken (no. of circuits over the steps)**

19       Sublingual GTN tablet before exercise = 80.9

20       Sublingual GTN tablet after exercise = 80.0

21       Mean change = +0.9 circuits

22       GTN spray before exercise = 83.5

23       GTN spray after exercise = 81.5

24       Mean change = + 2.0

25       Placebo aerosol before exercise: 83.0

26       Placebo after exercise: 80.9

27       Mean change: +2.1

28       p = non significant (reported by author)

29

30

1 **Time taken to develop angina (sec)**

2 Sublingual GTN tablet –Time taken for angina to develop (sec): mean change  
3 =+68.2 sec

4

5 GTN spray- Time taken for angina to develop (sec): Mean change = +14.5 sec

6 Placebo aerosol time taken for angina to develop (sec):+64.9 sec

7 p = non significant (reported by author)

8 **Duration of angina: (sec)**

9 Sublingual GTN tablet: 158.9 sec

10 GTN spray: 158.9 sec

11 Placebo aerosol: 218.0 sec

12 p = non significant (reported by author)

13

14 **Patient assessment:** Outpatient assessment showed that 10 patients regarded the  
15 active aerosol as more effective in relieving anginal pain, 11 chose the placebo  
16 aerosol, while two regarded active and placebo aerosols as equally effective. Only  
17 2 patients thought that tablets were better than aerosol. The only side effect  
18 encountered with the active aerosols was headache, which occurred in 6 patients.

19

20 **6.2.3 Economic evidence**

21 No economic studies were identified on this question. We calculated the range of cost per  
22 dose based on the unit cost reported in the BNF59[18].

23 **Table 6.4 Drug cost - short-acting drugs**

	<b>Specific drugs and doses</b>	<b>Cost per dose* (£)</b>
Short-acting nitrate tablets	Low = glyceryl trinitrate 300 micrograms	0.05
	High = glyceryl trinitrate 600 micrograms	0.28
Short-acting nitrate spray	Glyceryl trinitrate 400 micrograms	0.03

Short-acting nifedipine capsules	Low = nifedipine 5mg	0.07
	High = nifedipine 10mg	0.09

1 \* dose = 2 tablets or 2 sprays  
2

3 Overall the drug cost of short-acting nitrate spray is lower than the drug cost of  
4 sublingual nitrate tablets or nifedipine capsules.

5

6 **6.2.4 Evidence statements**

**Clinical Sublingual nifedipine versus placebo**

**Atterhog 1975[13]:** Evidence from one cross over RCT shows that compared to placebo, prophylactic sublingual nifedipine was associated with significantly higher mean total work time for stepped increase in load (mins) [MD 5.20 [0.81 to 9.59]]; estimated workload at breakpoint for stepped increase in load (kpm/min) (MD 146.00 [34.28 to 257.72] ;total work for stepped increase in load (kpm) (MD 3685.00 [880.29 to, 6489.71] ] ; mean total work time for continuous increase in load (mins) (MD 1.10 [0.00 to 2.20]); estimated workload at breakpoint for continuous increase in load (kpm/min) [MD 112.00 [0.09 to 223.91]; total work for continuous increase in load (kpm) (MD 1146.00 [403.17 to 1888.83]]; mean work capacity at angina threshold (mins of exercise): [MD 2.10 [0.85 to 3.35]] and maximal work capacity at maximal exercise level (mins of exercise): (MD 2.30 [0.93 to 3.67])).

**Sublingual nifedipine versus no treatment**

**Pupita 1993[14]:** Evidence from one cross over RCT shows that compared to no treatment, sublingual nifedipine significantly increased the mean exercise time to 1mm ST depression (sec): [MD 146.00 [34.87 to 257.13]

**Sublingual GTN versus sublingual nifedipine**

**Mooss 1989[15]:** Evidence from one parallel RCT shows that sublingual GTN was significantly more effective than sublingual nifedipine in reducing pain severity (mean pain intensity rating) at 2 minutes post treatment: MD -6.30 [-8.40 to -4.20] and at reducing pain severity (mean pain intensity rating) 4 minutes post treatment: MD -5.60 [-7.08 to -4.12]. By four minutes only 2 of 6 participants in the sublingual nifedipine group had >50% reduction in mean pain intensity.

**Mooss 1989[15]:** sublingual GTN was significantly more effective than sublingual nifedipine in providing complete pain resolution at 2 minutes

post treatment: [RR 9.63 [0.64 to 144.88]] and complete pain resolution at 4 minutes post treatment: [RR 9.63 [0.64 to 144.88]].

**Pupita 1993[14]:** There was no statistically significant difference between sublingual GTN and sublingual nifedipine in the mean exercise time to 1 mm ST depression (sec): [MD 90.00 [-14.07 to 194.07]].

**Economic** No economic evidence was found on this question. A simple cost analysis showed a small difference in drug costs between short-acting nitrates and nifedipine and between spray and sublingual short-acting nitrates; spray nitrates are the least costly.

1 **6.2.5 Recommendations and link to evidence**

<i>Recommendation</i>	<p><b>Offer a short-acting nitrate for preventing and treating episodes of angina. Advise people:</b></p> <ul style="list-style-type: none"> <li>• <b>how to administer the short-acting nitrate</b></li> <li>• <b>to use it immediately before any planned exercise or exertion</b></li> <li>• <b>that side effects such as flushing, headache and light-headedness may occur</b></li> <li>• <b>to sit down or find something to hold on to if feeling light-headed</b></li> <li>• <b>when treating episodes of angina, to repeat the dose after 5 minutes if the pain has not gone</b></li> <li>• <b>to call an emergency ambulance if the pain has not gone 5 minutes after taking a second dose of short-acting nitrate.</b></li> </ul>
-----------------------	---

**Relative values of different outcomes**

The outcome of interest was relief and prevention of episodes of angina.

**Trade off between clinical benefits and harms**

Evidence from three small randomised trials suggests that sublingual nifedipine increases measures of exercise capacity on a treadmill relative to placebo or to no treatment. Sublingual glyceryl trinitrate was more effective than sublingual nifedipine at reducing pain severity and providing complete symptom relief at two and four minutes after treatment. One trial reported that buccal glyceryl trinitrate tablet (held in cheek for 15 minutes) is more effective than

sublingual glyceryl trinitrate tablet at reducing the number of angina episodes requiring treatment and at preventing expected angina attacks.

One trial compared sublingual glyceryl trinitrate tablets with glyceryl trinitrate spray during daily exercise tests for six days and reported no significant differences in the amount of exercise or in the time to onset of anginal symptoms between the two treatment groups.

The GDG concluded that people with stable angina should be offered a short-acting drug to relieve episodes of angina. Weak evidence suggests that glyceryl trinitrate relieves episodes of angina more effectively than nifedipine.

**Economic considerations**

No economic evidence on the use of short-acting drugs was available for review. As glyceryl trinitrate is more effective at relieving episodes of angina and it does not increase costs compared to nifedipine, this drug is likely to be more cost-effective.

**Quality of evidence**

The trials in this review were very small and of poor quality.

No economic evidence was available.

**Other considerations**

The GDG noted that glyceryl trinitrate spray is easy to use and can be stored over long periods without loss of effect. After exposure to air glyceryl trinitrate tablets lose efficacy and should be discarded after eight weeks in use[18]. An advantage of glyceryl trinitrate tablets is that they can be discarded as soon as the angina episode is relieved to avoid the onset of adverse effects (including headache). The GDG did not however consider they could recommend one formulation of GTN over another and formulation should be chosen according to patient preferences and needs.

The GDG considered it important that patients are given adequate information regarding use of short acting nitrates and made a consensus recommendation about instructions for patients. These were informed by the current advice from the British Heart Foundation.

([http://www.bhf.org.uk/living\\_with\\_a\\_heart\\_condition/understanding\\_heart\\_conditions/types\\_of\\_heart\\_conditions/angina.aspx](http://www.bhf.org.uk/living_with_a_heart_condition/understanding_heart_conditions/types_of_heart_conditions/angina.aspx) )

1

2

1

## 2 **7 Beta blockers vs. calcium channel blockers**

### 3 **7.1 Introduction**

4 Anti-anginal drugs prevent attacks of angina by decreasing myocardial oxygen  
5 consumption (by lowering heart rate, blood pressure, myocardial loading, or  
6 myocardial contractility) and/or by increasing myocardial oxygen supply (by  
7 increasing coronary blood flow).

8 Evidence that monotherapy with single anti-anginal agents prevents attacks of angina  
9 has been reviewed previously. The quantity and quality of this evidence is limited but  
10 there is consensus that BBs and CCBs are effective in the treatment of people with  
11 stable angina [19-22].

12 The aim of this review was to determine whether BBs or CCBs offer advantages as  
13 first-line treatment for people with stable angina. The review includes evidence from  
14 nine trials of monotherapy with BBs versus monotherapy with CCBs.

#### 15 **Beta blockers**

16 Beta blockers reduce myocardial oxygen consumption by competitive inhibition of  
17 beta-adrenoceptors, which lowers heart rate, blood pressure, and myocardial  
18 contractility. The bradycardia prolongs diastole, thereby increasing the period of  
19 maximal coronary blood flow. Relative contra-indications to beta-blockade include  
20 obstructive airways disease, acute heart failure, and impaired atrioventricular  
21 conduction. Side effects of BBs include fatigue, altered carbohydrate metabolism,  
22 peripheral vasoconstriction, sexual dysfunction, and bronchoconstriction. Some BBs  
23 (e.g. atenolol, metoprolol, bisoprolol) are relatively cardioselective with greater  
24 inhibition of the cardiac beta<sub>1</sub> receptors than the beta<sub>2</sub> (bronchial) receptors and  
25 therefore have less effect on airways resistance. Atenolol, bisoprolol, and nadolol  
26 have a relatively long duration of action and are given once daily. Other BBs with  
27 shorter half-lives may be given in slow-release formulations.

28 In the United Kingdom the most frequently prescribed BBs are atenolol, bisoprolol,  
29 and propranolol. The cost of a BB for four weeks is low (e.g. £0.99 for atenolol 50mg  
30 daily, £1.33 for bisoprolol 10mg daily)[23].

31

1           **Calcium channel blockers**

2           Calcium channel blockers inhibit movement of calcium through slow calcium channels of  
3           cell membranes in the myocardium, cardiac conduction tissues, and vascular smooth  
4           muscle. Calcium channel blockers dilate peripheral and coronary arteries, and to a  
5           varying degree depress myocardial contractility and intra-cardiac conduction.  
6           Calcium channel blockers include dihydropyridines (e.g. amlodipine), benzothiapines  
7           (e.g. diltiazem), and phenylalkylamines (e.g. verapamil). Dihydropyridines may cause  
8           reflex tachycardia, flushing, headache, and ankle swelling. Diltiazem and verapamil  
9           depress cardiac conduction and cause bradycardia, and should not be given to  
10          people with heart block or treated with a BB. Verapamil may cause constipation.

11          In the United Kingdom the most frequently prescribed CCBs are amlodipine,  
12          nifedipine, felodipine, diltiazem, and verapamil. The costs of CCBs for four weeks are  
13          higher than the costs of atenolol (e.g. amlodipine 10mg daily £1.43; diltiazem MR  
14          60mg tds £2.93; verapamil 80mg tds £2.07)[23]. Slow release formulations of CCBs  
15          are more expensive.

16          **Generic beta blockers, calcium channel blockers included in evidence reviews**

17  
18          A large number of BBs and CCBs are available for clinical use in the UK and different  
19          BBs and CCBs have been used in different trials.

20          We looked at the prescription cost analysis table in the NHS Information Centre. We  
21          extracted the number of prescriptions in 2005 and 2008 for BB and CCB dispensed  
22          in the community. The GDG reviewed the lists and made a judgement about which  
23          drugs were currently used in stable angina and were known to be included in studies.

24          In this guideline we have considered evidence for the use of the following drugs:

- 25                 • BB: atenolol, propranolol, bisoprolol, metoprolol, nadolol,  
26                 • CCB: amlodipine, diltiazem, felodipine, nifedipine, verapamil

27          In this review we have assumed that the clinical effects are consistent within a class  
28          of drug (e.g. BB or CCB), across a range of doses, and in all trial participants.

29

30          **7.2 Beta blocker vs. calcium channel blocker**

31          **7.2.1 Clinical question**

32          What is the comparative clinical /cost effectiveness of standard antianginal drugs  
33          (BBs/CCBs) for the management of angina?

34

1 **7.2.2 Clinical evidence**

2 The “Review Protocol” for this topic can be found in Appendix C, the “Search  
3 Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
4 E1, the “Clinical Evidence Tables” in Appendix E2, and the “Forest Plots” in Appendix  
5 F.

6

1 **Table 7.1: BB vs. CCB for stable angina**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							BB	CCB	Relative (95% CI)	Absolute	
<b>Exercise duration (min) (metoprolol vs. diltiazem; propranolol vs. diltiazem; propranolol vs. nifedipine) (follow-up 6 weeks-6 months; better indicated by higher values)</b>											
van Dijk 1988[24]; O'Hara 1987[25]; Kawanishi 1992[26]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	88	83	-	MD 0.05 higher (0.82 lower to 0.92 higher)	⊕⊕○○ LOW
<b>Time to 1mm ST depression (sec) - metoprolol vs. nifedipine (follow-up 10 weeks; better indicated by higher values)</b>											
Savonitto 1996[27] (IMAGE)	randomised trials	serious (c)	no serious inconsistency	no serious indirectness	serious (b)	none	65	62	-	MD 12 higher (35.06 lower to 59.06 higher)	⊕⊕○○ LOW
<b>Time to onset of angina (min) (metoprolol vs. diltiazem; propranolol vs. nifedipine) (follow-up 6 weeks-6 months; better indicated by lower values)</b>											
van Dijk 1988[24]; Kawanishi 1992[26]	randomised trials	serious (d)	no serious inconsistency	no serious indirectness	serious (b)	none	54	49	-	MD 0.63 higher (0.27 lower to 1.53 higher)	⊕⊕○○ LOW
<b>Total mortality (atenolol vs. verapamil; metoprolol vs. verapamil; metoprolol vs. verapamil) (follow-up 2.7-9.1 years)</b>											
Pepine 2003[28] (INVEST); Rehnqvist 1996[29] (APSYS); Hjerdahl 2006[30] (APSYS)	randomised trials	serious (e)	no serious inconsistency	no serious indirectness	no serious imprecision	none	972/12121 (8%)	964/12073 (8%)	RR 1 (0.92 to 1.09)	0 fewer per 1000 (from 6 fewer to 7 more)	⊕⊕⊕○ MODERATE
<b>Cardiovascular death (atenolol vs. verapamil; atenolol vs. nifedipine; metoprolol vs. verapamil) (follow-up 2-3.4 years)</b>											
Pepine 2003[28] (INVEST); Dargie 1996[31] (TIBET); Rehnqvist 1996[29] (APSYS)	randomised trials	serious (f)	no serious inconsistency	no serious indirectness	no serious imprecision	none	453/11941 (3.8%)	456/11902 (3.8%)	OR 0.99 (0.87 to 1.12)	0 fewer per 1000 (from 5 fewer to 4 more)	⊕⊕⊕○ MODERATE
<b>Non fatal MI (atenolol vs. verapamil; atenolol vs. nifedipine; metoprolol vs. verapamil) (follow-up 2-3.4 years)</b>											
Pepine 2003[28] (INVEST); Dargie 1996[31] (TIBET); Hjerdahl 2006[30] (APSYS)	randomised trials	serious (g)	no serious inconsistency	no serious indirectness	no serious imprecision	none	184/11941 (1.5%)	185/11902 (1.6%)	RR 0.99 (0.81 to 1.22)	0 fewer per 1000 (from 3 fewer to 3 more)	⊕⊕⊕○ MODERATE
<b>CV related hospitalisation – (atenolol vs. verapamil) (follow-up mean 2.7 years)</b>											
Pepine 2003[28] (INVEST)	randomised trials	no serious limitations (h)	no serious inconsistency	no serious indirectness	no serious imprecision	none	709/11309 (6.3%)	726/11267 (6.4%)	OR 0.97 (0.88 to 1.08)	2 fewer per 1000 (from 7 fewer to 5 more)	⊕⊕⊕⊕ HIGH
<b>Non fatal CV events (combined) – (metoprolol vs. verapamil) (follow-up median 3.4 years)</b>											
Rehnqvist 1996[29] (APSYS)	randomised trials	serious (i)	no serious inconsistency	no serious indirectness	serious (j)	none	106/406 (26.1%)	98/403 (24.3%)	RR 1.07 (0.85 to 1.36)	17 more per 1000 (from 36 fewer to 88 more)	⊕⊕○○ LOW
<b>Angina episodes/week (atenolol vs. verapamil; metoprolol vs. diltiazem; propranolol vs. nifedipine; metoprolol vs. nifedipine) (follow-up 6 weeks-2.7 years; better indicated by lower values)</b>											

Pepine 2003[28] (INVEST); van Dijk 1988[24]; Kawanishi 1992[26]; Savonitto 1996[27] (IMAGE) (s)	randomised trials	serious (k)	no serious inconsistency	no serious indirectness	no serious imprecision	none	11424	11377	-	MD 0.11 higher (0.07 to 0.15 higher)	⊕⊕⊕○ MODERATE
<b>Prevalence of angina – (atenolol vs. verapamil) (follow-up mean 2.7 years)</b>											
Pepine 2003[28] (INVEST)	randomised trials	no serious limitations (h)	no serious inconsistency	no serious indirectness	serious (j)	none	228/11309 (2%)	261/11267 (2.3%)	RR 0.87 (0.73 to 1.04)	3 fewer per 1000 (from 6 fewer to 1 more)	⊕⊕⊕○ MODERATE
<b>Severity of angina assessed by investigator (moderate/markedly improved) – (nadolol vs. amlodipine) (follow-up 6 months)</b>											
Singh 1993[32]	randomised trials	serious (l)	no serious inconsistency	no serious indirectness	no serious imprecision	none	21/39 (53.8%)	29/39 (74.4%)	RR 0.72 (0.51 to 1.02)	208 fewer per 1000 (from 364 fewer to 15 more)	⊕⊕⊕○ MODERATE
<b>Severity of angina assessed by patients (moderate/severe) – (nadolol vs. amlodipine) (follow-up 6 months)</b>											
Singh 1993[32]	randomised trials	serious (l)	no serious inconsistency	no serious indirectness	no serious imprecision	none	16/40 (40%)	12/40 (30%)	RR 1.33 (0.73 to 2.45)	99 more per 1000 (from 81 fewer to 435 more)	⊕⊕⊕○ MODERATE
<b>Nitroglycerin use – (propranolol vs. nifedipine) (follow-up 6 months; better indicated by lower values)</b>											
Kawanishi 1992[26]	randomised trials	serious (m)	no serious inconsistency	no serious indirectness	no serious imprecision	none	21	16	-	MD 0 higher (0.94 lower to 0.94 higher)	⊕⊕⊕○ MODERATE
<b>Adverse effects (head ache) – (metoprolol vs. verapamil) (follow-up median 3.4 years)</b>											
Rehmqvist 1996[29] (APSYS)	randomised trials	serious (i)	no serious inconsistency	no serious indirectness	no serious imprecision	none	3/406 (0.7%)	4/403 (1%)	RR 0.74 (0.17 to 3.31)	3 fewer per 1000 (from 8 fewer to 23 more)	⊕⊕⊕○ MODERATE
<b>Adverse effects (GI events) – (metoprolol vs. verapamil) (follow-up median 3.4 years)</b>											
Rehmqvist 1996[29] (APSYS)	randomised trials	serious (i)	no serious inconsistency	no serious indirectness	serious (n)	none	10/406 (2.5%)	22/403 (5.5%)	OR 0.45 (0.22 to 0.94)	29 fewer per 1000 (from 3 fewer to 42 fewer)	⊕⊕○○ LOW
<b>Adverse effects (dizziness) – (atenolol vs. verapamil) (follow-up mean 2.7 years)</b>											
Pepine 2003[28] (INVEST)	randomised trials	no serious limitations (h)	no serious inconsistency	no serious indirectness	no serious imprecision	none	151/11309 (1.3%)	154/11267 (1.4%)	RR 0.98 (0.78 to 1.22)	0 fewer per 1000 (from 3 fewer to 3 more)	⊕⊕⊕⊕ HIGH
<b>Adverse effects (lightheadedness) – (atenolol vs. verapamil) (follow-up mean 2.7 years)</b>											
Pepine 2003[28] (INVEST)	randomised trials	no serious limitations (h)	no serious inconsistency	no serious indirectness	no serious imprecision	none	70/11309 (0.6%)	48/11267 (0.4%)	RR 1.45 (1.01 to 2.1)	2 more per 1000 (from 0 more to 5 more)	⊕⊕⊕⊕ HIGH
<b>Adverse effects (overall) (atenolol vs. amlodipine; metoprolol vs. verapamil; nadolol vs. amlodipine) (follow-up 10 weeks-3.4years)</b>											
Pehrsson 2000[33]; Rehmqvist 1996[29] (APSYS); Singh	randomised trials	serious (o)	no serious inconsistency	no serious indirectness	no serious imprecision	none	139/562 (24.7%)	146/559 (26.1%)	RR 0.95 (0.79 to	13 fewer per 1000 (from 55 fewer to	⊕⊕⊕○ MODERATE

1993[32] (q)									1.14)	37 more)	
<b>Adverse effects (constipation) – (atenolol vs. verapamil) (follow-up mean 2.7 years)</b>											
Pepine 2003[28] (INVEST)	randomised trials	no serious limitations (h)	no serious inconsistency	no serious indirectness	no serious imprecision	none	15/11309 (0.1%)	195/11267 (1.7%)	RR 0.08 (0.05 to 0.13)	16 fewer per 1000 (from 15 fewer to 16 fewer)	⊕⊕⊕⊕ HIGH
<b>Withdrawals due to adverse effects – (atenolol vs. nifedipine) (follow-up mean 2 years)</b>											
Dargie 1996[31] (TIBET) (r)	randomised trials	serious (p)	no serious inconsistency	no serious indirectness	serious (n)	none	60/226 (26.5%)	93/232 (40.1%)	RR 0.66 (0.51 to 0.87)	136 fewer per 1000 (from 52 fewer to 196 fewer)	⊕⊕○○ LOW
<b>Combined outcome (death, non fatal MI, non fatal stroke) (diabetes) - atenolol vs. verapamil (follow-up mean 2.7 years)</b>											
Pepine 2003[28] (INVEST)	randomised trials	no serious limitations (h)	no serious inconsistency	no serious indirectness	no serious imprecision	none	450/3231 (13.9%)	463/3169 (14.6%)	RR 0.95 (0.85 to 1.07)	7 fewer per 1000 (from 22 fewer to 10 more)	⊕⊕⊕⊕ HIGH
<b>Combined outcomes (death, non fatal MI, non fatal stroke) (females) - atenolol vs. verapamil (follow-up mean 2.7 years)</b>											
Pepine 2003[28] (INVEST)	randomised trials	no serious limitations (h)	no serious inconsistency	no serious indirectness	no serious imprecision	none	540/5920 (9.1%)	524/5850 (9%)	RR 1.02 (0.91 to 1.14)	2 more per 1000 (from 8 fewer to 13 more)	⊕⊕⊕⊕ HIGH
<b>Combined (death, non fatal MI, non fatal stroke) - subgroup age &gt;70 - atenolol vs. verapamil (follow-up mean 2.7 years)</b>											
Pepine 2003[28] (INVEST)	randomised trials	no serious limitations (h)	no serious inconsistency	no serious indirectness	no serious imprecision	none	664/3829 (17.3%)	596/3694 (16.1%)	RR 1.07 (0.97 to 1.19)	11 more per 1000 (from 5 fewer to 31 more)	⊕⊕⊕⊕ HIGH
<b>Quality of life (sleep disturbance) – (metoprolol vs. verapamil) (follow-up median 3.4 years; better indicated by lower values)</b>											
Rehqvist 1996[29] (APSYS)	randomised trials	serious (i)	no serious inconsistency	no serious indirectness	serious (b)	none	270	275	-	MD 0.4 lower (1.3 lower to 0.5 higher)	⊕⊕○○ LOW
<b>Quality of life (overall life satisfaction) –(metoprolol vs. verapamil) (follow-up median 3.4 years; better indicated by lower values)</b>											
Rehqvist 1996[29] (APSYS)	randomised trials	serious (i)	no serious inconsistency	no serious indirectness	serious (b)	none	268	275	-	MD 0.7 lower (5.07 lower to 3.67 higher)	⊕⊕○○ LOW
<b>Quality of life (psychosomatic symptoms) – (metoprolol vs. verapamil) (follow-up median 3.4 years; better indicated by lower values)</b>											
Rehqvist 1996[29] (APSYS)	randomised trials	serious (i)	no serious inconsistency	no serious indirectness	serious (b)	none	275	282	-	MD 1.3 lower (3.89 lower to 1.29 higher)	⊕⊕○○ LOW

- 1 (a) van Dijk 1988[24]; O'Hara 1987[25]; Kawanishi 1992[26]: All 3 studies randomised. Allocation concealment not reported in all 3 studies. All 3 studies double blind. ITT not reported in all 3 studies.
- 2
- 3 (b) 95% CI includes no effect and the upper and lower CI crosses the MID.
- 4 (c) Randomised. Double blind. Allocation concealment not reported. Baseline comparison made. Drop out <20% (11%). Intention to treat analysis not reported.

- 1 (d) van Dijk 1988[24]; Kawanishi 1992[26]: Both studies randomised. Allocation concealment not reported in both studies. ITT not reported in both studies. Both studies double  
2 blind.
- 3 (e) Pepine 2003[28]; Hjemdahl 2006[30] (APSYS); Rehnqvist 1996[29] (APSYS); All 3 randomised. Allocation concealment not reported in all 3 studies and ITT used in all 3  
4 the studies. All 3 studies double blind.
- 5 (f) Pepine 2003[28]; Rehnqvist 1996[29] (APSYS); Dargie 1996[31] (TIBET): All 3 studies randomised. Allocation concealment reported in 1 of the 3 studies. ITT reported in  
6 all 3 studies. All 3 studies double blind.
- 7 (g) Dargie 1996 (TIBET); Pepine 2003[28]; Hjemdahl 2006[30] (APSYS): All 3 studies randomised. Allocation concealment reported in 1 of 3 studies. ITT reported in all 3  
8 studies. All 3 studies double blind.
- 9 (h) Randomised. Allocation concealment reported. Double blind. Power calculation reported. Drop-out rate <20% (2.5%). Baseline comparisons made. Intention to treat  
10 analysis used.
- 11 (i) Double blind. Randomised. Allocation concealment not reported. Baseline comparisons made. Power calculation reported. Drop out <20%. Intention to treat analysis  
12 reported.
- 13 (j) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.
- 14 (k) Pepine 2003[28]; van Dijk 1988[24]; Kawanishi 1992[26]; Savonitto 1996[27]: All 4 studies randomised. Allocation concealment not reported in 3 of the 4 studies. ITT  
15 not reported in 3 of the 4 studies. All 4 studies double blind.
- 16 (l) Double blind. Randomised. Allocation concealment not reported. Baseline comparisons made. Drop out >20% 23% [(19/80) drop out; 20% (8/40) in the amlodipine  
17 group and 27% (11/40) in the nadolol group]. Intention to treat analysis not reported.
- 18 (m) Randomised. Double blind. Allocation concealment not reported. Baseline comparisons made. Drop out < 20% (2.8% in nifedipine group and 2.6% in propranolol group).  
19 Intention to treat analysis not reported.
- 20 (n) 95% CI around the pooled estimate of effect includes appreciable benefit or appreciable harm.
- 21 (o) Pehrsson 2000[33]; Rehnqvist 1996[29] (APSYS); Singh 1993[32]: Randomised all 3 studies. Allocation concealment not reported in all 3 of the studies. ITT not reported  
22 in 2 of the 3 studies. All 3 studies double blind.
- 23 (p) Randomised. Double blind. Allocation concealment not reported. Baseline comparisons made. Drop-out >20% [60(27%) for atenolol, 93 (40%) for nifedipine, 64  
24 (29%) for their combination]. Intention to treat analysis reported.
- 25 (q) Most commonly reported side effects with nadolol were bradycardia, dizziness, headache, nausea, dyspnoea, palpitations, and fatigue. Most frequently reported side  
26 effects with amlodipine were headache, oedema, palpitations, hypoesthesia, and flushing.
- 27 (r) Not reported what were the side effects.
- 28 (s) The comparisons here are metoprolol +placebo vs. nifedipine +placebo.
- 29

1        **Additional data:**

2        **Vliegen 1991[34]**

3        **Population:** n=56 (n=26 metoprolol, n=30 diltiazem).

4        People with stable effort induced angina pectoris for at least 3 months. The mean  
5        age in the diltiazem group was  $58\pm 9$  yrs and in the metoprolol group was  $64\pm 9$  yrs  
6        ( $p<0.05$ );

7        **Intervention:** metoprolol 100 mg b.i.d.

8        The treatment was preceded by a 2 week run-in period. If the patients were already  
9        taking antianginal medication (other than short acting nitrates) this was gradually  
10       discontinued. In the second week of the run-in period, only short acting nitrates were  
11       used by all patients. If the patients were not taking antianginal medication, the single  
12       blind run-in period was 1 week.

13       **Comparison:** diltiazem 120 mg b.i.d.

14       **Follow-up:** Follow-up 8 weeks, 20 weeks and 32 weeks.

15       **Results:**

- 16        • Exercise test (32 weeks): during treatment, mean changes in duration of  
17        exercise, time to angina pectoris, time to 1 mm ST segment depression,  
18        maximal ST segment depression were not significantly different between the  
19        patients on diltiazem and those on metoprolol. However at 20 weeks, exercise  
20        duration was longer in patients on diltiazem than in patients on metoprolol.
- 21        • Frequency of angina (8 weeks): the mean frequency of anginal attacks/ week  
22        decreased in diltiazem group from 5.9 at baseline to 3.5 during treatment  
23        ( $p<0.05$ ) and in the metoprolol group from 7.4 at baseline to 4.7 during  
24        treatment ( $p<0.01$ ). No differences were observed between the two  
25        treatment groups.

26        **Side effects:** no significant differences were found in incidence and severity of side  
27        effects between the 2 groups.

28        **Drug dosages in each study:**

- 29        1. Dargie 1996[31] (TIBET) - atenolol 50 mg twice daily, nifedipine (slow  
30        release) 20-40 mg twice daily
- 31        2. Pepine 2003[28] (INVEST) - Group 1: atenolol 50 mg twice daily +  
32        hydrochlorothiazide 25 mg twice daily + trandolapril 2mg/d ; Group 2:  
33        verapamil sustained release, 180 mg twice daily + hydrochlorothiazide, 25  
34        mg/d + trandolapril, 2 mg twice daily
- 35        3. Pehrsson 2000[33] - amlodipine 10 mg once daily, atenolol 100 mg once  
36        daily.
- 37

- 1 4. van Dijk 1988[24] - diltiazem 240 mg (60 mg four times daily), metoprolol  
2 200 mg (100 mg twice daily)
- 3 5. Savonitto 1996[27] (IMAGE)- metoprolol (controlled release, 200 mg once  
4 daily), nifedipine (retard, 20 mg tablets twice daily)
- 5 6. Rehnqvist 1996[29] (APSYS), Hjemdahl 2006[30] (APSYS)- metoprolol (Seloken  
6 ZOC 200 mg once daily), verapamil (Isoptin Retard 240 twice daily)
- 7 7. Singh 1993[32] - amlodipine 2.5-10 mg once daily, nadolol 40-160 mg once  
8 daily
- 9 8. O' Hara 1987[25] - diltiazem 360 mg once daily, propranolol 240 mg once  
10 daily
- 11 9. Kawanishi 1992[26] - nifedipine 10 mg four times daily vs. propranolol 20mg  
12 four times daily (not specified if it is long or short acting nifedipine)

13

14 **7.2.3 Economic evidence**

15 One study[35] included the relevant comparison. This is summarised in the economic  
16 evidence profile below. The base case results reported are for patients without any of the  
17 following comorbidities: ischaemic heart disease (excluding angina), hypertension,  
18 congestive cardiac failure, hypercholesterolaemia and cerebrovascular disease. We report  
19 the results for patients with comorbidity as a part of sensitivity analysis. See also Economic  
20 Evidence Tables in Appendix G.

21

22

**Table 7.2: BB vs. CCB - Economic study characteristics**

<b>Study</b>	<b>Limitations</b>	<b>Applicability</b>	<b>Other Comments</b>
<b>Borghi 2000[35]</b>	Potentially serious limitations (a)	Partial applicability (b)	Tenormin and Tildiem were respectively the BB and CCB evaluated. Resource use data were obtained from a database.

23 a) Based on a cross-sectional study; only one drug from each group was evaluated.

24 b) Not a full economic evaluation: only costs, not health effects.

25

1 **Table 7.3: BB vs. CCB - Economic summary of findings**

Study	Incremental cost per patient (£)	Incremental effects	ICER	Uncertainty
<b>Patients in their first year of antianginal treatment</b>				
<b>Borghi 2000[35]</b>	Saves 358 (a)	NR	NA	Subgroup analysis in patients with comorbidities: BB has an incremental cost of £580 per patient. The overall results do not change when: <ul style="list-style-type: none"> <li>- frequency of GP visits is varied</li> <li>- incidence of hospitalisation is varied (from 0 to double)</li> <li>- the cost of generic drugs is used.</li> </ul>
<b>Patients in the year following a change in previous medication</b>				
<b>Borghi 2000[35]</b>	97 (a)	NR	NA	Subgroup analysis in patients with comorbidities: costs had the same trend. The overall results do not change when: <ul style="list-style-type: none"> <li>- frequency of GP visits is varied</li> <li>- incidence of hospitalisation is varied (from 0 to double)</li> <li>- the cost of generic drugs is used.</li> </ul>
<b>Patients who had received the same treatment during the previous year</b>				
<b>Borghi 2000[35]</b>	Saves 16 (a)	NR	NA	Subgroup analysis in patients with comorbidities: costs had the same trend. The overall results do not change when: <ul style="list-style-type: none"> <li>- frequency of GP visits is varied</li> <li>- incidence of hospitalisation is varied (from 0 to double)</li> <li>- the cost of generic drugs is used.</li> </ul>

2 (a) 1997/1998 GBP. Costs included were cost of anti-anginal drugs, additional medication, GP-initiated  
3 tests, GP and practice nurse visits, outpatient visits, elective and emergency admissions. Resource costs  
4 were obtained NHS databases and UK cost studies.

5

6 **7.2.4 Evidence statements****Clinical BB vs. CCB****Clinical efficacy:**

**Pepine 2003[28] (INVEST), Van Dijk 1988[24], Kawanishi 1992[26], Savonitto 1996[27] (IMAGE):** Evidence from 4 RCT's shows that there were significantly fewer anginal episodes/week [MD 0.11 (0.07 to 0.15)] with CCB (verapamil, diltiazem, nifedipine) compared with BB (atenolol, metoprolol, propranolol) (follow-up 6 weeks-2.7 years).

**Van Dijk 1988[24], O'Hara 1987[25], Kawanishi 1992[26]:** Evidence from 3 RCTs shows that there was no significant difference between BB (metoprolol, propranolol) and CCB (diltiazem, nifedipine) for exercise duration (min) [MD 0.05 (-0.82 to 0.92)] (follow-up 6 weeks- 6 months).

**Savonitto 1996[27] (IMAGE):** Evidence from one RCT shows that there was no significant difference between BB (metoprolol) and CCB (nifedipine) for time to 1mm ST segment depression [MD 12 (-35.06 to 59.06)] (follow-up 10 weeks).

**Van Dijk 1988[24], Kawanishi 1992[26]:** Evidence from 2 RCTs shows that there was no significant difference between BB (metoprolol, propranolol) and CCB (diltiazem, nifedipine) for time to onset of angina (min) [MD 0.63 (-0.27 to 1.53)] (follow-up 6 weeks-6 months).

**Pepine 2003[28] (INVEST), Rehnqvist 1996[29] (APSYS), Hjemdahl 2006[30] (APSYS):** Evidence from 2 RCTs (3 papers) shows that there was no significant difference between BB (atenolol, metoprolol) and CCB (verapamil) for total mortality [RR 1 (0.92 to 1.09)]. (follow-up 2.7- 9.1 years)

**Pepine 2003[28] (INVEST), Dargie 1996[31] (TIBET), Rehnqvist 1996[29] (APSYS):** Evidence from 3 RCTs shows that there was no significant difference between BB (atenolol, metoprolol) and CCB (verapamil, nifedipine) for cardiovascular death [RR 0.99 (0.87 to 1.12)] (follow-up 2-3.4 years).

**Pepine 2003[28] (INVEST), Dargie 1996[31] (TIBET), Hjemdahl 2006[30] (APSYS):** Evidence from 3 RCTs shows that there was no significant difference between BB (atenolol, metoprolol) and CCB (verapamil, nifedipine) for non fatal MI [RR 0.99 (0.81 to 1.22)] (follow-up 2-3.4 years).

**Pepine 2003[28] (INVEST):** Evidence from one RCT shows that there was no significant difference between BB (atenolol) and CCB (verapamil) for cardiovascular related hospitalisation [RR 0.97 (0.88 to 1.08)] (follow-up mean 2.7 years).

**Rehnqvist 1996[29] (APSYS):** Evidence from one RCT shows that there was no significant difference between BB (metoprolol) and CCB (verapamil) for non fatal CV events (acute MI, incapacitating or unstable angina, cerebrovascular events or peripheral vascular events). [RR 1.07 (0.85 to 1.36)] (Follow-up median 3.4 years).

**Pepine 2003[28] (INVEST):** Evidence from one RCT shows that there was no significant difference between BB (atenolol) and CCB (verapamil) for prevalence of angina [RR 0.87 (0.73 to 1.04)] (follow-up mean 2.7 years).

**Singh 1993[32]:** Evidence from one RCT shows that there was no significant difference between BB (nadolol) and CCB (amlodipine) for severity of angina (assessed by investigators as moderate/markedly improved) [RR 0.72 (0.51 to 1.02)] (follow-up 6 months).

**Singh 1993[32]:** Evidence from one RCT shows that there was no significant difference between BB (nadolol) and CCB (amlodipine) for severity of angina (assessed by patients as moderate/severe) [RR 1.33 (0.73 to 2.45)] (follow-up 6 months).

**Kawanishi 1992[26]:** Evidence from one RCT shows that there was no significant difference between BB (propranolol) and CCB (nifedipine) for

use of nitroglycerin tablets/week [MD 0 (-0.94 to 0.94)] (follow-up 6 months).

**Rehnqvist 1996[29] (APSYS):** Evidence from one RCT shows that there was no significant difference between BB (metoprolol) and CCB (verapamil) for quality of life psychosomatic symptoms [MD -1.3 (-3.89 to 1.29)], overall life satisfaction [MD -0.7 (-5.07 to 3.67)], and sleep disturbance [MD -0.4 (-1.3 to 0.5)] [ (follow-up median 3.4 years).

**Pepine 2003[28] (INVEST):** Evidence from one RCT shows that there was no significant difference between BB and CCB for combined outcomes (death, non fatal MI, non fatal stroke) in sub group analyses conducted for age > 70 years [RR 1.07 (0.97 to 1.19)], female gender [RR 1.02 (0.91 to 1.14)] and people with diabetes [RR 0.95 (0.85 to 1.07)] (follow-up mean 2.7 years).

**Adverse effects:**

**Dargie 1996[31] (TIBET):** Evidence from one RCT shows that there were significantly more withdrawals due to adverse effects [RR 0.66 (0.51 to 0.87)] with CCB (nifedipine) compared to BB (atenolol) (follow-up mean 2 years).

**Pepine 2003[28] (INVEST):** Evidence from one RCT shows that there were significantly more adverse effects (constipation) [RR 0.08 (0.05 to 0.13)] with CCB (verapamil) compared to BB (atenolol) (follow-up mean 2.7 years).

**Pepine 2003[28] (INVEST):** Evidence from one RCT shows that there were significantly more adverse effects (light headedness) [RR 1.45 (1.01 to 2.1)] with BB (atenolol) compared to CCB (verapamil) (follow-up mean 2.7 years).

**Rehnqvist 1996[29] (APSYS):** Evidence from one RCT shows that there were significantly more adverse effects (GI events) [RR 0.45 (0.22 to 0.94)] with CCB (verapamil) compared to BB (metoprolol) (median 3.4 years).

**Pehrsson 2000[33], Rehnqvist 1996[29] (APSYS), Singh 1993:** Evidence from 3 RCTs shows that there was no significant difference between BB (atenolol, metoprolol, nadolol) and CCB (amlodipine, verapamil) for adverse effects (overall) [RR 0.95 (0.79 to 1.14)] (follow-up 10 weeks- 3.4 years).

**Pepine 2003[28] (INVEST):** Evidence from one RCT shows that there was no significant difference between BB (atenolol) and CCB (verapamil) in adverse effects (dizziness) [RR 0.98 (0.78 to 1.22)] (follow-up mean 2.7 years).

**Rehnqvist 1996[29] (APSYS):** Evidence from one RCT shows that there

was no significant difference between BB (metoprolol) and CCB (verapamil) for adverse effects (head ache) [RR 0.74 (0.17 to 3.31)] (follow-up median 3.4 years).

**Economic** Patients with and without co-morbidities were analysed separately. In patients without comorbidities BB generate fewer costs during the first year of treatment. BB and CCB have similar costs after the first year. In patients with comorbidities BB generate more costs also during the first year. This evidence has potentially serious limitations and partial applicability.

1 **7.2.5 Recommendations and link to evidence**

<b>Recommendation</b>	<p><b>Offer either a beta blocker or a calcium channel blocker as first-line treatment for stable angina. Decide which drug to use based on comorbidities, contraindications and the person's preference.</b></p> <p><b>If the person cannot tolerate the beta blocker or calcium channel blocker or if it is contraindicated, switch to the other option (calcium channel blocker or beta blocker).</b></p> <p><b>If the person's symptoms are not controlled, consider either switching to the other option (calcium channel blocker or beta blocker) or using a combination of the two*.</b></p> <p><b>Do not routinely offer anti-anginal drugs other than beta blockers or calcium channel blockers as first-line treatment for stable angina.</b></p> <p><i>*When combining a calcium channel blocker with a beta blocker, a dihydropyridine calcium channel blocker should be used</i></p>
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**Relative values of different outcomes**

The outcomes of most interest were long-term mortality (total and cardiovascular) and rates of major adverse cardiovascular events (myocardial infarction, stroke, myocardial revascularisation). Other outcomes included were measures of symptom severity (frequency of angina, exercise test outcomes).

**Trade off between clinical benefits and harms**

We found no evidence of a difference in total or cardiovascular mortality, or in risk of myocardial infarction or stroke, between people with stable angina treated with CCB

or BB. In one large trial the effect of treatment with CCB and BB on a combined endpoint (death, non-fatal myocardial infarction, non-fatal stroke) was consistent across subgroups, including women, and people with diabetes or aged over 70 years.

In one high quality trial the prevalence of angina two years after randomisation was similar amongst people treated with sustained release verapamil and amongst people treated with atenolol. On the other hand, evidence from four randomised controlled trials suggests that there are 0.11 fewer angina episodes per week amongst patients treated with CCB than amongst patients treated with BB. This difference equates to a single episode of angina every nine weeks and the GDG did not consider this to be of major clinical significance.

In one trial there was no difference in quality of life assessed with the Cornell Medical Index between patients treated with CCB or with BB.

There is no evidence of a consistent and clinically important difference in the rate of adverse events between patients treated with BB or CCB. In one large trial treatment with verapamil was associated with constipation but treatment with atenolol was associated with light-headedness.

The GDG concluded that there is no evidence on which to base the choice between BB and CCB for the initial treatment of people with stable angina.

Evidence to guide treatment if monotherapy with a BB or a CCB is not tolerated or does not control symptoms of angina is very limited. The GDG reached a consensus that if one class of anti-anginal drug is not tolerated or is ineffective a switch to the other class of anti-anginal drug can be considered.

**Economic considerations**

The cost of treatments with BB and CCB and their consequences is similar after the first year. The presence of comorbidities might influence the level of resource use (e.g. admissions) during the first year.

**Quality of evidence**

Randomised trials of BBs and CCBs in people with stable angina have mainly studied older drugs within each drug class (e.g. propranolol, atenolol, metoprolol, nifedipine, diltiazem, verapamil). The trials selectively recruited patients who were suitable for treatment with either a BB or CCB.

Information about the long term effects of BBs and CCBs in the treatment of people with stable angina is very limited. Most trials were not designed to study the effects of treatment on mortality or other major cardiovascular outcomes, are limited

by small study size, and only report short to medium term follow-up. One large trial was designed to detect a difference in the composite rate of death, non-fatal myocardial infarction, and non-fatal stroke at two years.[28]

The economic evidence has potentially serious limitations (it was based on a cross-sectional study and only one drug from each group was evaluated) and partial applicability (only costs, not health effects were measured).

### **Other considerations**

The GDG recognised the historical consensus that monotherapy with BB or CCB is effective for the prevention of attacks of angina. The GDG was also aware that monotherapy with organic nitrates is limited by the development of tolerance, and that evidence to support monotherapy with other antianginal drugs (nicorandil, ivabradine, ranolazine) is very limited. The GDG concluded that anti-anginal drugs other than BBs or CCBs should not be used as first line treatments for stable angina.

Previous guidelines[19-21] have suggested that BBs should be the first-line treatment for stable angina because of evidence that beta-blockade reduces mortality after acute myocardial infarction[36] and in people with chronic heart failure[37,38]. It has also been suggested that short-acting dihydropyridines may have deleterious effects in people with coronary artery disease[39]. We found no evidence to differentiate between the use of BB versus the use of CCB for first-line treatment of stable angina.

The GDG were also aware of a consensus that BBs and CCBs have a class effect on symptoms of angina, but that the potential for a particular drug to cause adverse effects may be influenced by its pharmacological profile (for example cardioselectivity for BBs and effects on intra-cardiac conduction for CCBs). The GDG considered that the available evidence did not support a recommendation to use a specific BB or CCB. Nevertheless, clinicians should be aware that evidence for anti-anginal efficacy is mainly confined to the use of a small number of older agents (e.g. propranolol, atenolol, metoprolol, nifedipine, diltiazem, verapamil), and clinicians should consider comorbidity, contra-indications and patient preference when selecting a first-line anti-anginal agent. The difference in cost between BBs and CCBs is small and was not considered significant by the GDG. The choice of initial treatment should therefore be determined by co-morbidity, contraindications and patient preference.

1

## 2 **8 Combination of beta blockers and calcium** 3 **channel blockers**

### 4 **8.1 Introduction**

5 This guideline identifies BBs and CCBs as first-line anti-anginal agents for the  
6 treatment of people with stable angina. In some people with angina monotherapy  
7 with a BB or CCB will control the symptoms but other people may continue to  
8 experience episodes of angina. In these people future treatment options include  
9 switching to an alternative anti-anginal drug, or addition of a second anti-anginal  
10 drug. The GDG were also interested to know whether there is long term benefit from  
11 using more than one drug even if symptoms are controlled.

12 In this section we review evidence that the addition of a BB to a CCB, or the addition  
13 of a CCB to a BB improves symptoms or clinical outcomes in people with stable  
14 angina.

15

### 16 **8.2 Beta blocker vs. beta blocker+calcium channel blocker**

#### 17 **8.2.1 Clinical question**

18 What is the comparative clinical/cost effectiveness of BB vs. BB+CCB for the  
19 management of angina?

20

#### 21 **8.2.2 Clinical evidence**

22 The "Review Protocol" for this topic can be found in Appendix C, the "Search  
23 Strategies" in Appendix D, the "List of Included and Excluded Studies" in Appendix  
24 E1, the "Clinical Evidence Tables" in Appendix E2, and the "Forest Plots" in Appendix  
25 F.

26

1 **Table 8.1: BB vs. BB + CCB for stable angina**

Quality assessment							Summary of findings				
							No of patients		Effect		Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	BB	BB +CCB	Relative (95% CI)	Absolute	
<b>Exercise time (min) (follow-up 10 weeks-6 months; better indicated by higher values) (propranolol vs. propranolol+nifedipine; propranolol vs. propranolol+diltazem; propranolol vs. propranolol+nifedipine)</b>											
Tweddel 1981[40]; O'Hara 1987[25]; Kawanishi 1992[26]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	73	41	-	MD 0.89 lower (1.67 to 0.11 lower)	⊕⊕○○ LOW
<b>Time to onset of angina (min) – (propranolol vs. propranolol+nifedipine) (follow-up 6 months; better indicated by highervalues)</b>											
Kawanishi 1992[26] (k)	randomised trials	serious (c)	no serious inconsistency	no serious indirectness	serious (b)	none	21	16	-	MD 0.2 higher (1.13 lower to 1.53 higher)	⊕⊕○○ LOW
<b>Angina attacks/week (follow-up 10weeks-6 months; better indicated by lower values) (propranolol vs. propranolol+nifedipine; metoprolol vs. metoprolol+nifedipine)</b>											
Kawanishi 1992[26]; Savonitto 1996[27] (IMAGE )	randomised trials	serious (d)	no serious inconsistency	no serious indirectness	no serious imprecision	none	82	77	-	MD 0.43 higher (0.56 lower to 1.41 higher)	⊕⊕⊕○ MODERATE
<b>Angina attacks/day –( propranolol vs. propranolol+nifedipine )(follow-up 10 weeks; better indicated by lower values)</b>											
Tweddel 1981[40]	randomised trials	serious (e)	no serious inconsistency	no serious indirectness	serious (b)	none	18	18	-	MD 3 higher (2.49 lower to 8.49 higher)	⊕⊕○○ LOW
<b>Nitroglycerin tablets/week –( propranolol vs. propranolol+nifedipine) (follow-up 6 months; better indicated by lower values)</b>											
Kawanishi 1992[26]	randomised trials	serious (c)	no serious inconsistency	no serious indirectness	no serious imprecision	none	21	16	-	MD 0.4 higher (0.15 lower to 0.95 higher)	⊕⊕⊕○ MODERATE
<b>Cardiac death – (atenolol vs. atenolol+nifedipine) (follow-up mean 2 years)</b>											
Dargie 1996[31] (TIBET)	randomised trials	serious (f)	no serious inconsistency	no serious indirectness	serious (g)	none	3/226 (1.3%)	4/224 (1.8%)	RR 0.74 (0.17 to 3.28)	5 fewer per 1000 (from 15 fewer to 41 more)	⊕⊕○○ LOW
<b>Non fatal MI – (atenolol vs. atenolol+nifedipine) (follow-up mean 2 years)</b>											
Dargie 1996[31] (TIBET)	randomised trials	serious (f)	no serious inconsistency	no serious indirectness	serious (g)	none	14/226 (6.2%)	7/224 (3.1%)	RR 1.98 (0.82 to 4.82)	31 more per 1000 (from 6 fewer to 119 more)	⊕⊕○○ LOW
<b>Withdrawals due to side effects – (atenolol vs. atenolol+nifedipine) (follow-up mean 2 years)</b>											
Dargie 1996[31] (TIBET) (j)	randomised trials	serious (f)	no serious inconsistency	no serious indirectness	serious (g)	none	60/226 (26.5%)	64/224 (28.6%)	RR 0.93 (0.69 to 1.25)	20 fewer per 1000 (from 89 fewer to 71 more)	⊕⊕○○ LOW
<b>Adverse effects (overall) – (atenolol vs. atenolol+amlodipine) (follow-up 10 weeks)</b>											
Pehrsson 2000[33]	randomised trials	serious (h)	no serious inconsistency	no serious indirectness	serious (g)	none	52/116 (44.8%)	59/119 (49.6%)	RR 0.9 (0.69 to 1.19)	50 fewer per 1000 (from 154 fewer to 94 more)	⊕⊕○○ LOW

											more)	
<b>Time to 1mm ST depression (sec)- (metoprolol vs. metoprolol+nifedipine) (follow-up 10 weeks; better indicated by higher values)</b>												
Savonitto 1996[27] (IMAGE)	randomised trials	serious (i)	no serious inconsistency	no serious indirectness	serious (b)	none	65	63	-	MD 59 lower (107.3 to 10.7 lower)	⊕⊕○○	LOW

- 1 (a) O'Hara 1987[25]: Randomised cross over trial. Double blind. Allocation concealment not reported. Baseline characteristics not reported. Drop out >20% (32%). Intention
- 2 to treat analysis not reported. Kawanishi 1992[26]: Randomised. Double blind. Allocation concealment not reported. Baseline comparisons made. Drop out < 20% (2.8%
- 3 in nifedipine group and 2.6% in propranolol group). Intention to treat analysis not reported. Tweddel 1981[40]: Randomised cross over trial. Double blind. Baseline
- 4 characteristics not reported. Allocation concealment not reported. Drop-out >20% (28%). Intention to treat analysis not reported.
- 5 (b) 95% CI includes no effect and the upper and lower CI crosses the MID.
- 6 (c) Randomised. Double blind. Allocation concealment not reported. Baseline comparisons made. Drop out < 20% (2.8% in nifedipine group and 2.6% in propranolol group).
- 7 Intention to treat analysis not reported.
- 8 (d) Kawanishi 1992[26]: Randomised. Double blind. Allocation concealment not reported. Baseline comparisons made. Drop out < 20% (2.8% in nifedipine group and 2.6%
- 9 in propranolol group). Intention to treat analysis not reported. Savonitto 1996[27]: Randomised. Double blind. Allocation concealment not reported. Baseline comparison
- 10 made. Drop out <20% (11%). Intention to treat analysis not reported.
- 11 (e) Randomised cross over trial. Double blind. Baseline characteristics not reported. Allocation concealment not reported. Drop-out >20% (28%). Intention to treat analysis
- 12 not reported.
- 13 (f) Randomised. Double blind. Allocation concealment not reported. Baseline comparisons made. Drop-out >20% [60(27%) for atenolol, 93 (40%) for nifedipine, 64
- 14 (29%) for their combination]. Intention to treat analysis reported.
- 15 (g) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.
- 16 (h) Randomised. Double blind. Allocation concealment not reported. Drop out <20% Baseline comparisons made. Intention to treat analysis not reported.
- 17 (i) Randomised. Double blind. Allocation concealment not reported. Baseline comparison made. Drop out <20% (11%). Intention to treat analysis not reported.
- 18 (j) Not reported what were the side effects
- 19 (k) At baseline (n= 74 participants): NYHA angina class I (4%), class II (73%), class III (23%)

1  
2  
3**Drug dosages in each study:**

- 4 1. Pehrsson 2000[33] - amlodipine 10 mg once daily, atenolol 100 mg once daily.  
 5 2. Dargie 1996[31] (TIBET) - atenolol 50 mg twice daily, nifedipine (slow release) 20-  
 6 40 mg twice daily  
 7 3. O' Hara[25] - 1987 - diltiazem 360 mg once daily, propranolol 240 mg once  
 8 daily,  
 9 4. Savonitto 1996[27] (IMAGE study)- metoprolol (controlled release, 200 mg once  
 10 daily), nifedipine (Retard, 20 mg tablets twice daily)  
 11 5. Kawanishi 1992[26] - nifedipine 10 mg four times daily, propranolol 20 mg four  
 12 times daily (not specified if it is long or short acting nifedipine)  
 13 6. Tweddel 1981[40] - nifedipine 10 mg three times daily, propranolol dose not  
 14 reported. After an initial placebo phase patients were commenced on propranolol,  
 15 with increasing doses at weekly intervals until a resting heart rate of less than 60  
 16 beats/min was obtained, and there was a 30% reduction in exercise tachycardia.  
 17 Patients were then randomly allocated to the addition of placebo or nifedipine in a  
 18 dose of 10mg, three times daily to their B-blocker therapy in a double blind cross  
 19 over fashion over two consecutive 3 week periods. Finally the B-blocker dose of  
 20 propranolol was gradually halved over a 2 week period. Patients continued on the  
 21 50% B-blocker dose and nifedipine for a further 2 weeks

22 **8.2.3 Economic evidence**

23 No economic studies were identified on this question. We calculated the range (low and  
 24 high) of daily and annual cost of adding CCB based on the unit cost reported in the  
 25 BNF59[18].

26 **Table 8.2: Cost of adding CCB**

	Specific drugs used for cost range	Additional cost per day (£)	Additional cost per year (£)
CCB	Low = amlodipine	0.04	15
	High = felodipine	0.15	55

27

28 The costs of future adverse effects and events were not estimated.

29 **8.2.4 Evidence statements****Clinical BB vs. BB+CCB****Clinical efficacy:**

**Tweddel 1981[40], O'Hara 1987[25], Kawanishi 1992[26]:** Evidence from 3 RCTs shows that exercise time (min) [MD -0.89 (-1.67 to -0.11)] was significantly higher with BB+CCB (propranolol+nifedipine, propranolol+diltiazem) compared to BB (propranolol) (follow-up 10

weeks to 6 months).

**Savonitto 1996[27] (IMAGE)**: Evidence from one RCT shows that time to 1 mm ST segment depression (sec) [MD -59 (-107.3 to -10.7)] was significantly higher with BB+CCB (metoprolol+nifedipine) than with BB (metoprolol) (follow-up 10 weeks).

**Kawanishi 1992[26]**: Evidence from one RCT shows that there was no significant difference between BB (propranolol) and BB+CCB (propranolol+nifedipine) for time to onset of angina (min) [MD 0.2 (-1.13 to 1.53)] (follow-up 6 months).

**Kawanishi 1992[26], Savonitto 1996[27] (IMAGE)**: Evidence from 2 RCTs shows that there was no significant difference between BB (propranolol, metoprolol) and BB+CCB (propranolol +nifedipine, metoprolol+nifedipine) for angina attacks/week. [MD 0.43 (-0.56 to 1.41)] (follow-up 10 weeks-6 months).

**Tweddel 1981[40]**: Evidence from one RCT shows that there was no significant difference between BB (propranolol) and BB+CCB (propranolol+nifedipine) for no. of angina attacks/day. [MD 3 (-2.49 to 8.49)] (follow-up 10 weeks).

**Kawanishi 1992[26]**: Evidence from one RCT shows that there was no significant difference between BB (propranolol) and BB+CCB (propranolol+nifedipine) for use of nitroglycerin tablets/week. [MD 0.4 (-0.15 to 0.95)] (follow-up 6 months)

**Dargie 1996[31] (TIBET)**: Evidence from one RCT shows that there was no significant difference between BB (atenolol) and BB+CCB (atenolol+nifedipine) for cardiac death. [RR 0.74 (0.17 to 3.28)] (follow-up mean 2 years).

**Dargie 1996[31] (TIBET)**: Evidence from one RCT shows that there was no significant difference between BB (atenolol) and BB+CCB (atenolol+nifedipine) for non fatal MI. [RR 1.98 (0.82 to 4.82)] (follow-up mean 2 years).

**Adverse effects:**

**Dargie 1996[31] (TIBET)**: Evidence from one RCT shows that there was no significant difference between BB (atenolol) and BB+CCB (atenolol +nifedipine) for withdrawal due to side effects [RR 0.93 (0.69 to 1.25)] (follow-up mean 2 years).

**Pehrsson 2000[33]**: Evidence from one RCT shows that there was no significant difference between BB (atenolol) and BB+CCB (atenolol +amlodipine) for adverse effects (overall) [RR 0.9 (0.69 to 1.19)] (follow-up 10 weeks).

**Economic** No economic evidence was found on this question. A simple cost analysis showed a small increase in drug costs when a CCB is added to the therapy.

1 **8.3 Calcium channel blocker vs. beta blocker + calcium channel blocker**

2 **8.3.1 Clinical question**

3 What is the comparative clinical/cost effectiveness of CCB vs. BB+CCB for the  
4 management of angina?

5

6 **8.3.2 Clinical evidence**

7 The “Review Protocol” for this topic can be found in Appendix C, the “Search Strategies” in  
8 Appendix D, the “List of Included and Excluded Studies” in Appendix E1, the “Clinical  
9 Evidence Tables” in Appendix E2, and the “Forest Plots” in Appendix F

1 **Table 8.3: CCB vs. BB + CCB for stable angina**

Quality assessment							Summary of findings					Importance
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality	
							CCB	BB +CCB	Relative (95% CI)	Absolute		
<b>Exercise time (min) (follow-up 18 weeks -6 months; better indicated by higher values) (diltiazem vs. propranolol +diltiazem; nifedipine vs. propranolol +nifedipine)</b>												
O'Hara 1987[25]; Kawanishi 1992[26]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	50	26	-	MD 1.91 lower (2.87 to 0.95 lower)	⊕⊕⊕○ MODERATE	
<b>Cardiac death (nifedipine vs. atenolol+nifedipine) (follow-up mean 2 years)</b>												
Dargie 1996[31] (TIBET)	randomised trials	serious (b)	no serious inconsistency	no serious indirectness	serious (c)	None	6/232 (2.6%)	4/224 (1.8%)	RR 1.45 (0.41 to 5.06)	8 more per 1000 (from 11 fewer to 72 more)	⊕⊕○○ LOW	
<b>Non fatal MI - nifedipine vs. atenolol+nifedipine (follow-up mean 2 years)</b>												
Dargie 1996[31] (TIBET)	randomised trials	serious (b)	no serious inconsistency	no serious indirectness	serious (c)	None	15/232 (6.5%)	7/224 (3.1%)	RR 2.07 (0.86 to 4.98)	33 more per 1000 (from 4 fewer to 124 more)	⊕⊕○○ LOW	
<b>Withdrawals due to side effects - nifedipine vs. atenolol+nifedipine (follow-up mean 2 years)</b>												
Dargie 1996[31] (TIBET) (h)	randomised trials	serious (b)	no serious inconsistency	no serious indirectness	serious (d)	None	93/232 (40.1%)	64/224 (28.6%)	RR 1.4 (1.08 to 1.82)	114 more per 1000 (from 23 more to 234 more)	⊕⊕○○ LOW	
<b>Adverse effects (overall) - amlodipine vs. atenolol+amlodipine (follow-up 10 weeks)</b>												
Pehrsson 2000[33]	randomised trials	serious (e)	no serious inconsistency	no serious indirectness	serious (c)	None	60/116 (51.7%)	59/119 (49.6%)	RR 1.04 (0.81 to 1.34)	20 more per 1000 (from 94 fewer to 169 more)	⊕⊕○○ LOW	
<b>Time to onset of angina (min) - nifedipine vs. propranolol+nifedipine (follow-up 6 months; better indicated by higher values)</b>												
Kawanishi 1992[26] (j)	randomised trials	serious (f)	no serious inconsistency	no serious indirectness	serious (g)	None	16	19	-	MD 0.5 lower (1.93 lower to 0.93 higher)	⊕⊕○○ LOW	
<b>Angina episodes/week (follow-up 10 weeks-6 months; better indicated by lower values) (nifedipine vs. propranolol+nifedipine; nifedipine vs. metoprolol+nifedipine)</b>												
Kawanishi 1992[26]; Savonitto 1996[27] (IMAGE) (i)	randomised trials	serious (h)	no serious inconsistency	no serious indirectness	no serious imprecision	None	77	76	-	MD 0.1 higher (1.62 lower to 1.82 higher)	⊕⊕⊕○ MODERATE	
<b>Nitroglycerin tablets/week - nifedipine vs. propranolol+nifedipine (follow-up 6 months; better indicated by lower values)</b>												
Kawanishi 1992[26]	randomised trials	serious (i)	no serious inconsistency	no serious indirectness	no serious imprecision	None	16	19	-	MD 0.4 lower (1.66 lower to 0.86 higher)	⊕⊕⊕○ MODERATE	
<b>Time to 1 mm ST segment depression (sec) - nifedipine vs. metoprolol+nifedipine (follow-up 10 weeks; better indicated by lower values)</b>												
Savonitto 1996[27]	randomised	serious <sup>10</sup>	no serious	no serious	serious <sup>7</sup>	None	62	59	-	MD 70 lower	⊕⊕○○	

(IMAGE)	trials		inconsistency	indirectness						(125.13 to 14.87 lower)	LOW	
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- 1 (a) *Kawanishi 1992[26]: Randomised. Double blind. Allocation concealment not reported. Baseline comparisons made. Drop out < 20% (2.8% in nifedipine group and 2.6% in propranolol group). Intention to treat analysis not reported; O'Hara 1987[25]: Randomised cross over trial. Double blind. Allocation concealment not reported. Baseline characteristics not reported. Drop out >20% (32%). Intention to treat analysis not reported.*
- 2
- 3
- 4 (b) *Randomised. Double blind. Allocation concealment not reported. Baseline comparisons made. Drop-out >20% [60(27%) for atenolol, 93 (40%) for nifedipine, 64 (29%) for their combination]. Intention to treat analysis reported.*
- 5
- 6 (c) *95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.*
- 7 (d) *95% CI around the pooled estimate of effect includes appreciable benefit or appreciable harm.*
- 8 (e) *Randomised. Double blind. Allocation concealment not reported. Drop out <20% Baseline comparisons made. Intention to treat analysis not reported.*
- 9 (f) *Randomised. Double blind. Allocation concealment not reported. Baseline comparisons made. Drop out < 20% (2.8% in nifedipine group and 2.6% in propranolol group). Intention to treat analysis not reported.*
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- 11 (g) *95% CI includes no effect and the upper and lower CI crosses the MID.*
- 12 (h) *Kawanishi 1992[26]: Randomised. Double blind. Allocation concealment not reported. Baseline comparisons made. Drop out < 20% (2.8% in nifedipine group and 2.6% in propranolol group). Intention to treat analysis not reported; Savonitto 1996[27]: Randomised. Double blind. Allocation concealment not reported. Baseline comparison made. Drop out <20% (11%). Intention to treat analysis not reported.*
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- 14
- 15 (i) *Randomised. Double blind. Allocation concealment not reported. Baseline comparisons made. Drop out < 20% (2.8% in nifedipine group and 2.6% in propranolol group). Intention to treat analysis not reported.*
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- 17 (j) *Randomised. Double blind. Allocation concealment not reported. Baseline comparison made. Drop out <20% (11%). Intention to treat analysis not reported.*
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**Drug dosages in each study:**

1. Pehrsson 2000[33] - amlodipine 10 mg once daily, atenolol 100 mg once daily.
2. Dargie 1996[31] (TIBET) - atenolol 50 mg twice daily, nifedipine (slow release) 20-40 mg twice daily
3. O' Hara[25] - 1987 - diltiazem 360 mg once daily, propranolol 240 mg once daily
4. Savonitto 1996[27] (IMAGE)- metoprolol (controlled release, 200 mg once daily), nifedipine (Retard, 20 mg tablets twice daily)
5. Kawanishi 1992[26] - nifedipine 10 mg four times daily, propranolol 20 mg four times daily (not specified if it is long or short acting nifedipine)
6. Tweddel 1981[40] - nifedipine 10 mg three times daily, propranolol dose not reported.

1

2 **8.3.3 Economic evidence**

3 No economic studies were identified on this question. For drug cost of adding CCB see  
4 8.2.3.

5

6 **8.3.4 Evidence statements**

**Clinical**                      **CCB vs. BB+CCB**

**Clinical efficacy:**

**O'Hara 1987[25], Kawanishi 1992[26]:** Evidence from 2 RCTs shows that exercise time (min) [MD -1.91 (-2.87 to -0.95)] was significantly higher with BB+CCB (propranolol +diltiazem, propranolol+ nifedipine) compared to CCB (diltiazem, nifedipine) (follow-up 18 weeks-6 months).

**Savonitto 1996[27] (IMAGE):** Evidence from one RCT shows that time to 1mm ST segment depression (sec) [MD -70 (-125.13 to -14.87)] was significantly higher in the BB+CCB (metoprolol +nifedipine) compared to CCB (nifedipine) (follow-up 10 weeks).

**Dargie 1996[31] (TIBET):** Evidence from one RCT shows that there was no significant difference between CCB (nifedipine) and BB+CCB (atenolol +nifedipine) for cardiac death [RR 1.45 (0.41 to 5.06)] (follow-up mean 2 years).

**Dargie 1996[31] (TIBET):** Evidence from one RCT shows that there was no significant difference between CCB (nifedipine) and BB+CCB (atenolol +nifedipine) for non fatal MI [RR 2.07 (0.86 to 4.98)] (follow-up mean 2 years).

**Kawanishi 1992[26]:** Evidence from one RCT shows that there was no significant difference between CCB (nifedipine) and BB+CCB (propranolol +nifedipine) for time to onset of angina (min) [MD -0.5 (-1.93 to 0.93)] (follow-up 6 months).

**Kawanishi 1992[26], Savonitto 1996[27] (IMAGE):** Evidence from 2 RCTs shows that there was no significant difference between CCB (nifedipine) and BB+CCB (propranolol +nifedipine, metoprolol +nifedipine) for angina episodes/week [MD 0.1 (-1.62 to 1.82)] (follow-up 10 weeks-6 months).

**Kawanishi 1992[26]:** Evidence from one RCT shows that there was no significant difference between CCB (nifedipine) and BB+CCB (propranolol +nifedipine) for no. of nitroglycerin tablets/week [MD -0.4 (-1.66 to 0.86)] (follow-up 6 months).

**Adverse effects:**

**Dargie 1996[31] (TIBET):** Evidence From one RCT shows that there were significantly more withdrawals due to side effects [RR 1.4 (1.08 to 1.82)] in CCB (nifedipine) compared to BB+CCB group (atenolol +nifedipine) (follow-up mean 2 years).

**Pehrsson 2000[33]:** Evidence from one RCT shows that there was no significant difference between CCB (amlodipine) and BB+CCB (atenolol +amlodipine) for adverse effects (overall) [RR 1.04 (0.81 to 1.34)] (follow-up 10 weeks).

**Economic**

No economic evidence was found on this question. A simple cost analysis showed a small increase in drug costs when a CCB is added to the therapy.

1 **8.4 Addition of CCB to basic (or standard) anti-anginal treatment**

2 **8.4.1 Clinical question**

3 What is the comparative clinical /cost effectiveness of adding CCB to basic (or  
4 standard) anti-anginal treatment for the management of angina?

5 **8.4.2 Clinical Evidence**

6 The ACTION trial reports the effects of adding CCB nifedipine to usual anti-anginal  
7 treatment. Although not designed to specifically examine the question of the addition  
8 of CCB to BB the GDG considered that the trial should be included as significant  
9 proportions of patients (80%) were on a BB and the trial provided useful information  
10 on long term safety of CCBs. The information from this trial also influenced the GDG  
11 in their consideration of the use of three anti-anginal drugs.

1 **Table 8.4: CCB +basic regimen vs. placebo +basic regimen**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							CCB +basic regimen	Placebo +basic regimen	Relative (95% CI)	Absolute	
<b>All cause mortality (follow-up mean 4.9 patient-years)</b>											
Poole-Wilson 2004[41] (ACTION) (b,d)	randomised trial	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	310/3825 (8.1%)	291/3840 (7.6%)	RR 1.07 (0.92 to 1.25)	5 more per 1000 (from 6 fewer to 19 more)	⊕⊕⊕⊕ HIGH
<b>Cardiovascular or unknown death (follow-up mean 4.9 years)</b>											
Poole-Wilson 2004[41] (ACTION)	randomised trial	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	178/3825 (4.7%)	177/3840 (4.6%)	RR 1.01 (0.82 to 1.24)	0 more per 1000 (from 8 fewer to 11 more)	⊕⊕⊕⊕ HIGH
<b>MI (follow-up mean 4.9 years)</b>											
Poole-Wilson 2004[41] (ACTION)	randomised trial	no serious limitations (a)	no serious inconsistency	no serious indirectness	serious imprecision (c)	none	320/3825 (8.4%)	296/3840 (7.7%)	RR 1.09 (0.93 to 1.26)	7 more per 1000 (from 5 fewer to 20 more)	⊕⊕⊕○ MODERATE
<b>Withdrawal due to adverse effects (follow-up mean 4.9 years)</b>											
Poole-Wilson 2004[41] (ACTION)	randomised trial	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	389/3825 (10.2%)	172/3840 (4.5%)	RR 2.27 (1.91 to 2.7)	57 more per 1000 (from 41 more to 77 more)	⊕⊕⊕⊕ HIGH
<b>Combined outcome (death from any cause, acute MI, refractory angina, new overt heart failure, debilitating stroke and peripheral revascularisation) (subgroup age &gt;65yrs) (follow-up mean 4.9 years)</b>											
Poole-Wilson 2004[41] (ACTION)	randomised trial	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	467/1772 (26.4%)	466/1776 (26.2%)	RR 1 (0.9 to 1.12)	0 fewer per 1000 (from 26 fewer to 31 more)	⊕⊕⊕⊕ HIGH
<b>Combined outcome (death from any cause, acute MI, refractory angina, new overt heart failure, debilitating stroke and peripheral revascularisation) (subgroup females) (follow-up mean 4.9 years)</b>											
Poole-Wilson 2004[41] (ACTION)	randomised trial	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	166/784 (21.2%)	147/797 (18.4%)	RR 1.15 (0.94 to 1.4)	28 more per 1000 (from 11 fewer to 74 more)	⊕⊕⊕⊕ HIGH
<b>Combined outcome (death from any cause, acute MI, refractory angina, new overt heart failure, debilitating stroke and peripheral revascularisation) (subgroup diabetes) (follow-up mean 4.9 years)</b>											
Poole-Wilson 2004[41] (ACTION)	randomised trial	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	164/565 (29%)	170/545 (31.2%)	RR 0.93 (0.78 to 1.11)	22 fewer per 1000 (from 69 fewer to 34 more)	⊕⊕⊕⊕ HIGH

Combined outcome (death from any cause, acute MI, refractory angina, new overt heart failure, debilitating stroke and peripheral revascularisation) (subgroup age <65 years) (follow-up mean 4.9 years)											
Poole-Wilson 2004[41] (ACTION)	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	337/2053 (16.4%)	362/2064 (17.5%)	RR 0.94 (0.82 to 1.07)	11 fewer per 1000 (from 32 fewer to 12 more)	⊕⊕⊕⊕ HIGH
Combined outcome (death from any cause, acute MI, refractory angina, new overt heart failure, debilitating stroke and peripheral revascularisation) (sub group males) (follow-up mean 4.9 years)											
Poole-Wilson 2004[41] (ACTION)	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	638/3041 (21%)	681/3043 (22.4%)	RR 0.94 (0.85 to 1.03)	13 fewer per 1000 (from 34 fewer to 7 more)	⊕⊕⊕⊕ HIGH
Combined outcome (death from any cause, acute MI, refractory angina, new overt heart failure, debilitating stroke and peripheral revascularisation) (sub group no diabetes) (follow-up mean 4.9 years)											
Poole-Wilson 2004[41] (ACTION)	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	640/3260 (19.6%)	658/3295 (20%)	RR 0.98 (0.89 to 1.08)	4 fewer per 1000 (from 22 fewer to 16 more)	⊕⊕⊕⊕ HIGH

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- (a) Sample size calculation reported. Baseline comparison made. Allocation concealment reported. Blocked randomisation. Double blind. Drop-out <20% (12.8% in the nifedipine group and 12.2% in the placebo group). Intention to treat analysis reported.
- (b) Drug dosage: nifedipine GITS 30 mg once daily, increasing to 60 mg once daily within 6 weeks if no evidence of intolerance seen.
- (c) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.
- (d) Concomitant treatments at baseline:
  - Anti anginal drug: nifedipine +Basic regimen (n=3825): placebo +basic regimen (n=3840)
  - B-blocker- 3032 (79%): 3066 (80%)
  - Organic nitrate, as needed- 2157 (56%): 2175 (57%)
  - Organic nitrate, daily maintenance- 1455 (38%): 1417 (37%)
  - Other vasodilator- 158 (4%): 148 (4%)
  - Any of the above- 3775 (99%):3784 (99%)
  - Any two of the above- 1888 (49%): 1960 (51%)
  - Any three or four of the above- 563 (15%): 520 (14%)
  - Lipid lowering:
    - Statin- 2409 (63%): 2389 (62%)
    - Fibrate 242 (6%): 246 (6%)
    - Other- 45 (1%): 68 (2%)
    - Any of the above- 2607 (68%): 2591 (67%)
  - Blood pressure lowering:
    - ACE inhibitor – 771 (20%): 792 (21%)
    - Angiotensin II antagonist- 90 (2%):93 (2%)

## DRAFT FOR CONSULTATION

1                    *Diuretic – 432 (1%): 447 (12%)*  
2                    *Other- 113 (3%): 81 (2%)*  
3                    *Any of the above- 1165 (30%): 1166 (30%)*  
4

5                    **Subgroup interaction:**

6                    *There was no significant difference between sub group age > 65 years and > 65 years for combined outcomes [p=0.42]*

7                    *There was no significant difference between sub group males and females for combined outcomes [p=0.070]*

8                    *There was no significant difference between subgroup diabetes and no diabetes for combined outcomes [p=0.59]*

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2 **8.4.3 Economic evidence**

3 No economic studies were identified on this question. For drug cost of adding CCB see  
4 8.2.3.

5 **8.4.4 Evidence statements**

**Clinical** **Addition of CCB to basic (or standard) anti-anginal treatment**

**Clinical efficacy:**

**Poole-Wilson 2004[41] (ACTION):** Evidence from one RCT shows that there was no significant difference between CCB and placebo when added to usual anti-anginal treatment for all cause mortality [RR 1.07 (0.92 to 1.25)], cardiovascular or unknown death [RR 1.01 (0.82 to 1.24)] and MI [RR 1.09 (0.93 to 1.26)] (follow-up mean 4.9 patient-years).

**Poole-Wilson 2004[41] (ACTION):** Evidence from one RCT shows that there was no significant difference between CCB and placebo when added to usual treatment for combined outcomes (death from any cause, acute MI, refractory angina, new overt heart failure, debilitating stroke and peripheral revascularisation) for subgroup of patients >65 yrs [RR 1 (0.9 to 1.12)] sub group patients <65 yrs [RR 0.94 (0.82 to 1.07) ], sub group of female patients [RR 1.15 (0.94 to 1.4)], subgroup male patients [RR 0.94 (0.85 to 1.03)], and subgroup of diabetic patients [RR 0.93 (0.78 to 1.11)] and people with no diabetes [RR 0.98 (0.89 to 1.08)] (follow-up mean 4.9 patient-years).

Sub group interaction: There was no significant interaction between the rate of the combined outcome in the two treatment groups and age >65 years [p=0.42], gender [p=0.070], or presence of diabetes [p=0.59].

**Adverse effects:**

**Poole-Wilson 2004[41] (ACTION):** Evidence from one RCT shows that there was significantly more withdrawal due to adverse effects in the CCB group compared to placebo [RR 2.27 (1.91 to 2.7)] (follow-up mean 4.9 patient-years).

**Economic**

No economic evidence was found on this question. A simple cost analysis showed a small increase in drug costs when a CCB is added to the therapy.

1 **8.5 Recommendations and link to evidence**

<b>Recommendation</b>	<p><b>If the person’s symptoms are not controlled, consider either switching to the other option (calcium channel blocker or beta blocker)* or using a combination of the two**.</b></p> <p><b>Do not routinely offer anti-anginal drugs other than beta blockers or calcium channel blockers as first-line treatment for stable angina.</b></p> <p><i>* Evidence on the use of BBs or CCBs as monotherapy, is presented in chapter 7</i></p> <p><i>**When combining a calcium channel blocker with a beta blocker, a dihydropyridine calcium channel blocker should be used</i></p>
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**Relative values of different outcomes**

Outcomes of interest included long-term mortality (total and cardiovascular) and rates of major adverse cardiovascular events (myocardial infarction, stroke, myocardial revascularisation). Additional outcomes of interest included measures of symptom severity (frequency of angina, exercise test outcomes).

**Trade off between clinical benefits and harms**

There is no evidence of a difference in cardiac mortality or rate of non-fatal myocardial infarction between patients treated with the combination of BB and CCB compared with BB or CCB alone.

There is evidence that during exercise testing the combination of BB and CCB increases exercise time and time to 1 mm ST segment depression in the short term when compared with BB or CCB alone. This beneficial effect of combination treatment was not matched by evidence of improved symptom control, as assessed by the frequency of episodes of angina and the use of nitroglycerine.

One trial reported more treatment withdrawals amongst patients treated with nifedipine and atenolol compared with patients treated with atenolol. One other trial reported no difference in the rate of adverse events amongst patients treated with amlodipine versus patients treated with amlodipine and atenolol in combination. Overall there was no consistent evidence of an increased risk of adverse events amongst patients treated with combination therapy when compared with BB or CCB alone.

The GDG concluded that evidence that combination therapy with a BB and a CCB is superior to a BB or CCB alone is weak, and mainly confined to a modest increase in exercise time

during formal exercise testing.

**Economic considerations**

No economic evidence on the use of BBs in combination with CCBs versus CCBs or BBs alone for the first-line treatment of stable angina was available for review.

**Quality of evidence**

Trials comparing the combination of BB and CCB with BB or CCB alone were relatively small with limited statistical power to detect differences in mortality or other major adverse clinical outcomes and only short-term follow-up data were available.

No economic evidence was available.

**Other considerations**

The GDG concluded that there is no evidence to recommend addition of a BB to a CCB, or CCB to a BB for patients whose symptoms are controlled on one drug alone. There is some evidence of short-term improvement in exercise tolerance with combination therapy and the GDG considered that patients not controlled on one drug class should be offered a change to the other drug class, or combination therapy with both drug classes. A dihydropyridine CCB should be used when a CCB is combined with a BB.

<b>Recommendation</b>	<p><b>Do not offer a third anti-anginal drug* to people whose stable angina is controlled with two anti-anginal drugs.</b></p> <p><b>Consider adding a third anti-anginal drug* when:</b></p> <ul style="list-style-type: none"> <li>• <b>the person's symptoms are not controlled with two anti-anginal drugs and</b></li> <li>• <b>the person is waiting for revascularisation or it is not appropriate or acceptable.</b></li> </ul> <p><b>Decide which drug* to use based on comorbidities, contraindications, the person's preference and costs.</b></p> <p><i>*These recommendations also draw on the evidence reviews of nicorandil, ranolazine and ivabradine</i></p>
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**Relative values of different outcomes**

Outcomes of interest included long-term mortality (total and cardiovascular), rates of major adverse cardiovascular events (myocardial infarction, stroke, myocardial revascularisation), and measures of symptom severity (frequency of angina, exercise test outcomes).

**Trade off between clinical benefits and harms**

We found no evidence that directly addressed the use of three classes of anti-anginal drug (BB, CCB, long acting nitrate, or a new anti-anginal drug [nicorandil, ivabradine, ranolazine]) in

combination (versus one or two classes of anti-anginal drug) in people with stable angina.

In one large trial (ACTION) there was no evidence that the addition of long-acting nifedipine GITS to standard anti-anginal treatment (with BB and/or organic nitrate) reduces the risk of death or myocardial infarction in people with stable angina. There is no evidence of an advantage of nifedipine GITS in women, people aged over 65 years, or people with diabetes.

Impact of nifedipine GITS on symptoms of angina was not reported, but nifedipine GITS was associated with a lower rate of coronary arteriography and coronary artery bypass surgery than placebo.

Treatment withdrawal due to adverse effects was increased with nifedipine GITS.

The GDG concluded that routine addition of nifedipine GITS to standard anti-anginal treatment (with BB and/or long-acting nitrate) does not confer any major clinical benefit.

**Economic considerations**

There is no clinical evidence that adding a third drug to standard antianginal treatment generates any clinical benefit. It could therefore increase costs with no additional benefit.

**Quality of evidence**

There was no evidence on the use of three classes of anti-anginal drug in people with stable angina. A large high quality randomised controlled trial provided evidence on the use of nifedipine GITS in addition to treatment with BB and/or long-acting nitrate.

**Other considerations**

The GDG concluded that there is no evidence that routine addition of a third class of antianginal drug provides benefit in people with stable angina already treated with two classes of antianginal drug. The GDG did not consider it appropriate that patients should have repeated trials of different antianginal drug combinations given the lack of evidence for use of more than two drugs.

The GDG considered that a therapeutic trial of a third class of anti-anginal drug could be considered in people with stable angina whose symptoms are not controlled by two classes of anti-anginal drug, and when awaiting revascularisation or when revascularisation is not appropriate or desirable. The response to the addition of a third class of antianginal drug should be reviewed after 2-4 weeks and the drug should be continued only if the person's angina is improved.

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## 2 **9 Long acting nitrates**

### 3 **9.1 Introduction**

4 Long-acting organic nitrates are indicated for the prophylaxis and treatment of  
5 angina. The therapeutic effects of organic nitrates are mediated through dilatation of  
6 capacitance veins and conductive coronary and peripheral arteries. These  
7 haemodynamic changes reduce ventricular preload, and to a lesser extent ventricular  
8 afterload, thereby lowering myocardial oxygen demand and improving  
9 subendocardial blood flow.

10 In many people with stable angina continuous use of organic nitrates induces  
11 tolerance, with reduced therapeutic effect. Tolerance can be avoided by a nitrate-  
12 free interval each day, but this may lower the threshold for episodes of angina. The  
13 pathophysiology of tolerance is incompletely understood but continuous treatment with  
14 organic nitrates causes sympathetic activation, increases oxidative stress, and induces  
15 endothelial dysfunction. Other unwanted effects of nitrates include flushing, headache,  
16 and postural hypotension. Phosphodiesterase type 5 inhibitors should not be used  
17 within 24 hours of long acting nitrate administration because of the risk of severe  
18 hypotension.

19 The GDG were interested in whether there was evidence for the addition of a long-  
20 acting nitrate to treatment with a BB or CCB.

#### 21 **9.1.1 Clinical question**

22 What is the clinical and cost effectiveness of adding long acting nitrates to BB and/or  
23 CCBs?

#### 24 **9.1.2 Clinical evidence**

25 The “Review Protocol” for this topic can be found in Appendix C, the “Search  
26 Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
27 E1, the “Clinical Evidence Tables” in Appendix E2, and the “Forest Plots” in Appendix  
28 F.

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1 **Table 9.1: BB+nitrates vs. BB+CCB for stable angina**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							BB+nitrates	BB+CCB	Relative (95% CI)	Absolute	
<b>Exercise time (sec) (follow-up 12 weeks; better indicated by higher values)</b>											
de Vries[42] 1994 (d)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	None	46	46	-	MD 10 lower (41.14 lower to 21.14 higher)	⊕⊕○○ LOW
<b>Time to onset of angina (sec) (follow-up 12 weeks; better indicated by higher values)</b>											
de Vries[42] 1994	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	None	46	46	-	MD 31 lower (78.08 lower to 16.08 higher)	⊕⊕○○ LOW
<b>Time to ST segment depression (sec) (better indicated by higher values)</b>											
de Vries[42] 1994	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	None	46	46	-	MD 47 lower (102.4 lower to 8.4 higher)	⊕⊕○○ LOW
<b>Adverse effects (overall) (follow-up 12 weeks) (d)</b>											
de Vries[42] 1994	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (c)	None	22/46 (47.8%)	14/43 (32.6%)	RR 1.47 (0.87 to 2.48)	153 more per 1000 (from 42 fewer to 482 more)	⊕⊕○○ LOW
<b>Stopping treatment due to adverse events (follow-up 12 weeks)</b>											
de Vries[42] 1994	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (c)	None	8/46 (17.4%)	2/43 (4.7%)	RR 3.74 (0.84 to 16.64)	127 more per 1000 (from 7 fewer to 727 more)	⊕⊕○○ LOW
<b>Headache (follow-up 12 weeks)</b>											
de Vries[42] 1994	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (c)	None	10/46 (21.7%)	4/43 (9.3%)	RR 2.34 (0.79 to 6.9)	125 more per 1000 (from 20 fewer to 549 more)	⊕⊕○○ LOW

- (a) Randomised, double blind, cross over, single centre, sample size calculation reported, 4/46 (8.6%) lost to follow-up. Allocation concealment not reported, Intention to treat analysis not reported.
- (b) 95% CI includes no effect and the upper and lower CI crosses the MID.
- (c) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.
- (d) Drug dosages: felodipine extended release 5 mg daily, isosorbide mononitrate 10 mg or 20 mg thereafter twice daily, optimal B-blockade, fixed dose (exact dose not reported).
- (e) Adverse events: headache, peripheral oedema, tiredness, cerebrovascular disorder, flushing.

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1 **Table 9.2: BB+nitrates vs. BB+CCB for stable angina** (Results from one study - data reported graphically .Data reported as in the text of the paper.

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							BB+nitrates	BB+CCB	Relative (95% CI)	Absolute	
<b>Anginal attacks (Follow-up 15 weeks)</b>											
Morse 1985[43] (b)	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	27	27	Data reported graphically	BB+nitrates and BB+CCB resulted in significant reduction in anginal frequency. BB+CCB were superior to BB+nitrates in reducing the frequency of angina episodes. (p=0.03)	LOW
<b>Nitroglycerin consumption (follow-up 15 weeks)</b>											
Morse 1985[43]	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	27	27	Data reported graphically	No sig. difference for nitroglycerin consumption between BB+CCB and BB+nitrates	LOW
<b>Total Exercise time (sec) (follow-up 15 weeks)</b>											
Morse 1985[43]	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	27	27	Data reported graphically	BB+CCB resulted in significant increase in total exercise time compared to BB+nitrates (p<0.02)	LOW
<b>Time to onset of pain (sec) (follow-up 15 weeks)</b>											
Morse 1985[43]	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	27	27	Data reported graphically	Time to onset of angina was significantly prolonged in BB+CCB compared to BB+nitrates (p=0.003)	LOW

- 2 (a) Randomised cross over, double blind, drop out 10%, small sample size, allocation concealment not reported, intention to treat analysis not reported, data cannot be  
3 analysed as results reported graphically.  
4 (b) Drug dosages: nifedipine was 77.0 mg/day, isosorbide mononitrate 90.4 mg/day, propranolol median dose was 120 mg/day (range 60 to 240.

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2 **9.1.3 Economic evidence**

3 No economic studies were identified on this question. We calculated the range (low and  
4 high) of daily and annual cost of adding long-acting nitrates based on the unit cost  
5 reported in the BNF59[18].

6 **Table 9.3: Cost of adding long-acting nitrates**

	Specific drugs used for cost range	Cost per day (£)	Cost per year (£)
Long-acting nitrates	Low = isosorbide mononitrate	0.052	19
	High = glyceryl nitrate	0.55	201

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8 The cost of adverse effects was not estimated.

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10 **9.1.4 Evidence statements****Clinical** **Addition of nitrates****Clinical efficacy:**

**De Vries 1994[42]:** Evidence from one RCT shows that there was no significant difference between BB+nitrates and BB+CCB for exercise time (sec) [MD -10 (-41.14 to 21.14)], time to onset of angina (sec) [MD -31 (-78.08 to 16.08)] and time to ST segment depression (Sec) [MD -47 (-102.4 to 8.4)] (follow-up 12 weeks).

**Adverse effects:**

**De Vries 1994[42]:** Evidence from one RCT shows that there was no significant difference between BB+nitrates and BB+CCB for adverse effects overall [RR 1.47 (0.87 to 2.48)], stopping due to adverse events [RR 3.74 (0.84 to 16.64)] and headache [RR 2.34 (0.79 to 6.9)] (follow-up 12 weeks).

**Economic** No economic evidence was found on this question. A simple cost analysis showed the annual cost of adding long-acting nitrates to range between £19 and £201.

1 9.1.5 Recommendations and link to evidence

<b>Recommendation</b>	<p><b>If the person cannot tolerate beta blockers and calcium channel blockers or they are contraindicated, consider monotherapy with one of the following drugs*:</b></p> <ul style="list-style-type: none"> <li>• a long-acting nitrate</li> <li>• ivabradine</li> <li>• nicorandil** or</li> <li>• ranolazine.</li> </ul> <p><b>Decide which drug to use based on comorbidities, contraindications, the person's preference and costs.</b></p> <p><b>For people on beta blocker or calcium channel blocker monotherapy whose symptoms are not controlled and the other option (calcium channel blocker or beta blocker) is contraindicated or not tolerated, consider one of the following as an additional drug*:</b></p> <ul style="list-style-type: none"> <li>• a long-acting nitrate</li> <li>• ivabradine***</li> <li>• nicorandil** or</li> <li>• ranolazine</li> </ul> <p><b>Decide which drug to use based on comorbidities, contraindications, the person's preference and costs.</b></p> <p><i>* Evidence on ivabradine, nicorandil and ranolazine is presented in chapter 10 (Other anti-anginal drugs)</i></p> <p><i>**At the time of consultation (December, 2010), nicorandil did not have UK marketing authorisation for use in this indication. Informed consent should be obtained and documented</i></p> <p><i>*** Ivabradine should only be combined with a dihydropyridine CCB</i></p>
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**Relative values of different outcomes**

Outcomes of interest included long-term mortality (total and cardiovascular), rates of major adverse cardiovascular events (myocardial infarction, stroke, myocardial revascularisation), and measures of symptom severity (frequency of angina, exercise test outcomes).

**Trade off between clinical benefits and harms**

We found no evidence to confirm the safety or efficacy of long-term use of organic nitrate as an additional anti-anginal agent in patients already taking a BB or CCB.

In patients aged over 65 years addition of either isosorbide mononitrate or felodipine to treatment with a BB had comparable short-term effects on exercise time, and time to onset of angina or ST segment depression during exercise stress testing. There was no difference in adverse effects of treatment between the two groups.

In one small study of poor quality the combination of propranolol and nifedipine resulted in greater reduction in angina frequency and longer exercise times than the combination of isosorbide dinitrate and propranolol.

**Economic considerations**

No economic evidence on the use of long-acting organic nitrates for the treatment of stable angina was available for review. The drug cost ranges from £19 to £201 per year.

**Quality of evidence**

Evidence to support the use of long-acting nitrates in combination with BB in people with stable angina is of poor quality and available trials are limited by small sample size and short duration of follow-up.

No trials of nitrates in combination with CCBs were identified.

No economic evidence was available.

**Other considerations**

The GDG concluded that evidence to support the addition of long-acting nitrate to monotherapy with BB or CCB in people with stable angina is very weak.

The GDG recognized that organic nitrates have been used for the relief of attacks of angina for over 100 years. In addition there is consensus that monotherapy with long-acting nitrates is effective in the treatment of stable angina in the short-term, but that the efficacy of long acting nitrates may be limited by the development of tolerance.

The GDG made a consensus recommendation that long acting nitrates can be considered for monotherapy if BBs and CCBs are not tolerated or are contraindicated. The GDG also agreed that addition of a long-acting nitrate can be considered in people whose symptoms are not controlled by monotherapy with either BB or CCB if the combination of BB and CCB is not appropriate.

The cost of long acting nitrate varies widely between different

formulations but is less than the cost of newer antianginal drugs (e.g. nicorandil, ivabradine, ranolazine – see chapter 10).

Nevertheless the GDG concluded that there was insufficient evidence to make a firm recommendation about the choice of antianginal drug as monotherapy or as an additional antianginal drug if a CCB or BB is not tolerated or is contraindicated.

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## 2 **10 Other anti anginal drugs and general drug**

### 3 **recommendations**

#### 4 **10.1 Introduction**

5 Ivabradine, nicorandil, and ranolazine are anti-anginal drugs that are licensed for  
6 use in the treatment of stable angina. The GDG were interested in evidence for the  
7 use of these drugs either as monotherapy or in combination with other anti-anginal  
8 drugs, and their place in the pathway for people with stable angina.

9 Ivabradine is licensed for the treatment of angina in patients in sinus rhythm in  
10 combination with a BB, or when a BB is contra-indicated or not tolerated. Nicorandil  
11 has been available for longer than the other drugs considered in this chapter and  
12 although it does not have a licence to be used in combination with other antianginal  
13 drugs it is regularly used this way in practice. Ranolazine is licensed as adjunctive  
14 therapy in patients who are inadequately controlled or intolerant of first-line  
15 antianginal drugs.

16 The costs of drugs at standard doses are listed below and compared with the cost of  
17 long acting nitrates.

18 **Table 10: Cost of drugs**

	Specific drugs used for range	Cost per day (£)	Cost per year (£)
Long-acting nitrates	Low = isosorbide mononitrate	0.05	19
	High = glyceryl nitrate	0.55	201
Ivabradine, 5 mg or 7.5 mg twice daily	Low = high	1.39	507
Ranolazine, 375 mg, 500 mg or 750 mg twice daily	Low = high	1.63	595

Nicorandil	Low = 10 mg tablets twice daily	0.27	99
	High = 20 mg tablets twice daily	0.52	190

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3 **10.2 Ivabradine**

4 Ivabradine is a heart rate-lowering agent that acts by selectively inhibiting the If  
5 current, an ionic current across the sarcolemma in cells of the sino-atrial node that is  
6 involved in pacemaker activity. Ivabradine reduces the slope of spontaneous diastolic  
7 depolarization in sino-atrial cells, and lowers heart rate at rest and during exercise.  
8 Side effects of ivabradine include bradycardia, heart block, and visual disturbances  
9 (phosphenes and blurred vision).

10 **10.2.1 Clinical question**

11 What is the clinical /cost effectiveness of ivabradine for the management of stable  
12 angina?

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14 **10.2.2 Clinical evidence**

15 The “Review Protocol” for this topic can be found in Appendix C, the “Search  
16 Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
17 E1, the “Clinical Evidence Tables” in Appendix E2, and the “Forest Plots” in Appendix  
18 F.

19 The evidence review included evidence for the use of ivabradine as monotherapy or  
20 in combination with BB to control symptoms and improve outcome in people with  
21 stable angina.

1 **Table 10.1: Ivabradine vs. placebo**

Quality assessment							Summary of findings				
							No of patients		Effect		Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Ivabradine	placebo	Relative (95% CI)	Absolute	
<b>Time to angina onset (sec) (trough change from baseline) (follow-up 14 days; better indicated by higher values) (g)</b>											
Borer 2003[44] (e)	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	serious (b)	None	59	68	-	MD 14.1 higher (11.73 lower to 39.93 higher)	⊕⊕⊕O MODERATE
<b>Time to angina onset (sec) (peak change from baseline) (follow-up 14 days; better indicated by higher values) (h)</b>											
Borer[44] 2003	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	serious (b)	None	59	68	-	MD 43.2 higher (16.75 to 69.65 higher)	⊕⊕⊕O MODERATE
<b>Time to 1mm ST depression (sec) (at peak of drug activity) (follow-up 14 days; better indicated by higher values)</b>											
Borer[44] 2003	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	serious (b)	None	59	68	-	MD 52.90 higher (26.85 to 78.95 higher)	⊕⊕⊕O MODERATE
<b>Time to 1mm ST depression (sec) (at trough) (follow-up 14 days; better indicated by higher values)</b>											
Borer[44] 2003	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	serious (b)	None	59	68	-	MD 35.10 higher (9.68 to 60.52 higher)	⊕⊕⊕O MODERATE
<b>Patients with limiting angina (j) - CV death or hospitalisation for MI or HF - (follow-up median 18 months)</b>											
Fox 2009[45] (BEAUTIFUL) (f)	randomised trials	no serious limitation (c)	no serious inconsistency	no serious indirectness	serious (d)	None	88/734 (12%)	120/773 (15.5%)	RR 0.77 (0.6 to 1)	36 fewer per 1000 (from 62 fewer to 0 more)	⊕⊕⊕O MODERATE
<b>Patients with limiting angina - all cause mortality - (follow-up median 18 months)</b>											
Fox 2009[45] (BEAUTIFUL)(i)	randomised trials	no serious limitation (c)	no serious inconsistency	no serious indirectness	serious (d)	None	64/734 (8.7%)	77/773 (10%)	RR 0.88 (0.64 to 1.2)	12 fewer per 1000 (from 36 fewer to 20 more)	⊕⊕⊕O MODERATE
<b>Patients with limiting angina - cardiac death - (follow-up median 18 months)</b>											
Fox 2009[45] (BEAUTIFUL)	randomised trials	no serious limitation (c)	no serious inconsistency	no serious indirectness	serious (d)	None	11/734 (1.5%)	16/773 (2.1%)	RR 0.72 (0.34 to 1.55)	6 fewer per 1000 (from 14 fewer to 11 more)	⊕⊕⊕O MODERATE
<b>Patients with limiting angina - hospitalisation for HF - (follow-up median 18 months)</b>											
Fox 2009[45] (BEAUTIFUL)	randomised trials	no serious limitation (c)	no serious inconsistency	no serious indirectness	serious (d)	None	33/734 (4.5%)	41/773 (5.3%)	RR 0.85 (0.54 to 1.33)	8 fewer per 1000 (from 24 fewer to 18 more)	⊕⊕⊕O MODERATE
<b>Patients with limiting angina - hospitalisation for MI or unstable angina - (follow-up median 18 months)</b>											

Fox 2009[45] (BEAUTIFUL)	randomised trials	no serious limitation (c)	no serious inconsistency	no serious indirectness	serious (d)	None	56/734 (7.6%)	65/773 (8.4%)	RR 0.9 (0.64 to 1.28)	8 fewer per 1000 (from 30 fewer to 24 more)	⊕⊕⊕○ MODERATE
<b>Patients without limiting angina - CV death or hospitalisation for MI or heart failure (follow-up 18 months)</b>											
Fox 2009[45] (BEAUTIFUL)	randomised trials	no serious limitation (c)	no serious inconsistency	no serious indirectness	no serious imprecision	None	756/4745 (15.9%)	712/4665 (15.3%)	RR 1.04 (0.95 to 1.15)	6 more per 1000 (from 8 fewer to 23 more)	⊕⊕⊕⊕ HIGH
<b>Patients without limiting angina - all cause mortality (follow-up median 18 months)</b>											
Fox 2009[45] (BEAUTIFUL)	randomised trials	no serious limitation (c)	no serious inconsistency	no serious indirectness	no serious imprecision	None	508/4745 (10.7%)	470/4665 (10.1%)	RR 1.06 (0.94 to 1.2)	6 more per 1000 (from 6 fewer to 20 more)	⊕⊕⊕⊕ HIGH
<b>Patients without limiting angina - cardiac death (follow-up median 18 months)</b>											
Fox 2009[45] (BEAUTIFUL)	randomised trials	no serious limitation (c)	no serious inconsistency	no serious indirectness	serious (d)	None	125/4745 (2.6%)	135/4665 (2.9%)	RR 0.91 (0.72 to 1.16)	3 fewer per 1000 (from 8 fewer to 5 more)	⊕⊕⊕○ MODERATE
<b>Patients without limiting angina - hospitalisation for heart failure (follow-up median 18 months)</b>											
Fox 2009[45] (BEAUTIFUL)	randomised trials	no serious limitation (c)	no serious inconsistency	no serious indirectness	no serious imprecision	None	393/4745 (8.3%)	386/4665 (8.3%)	RR 1 (0.87 to 1.15)	0 fewer per 1000 (from 11 fewer to 12 more)	⊕⊕⊕⊕ HIGH
<b>Patients without limiting angina - hospitalisation for MI or unstable angina (follow-up median 18 months)</b>											
Fox 2009[45] (BEAUTIFUL)	randomised trials	no serious limitation (c)	no serious inconsistency	no serious indirectness	no serious imprecision	None	247/4745 (5.2%)	252/4665 (5.4%)	RR 0.96 (0.81 to 1.14)	2 fewer per 1000 (from 10 fewer to 8 more)	⊕⊕⊕⊕ HIGH
<b>All serious adverse events (follow-up median 18 months)</b>											
Fox 2009[45] (BEAUTIFUL)	randomised trials	no serious limitation (c)	no serious inconsistency	no serious indirectness	no serious imprecision	None	135/734 (18.4%)	144/773 (18.6%)	RR 0.99 (0.92 to 1.06)	2 fewer per 1000 (from 37 fewer to 41 more)	⊕⊕⊕⊕ HIGH

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(a) Randomisation, allocation concealment, blinding and ITT reported.

(b) The upper and lower CI crosses the MID.

(c) Randomisation, allocation concealment, blinding and ITT reported.

(d) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.

(e) Ivabradine 5 mg bid vs. placebo

(f) Ivabradine 7.5 mg bid vs. placebo

(g) Trough = 12 hours after administration of ivabradine

(h) Peak = 4 hours after administration of ivabradine

(i) In this post hoc analysis, the BEAUTIFUL population was divided according to the presence of limiting angina symptoms at baseline using the New York Heart Association (NYHA) functional classification. Patients were questioned at the inclusion visit regarding the presence of symptoms limiting activity, and whether they were related to anginal pain or due to presence of heart failure (fatigue, palpitations or dyspnoea).

1 (j) Limiting angina symptoms were identified in 13.8% of the BEAUTIFUL population at baseline (1507 out of 10917 patients). Of these, 734 were randomised to ivabradine  
 2 treatment and 773 to placebo.

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4 **Table 10.2: Ivabradine vs. atenolol**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Ivabradine	Atenolol	Relative (95% CI)	Absolute	
<b>Total exercise duration (sec) (trough change from baseline) (follow-up 16 weeks; better indicated by higher values)</b>											
Tardif 2005[46] (d)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	300	286	-	MD 8 higher (13.26 lower to 29.26 higher)	⊕⊕⊕O MODERATE
<b>Time to angina onset (sec) (trough change from baseline) (follow-up 16 weeks; better indicated by higher values)</b>											
Tardif 2005[46]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	None	300	286	-	MD 10 higher (14.96 lower to 34.96 higher)	⊕⊕OO LOW
<b>Weekly number of angina attacks (follow-up 16 weeks; better indicated by lower values)</b>											
Tardif 2005[46]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	307	294	-	MD 0.5 higher (0.99 lower to 1.99 higher)	⊕⊕⊕O MODERATE
<b>Short-acting nitrate consumption units/week (follow-up 16 weeks; better indicated by lower values)</b>											
Tardif 2005[46]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	307	294	-	MD 0.4 lower (1 lower to 0.2 higher)	⊕⊕⊕O MODERATE
<b>Withdrawal due to adverse events (follow-up 16 weeks)</b>											
Tardif 2005[46]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (c)	None	28/315 (8.9%)	17/307 (5.5%)	RR 1.61 (0.9 to 2.87)	34 more per 1000 (from 6 fewer to 104 more)	⊕⊕OO LOW

- 5 (a) Allocation concealment not reported. Randomisation, blinding and ITT reported.  
 6 (b) 95% CI includes no effect and the upper and lower CI crosses the MID.  
 7 (c) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.  
 8 (d) Ivabradine 5 mg bid for 4 weeks and then 7.5 bid for 12 weeks or atenolol 50 mg od for 4 weeks and then 100 mg od for 12 weeks.  
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**Table 10.3: Ivabradine + atenolol vs. atenolol**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Ivabradine plus atenolol	Atenolol	Relative (95% CI)	Absolute	
<b>Total exercise duration (sec) (change from baseline) (follow-up 2 months; better indicated by higher values) (e)</b>											
Tardif 2009[47] (d)	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	441	434	-	MD 8.7 higher (0.98 to 16.42 higher)	⊕⊕⊕⊕ HIGH
<b>Time to angina onset (sec) (change from baseline) (follow-up 2 months; better indicated by higher values)</b>											
Tardif 2009[47]	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	441	434	-	MD 13 higher (3.43 to 22.57 higher)	⊕⊕⊕⊕ HIGH
<b>Time to 1mm ST depression (sec) (change from baseline) (follow-up 2 months; better indicated by higher values)</b>											
Tardif 2009[47]	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	serious (b)	None	441	434	-	MD 27.2 higher (16.15 to 38.25 higher)	⊕⊕⊕○ MODERATE
<b>Total exercise duration (sec) (change from baseline) (follow-up 4 months; better indicated by higher values)</b>											
Tardif 2009[47]	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	441	434	-	MD 16.6 higher (8.05 to 25.15 higher)	⊕⊕⊕⊕ HIGH
<b>Time to onset of angina(sec) (change from baseline) (follow-up 4 months; better indicated by higher values)</b>											
Tardif 2009[47]	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	serious (b)	None	441	434	-	MD 26.4 higher (15.64 to 37.16 higher)	⊕⊕⊕○ MODERATE
<b>Time to 1 mm ST depression (sec) (change from baseline) (follow-up 4 months; better indicated by higher values)</b>											
Tardif 2009[47]	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	serious (b)	None	441	434	-	MD 30.3 higher (18.4 to 42.2 higher)	⊕⊕⊕○ MODERATE
<b>Adverse events (follow-up 4 months)</b>											
Tardif 2009[47]	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	serious (c)	None	13/441 (2.9%)	4/434 (0.9%)	RR 3.2 (1.05 to 9.73)	20 more per 1000 (from 0 more to 80 more)	⊕⊕⊕○ MODERATE

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- (a) Randomisation, allocation concealment, blinding and ITT reported.
- (b) The upper and lower CI crosses the MID.
- (c) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.
- (d) Patients receiving atenolol 50 mg/day were randomised to receive ivabradine 5 mg b.i.d for 2 months, increased to 7.5 mg b.i.d for a further 2 months.
- (e) 12 hours after last dose ivabradine, 24 hours after last dose atenolol

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**Table 10.4: Ivabradine vs. amlodipine**

Quality assessment							Summary of findings				
							No of patients		Effect		Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Ivabradine	Amlodipine	Relative (95% CI)	Absolute	
<b>Total exercise duration (sec) (follow-up 3 months; better indicated by higher values)</b>											
Ruzylo 2007[48] (c)	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	381	398	-	MD 3.6 lower (16.5 lower to 9.3 higher)	⊕⊕⊕⊕ HIGH
<b>Time angina onset (sec) (follow-up 3 months; better indicated by higher values)</b>											
Ruzylo 2007[48]	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	381	398	-	MD 1.9 lower (16.24 lower to 12.44 higher)	⊕⊕⊕⊕ HIGH
<b>Short-acting nitrate use (units/week) (follow-up 3 months; better indicated by lower values)</b>											
Ruzylo 2007[48]	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	389	398	-	MD 0.8 higher (0.04 to 1.56 higher)	⊕⊕⊕⊕ HIGH
<b>Frequency of angina attacks/week - (follow-up 3 months; better indicated by lower values)</b>											
Ruzylo 2007[48]	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	389	398	-	MD 0 higher (0.77 lower to 0.77 higher)	⊕⊕⊕⊕ HIGH
<b>Adverse events (follow-up 3 months)</b>											
Ruzylo 2007[48]	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	serious (b)	None	181/400 (45.3%)	152/404 (37.6%)	RR 1.2 (1.02 to 1.42)	75 more per 1000 (from 8 more to 158 more)	⊕⊕⊕○ MODERATE

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- (a) Randomisation, allocation concealment, blinding and ITT reported.
- (b) 95% CI around the pooled estimate of effect includes appreciable benefit or appreciable harm.
- (c) Ivabradine 7.5 mg twice daily vs. amlodipine 10 mg once daily

1 **10.2.3 Economic evidence**

2 No economic studies were identified on this question. We calculated the daily and  
3 annual cost of ivabradine treatment based on the unit cost reported in the BNF59[18].

4 **Table 10.5: Drug cost of Ivabradine**

	Cost per day (£)	Cost per year (£)
Ivabradine, 5 mg or 7.5mg twice daily	1.39	507

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6 The costs adverse effects were not estimated.

7 **10.2.4 Evidence statements**

**Clinical Ivabradine versus placebo**

**Borer 2003[44]:** Evidence from one RCT shows that there was no significant difference between ivabradine and placebo for time to angina onset (sec) (at trough) [MD 14.1 (-11.73 to 39.93)]. Time to angina onset (sec) (at peak) [MD 43.2 (16.75 to 69.65)], time to 1 mm ST depression (sec) at peak [MD 52.90 (26.85 to 78.95)], time to 1mm ST depression (sec) at trough [MD 35.10 (9.68 to 60.52)] was significantly higher in the ivabradine group (5 mg) compared to placebo (follow-up 14 days).

**Fox 2009[45]:** Evidence from one RCT shows that there was no statistically significant difference between ivabradine (7.5 mg) and placebo in patients with limiting angina for CV death or hospitalisation for MI or HF [RR 0.77 (0.6 to 1.0)], all cause mortality [RR 0.88 (0.64 to 1.2)], cardiac death [RR 0.72 (0.34 to 1.55)], hospitalisation for heart failure [RR 0.85 (0.54 to 1.33)], hospitalisation for heart failure or unstable angina [RR 0.9 (0.64 to 1.28)].

Evidence from RCT shows that there was no statistically significant difference between ivabradine (7.5 mg) and placebo in patients without limiting angina for CV death or hospitalisation for MI or HF [RR 1.04 (0.95 to 1.15)], all cause mortality [RR 1.06 (0.94 to 1.2)], cardiac death [RR 0.91 (0.72 to 1.16)], hospitalisation for heart failure [RR 1 (0.87 to 1.15)] and hospitalisation for heart failure or unstable angina [RR 0.96 (0.81 to 1.14)] and serious adverse events [RR 0.99 (0.92 to 1.06)] (median follow-up 18 months).

**Ivabradine versus atenolol**

**Tardif 2005[46]:** Evidence from one RCT shows that there was no statistically significant difference between ivabradine (5 mg bid for 4 weeks and then 7.5 mg bid for 12 weeks) and atenolol (50 mg) for total exercise duration at trough (sec) [MD 8.00 (-13.26 to 29.26)], time to

angina onset at trough (sec) [MD 10.00 (-14.96 to 34.96)], weekly number of angina attacks [MD 0.50 (0.99 to 1.99)], short-acting nitrate consumption (units/week) [MD -0.40 (1.00 to 0.20)], and withdrawal due to adverse events [RR1.61 (0.90 to 2.87)] (follow-up 16 weeks).

#### **Ivabradine plus atenolol versus atenolol**

**Tardif 2009[47]** (sub group diabetes): Evidence from one RCT shows that total exercise duration at trough (sec) [MD 8.70 (0.98 to 16.42)], time to angina onset at trough (sec) [MD 13.00 (3.43 to 22.57)] and time to 1mm ST segment depression (sec) [MD 27.2 (16.15 to 38.25)] at 2 months and total exercise duration at trough (sec) [MD 16.6 (8.05 to 25.15)], time to angina onset at trough (sec) [MD 26.4 (15.64 to 37.16)] [and time to 1mm ST segment depression (sec) [MD 30.3 (18.4 to 42.2)] at 4 months was significantly higher in the ivabradine plus atenolol (ivabradine 5 mg b.i.d for 2 months, increased to 7.5 mg b.i.d for a further 2 months) group compared to atenolol.

The rate of adverse events was significantly higher [(RR3.20 (1.05 to 9.73)] in the ivabradine plus atenolol group compared to atenolol alone (follow-up 2 months and 4 months).

#### **Ivabradine versus amlodipine**

**Ruzylo 2007[48]**: Evidence from one RCT shows that there were no statistically significant differences between ivabradine (7.5 mg bid) and amlodipine (10 mg/daily) for total exercise duration at trough (sec) [MD -3.60 (-16.5 to 9.3)], time to angina onset at trough (sec) [MD -1.90 (-16.24 to 12.44)], weekly number of angina attacks [MD 0.0 (-0.77 to 0.77)] or short-acting nitrate consumption (units/week) [MD 0.80 (0.04 to 1.56)]. There was significantly higher risk of adverse events with in the ivabradine group compared with amlodipine (RR1.20 (1.02 to 1.42) (follow-up 12 weeks).

**Economic** No economic evidence was found on this question. A simple cost analysis showed a significant drug cost of ivabradine.

1 10.2.5 Recommendations and link to evidence

<p><b>Recommendation</b></p>	<p><b>If the person cannot tolerate beta blockers and calcium channel blockers or they are contraindicated, consider monotherapy with one of the following drugs*:</b></p> <ul style="list-style-type: none"> <li>• a long-acting nitrate</li> <li>• ivabradine</li> <li>• nicorandil** or</li> <li>• ranolazine</li> </ul> <p><b>Decide which drug to use based on comorbidities, contraindications, the person's preference and costs.</b></p> <p><b>For people on beta blocker or calcium channel blocker monotherapy whose symptoms are not controlled and the other option (calcium channel blocker or beta blocker) is contraindicated or not tolerated, consider one of the following as an additional drug*:</b></p> <ul style="list-style-type: none"> <li>• a long-acting nitrate</li> <li>• ivabradine***</li> <li>• nicorandil** or</li> <li>• ranolazine</li> </ul> <p><b>Decide which drug to use based on comorbidities, contraindications, the person's preference and costs.</b></p> <p><i>* Evidence on long acting nitrates is presented in chapter 9. Evidence on nicorandil and ranolazine is presented in sections 10.3 and 10.4 respectively of this chapter.</i></p> <p><i>**At the time of consultation (December, 2010), nicorandil did not have UK marketing authorisation for this indication. Informed consent should be obtained and documented</i></p> <p><i>*** Ivabradine should only be combined with a dihydropyridine CCB</i></p>
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**Relative values of different outcomes**

Outcomes of interest included long-term mortality (total and cardiovascular), rates of major adverse cardiovascular events (myocardial infarction, stroke, myocardial revascularisation), and measures of symptom severity (frequency of angina, exercise test outcomes).

**Trade off between clinical benefits and harms**

Short-term trials of monotherapy with ivabradine versus monotherapy with atenolol or amlodipine demonstrated similar increases in total exercise duration, and similar reductions in the frequency of angina episodes in both treatment groups.

One short-term trial reported that addition of ivabradine to monotherapy with atenolol resulted in small increases in total exercise duration (16.3s) and time to angina on the treadmill, but did not reduce the frequency of episodes of angina.

These data suggest that ivabradine is an effective anti-anginal agent with comparable short-term efficacy to atenolol and amlodipine. In addition there is a statistically significant incremental benefit of adding ivabradine to atenolol in people with angina, but the magnitude of the benefit is small and of uncertain clinical significance.

There were trends to higher rates of adverse events in ivabradine treated patients, partly due to visual disturbance (phosphenes and blurred vision).

**Economic considerations**

No economic evidence on the use of ivabradine for the treatment of stable angina was available for review. The cost of ivabradine is substantially higher than the costs of other standard treatments (including BB, CCB, and long-acting nitrate).

**Quality of evidence**

The trials assessing the short-term anti-anginal efficacy of ivabradine were relatively large, well-designed studies. Evidence confirming the long-term efficacy and safety of ivabradine is limited.

The BEAUTIFUL trial assessed the effect of ivabradine in people with coronary artery disease and impaired left ventricular function. In a subgroup analysis of patients whose limiting baseline symptom was angina, ivabradine was associated with a reduction in the composite rate of the primary endpoint (cardiovascular death and hospitalization for myocardial infarction or heart failure) of borderline statistical significance. The rate of hospitalisation for myocardial infarction was lower in the ivabradine treated patients (RR 0.58, 95%CI 0.37–0.92, p=0.021)[45]. The subgroup was defined retrospectively, only includes 13.8% of the total trial population, and lacks statistical power for the primary endpoint. The GDG considered this analysis to be exploratory, rather than providing definitive evidence of benefit of ivabradine in people with stable angina and impaired left ventricular systolic function.

No economic evidence was found on this question.

**Other considerations**

There is some evidence for the use of ivabradine as monotherapy or in combination with BB, but no evidence for use of ivabradine in combination with CCB was found. Concomitant use of ivabradine with heart rate reducing CCB such as verapamil or diltiazem is not recommended by the manufacturers.

Ivabradine is a relatively new drug with limited information about long-term safety and efficacy. The cost of ivabradine is comparable with the costs of nicorandil and ranolazine but more than the cost of long-acting nitrate. Nevertheless the GDG considered that there was insufficient evidence to make a firm recommendation about the choice of antianginal drug as monotherapy or as an additional antianginal drug if a CCB or BB is not tolerated or is contraindicated.

The GDG concluded that monotherapy with ivabradine should not be used as an alternative to monotherapy with a BB or CCB. Monotherapy with ivabradine can be considered in people with stable angina in whom BB and CCB are contraindicated or not tolerated.

The GDG concluded that ivabradine can be considered as an additional drug for people whose symptoms are not controlled by monotherapy with a BB and the addition of CCB is contraindicated or not tolerated. Ivabradine should only be combined with dihydropyridine CCB

1 **10.3 Nicorandil**

2 Nicorandil is a nitrate derivative of nicotinamide that is licensed for the prevention  
3 and long-term treatment of angina. Nicorandil is believed to have a dual mechanism  
4 of action. Specifically nicorandil provides a nitrate moiety that dilates epicardial  
5 coronary arteries and systemic venous capacitance vessels. In addition, nicorandil  
6 opens ATP-sensitive potassium channels ( $K_{ATP}$ ) in vascular smooth muscle cells, thereby  
7 dilating arterial resistance vessels in the peripheral and coronary circulations. In  
8 humans nicorandil decreases ventricular filling pressure, coronary vascular resistance,  
9 and mean arterial pressure, and these combined effects increase coronary blood flow  
10 and reduce myocardial work.

11  $K_{ATP}$  channels are an important mediator of ischaemic preconditioning. The molecular  
12 mechanisms have not been fully elucidated but activation of the  $K_{ATP}$  channel has a  
13 cardioprotective effect similar to ischaemic preconditioning, while  $K_{ATP}$  channel  
14 blockade prevents preconditioning. Experimental and clinical studies of myocardial  
15 ischaemia provide evidence that pretreatment with nicorandil reduces ischaemic  
16 myocardial injury. It has therefore been suggested that in addition to relieving  
17 symptoms of ischaemia nicorandil may have a clinically relevant cardioprotective  
18 effect.

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2 **10.3.1 Clinical question**

3 What is the clinical /cost effectiveness of nicorandil for the management of stable  
4 angina?

5 **10.3.2 Clinical evidence**

6 The “Review Protocol” for this topic can be found in Appendix C, the “Search  
7 Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
8 E1, the “Clinical Evidence Tables” in Appendix E2, and the “Forest Plots” in Appendix  
9 F.

1 **Table 10.6: Nicorandil +usual treatment versus Placebo + usual treatment**

Quality assessment							Summary of findings				
							No of patients		Effect		Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Nicorandil	Placebo	Relative (95% CI)	Absolute	
<b>CHD death (follow-up 1.6 years)</b>											
Dargie 2002[49] (IONA) (d)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	None	60/2565 (2.3%)	73/2561 (2.9%)	RR 0.82 (0.59 to 1.15)	5 fewer per 1000 (from 12 fewer to 4 more)	⊕⊕○○ LOW
<b>Non fatal MI (follow-up 1.6 years)</b>											
Dargie 2002[49] (IONA)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	None	56/2565 (2.2%)	72/2561 (2.8%)	RR 0.78 (0.55 to 1.1)	6 fewer per 1000 (from 13 fewer to 3 more)	⊕⊕○○ LOW
<b>Unstable angina (follow-up 1.6 years)</b>											
Dargie 2002[49] (IONA)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	None	115/2565 (4.5%)	127/2561 (5%)	RR 0.9 (0.71 to 1.16)	5 fewer per 1000 (from 14 fewer to 8 more)	⊕⊕○○ LOW
<b>All cause mortality (follow-up 1.6 years)</b>											
Dargie 2002[49] (IONA)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	None	111/2565 (4.3%)	129/2561 (5%)	RR 0.86 (0.67 to 1.1)	7 fewer per 1000 (from 17 fewer to 5 more)	⊕⊕○○ LOW
<b>Worsening of angina status (follow-up 1.6 years)</b>											
Dargie 2002[49] (IONA)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	569/2565 (22.2%)	602/2561 (23.5%)	RR 0.94 (0.85 to 1.04)	14 fewer per 1000 (from 35 fewer to 9 more)	⊕⊕⊕○ MODERATE
<b>GI disturbances (follow-up 1.6 years)</b>											
Dargie 2002[49] (IONA)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (c)	None	194/2565 (7.6%)	132/2561 (5.2%)	RR 1.47 (1.18 to 1.82)	24 more per 1000 (from 9 more to 42 more)	⊕⊕○○ LOW
<b>Combined outcome CHD death, non-fatal MI or hospital admission for chest pain (diabetes subgroup) (follow-up 1.6 years)</b>											
IONA Study Group 2004[50] (IONA)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	None	27/197 (13.7%)	40/232 (17.2%)	RR 0.79 (0.51 to 1.25)	36 fewer per 1000 (from 84 fewer to 43 more)	⊕⊕○○ LOW
<b>Combined outcomes CHD death, non-fatal MI or hospital admission for chest pain (age subgroup &gt;70 yrs) (follow-up 1.6 years)</b>											
IONA Study Group 2004[50] (IONA)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	None	131/927 (14.1%)	167/948 (17.6%)	RR 0.8 (0.65 to 0.99)	35 fewer per 1000 (from 2 fewer to 62 fewer)	⊕⊕○○ LOW
<b>Combined outcomes CHD death, non-fatal MI or hospital admission for chest pain (female subgroup) (follow-up 1.6 years)</b>											
IONA Study Group 2004[50] (IONA)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	86/603 (14.3%)	87/613 (14.2%)	RR 1 (0.76 to 1.32)	0 fewer per 1000 (from 34 fewer to 45 more)	⊕⊕○○ LOW
<b>Composite (CHD death, non fatal MI or hospital admission. for chest pain) (follow-up 1.6 years)</b>											
Dargie 2002[49]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (c)	None	337/2565 (13.1%)	398/2561 (15.5%)	RR 0.85 (0.74 to 0.97)	23 fewer per 1000 (from 5 fewer to 40)	⊕⊕○○

(IONA)										fewer)	LOW
<b>Headache (follow-up 1.6 years)</b>											
Dargie 2002[49] (IONA)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	364/2565 (14.2%)	81/2561 (3.2%)	RR 4.49 (3.55 to 5.67)	110 more per 1000 (from 81 more to 148 more)	⊕⊕⊕○ MODERATE

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- (a) Randomisation process was reported; allocation concealment was not reported; study was double blind; Number of drop outs were reported and >20%; Intention to treat analysis was reported; the study was powered for primary outcome (CHD death, non fatal MI, or unplanned hospitalisation).
- (b) 95% CI around the pooled estimate of effect includes both 1) no effect 2) appreciable benefit or appreciable harm
- (c) 95% CI around the pooled estimate of effect includes appreciable benefit or appreciable harm
- (d) Canadian Cardiovascular Society Functional classification of angina at the end of the study (follow-up mean 1.6 years):  
 Class I - Nicorandil 985 (43%); placebo 989 (43%)  
 Class II- Nicorandil 1159 (50%); placebo 1124 (49%)  
 Class III- Nicorandil 162 (7%); placebo 163 (7%)  
 Class IV- Nicorandil 9 (<1%) ; placebo 15 (1%)

1 **Table 10.7: Nicorandil versus diltiazem**

Quality assessment							Summary of findings				
							No of patients		Effect		Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Nicorandil	Diltiazem	Relative (95% CI)	Absolute	
<b>Exercise capacity (work to peak exercise) (KJ) (follow-up 90 days: better indicated by more)</b>											
Guermontprez 1993[51]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	None	50	56	-	MD 2.4 higher (60.15 lower to 64.95 higher)	⊕⊕○○ LOW
<b>Exercise capacity (work to onset of angina) (KJ) (follow-up 90 days: better indicated by more)</b>											
Guermontprez 1993[51]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	None	50	56	-	MD 3.40 higher (58.91 lower to 65.71 higher)	⊕⊕○○ LOW
<b>Adverse events (combined) (follow-up 90 days)</b>											
Guermontprez 1993[51]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (c)	None	19/60 (31.7%)	19/63 (30.2%)	RR 1.05 (0.62 to 1.78)	15 more per 1000 (from 115 fewer to 235 more)	⊕⊕○○ LOW

(a) Allocation concealment was not reported; study was double blind; number of drop-outs were reported and < 20%; intention to treat analysis was not reported.

(b) 95% CI includes no effect and the upper and the lower confidence limit crosses the MID.

(c) 95% CI around the pooled estimate of effect includes both 1) no effect 2) appreciable benefit or appreciable harm

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1 **Table 10.8: Nicorandil vs. diltiazem** \* (Data for this outcome not able to analyse. Results reported as in the paper)

<b>Outcome</b>	<b>Number of studies</b>	<b>Design</b>	<b>Limitations</b>	<b>Inconsistency</b>	<b>Directness</b>	<b>Imprecision</b>
Frequency of anginal attacks per week	1 (Guermonprez)	RCT (double blind)	Serious <sup>1</sup>	No serious Inconsistency	No serious indirectness	No serious imprecision

<b>Outcome</b>	<b>Nicorandil</b>	<b>Placebo</b>	<b>Relative risk</b>	<b>Absolute effect</b>	<b>Quality</b>
Follow-up 90 days					
Frequency of anginal attacks per week	0.7 (mean) <sup>2</sup>	-	-	SD not reported. P=0.56 (Difference between groups not significant).	MODERATE

2 <sup>1</sup> Allocation concealment was not reported; study was double blind; Number of drop-outs were reported and < 20%; Intention to treat analysis was not reported.

3 <sup>2</sup> Mean value reported for both groups together. No standard deviation (SD) reported

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**Table 10.9: Nicorandil versus amlodipine**

Quality assessment							Summary of findings				Quality
							No of patients		Effect		
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Nicorandil	Amlodipine	Relative (95% CI)	Absolute	
<b>ETT (Total exercise duration) (min) (follow-up 8 weeks; better indicated by higher values)</b>											
Chatterjee 1999[52]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	56	62	-	MD 0.7 lower (1.69 lower to 0.29 higher)	⊕⊕○○ LOW
<b>ETT (Time to ST-segment depression) (follow-up 8 weeks; better indicated by higher values)</b>											
Chatterjee 1999[52]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	56	62	-	MD 0.6 lower (1.45 lower to 0.25 higher)	⊕⊕○○ LOW
<b>ETT (Time to onset of anginal pain) (follow-up 8 weeks; better indicated by higher values)</b>											
Chatterjee 1999[52]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	56	62	-	MD 0.9 lower (2 lower to 0.2 higher)	⊕⊕○○ LOW
<b>Sum of weekly anginal attacks (follow-up 8 weeks; better indicated by lower values)</b>											
Chatterjee 1999[52]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (c)	none	56	62	-	MD 1.2 higher (0.54 to 1.86 higher)	⊕⊕○○ LOW
<b>Adverse events (combined) (follow-up 8 weeks)</b>											
Chatterjee 1999[52]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (d)	none	20/57 (35.1%)	20/64 (31.3%)	RR 1.12 (0.68 to 1.86)	38 more per 1000 (from 100 fewer to 269 more)	⊕⊕○○ LOW

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- (a) Allocation concealment was not reported; study was double blind; number of drop-outs were reported and < 20%; intention to treat analysis was reported.
- (b) 95% CI includes no effect and the upper and the lower confidence limit crosses the MID.
- (c) The upper and the lower confidence limit crosses the MID.
- (d) 95% CI around the pooled estimate of effect includes both 1) no effect 2) appreciable benefit or appreciable harm

1 **Table 10.10: Nicorandil vs. nifedipine for stable angina**

Quality assessment							Summary of findings				
							No of patients		Effect		Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Nicorandil	Nifedipine	Relative (95% CI)	Absolute	
<b>Weekly anginal attack rate (follow-up after 8 weeks of treatment; better indicated by lower values)</b>											
Ulvenstam 1992[53]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	27	23	-	MD 5.3 lower (11.48 lower to 0.88 higher)	⊕⊕⊕⊕ LOW
<b>Exercise duration (min) (follow-up after 8 weeks of treatment; better indicated by higher values)</b>											
Ulvenstam 1992[53]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	25	23	-	MD 1 higher (0.59 lower to 2.59 higher)	⊕⊕⊕⊕ LOW
<b>Time to onset of angina pectoris (min) (follow-up after 8 weeks of treatment; better indicated by higher values)</b>											
Ulvenstam 1992[53]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	23	22	-	MD 1.1 higher (0.75 lower to 2.95 higher)	⊕⊕⊕⊕ LOW
<b>Time to 1mm ST-depression (min) (follow-up after 8 weeks of treatment; better indicated by higher values)</b>											
Ulvenstam 1992[53]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	23	20	-	MD 1.6 higher (0.02 lower to 3.22 higher)	⊕⊕⊕⊕ LOW
<b>ST depression on maximal identical workload (mm) (follow-up after 8 weeks of treatment; better indicated by higher values)</b>											
Ulvenstam 1992[53]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	24	20	-	MD 0.2 higher (0.28 lower to 0.68 higher)	⊕⊕⊕⊕ LOW
<b>Adverse events (combined) follow-up after 8 weeks of treatment; better indicated by lower values)</b>											
Ulvenstam 1992[53]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	25/29 (86.2%)	28/29 (96.6%)	RR 0.89 (0.76 to 1.05)	106 fewer per 1000 (from 232 fewer to 48 more)	⊕⊕⊕⊕ MODERATE

(a) Double-blind, randomised, multicentre study. 55/58 completed the study. Allocation concealment not reported. ITT not reported.  
 (b) 95% CI includes no effect and the upper and the lower confidence limit crosses the MID.

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1 **Table 10.11: Nicorandil versus isosorbide mononitrate**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Nicorandil	ISMN	Relative (95% CI)	Absolute	
<b>ETT (Total exercise time) (sec) (follow-up 2 weeks; better indicated by higher values)</b>											
Zhu 2007[54]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious imprecision(b)	None	115	117	-	MD 3.2 lower (37.26 lower to 30.86 higher)	⊕⊕⊕○ LOW
<b>ETT (Time to ST depression) (follow-up 2 weeks; better indicated by higher values)</b>											
Zhu 2007[54]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious imprecision (b)	None	114	116	-	MD 2.4 higher (37.98 lower to 42.78 higher)	⊕⊕⊕○ LOW
<b>Adverse event (Headache) (follow-up 2 weeks)</b>											
Zhu 2007[54]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (c)	None	15/123 (12.2%)	18/123 (14.6%)	RR 0.83 (0.44 to 1.58)	25 fewer per 1000 (from 82 fewer to 85 more)	⊕⊕○○ LOW

(a) Allocation concealment was not reported; study was double blind; number of drop-outs were reported and < 20%; intention to treat analysis was not reported.

(b) 95% CI includes no effect and the upper and the lower confidence limit crosses the MID.

(c) 95% CI around the pooled estimate of effect includes both 1) no effect 2) appreciable benefit or appreciable harm

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1 **Table 10.12: Nicorandil versus propranolol**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Nicorandil	Propranolol	Relative (95% CI)	Absolute	
<b>Angina free in daily life (%) (follow-up 6 weeks ;better indicated by higher values)</b>											
Meeter 1992[55]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision (b)	None	11/32 (34.4%)	13/37 (35.1%)	RR 0.98 (0.51 to 1.87)	7 fewer per 1000 (from 172 fewer to 306 more)	⊕⊕⊕O MODERATE
<b>12 hrs after medication - change in maximal work load (follow-up 3 weeks; better indicated by higher values)</b>											
Meeter 1992[55]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (c)	None	32	37	-	MD 6 lower (14.77 lower to 2.77 higher) <sup>4</sup>	⊕⊕OO LOW
<b>12 hrs after medication - change in maximal work load (W) (follow-up 6 weeks; better indicated by higher values)</b>											
Meeter 1992[55]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (e)	None	32	37	-	MD 5 lower (15.72 lower to 5.72 higher)	⊕⊕OO LOW
<b>12 hrs after treatment - change in time to angina (follow-up 3 weeks; better indicated by higher values)</b>											
Meeter 1992[55]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (c)	None	32	37	-	MD 0.10 lower (1.05 lower to 0.85 higher)	⊕⊕OO LOW
<b>12 hrs after treatment - change in time to angina (follow-up 6 weeks; better indicated by lower values)</b>											
Meeter 1992[55]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (c)	None	32	37	-	MD 0.40 lower (1.35 lower to 0.55 higher)	⊕⊕OO LOW
<b>2 hrs after treatment - change in maximal work load (follow-up 3 weeks; better indicated by higher values)</b>											
Meeter 1992[55]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	32	37	-	MD 5.00 lower (13.07 lower to 3.07 higher)	⊕⊕⊕O MODERATE
<b>2 hrs after treatment - change in maximal work load (W) (follow-up 6 weeks; better indicated by higher values)</b>											
Meeter 1992[55]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (c)	None	32	37	-	MD 5.00 lower (14.47 lower to 4.47 higher)	⊕⊕OO LOW
<b>2 hrs after treatment - change in time to angina (follow-up 3 weeks; better indicated by lower values)</b>											
Meeter 1992[55]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (c)	None	32	37	-	MD 0.20 higher (0.53 lower to 0.93 higher)	⊕⊕OO LOW
<b>2 hrs after medication - change in time to angina (follow-up 6 weeks; better indicated by higher values)</b>											
Meeter 1992[55]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision (c)	None	32	37	-	MD 0.60 higher (0.35 lower to 1.55 higher)	⊕⊕⊕O MODERATE

- 2 (a) 1/1 Allocation concealment not reported; 1/1 double blind; 1/1 drop-out rate reported and < 20%; Intention to treat analysis not reported.
- 3 (b) 95% C around the pooled estimate of effect includes both 1) no effect 2) appreciable benefit or appreciable harm.
- 4 (c) The upper and lower limits of 95% CI crosses the MI.

1 **10.3.3 Economic evidence**

2 No economic evidence was found on the use of nicorandil as monotherapy. Based on the  
3 unit cost reported in the BNF59[18] the annual drug cost ranges from £99 and £190.

4 We found one study[56] comparing the addition of nicorandil to usual care with placebo.  
5 This is summarised in the economic evidence profile below. See also Economic Evidence  
6 Tables in Appendix G.

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8 **Table 10.13: Nicorandil+usual care vs. placebo+usual care - Economic study characteristics**

Study	Limitations	Applicability	Other Comments
Walker 2006[56]	Potentially serious limitations (a)	Partial applicability (b)	Intervention was nicorandil 20mg bd + usual care (57% BB, 55% CCB, 87% nitrates, 88% aspirin). Based on the IONA trial[49] included in the clinical review.

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- c) Short follow-up (up to 1.6 years). Sensitivity analysis was quite limited and was applied only to the primary analysis (cost of care after discharge excluded). Morbidity associated with gastro-intestinal events is not included. Effectiveness data were reported only in the incremental analysis.
- d) QALYs were not estimated.

**Table 10.14: Nicorandil+usual care vs. placebo+usual care - Economic summary of findings**

Study	Incremental cost per patient (£)	Incremental effects per patient (primary end-point averted)	ICER	Uncertainty
Walker 2006[56]	Saves £0.12 (a)	0.024 (b)	Dominant	Nicorandil is not cost-saving when: - cost of care after discharge is included - either cost of cardiology, cardiac surgery or ICU is reduced by 20%. Results were similar when the measure of effectiveness considered was the number of event-free survivors (events were cardiac death, non-fatal MI, unstable and stable angina, stroke, hospital admission for TIA) or the number of cases of definite acute coronary syndromes (coronary heart disease death, non-fatal myocardial infarction or unstable angina).

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- (a) 2002 GBP. Costs included were cost of Nicorandil (including 10% dispensing fee and two additional physician visits), adverse events related to Nicorandil, hospital admissions, surgical procedures. The cost of post-discharge care was not included in the base case analysis.
- (b) Primary end-points considered in the analysis were cardiac death, non-fatal MI, hospital admission for cardiac chest pain.

## 1 10.3.4 Evidence statements

## Clinical

**A. Clinical Efficacy****Nicorandil versus Placebo**

**Dargie 2002[49] and IONA Study Group 2004[50]:** Evidence from one RCT shows that the composite outcomes (CHD death, non-fatal MI, or unplanned hospital admission for chest pain) for the entire group [RR 0.85 (0.74 to 0.97)] and for people aged over 70 years [RR 0.80 (0.65 to 0.99)] were significantly reduced in the nicorandil group compared to placebo (mean follow-up 1.6 years).

**Dargie 2002[49] and IONA Study Group 2004[50]:** Evidence from one RCT shows that there were no statistically significant differences between nicorandil and placebo for CHD death [RR 0.82 (0.59 to 1.15)], non fatal MI [RR 0.78 (0.55 to 1.10)], all cause mortality [RR (0.86 (0.67 to 1.10)], unstable angina [RR 0.90 (0.71 to 1.16)], and worsening of angina status [RR 0.94 (0.85 to 1.04)]. There were no statistically significant differences between treatment groups for a composite morbidity/mortality outcome (CHD death, non-fatal MI, or unplanned hospital admission for chest pain) in subgroup analyses of results for women [RR 1.00 [0.76 to 1.31], and people with diabetes [RR 0.79 (0.51 to 1.25)] (mean follow-up 1.6 years).

**Nicorandil versus diltiazem**

**Guermontez 1992[51]:** Evidence from one RCT shows that there was no significant difference between nicorandil and diltiazem for exercise capacity (work to peak exercise) [MD 2.4 (-60.15 to 64.95) and exercise capacity (work required to reach onset of angina) [MD 3.40 (-58.91 to 64.95)] (follow-up 90 days).

**Nicorandil versus amlodipine**

**Chatterjee 1999[52]:** Evidence from one RCT shows that there were no significant differences between nicorandil and amlodipine for total exercise duration (min), MD -0.70 [-1.69 to 0.29] , ETT (Time to onset of anginal pain) MD -0.9 (-2 to 0.2 ) , and ETT (Time to ST-segment depression) [MD -0.6 (-1.45 to 0.25 higher) and sum of weekly anginal attacks, [MD 1.20 [0.54 to 1.86] (follow-up 8 weeks).

**Nicorandil versus nifedipine**

**Ulvenstam 1992[53]:** Evidence from one RCT shows that there was no statistically significant differences between nicorandil and nifedipine for weekly anginal attack rate [MD -5.3 (-11.48 to 0.88)], exercise duration (min) [MD 1 higher (-0.59 to 2.59)], time to onset of angina pectoris (min) [MD 1.1 (-0.75 to 2.95)], time to 1mm ST-depression (min) [MD 1.6 (-0.02 to 3.22)], and ST depression on maximal identical workload (mm) [MD 0.2 (-0.28 to 0.68)] (follow-up after 8 weeks of treatment).

**Nicorandil versus isosorbide mononitrate**

**Zhu 2007[54]:** Evidence from one RCT shows that there was no

significant difference between nicorandil and isosorbide mononitrate for total exercise time (sec) [MD -3.20 [-37.26 to 30.86]] and ETT (time to ST-depression) [MD 2.4 (-37.98 to 42.78)] (follow-up 2 weeks).

#### **Nicorandil versus propranolol**

**Meeter 1992[55]:** Evidence from one RCT shows that there was no significant difference between nicorandil and propranolol for frequency of anginal attacks [RR 0.98 (0.51 to 1.87)] (follow-up 6 weeks).

**Meeter 1992[55]:** Evidence from one RCT shows that there was no significant difference between nicorandil and propranolol for change in maximal workload 12 hrs after medication at 3 weeks [MD -6 (-14.77 to 2.77)] and 6 weeks [MD -5 (-15.72 to 5.72)], change in time to angina decimal min 12 hrs after medication at 3 weeks [MD -5.40 (-6.35 to -4.45)] and 6 weeks [MD -0.40 (-1.35 to 0.55)], change in maximal workload 2 hrs after treatment at 3 weeks [MD -5.00 (-13.07 to 3.07)] and 6 weeks [MD -5.00 (-14.47 to 4.47)], change in time to angina 2 hrs after treatment at 3 weeks [MD 0.20 (-0.53 to 0.93)] and 6 weeks [MD 0.60 (-0.35 to 1.55)] (follow-up 6 weeks).

### **B. Adverse events**

#### **Nicorandil versus placebo**

**Dargie 2002[49] (IONA):** Evidence from one RCT shows that there were significantly greater GI disturbances [RR 1.47 (1.18 to 1.82)] and headaches in the nicorandil compared to placebo [RR 4.49 (3.55 to 5.67)] (mean follow-up 1.6 years) (follow-up mean 1.6 years).

#### **Nicorandil versus diltiazem**

**Guermontprez 1993[51]:** Evidence from one RCT shows that there were no statistically significant differences between nicorandil and diltiazem for adverse effects (combined) [RR 1.05 (0.62 to 1.78)] (follow-up 90 days).

#### **Nicorandil versus amlodipine**

**Chatterjee 1999[52]:** Evidence from one RCT suggests that there were no statistically significant differences between nicorandil and amlodipine for adverse effects (combined) [RR 1.1 (0.68 to 1.86)] (follow-up 8 weeks).

#### **Nicorandil vs. nifedipine**

**Ulvenstam 1992[53]:** Evidence from one RCT suggests that there were no statistically significant differences between nicorandil and nifedipine for adverse events (combined) [RR 0.89 (0.76 to 1.05)] (follow-up after 8 weeks of treatment).

#### **Nicorandil versus isosorbide mononitrate**

**Zhu 2007[54]:** Evidence from one RCT suggests that there were no statistically significant difference between nicorandil and isosorbide mononitrate for adverse effects (headache) [RR 0.83 (0.44 to 1.58)]

(follow-up 2 weeks).

**Nicorandil versus propranolol**

**Meeter 1992[55]:** Adverse effects not reported (follow-up 6 weeks).

**Economic** Nicorandil is cost-neutral when post discharge care is not included and over a short time (1.6 years). It could be less cost effective when post-discharge care is included. This evidence has potentially serious limitations and partial applicability.

1 **10.3.5 Recommendations and link to evidence**

<b>Recommendation</b>	<p><b>If the person cannot tolerate beta blockers and calcium channel blockers or they are contraindicated, consider monotherapy with one of the following drugs*:</b></p> <ul style="list-style-type: none"> <li>• a long-acting nitrate</li> <li>• ivabradine</li> <li>• nicorandil** or</li> <li>• ranolazine</li> </ul> <p><b>Decide which drug to use based on comorbidities, contraindications, the person's preference and costs.</b></p> <p><i>* Evidence on long acting nitrates is presented in chapter 9. Evidence on ivabradine and ranolazine is presented in sections 10.2 and 10.4 respectively of this chapter.</i></p> <p><i>**At the time of consultation (December, 2010), nicorandil did not have UK marketing authorisation for this indication. Informed consent should be obtained and documented.</i></p>
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**Relative values of different outcomes**

Outcomes of interest included long-term mortality (total and cardiovascular), rates of major adverse cardiovascular events (myocardial infarction, stroke, myocardial revascularisation), and measures of symptom severity (frequency of angina, exercise test outcomes).

**Trade off between clinical benefits and harms**

No evidence was found to assess the effects of monotherapy with nicorandil on long term mortality or rates of major adverse cardiovascular events in people with stable angina.

Short-term trials of monotherapy with nicorandil versus monotherapy with other anti-anginal drugs (diltiazem, amlodipine, or propranolol) demonstrated similar reductions in

the frequency of episodes of angina in both treatment groups.

These trials also reported similar increases in total exercise capacity during monotherapy with nicorandil and monotherapy with diltiazem, amlodipine, propranolol, or isosorbide mononitrate.

No difference in the short-term rate of adverse effects was reported between nicorandil and diltiazem, amlodipine, or isosorbide mononitrate.

**Economic considerations**

No economic evidence on the use of nicorandil in monotherapy was found. The annual drug cost of nicorandil ranges from £99 to £190.

**Quality of evidence**

Low quality evidence from trials with small sample size and short duration of follow-up. In one trial [55] no intention to treat analysis was carried out.

No economic evidence was found.

**Other considerations**

The GDG concluded that there is insufficient evidence to recommend monotherapy with nicorandil in preference to monotherapy with a BB or CCB as first line treatment for angina. Nicorandil can be considered as monotherapy for the treatment of stable angina if BB and CCB are not tolerated or contraindicated.

Adverse effects of nicorandil include headache (especially on initiation of treatment), flushing, dizziness, reduction in blood pressure and/or increase in heart rate, and gastrointestinal side effects including mucosal ulceration. In the IONA trial routine treatment with nicorandil was associated with a higher risk of gastrointestinal side effects and GDG members have experience of patients who developed gastrointestinal ulceration during treatment with nicorandil.

<b>Recommendation</b>	<p><b>For people on beta blocker or calcium channel blocker monotherapy whose symptoms are not controlled and the other option (calcium channel blocker or beta blocker) is contraindicated or not tolerated, consider one of the following as an additional drug:</b></p> <ul style="list-style-type: none"> <li>• <b>a long-acting nitrate</b></li> </ul>
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	<ul style="list-style-type: none"> <li>• <b>ivabradine</b></li> <li>• <b>nicorandil<sup>3</sup> or</b></li> <li>• <b>ranolazine.</b></li> </ul> <p><i>* Evidence on long acting nitrates is presented in chapter 9. Evidence on ivabradine and ranolazine is presented in sections 10.2 and 10.4 respectively of this chapter.</i></p> <p><i>**At the time of consultation (December, 2010), nicorandil did not have UK marketing authorisation to be combined with any other drugs. Informed consent should be obtained and documented.</i></p>
<p><b>Relative values of different outcomes</b></p>	<p>Outcomes of interest included long-term mortality (total and cardiovascular), rates of major adverse cardiovascular events (myocardial infarction, stroke, myocardial revascularisation), and measures of symptom severity (frequency of angina, exercise test outcomes).</p>
<p><b>Trade off between clinical benefits and harms</b></p>	<p>In a large trial addition of nicorandil to standard antianginal treatment (BB 56%, CCB 55%, nitrate 87%) in people with stable angina reduced the composite of coronary heart disease death, myocardial infarction, and unplanned hospitalisation for chest pain. There were trends for lower rates of all events included in the composite primary endpoint in the nicorandil group, but these were not statistically significant. At the end of the study (1.6 years) the Canadian Cardiovascular Society angina class did not differ between the two groups. Headache, gastrointestinal disturbance, and treatment withdrawal because of adverse effects were more frequent in the nicorandil group.</p> <p>The GDG concluded that the 2.4% absolute reduction in the rate of the primary composite endpoint in IONA did not justify the routine use of nicorandil as add-on therapy to standard antianginal treatment in people with stable angina, particularly as the drug is associated with an excess risk of adverse events, including headache and gastrointestinal disturbance.</p>
<p><b>Economic considerations</b></p>	<p>When symptoms are not controlled with standard treatment, adding nicorandil could be a cost-effective option. Nicorandil is cost-neutral when post discharge care is not included and</p>

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<sup>3</sup> At the time of consultation (November 2010), nicorandil did not have UK marketing authorisation for use in this indication. Informed consent should be obtained and documented.

over a short time (1.6 years). It could be less cost effective when post-discharge care is included.

**Quality of evidence**

Moderate quality evidence from a large multicentre trial powered to detect a 20% reduction in the primary endpoint. Allocation concealment was not reported and treatment withdrawal was >20% in both groups.

The economic evidence has potentially serious limitations and partial applicability.

**Other considerations**

The GDG concluded that addition of nicorandil is an option for people whose symptoms of angina are not controlled by a BB or CCB. Nicorandil is slightly cheaper than ivabradine and ranolazine but more than the cost of long-acting nitrate. Nicorandil is currently not licensed for use in combination treatment. Nevertheless the GDG concluded that there was insufficient evidence to make a firm recommendation about the choice of an additional antianginal drug if a BB or CCB is not tolerated or is contraindicated.

1 **10.4 Ranolazine**

2 The mechanism of action of ranolazine has not been fully elucidated, but it is believed  
3 to act by selective inhibition of late sodium influx across the sarcolemma, which  
4 attenuates the abnormalities of ventricular repolarisation and contractility associated  
5 with myocardial ischaemia. Reported side-effects include dizziness, constipation, and  
6 nausea. Ranolazine has the potential to prolong the QT interval and is  
7 contraindicated in people with pre-existing QT prolongation. Ranolazine should be  
8 avoided in severe hepatic or renal impairment. Ranolazine is available in a sustained  
9 release formulation, with an elimination half-life of about seven hours.

10 This section reviews evidence for the use of ranolazine as adjunctive therapy to  
11 control symptoms and improve outcome in people with stable angina.

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13 **10.4.1 Clinical question**

14 What is the clinical/cost effectiveness of ranolazine for the management of stable  
15 angina?

16 **10.4.2 Clinical evidence**

17 The "Review Protocol" for this topic can be found in Appendix C, the "Search  
18 Strategies" in Appendix D, the "List of Included and Excluded Studies" in Appendix  
19 E1, the "Clinical Evidence Tables" in Appendix E2, and the "Forest Plots" in Appendix  
20 F. .

1 **Table 10.15: Ranolazine (750 mg bid ) + antianginal treatment vs. placebo + antianginal treatment (follow-up 12 weeks)**

Quality assessment							Summary of findings				
							No of patients		Effect		Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Ranolazine (750 mg bid ) + antianginal	Placebo + antianginal	Relative (95% CI)	Absolute	
<b>Exercise duration (sec) (trough - change from baseline) - (follow-up 12 weeks; better indicated by higher values)</b>											
Chaitman 2004[57] (CARISA) (c)	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	serious imprecision (d)	none	272	258	-	MD 23.7 higher (1.11 to 46.29 higher)	⊕⊕⊕⊕ MODERATE
<b>Time to onset of angina (sec) (trough - change from baseline) - (follow-up 12 weeks; better indicated by higher values)</b>											
Chaitman 2004[57] (CARISA)	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	serious imprecision (d)	none	272	258	-	MD 29.7 higher (4.62 to 54.78 higher)	⊕⊕⊕⊕ MODERATE
<b>Exercise duration (sec) (peak - change from baseline) - (follow-up 12 weeks; better indicated by higher values)</b>											
Chaitman 2004[57] (CARISA)	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	serious imprecision (d)	none	270	256	-	MD 34 higher (11.96 to 56.04 higher)	⊕⊕⊕⊕ MODERATE
<b>Time to onset of angina (sec) (peak - change from baseline) - (follow-up 12 weeks; better indicated by higher values)</b>											
Chaitman 2004[57] (CARISA)	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	serious imprecision (d)	none	272	256	-	MD 38 higher (13.91 to 62.09 higher)	⊕⊕⊕⊕ MODERATE
<b>Adverse events (follow-up 12 weeks)</b>											
Chaitman 2004[57] (CARISA)	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	serious (b)	none	82/279 (29.4%)	71/269 (26.4%)	RR 1.11 (0.85 to 1.46)	29 more per 1000 (from 40 fewer to 121 more)	⊕⊕⊕○ MODERATE
<b>Angina attacks per week (follow-up 12 weeks; better indicated by lower values)</b>											
Chaitman 2004[57] (CARISA)	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	272	258	-	MD 0.8 lower (1.52 to 0.08 lower)	⊕⊕⊕⊕ HIGH

(a) Randomised. Allocation concealment reported. ITT reported.

(b) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.

(c) Ranolazine 750 mg twice plus anti-anginal drugs including atenolol 50 mg (45% patients), amlodipine 5 mg (30%) and diltiazem 180 mg (26%) vs. placebo plus antianginal drugs

(d) 95% CI includes no effect and the upper and lower CI crosses the MID.

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1 **Table 10.16: Ranolazine (750 mg bid) + antianginal treatment vs. placebo+antianginal treatment – Subgroup diabetes (follow-up 12 weeks)**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Ranolazine (750 mg bid) + antianginal treatment	Placebo+antianginal treatment - diabetic patients	Relative (95% CI)	Absolute	
<b>Exercise duration sec (trough change from baseline) - 12 wks (follow-up 12 weeks; better indicated by higher values)</b>											
Timmis 2006[58] (CARISA) (c)	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	serious (b)	none	68	57	-	MD 28.7 higher (50.9 lower to 108.3 higher)	⊕⊕⊕O MODERATE
<b>Time to onset of angina sec (trough change from baseline) - 12 wks (follow-up 12 weeks; better indicated by higher values)</b>											
Timmis 2006[58] (CARISA)	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	serious (b)	none	68	57	-	MD 50.8 higher (37.56 lower to 139.16 higher)	⊕⊕⊕O MODERATE
<b>Angina episodes per week - 12 wks (follow-up 12 weeks; better indicated by lower values)</b>											
Timmis 2006[58] (CARISA)	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	serious (b)	none	68	57	-	MD 0.91 lower (3.25 lower to 1.43 higher)	⊕⊕⊕O MODERATE
<b>Nitroglycerin consumption per week - 12 wks (follow-up 12 weeks; better indicated by lower values)</b>											
Timmis 2006[58] (CARISA)	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	serious (b)	none	68	57	-	MD 2.32 lower (7.18 lower to 2.54 higher)	⊕⊕⊕O MODERATE

(a) Randomised. Allocation concealment reported. ITT reported.

(b) 95% CI includes no effect and the upper and lower CI crosses the MID.

(c) Ranolazine 750 mg twice plus anti-anginal drugs including atenolol 50 mg (45% patients), amlodipine 5 mg (30%) and diltiazem 180 mg (26%) vs. placebo plus antianginal drugs

**Sub-group interaction between diabetic and non-diabetic patients:** There was no significant treatment by subgroup interaction for exercise duration ( $p=0.89$ ) and time to onset of angina ( $p=0.54$ ) between diabetic and non diabetic patients. Statistical tests for interaction between diabetes status and treatment effect showed no evidence that the effects of ranolazine differed between diabetic and non-diabetic patients either in the number of angina episodes per week ( $p=0.81$ ) or nitroglycerin usage ( $p=0.063$ ); and therefore no evidence that the treatment effect differed between diabetic and non-diabetic patients.

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1 **Table 10.17: Ranolazine (1000 mg bid) + antianginal treatment vs. placebo +antianginal treatment- Subgroup age (follow-up 6 weeks)**

Quality assessment							Summary of findings				
							No of patients		Effect		Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Ranolazine (1000 mg bid) + antianginal treatment	Placebo +antianginal treatment- age	Relative (95% CI)	Absolute	
<b>Weekly angina attacks &lt; 70 yrs (follow-up 6 weeks; better indicated by lower values)</b>											
Rich 2007[59] (CARISA)	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	403	409	-	MD 0.5 lower (1.1 lower to 0.1 higher)	⊕⊕⊕⊕ HIGH
<b>Weekly angina attacks &gt; 71 yrs (follow-up 6 weeks; better indicated by lower values)</b>											
Rich 2007[59] (CARISA)	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	135	130	-	MD 1.13 lower (2.05 to 0.21 lower)	⊕⊕⊕⊕ HIGH
<b>Nitroglycerin consumption &lt; 70 yrs (follow-up 6 weeks; better indicated by lower values)</b>											
Rich 2007[59] (CARISA)	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	403	409	-	MD 0.97 lower (1.64 to 0.3 lower)	⊕⊕⊕⊕ HIGH
<b>Nitroglycerin consumption &gt; 71 yrs (follow-up 6 weeks; better indicated by lower values)</b>											
Rich 2007[59] (CARISA)	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	135	130	-	MD 0.94 lower (1.74 to 0.14 lower)	⊕⊕⊕⊕ HIGH
<b>Adverse events &lt;70 years (follow-up 6 weeks) (c)</b>											
Rich 2007[59] (CARISA)	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	194/604 (32.1%)	131/420 (31.2%)	RR 1.03 [0.86, 1.24]	9 more per 1000 (from 44 fewer to 75 more )	⊕⊕⊕⊕ HIGH
<b>Adverse events &gt; 70 years (follow-up 6 weeks) (c)</b>											
Rich 2007[59] (CARISA)	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	serious (b)	none	102/231 (44.2%)	43/132 (32.6%)	RR 1.36 [1.02, 1.80]	117 more per 1000 (from 7 more to 261 more)	⊕⊕⊕⊕ MOAEPATE

(a) Randomised. Allocation concealment reported. ITT reported.

(b) 95% CI around the pooled estimate of effect includes appreciable benefit or appreciable harm.

(c) Adverse events- cardiac adverse events, constipation, nausea, dyspepsia, dizziness, headache, peripheral edema asthenia, serious adverse events such as MI, syncope,, transient ischemic attack. The most common events resulting in discontinuation of study drug were related to the gastrointestinal, nervous, and cardiac organ systems.

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1 **Table 10.18: Ranolazine (1000 mg bid) plus amlodipine (10 mg) vs. amlodipine (10mg) (follow-up 6 weeks)**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Ranolazine (1000 mg bid) plus amlodipine (10 mg)	amlodipine (10mg)	Relative (95% CI)	Absolute	
<b>Adverse events (follow-up 6 weeks)</b>											
Stone 2006[60] (ERICA) (c)	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	serious (b)	none	112/281 (39.9%)	100/284 (35.2%)	RR 1.13 (0.91 to 1.4)	46 more per 1000 (from 32 fewer to 141 more)	⊕⊕⊕○ MODERATE
<b>Weekly angina frequency - (follow-up 6 weeks; better indicated by lower values)</b>											
Stone 2006[60] (ERICA)	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	277	281	-	MD 0.43 lower (1 lower to 0.14 higher)	⊕⊕⊕⊕ HIGH
<b>Weekly nitroglycerin consumption - (follow-up 6 weeks; better indicated by lower values)</b>											
Stone 2006[60] (ERICA)	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	277	281	-	MD 0.65 lower (1.23 to 0.07 lower)	⊕⊕⊕⊕ HIGH

2 <sup>1</sup> Randomised. Allocation concealment reported. Blinding of outcome assessors reported. ITT reported.<sup>2</sup> 95% CI around the pooled estimate of effect includes both: 1) no effect and  
 3 2) appreciable benefit or appreciable harm.<sup>3</sup>ERICA - Ranolazine 1000 mg twice daily plus amlodipine 10 mg/daily vs. amlodipine 10 mg/daily  
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### 1 10.4.3 Economic evidence

2 No economic studies were identified on this question. We calculated the daily and  
3 annual cost of ranolazine treatment based on the unit cost reported in the BNF59[18].

4 **Table 10.19: Drug cost of Ranolazine**

	Cost per day (£)	Cost per year (£)
Ranolazine, 375 mg, 500 mg or 750 mg twice daily	1.63	595

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6 The costs of adverse effects were not estimated.

7

### 8 10.4.4 Evidence statements

#### Clinical A. Clinical Efficacy

##### Ranolazine plus antianginal treatment versus placebo plus antianginal treatment

**Chaitman 2004[57] (CARISA):** Evidence from one RCT shows that exercise duration at trough (sec) [MD 23.70 (1.11 to 46.29)], time to onset of angina at trough (sec) [MD 29.70 (4.62 to 54.76)], exercise duration at peak (sec) [MD 34 (11.96 to 56.04)] and time to onset of angina at peak(sec) [MD 38 (13.91 to 62.09)] were significantly higher in the ranolazine plus antianginal treatment compared with placebo plus antianginal treatment [follow-up 12 weeks]. There were no statistically significant differences between ranolazine plus antianginal treatment and placebo plus antianginal treatment for the outcome of adverse events [RR 1.11 (0.85 to 1.46)] (follow up 12 weeks).

**Timmis 2006[58] (CARISA):** Evidence from a post-hoc sub-group analyses of one RCT shows that there were no statistically significant differences in the outcomes of exercise duration (sec) [MD 28.70 (-50.90 to 108.30)], time to onset of angina (sec) [MD 50.80 (-37.56 to 139.16)], frequency of angina attacks [MD -0.91 (-3.25 to 1.43)] and nitroglycerin consumption [MD -2.32 (-7.18 to 2.54)] between ranolazine plus anti anginal treatment and placebo plus anti-anginal treatment in people with diabetes (follow-up 12 weeks).

**Rich 2007[59] (CARISA):** Evidence from one post-hoc sub-group analysis of a RCT shows that in patients younger than 70 years ranolazine plus anti anginal treatment resulted in a statistically significant reduction in nitroglycerine consumption [MD -0.97 (-1.64 to -0.30)] but no significant difference in weekly angina attacks [MD -0.50 (-1.10 to 0.10)] or adverse events [RR1.03 [0.86, 1.24]] when compared

with placebo plus anti-anginal treatment [follow-up 6 weeks]. In patients older than 70 years ranolazine plus anti-anginal treatment resulted in statistically significant reductions in weekly angina attacks [MD -1.13 (-2.05 to -0.21)] and nitroglycerin consumption [MD -0.94 (-1.74 to -0.14)] but a statistically significant increase in adverse events [RR 1.36 [1.02, 1.80]] when compared with placebo plus anti-anginal treatment. (follow-up six weeks).

**Ranolazine plus amlodipine versus amlodipine**

**Stone 2006[60] (ERICA):** Evidence from one RCT shows that weekly nitroglycerin consumption was significantly lower with ranolazine plus amlodipine compared to amlodipine alone [MD -0.65 (-1.23 to -0.07)]. There were no statistically significant differences between ranolazine plus amlodipine and amlodipine for weekly angina frequency [MD -0.43 (-1.00 to 0.14)] and adverse events (RR 1.13 (0.91 to 1.40)) (follow-up 6 weeks).

**Economic** No economic evidence was found on this question. A simple cost analysis showed a significant drug cost of ranolazine.

1 10.4.5 Recommendations and link to evidence

<b>Recommendation</b>	<p><b>If the person cannot tolerate beta blockers and calcium channel blockers or they are contraindicated, consider monotherapy with one of the following drugs*:</b></p> <ul style="list-style-type: none"> <li>• a long-acting nitrate</li> <li>• ivabradine</li> <li>• nicorandil** or</li> <li>• ranolazine</li> </ul> <p><b>Decide which drug to use based on comorbidities, contraindications, the person's preference and costs.</b></p> <p><b>For people on beta blocker or calcium channel blocker monotherapy whose symptoms are not controlled and the other option (calcium channel blocker or beta blocker) is contraindicated or not tolerated, consider one of the following as an additional drug*:</b></p> <ul style="list-style-type: none"> <li>• a long-acting nitrate</li> <li>• ivabradine***</li> <li>• nicorandil** or</li> <li>• ranolazine</li> </ul> <p><b>Decide which drug to use based on comorbidities, contraindications, the person's preference and costs.</b></p> <p><i>* Evidence on long acting nitrates is presented in chapter 9. Evidence on ivabradine and nicorandil is presented in sections 10.2 and 10.3 respectively of this chapter.</i></p> <p><i>** At the time of consultation (December, 2010), nicorandil did not have UK marketing authorisation for use in this indication. Informed consent should be obtained and documented.</i></p> <p><i>*** Ivabradine should only be combined with a dihydropyridine CCB</i></p>
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**Relative values of different outcomes**

Outcomes of interest included long-term mortality (total and cardiovascular), rates of major adverse cardiovascular events (myocardial infarction, stroke, myocardial revascularisation), and measures of symptom severity (frequency of angina, exercise test outcomes).

**Trade off between clinical benefits and harms**

We found no evidence about the effects of ranolazine monotherapy or ranolazine in combination with other anti-anginal drugs on long-term outcome in people with stable angina.

In one randomised trial addition of ranolazine to standard anti-anginal treatment for twelve weeks increased exercise duration (by 20 to 30 seconds) and time to angina at trough (and at peak). Ranolazine reduced the frequency of angina attacks and nitroglycerine use by about one per week. These effects were consistent in people with diabetes and in people aged over 70 years.

In one randomised trial addition of ranolazine to amlodipine reduced nitroglycerine consumption (by 0.65 doses per week) but not weekly angina frequency after six weeks follow-up.

Ranolazine did not increase the risk of adverse events.

**Economic considerations**

No economic evidence on the use of ranolazine for the treatment of stable angina was available for review. The cost of ranolazine is substantially higher than the costs of first-line anti-anginal drugs (BBs and CCBs) and long-acting nitrates.

**Quality of evidence**

Randomised trials of ranolazine are of modest size and were not designed to assess the long-term effects of ranolazine on mortality or other major adverse cardiac events. The improvements in exercise time and symptom severity associated with short-term ranolazine treatment are modest and of uncertain clinical significance.

No economic evidence was available on this question.

**Other considerations**

Evidence to support the long-term use of ranolazine as adjunctive anti-anginal therapy is very limited. The GDG concluded that there is insufficient evidence to recommend routine use of ranolazine, but ranolazine may have a role in people with stable angina who are inadequately controlled or intolerant of first-line anti-anginal therapies.

The cost of ranolazine is comparable with the costs of ivabradine but more than the cost of long-acting nitrate. Ranolazine has a licence for use in combination treatment. Nevertheless the GDG concluded that there was insufficient evidence to make a firm recommendation about the choice of an additional anti-anginal drug if a BB or CCB is not tolerated or is contraindicated.

1 **10.5 General drug recommendations**

Recommendation	<b>Offer people optimal drug treatment for the initial management of stable angina. Optimal drug treatment consists of one or two anti-anginal drugs as necessary plus drugs for secondary prevention of cardiovascular disease.</b>
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Other considerations      The evidence reviews indicated benefit from secondary prevention treatment and anti-anginal treatment. The GDG considered it important to emphasise the importance for patients to receive optimal medical treatment and made a consensus recommendation for this.

2

Recommendation	<b>Advise people that the aim of anti-anginal drug treatment is to prevent episodes of angina and the aim of secondary prevention treatment is to prevent cardiovascular events such as heart attack and stroke.</b>
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Other considerations      The GDG were aware of the importance of patient adherence to secondary prevention treatment. They also considered it important that patients understand that the purpose of anti-anginal drugs is to improve symptoms. The GDG made a consensus recommendation to ensure that professionals explain these points adequately to patients.

3

Recommendation	<p><b>Review the person's response to treatment, including any side effects, 2–4 weeks after starting or changing drug treatment.</b></p> <p><b>Titrate the drug dosage against symptoms up to the maximum tolerable dosage.</b></p> <p><b>Discuss how side effects of drug treatment might affect the person's daily activities and explain why it is important to take drug treatment regularly.</b></p>
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Other considerations      The GDG debated the need to review the response to treatment after starting or changing any anti-anginal medication. The GDG reached a consensus that response to treatment, including any side effects, should be reviewed 2-4 weeks after starting or changing any anti-anginal drug. If the person's angina is not controlled, the dose of the anti-anginal drug should be titrated up to the maximum tolerable dose

(within the licensed dose range) with the objective of achieving control of symptoms of stable angina. The GDG also considered it important that patients do not remain on drugs that are not providing benefit to them and that health care professionals should stop anti-anginal drugs that are not providing symptomatic benefit.

Recommendation	<p><b>Do not offer a third anti-anginal drug to people whose stable angina is controlled with two anti-anginal drugs.</b></p> <p><b>Consider adding a third anti-anginal drug when:</b></p> <ul style="list-style-type: none"><li>• <b>the person's symptoms are not controlled with two anti-anginal drugs and</b></li><li>• <b>the person is waiting for revascularisation or it is not considered appropriate or acceptable.</b></li></ul> <p><b>Decide which drug to use based on comorbidities, contraindications, the person's preference and costs.</b></p>
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See section 8.5 for link to evidence for these recommendations

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## 2 10.6 Research recommendation

3 The GDG recommended the following research question:

4 ➤ **Research question:** What is the clinical and cost effectiveness of adding a newer  
5 anti-anginal drug (nicorandil, ivabradine or ranolazine) to a calcium channel blocker  
6 for treating stable angina?

7 ➤ **Why this is important:** We do not know the clinical and cost effectiveness of  
8 adding a newer anti-anginal drug to a calcium channel blocker in people with  
9 stable angina. We propose a double-blind placebo-controlled randomised trial  
10 comparing the addition of a newer anti-anginal drug to a calcium channel blocker  
11 with a calcium channel blocker alone in people with stable angina whose symptoms  
12 are not being controlled. Endpoints would include symptom severity, quality of life,  
13 long-term morbidity and mortality, and cost effectiveness. The results of the trial  
14 would influence clinical practice and inform future updates of key recommendations  
15 in this guideline.

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## 2 **11 Medical versus revascularisation**

### 3 **interventions**

#### 4 **11.1 Introduction**

5 This chapter compares the effectiveness of medical treatment to revascularisation (PCI  
6 or CABG) for treating people with stable angina.

7 Coronary artery bypass surgery has been used to treat people with stable angina  
8 since the 1970s. Until recently coronary surgery required extracorporeal circulation,  
9 but new techniques have facilitated 'off-pump' surgery without circulatory bypass[61].  
10 During surgery reversed saphenous vein, and internal mammary or other arterial  
11 conduits are used to bypass areas of coronary arterial obstruction.

12 Percutaneous transluminal coronary (balloon) angioplasty (PTCA) was established as a  
13 routine treatment for stable angina in the 1980's. The results of coronary balloon  
14 angioplasty were limited by peri-procedural occlusion of the treated artery, and by  
15 recurrence of the arterial stenosis ('restenosis') within a few months in around one third  
16 of patients. The introduction of metallic ('bare metal') coronary artery stents in the  
17 1990's improved the results of percutaneous coronary intervention, but was  
18 associated with the new problems of thrombotic stent occlusion ('stent thrombosis') and  
19 in-stent restenosis. In the last decade the development of drug-eluting stents has  
20 facilitated focal inhibition of the intimal proliferative response to arterial wall injury,  
21 resulting in a reduced risk of in-stent restenosis but a small but important risk of late  
22 stent thrombosis. Meta-analyses of randomised trials confirm that bare metal and  
23 drug eluting coronary stents reduce the risk of restenosis and need for repeat  
24 revascularisation procedures, but have no impact on mortality[62-64].

25 The role of coronary arteriography and myocardial revascularisation in people with  
26 coronary artery disease has been investigated in numerous randomised trials.  
27 Nevertheless, after several decades of research there is persisting uncertainty about  
28 the indications for, and optimal timing of invasive investigation and myocardial  
29 revascularisation in people with stable angina. The trials in this review compared an  
30 initial treatment strategy of continued medical therapy versus an initial treatment  
31 strategy of continued medical therapy and myocardial revascularisation (with  
32 coronary artery bypass surgery or percutaneous coronary intervention).

1           **Evidence review - studies included**

2           The focus of this guideline is the management of stable angina and we only included  
3           studies that had more than 60% stable angina patients.

4           The evidence review includes evidence from RCTs and from individual patient data  
5           (IPD) meta-analyses of medical treatment vs. surgery [65].

6           The RCT evidence addressed three main comparisons:

- 7           • Medical vs. CABG
- 8           • Medical vs. PCI
- 9           • Medical vs. PCI or CABG

10          Some trials selectively recruited patients with single vessel coronary artery disease  
11          but other trials recruited patients with single or multi-vessel disease and/or presented  
12          subgroup analyses by the number of diseased vessels. Definitions for these subgroups  
13          are not universally agreed and results for patients with single and multi-vessel disease  
14          are not reported consistently across the trials. In the evidence reviews we have  
15          combined evidence from trials that included patients with multi-vessel disease, but  
16          results for patients with single vessel disease are considered separately. We also  
17          consider subgroups of older patients, those with two or three vessel disease, and those  
18          with involvement of the left anterior descending artery or with left main stem disease.  
19          Results are presented for three time periods, short term (1 yr), medium term (2-4 yrs),  
20          and longer term follow-up (>4yrs).

21          **Evidence review - outcomes**

22          The main outcomes analysed were:

- 23          • Death (all causes)
- 24          • Cardiac death
- 25          • MI/non fatal MI
- 26          • Stroke
- 27          • Non protocol revascularisation (PCI and/or CABG)
- 28          • Freedom from angina

29          **Evidence review- presentation of results**

30          The results of the review are presented as follows:

31          **A. Medical vs. CABG**

- 32          • Multi-vessel disease – short term follow-up (1 year)

- 1 • Multi-vessel disease - medium term follow-up (2 to 4 years)
- 2 • Multi-vessel disease - long term follow-up (>4 years)
- 3 • Single vessel disease - medium term follow-up (2 to 4 years)
- 4 • Single vessel disease - long term follow-up (>4 years)
- 5 • Left main stem disease - medium term follow-up (2 to 4 years)
- 6 • Left main stem disease - long term follow-up (>4 years)
- 7 • Left anterior descending artery - long term follow-up (>4 years)

8 **B. Medical vs. PCI**

- 9 • Multi-vessel disease - short term follow-up (1 year)
- 10 • Multi-vessel disease - medium term follow-up (2 to 4 years)
- 11 • Multi-vessel disease - long term follow-up (> 4 years follow-up)
- 12 • Single vessel disease - medium term follow-up (2 -4 years)
- 13 • Single vessel disease - long term follow-up (>4 years)

14 **C. Medical vs. PCI or CABG**

- 15 • Multi-vessel disease - short term follow-up (1 year)
- 16 • Multi-vessel disease - medium term follow-up (2 to 4 years)
- 17 • Multi-vessel disease - long term follow-up (>4 years)

18 The narrative summary of the outcome 'Quality of life' is presented separately for  
19 each of the above comparisons (as data was not analysed for this outcome).

20

21 **11.2 Medical interventions versus CABG**

22 **11.2.1 Clinical question**

23 What is the clinical and cost effectiveness of medical interventions versus CABG in  
24 people with stable angina?

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1    **11.2.2    Clinical evidence**

2            The “Review Protocol” for this topic can be found in Appendix C, the “Search  
3            Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
4            E1, the “Clinical Evidence Tables” in Appendix E2, and the “Forest Plots” in Appendix  
5            F.

6

1 **Table 11.1: Multi-vessel disease - Short term follow-up (1 year) for stable angina**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Multi vessel disease- Short term follow-up	CABG	Relative (95% CI)	Absolute	
<b>Death (follow-up 1 year)</b>											
Hueb 2004[66] (MASS-II)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	None	3/203 (1.5%)	8/203 (3.9%)	RR 0.38 (0.1 to 1.39)	24 fewer per 1000 (from 35 fewer to 15 more)	⊕⊕○○ LOW
<b>Q wave MI (follow-up 1 year)</b>											
Hueb 2004[66] (MASS-II)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	None	10/203 (4.9%)	4/203 (2%)	RR 2.5 (0.8 to 7.84)	30 more per 1000 (from 4 fewer to 135 more)	⊕⊕○○ LOW
<b>Stroke (follow-up 1 year)</b>											
Hueb 2004[66] (MASS-II)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	None	3/203 (1.5%)	3/203 (1.5%)	RR 1 (0.2 to 4.9)	0 fewer per 1000 (from 12 fewer to 58 more)	⊕⊕○○ LOW
<b>Non protocol revascularisation (follow-up 1 year)</b>											
Hueb 2004[66] (MASS-II)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	16/203 (7.9%)	1/203 (0.5%)	RR 16 (2.14 to 119.52)	74 more per 1000 (from 6 more to 584 more)	⊕⊕⊕○ MODERATE
<b>Free of angina (follow-up 1 year)</b>											
Hueb 2004[66] (MASS-II)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (c)	None	74/203 (36.5%)	120/203 (59.1%)	RR 0.62 (0.5 to 0.76)	225 fewer per 1000 (from 142 fewer to 296 fewer)	⊕⊕○○ LOW
<b>Death- subgroup diabetes (follow-up 1 year)</b>											
Soares 2006[67] (MASS-II)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	None	2/75 (2.7%)	4/59 (6.8%)	RR 0.39 (0.07 to 2.07)	41 fewer per 1000 (from 63 fewer to 73 more)	⊕⊕○○ LOW
<b>Death- subgroup no diabetes (follow-up 1 year)</b>											
Soares 2006[67] (MASS-II)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	None	2/128 (1.6%)	7/144 (4.9%)	RR 0.32 (0.07 to 1.52)	33 fewer per 1000 (from 45 fewer to 25 more)	⊕⊕○○ LOW

- 2 (a) Randomised. ITT reported. Allocation concealment unclear.  
 3 (b) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.  
 4 (c) 95% CI around the pooled estimate of effect includes appreciable benefit or appreciable harm.

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**Table 11.2: Multi-vessel disease - Medium term follow-up (2 to 4 years) for stable angina**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Medical	CABG	Relative (95% CI)	Absolute	
<b>Death (follow-up 2-4 years)</b>											
Read 1978[68] (VA); Varnauskas 1980[69] (ECSS)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	89/727 (12.2%)	67/726 (9.2%)	RR 1.29 (0.96 to 1.74)	27 more per 1000 (from 4 fewer to 68 more)	⊕⊕○○ LOW
<b>Cardiac death (follow-up 2 years)</b>											
Varnauskas 1980[69] (ECSS)	randomised trials	serious (c)	no serious inconsistency	no serious indirectness	no serious imprecision	none	27/373 (7.2%)	10/394 (2.5%)	RR 2.85 (1.4 to 5.81)	47 more per 1000 (from 10 more to 122 more)	⊕⊕⊕○ MODERATE
<b>MI (follow-up 2-2.8 years)</b>											
Guinn 1976[70]	randomised trials	serious (d)	no serious inconsistency	no serious indirectness	serious (b)	none	11/60 (18.3%)	5/56 (8.9%)	RR 2.05 (0.76 to 5.54)	94 more per 1000 (from 21 fewer to 405 more)	⊕⊕○○ LOW
<b>Free of angina (follow-up 2-2.8 years)</b>											
Guinn 1976[70]; Varnauskas 1980[69] (ECSS)	randomised trials	serious (e)	serious (f)	no serious indirectness	no serious imprecision	none	180/433 (41.6%)	353/450 (78.4%)	RR 0.53 (0.47 to 0.60)	369 fewer per 1000 (from 314 fewer to 416 fewer)	⊕⊕○○ LOW
<b>Death - sub group 2 vessel disease (follow-up 2 years)</b>											
Varnauskas 1980[69] (ECSS)	randomised trials	serious (c)	no serious inconsistency	no serious indirectness	serious (b)	none	6/154 (3.9%)	10/147 (6.8%)	RR 0.57 (0.21 to 1.54)	29 fewer per 1000 (from 54 fewer to 37 more)	⊕⊕○○ LOW
<b>Death - sub group 3 vessel disease (follow-up 2-4 years)</b>											
Detre 1977[71] (VA); Varnauskas 1980[69] (ECSS)	randomised trials	serious (g)	no serious inconsistency	no serious indirectness	serious (b)	none	46/346 (13.3%)	28/354 (7.9%)	RR 1.57 (1.02 to 2.44)	45 more per 1000 (from 2 more to 114 more)	⊕⊕○○ LOW
<b>Non protocol revascularisation (follow-up 2.8 years)</b>											
Guinn 1976[70]	randomised trials	serious (h)	no serious inconsistency	no serious indirectness	serious (b)	none	4/60 (6.7%)	1/56 (1.8%)	RR 3.73 (0.43 to 32.4)	49 more per 1000 (from 10 fewer to 561 more)	⊕⊕○○ LOW

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(a) Randomised, ITT reported in all studies. Allocation not reported in all studies. No heterogeneity  $I^2=0\%$   
 (b) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.  
 (c) Strengths: randomised. Low attrition bias. Intention to treat analysis used. Weaknesses: reporting of outcome is not always very clear; crossover 26/394 (6.5%) of patients assigned to surgery did not complete treatment; medical group 50/373 (13%) had surgery; unclear allocation concealment

- 1 (d) *Strengths: Randomised. No loss to follow-up. Baseline comparisons made. Intention to treat analysis reported. Limitations: allocation concealment not reported. No*  
2 *heterogeneity  $I^2=0\%$*   
3 (e) *Randomised, ITT reported in all. Allocation concealment not reported in both studies.*  
4 (f) *High heterogeneity -  $I^2=93\%$*   
5 (g) *Randomised, ITT used in both the studies. Allocation concealment not reported in both.*  
6 (h) *Strengths: Randomised. No loss to follow-up. Baseline comparisons made. Intention to treat analysis reported. Limitations: allocation concealment not reported.*  
7

8 **Sub group interaction**

9 *There was no significant difference between sub group of patients with 2 vessel or 3 vessel disease for death ( $p=0.07$ ) at medium term follow-up (2- to 4 years).*

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**Table 11.3: Multi-vessel disease -Long term follow-up (>4 years) for stable angina**

Quality assessment							Summary of findings				
							No of patients		Effect		Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Medical	CABG	Relative (95% CI)	Absolute	
<b>Death (follow-up 5-22 years)</b>											
Alderman 1990[72] (CASS); Frick 1985[73]; Kloster 1979[74]; Peduzzi 1998[75] (VA); Varnauskas 1988[76] (ECSS); Hueb 2010[77] (MASS-II)	randomised trials	serious (a)	serious (b)	no serious indirectness	no serious imprecision	none	533/1419 (37.6%)	484/1415 (34.2%)	RR 1.08 (0.99 to 1.17)	27 more per 1000 (from 3 fewer to 58 more)	⊕⊕○○ LOW
<b>Cardiac death (follow-up 12 years)</b>											
Bhayana 1980[78] (VA); Varnauskas 1988[76] (ECSS)	randomised trials	serious (c)	serious (d)	no serious indirectness	serious (e)	none	112/448 (25%)	79/465 (17%)	RR 1.44 (1.12 to 1.84)	75 more per 1000 (from 20 more to 143 more)	⊕○○○ VERY LOW
<b>MI (follow-up 5-22 years)</b>											
Fisher 1985[79] (CASS); Kloster 1979[74]; Peduzzi 1998[75] (VA); Hueb 2010[77] (MASS-II)	randomised trials	serious (f)	serious (g)	no serious indirectness	no serious imprecision	none	216/996 (21.7%)	221/976 (22.6%)	RR 0.94 (0.80 to 1.10)	14 fewer per 1000 (from 45 fewer to 23 more)	⊕⊕○○ LOW
<b>Free of angina (follow-up 5-15 years)</b>											
Peduzzi 1992[80] (VA); Rogers 1990[81] (CASS); Varnauskas 1982[82] (ECSS); Hueb 2010[77] (MASS-II)	randomised trials	serious (h)	serious (i)	no serious indirectness	serious (j)	none	365/1320 (27.7%)	507/1319 (38.4%)	RR 0.73 (0.66 to 0.81)	104 fewer per 1000 (from 73 fewer to 131 fewer)	⊕○○○ VERY LOW
<b>Stroke (follow-up 10 years)</b>											
Hueb 2010[77] (MASS-II)	randomised trials	serious (k)	no serious inconsistency	no serious indirectness	serious (e)	none	14/203 (6.9%)	17/203 (8.4%)	RR 0.82 (0.42 to 1.63)	15 fewer per 1000 (from 49 fewer to 53 more)	⊕⊕○○ LOW
<b>Non protocol revascularisation (follow-up 10-22 years)</b>											
Peduzzi 1998[75] (VA); Rogers 1990[81] (CASS); Hueb 2010[77] (MASS-II)	randomised trials	serious (l)	serious (m)	no serious indirectness	no serious imprecision	none	442/947 (46.7%)	142/925 (15.4%)	RR 3.02 (2.56 to 3.55)	310 more per 1000 (from 239 more to 391 more)	⊕⊕○○ LOW

Death- sub group 2 vessel disease (follow-up 5-12 years)											
Alderman 1990[72] (CASS); Kloster 1979[74]; Varnauskas 1982[82] (ECSS)	randomised trials	serious (n)	no serious inconsistency	no serious indirectness	serious (j)	none	53/321 (16.5%)	33/324 (10.2%)	RR 1.64 (1.1 to 2.45)	65 more per 1000 (from 10 more to 148 more)	⊕⊕○○ LOW
Death- sub group 3 vessel disease (follow-up 5-12 years)											
Alderman 1990[72] (CASS); Kloster 1979[74]; Varnauskas 1982[82] (ECSS)	randomised trials	serious (n)	serious (o)	no serious indirectness	serious (j)	none	71/343 (20.7%)	49/368 (13.3%)	RR 1.48 (1.07 to 2.06)	64 more per 1000 (from 9 more to 141 more)	⊕○○○ VERY LOW
Death age >53 yrs (follow-up 10 years)											
Alderman 1990[72] (CASS)	randomised trials	serious (p)	no serious inconsistency	no serious indirectness	serious (e)	none	46/163 (28.2%)	39/163 (23.9%)	RR 1.18 (0.82 to 1.7)	43 more per 1000 (from 43 fewer to 167 more)	⊕⊕○○ LOW
Death - age <47 years (follow-up 10 years)											
Alderman 1990[72] (CASS)	randomised trials	serious (p)	no serious inconsistency	no serious indirectness	serious (e)	none	16/101 (15.8%)	17/92 (18.5%)	RR 0.86 (0.46 to 1.6)	26 fewer per 1000 (from 100 fewer to 111 more)	⊕⊕○○ LOW
Death - age 47-53 years (follow-up 10 years)											
Alderman 1990[72] (CASS)	randomised trials	serious (p)	no serious inconsistency	no serious indirectness	serious (e)	none	23/126 (18.3%)	16/135 (11.9%)	RR 1.54 (0.85 to 2.78)	64 more per 1000 (from 18 fewer to 211 more)	⊕⊕○○ LOW

- 1 (a) Randomised, ITT used in all studies. Allocation concealment not reported in all 7 studies.
- 2 (b) Considerable heterogeneity  $I^2=71\%$
- 3 (c) Randomised, unclear allocation concealment in both the studies. ECSS- ITT used. Loss to follow-up not reported (Bhayana 1980)[78].
- 4 (d) Substantial heterogeneity  $-I^2=79\%$
- 5 (e) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.
- 6 (f) Randomised, ITT reported in all 4 studies. Allocation concealment not reported in all 4 studies. Loss to follow-up not reported (Kloster 1979)[74].
- 7 (g)  $I^2=73\%$
- 8 (h) Randomised, ITT used in all 4 studies. Allocation concealment not reported in all 4 studies.
- 9 (i) Substantial heterogeneity  $-I^2=70\%$
- 10 (j) 95% CI around the pooled estimate of effect includes appreciable benefit or appreciable harm.
- 11 (k) Randomised. ITT reported. Allocation concealment unclear.
- 12 (l) Randomised, ITT used in both studies. Allocation concealment not reported in both studies.
- 13 (m) Substantial heterogeneity  $-I^2=82\%$
- 14 (n) Randomised in all studies. Loss to follow-up and ITT not reported in one study (Kloster 1979)[74]. Allocation concealment not reported in all 3 studies.
- 15 (o) Substantial heterogeneity  $-I^2=75\%$

## DRAFT FOR CONSULTATION

1 (p) *Strengths: randomised (stratified randomisation). Baseline comparisons made. Intention to treat analysis reported. Limitations: Allocation concealment not reported.*

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3 **Sub group interaction:**

4 *There was no significant difference between sub groups 2 vessel and 3 vessel disease for death ( $p=0.70$ ) at long term follow-up (5-12 years).*

5 *There was no significant difference between sub groups age <47 years, 47-53 years and >53 years for death ( $p= 0.41$ ) at long term follow-up (10 years)*

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1 **Table 11.4: Single vessel disease – Medium term follow-up (2-4 years) for stable angina**

Quality assessment							Summary of findings				
							No of patients		Effect		Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Single vessel disease- medium term follow-up Medical	CABG	Relative (95% CI)	Absolute	
<b>Death (follow-up 3 years)</b>											
Hueb 1995[83] (MASS- I)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	None	0/72 (0%)	1/70 (1.4%)	RR 0.32 (0.01 to 7.83)	10 fewer per 1000 (from 14 fewer to 98 more)	⊕⊕○○ LOW
<b>Stroke (follow-up 3 years)</b>											
Hueb 1995[83] (MASS- I)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	0/72 (0%)	0/70 (0%)	not pooled	not pooled	⊕⊕⊕○ MODERATE
<b>MI (follow-up 3 years)</b>											
Hueb 1995[83] (MASS- I)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	None	2/72 (2.8%)	1/70 (1.4%)	RR 1.94 (0.18 to 20.96)	13 more per 1000 (from 12 fewer to 285 more)	⊕⊕○○ LOW
<b>Non protocol revascularisation (follow-up 3 years)</b>											
Hueb 1995[83] (MASS- I)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	None	7/72 (9.7%)	0/70 (0%)	RR 14.59 (0.85 to 250.71)	100 more per 1000 (from 20 more to 170 more)	⊕⊕○○ LOW
<b>Free of angina (follow-up 3 years)</b>											
Hueb 1995[83] (MASS- I)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	23/72 (31.9%)	68/70 (97.1%)	RR 0.33 (0.23 to 0.46)	651 fewer per 1000 (from 525 fewer to 748 fewer)	⊕⊕⊕○ MODERATE

- (a) Strengths: Randomised. Baseline comparisons made. Number of patients lost to follow-up not reported. ITT reported. Limitations: allocation concealment not reported. Blinding of outcome assessors not reported.
- (b) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm

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1 **Table 11.5: Single vessel disease - Long term follow-up (>4 years) for stable angina**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Single vessel disease -Long term follow-up Medical	CABG	Relative (95% CI)	Absolute	
<b>Death (follow-up 5-10 years)</b>											
Alderman 1990[72] (CASS); Kloster 1979[74]; Hueb 1999[84] (MASS-I)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	None	26/189 (13.8%)	18/185 (9.7%)	RR 1.41 (0.81 to 2.46)	40 more per 1000 (from 18 fewer to 142 more)	⊕⊕○○ LOW
<b>Cardiac death (follow-up 5 years)</b>											
Hueb 1999[84] (MASS-I)	randomised trials	serious (c)	no serious inconsistency	no serious indirectness	serious (b)	None	2/72 (2.8%)	2/70 (2.9%)	RR 0.97 (0.14 to 6.71)	1 fewer per 1000 (from 25 fewer to 163 more)	⊕⊕○○ LOW
<b>MI (follow-up 5 years)</b>											
Hueb 1999[84] (MASS-I)	randomised trials	serious (c)	no serious inconsistency	no serious indirectness	serious (b)	None	3/72 (4.2%)	3/70 (1.4%)	RR 0.97 (0.20 to 4.66)	27 more per 1000 (from 10 fewer to 377 more)	⊕⊕○○ LOW
<b>Stroke (follow-up 5 years)</b>											
Hueb 1999[84] (MASS-I)	randomised trials	serious (c)	no serious inconsistency	no serious indirectness	serious (b)	None	1/72 (1.4%)	1/70 (1.4%)	RR 0.97 (0.06 to 15.24)	0 fewer per 1000 (from 13 fewer to 203 more)	⊕⊕○○ LOW
<b>Non protocol revascularisation (follow-up 5 years)</b>											
Hueb 1999[84] (MASS-I)	randomised trials	serious (c)	no serious inconsistency	no serious indirectness	no serious imprecision	None	12/72 (16.7%)	0/70 (0%)	RR 24.32 (1.47 to 402.97)	170 more per 1000 (from 80 more to 260 more).	⊕⊕⊕○ MODERATE
<b>Free of angina (follow-up 5 years)</b>											
Hueb 1999[84] (MASS-I)	randomised trials	serious (c)	no serious inconsistency	no serious indirectness	no serious imprecision	None	17/72 (23.6%)	48/70 (68.6%)	RR 0.34 (0.22 to 0.54)	453 fewer per 1000 (from 315 fewer to 535 fewer)	⊕⊕⊕○ MODERATE

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- (a) All 3 Randomised. ITT not reported in Kloster 1979[74]. Allocation concealment not reported in all 3 papers.
- (b) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.
- (c) Randomised, Baseline comparisons made. Number of patients lost to follow-up not reported. ITT reported. Limitations: allocation concealment not reported. Blinding of outcome assessors not reported.

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**Table 11.6: Left main stem disease - Medium term follow-up (2 to 4 years) for stable angina**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Left main stem disease-Medium term follow-up (2 to 4 years) Medical	CABG	Relative (95% CI)	Absolute	
<b>Death (follow-up 2-4 years)</b>											
Detre 1977[71] (VA); Varnauskas 1980[69] (ECSS)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	20/75 (26.7%)	5/74 (6.8%)	RR 4 (1.6 to 10.03)	203 more per 1000 (from 41 more to 610 more)	⊕⊕⊕O MODERATE

(a) Randomised, ITT used in both studies. Allocation concealment not reported in both studies. Low heterogeneity  $I^2=19\%$

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**Table 11.7: Left main stem disease - Long term follow-up (>4 years) for stable angina**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Left main stem disease-Long term follow-up (>4 years) Medical	CABG	Relative (95% CI)	Absolute	
<b>Death (follow-up 10-22 years)</b>											
Alderman 1990[72] (CASS); Peduzzi 1998[75] (VA); Varnauskas 1982[82] (ECSS)	randomised trials	serious (a)	serious (b)	no serious indirectness	serious (c)	None	51/80 (63.8%)	47/84 (56%)	RR 1.18 (0.97 to 1.43)	101 more per 1000 (from 17 fewer to 241 more)	⊕○○○ VERY LOW
<b>MI (follow-up 22 years)</b>											
Peduzzi 1998[75] (VA)	randomised trials	serious (d)	no serious inconsistency	no serious indirectness	serious (c)	None	16/43 (37.2%)	21/48 (43.8%)	RR 0.85 (0.51 to 1.41)	66 fewer per 1000 (from 214 fewer to 179 more)	⊕⊕○○ LOW

(a) Randomised, ITT reported in all 3 studies. Allocation concealment not reported all 3 studies.

(b) Substantial heterogeneity  $I^2=79\%$

(c) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.

(d) Strengths: Randomised, baseline comparisons made. Intention to treat analysis reported. Limitations: allocation concealment not reported

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1 **Table 11.8: Left anterior descending artery - Long term follow-up (>4 years) for stable angina**

Quality assessment							Summary of findings				Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		
							Left anterior descending artery - Long term follow-up (>4 years) Medical	CABG	Relative (95% CI)	Absolute	
<b>Death (follow-up 10-12 years)</b>											
Alderman 1990[72] (CASS); Varnauskas 1988[76] (ECSS)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	None	144/515 (28%)	113/539 (21%)	RR 1.34 (1.09 to 1.66)	71 more per 1000 (from 19 more to 138 more)	⊕⊕⊕⊕ LOW

(a) Randomised, ITT used in both studies. Allocation concealment not reported in both studies. No heterogeneity  $I^2=0\%$

(b) 95% CI around the pooled estimate of effect includes appreciable benefit or appreciable harm.

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1 **Table 11.9: IPD meta analyses [Medical vs. CABG] – Multivessel disease – Long term follow-up**

Quality assessment							Summary of findings				Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		
							IPD meta analyses Medical	CABG	Odds ratio (95% CI)	Absolute	
<b>Total mortality (follow-up 10 years)</b>											
Yusuf 1994[65] (c)	randomised trial	serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	404/1325 (30.5%)	350/1324 (26.4%)	OR 0.83 (0.70 to 0.98)	38 fewer per 1000 (from 4 fewer to 69 fewer)	⊕⊕⊕O MODERATE
<b>Mortality - sub group one vessel disease (follow-up 5 years)</b>											
Yusuf 1994[65]	randomised trial	serious limitations (b)	no serious inconsistency	no serious indirectness	no serious imprecision	None	Not reported	Not reported	OR 0.54 (0.22 to 1.33)	Cannot be calculated	⊕⊕⊕O MODERATE
<b>Mortality - subgroup 2 vessels (follow-up 5 years)</b>											
Yusuf 1994[65]	randomised trial	serious limitations (b)	no serious inconsistency	no serious indirectness	no serious imprecision	None	Not reported	Not reported	OR 0.84 (0.54 to 1.32)	Cannot be calculated	⊕⊕⊕O MODERATE
<b>Mortality- sub group 3 vessels</b>											
Yusuf 1994[65]	randomised trial	serious limitations (b)	no serious inconsistency	no serious indirectness	no serious imprecision	None	Not reported	Not reported	OR 0.58 (0.42 to 0.80)	Cannot be calculated	⊕⊕⊕O MODERATE
<b>Mortality- sub group Left main artery (follow-up 5 years)</b>											
Yusuf 1994[65]	randomised trial	no serious limitations (b)	no serious inconsistency	no serious indirectness	no serious imprecision	None	Not reported	Not reported	OR 0.32 (0.15 to 0.70)	Cannot be calculated	⊕⊕⊕O MODERATE
<b>Mortality- LAD disease present</b>											
Yusuf 1994[65]	randomised trial	serious limitations (b)	no serious inconsistency	no serious indirectness	no serious imprecision	None	Not reported	Not reported	OR 0.50 (0.43 to 0.77)	Cannot be calculated	⊕⊕⊕O MODERATE
<b>Mortality- sub group normal LV function (follow-up 5 years)</b>											
Yusuf 1994[65]	randomised trial	serious limitations (b)	no serious inconsistency	no serious indirectness	no serious imprecision	None	Not reported	Not reported	OR 0.61 (0.46 to 0.81)	Cannot be calculated	⊕⊕⊕O MODERATE
<b>Mortality- sub group abnormal LV function (follow-up 5 years)</b>											
Yusuf 1994[65]	randomised trial	serious limitations (b)	no serious inconsistency	no serious indirectness	no serious imprecision	None	Not reported	Not reported	OR 0.59 (0.39 to 0.91)	Cannot be calculated	⊕⊕⊕O MODERATE

Mortality- subgroup class 0,I,II (follow-up 5 years)											
Yusuf 1994[65]	randomised trial	serious limitations (b)	no serious inconsistency	no serious indirectness	no serious imprecision	None	Not reported	Not reported	OR 0.63 (0.46 to 0.87)	Cannot be calculated	⊕⊕⊕○ MODERATE
Mortality- subgroup class III,IV (follow-up 5 years)											
Yusuf 1994[65]	randomised trial	serious limitations (b)	no serious inconsistency	no serious indirectness	no serious imprecision	None	Not reported	Not reported	OR 0.57 (0.40 to 0.81)	Cannot be calculated	⊕⊕⊕○ MODERATE

- 1 (a) This is an IPD (Individual patient data) meta analyses. This review addresses an appropriate and clearly focused question. The review included only RCTs which was relevant  
2 to the review question. There was adequate description of the methodology used in the meta analyses. The mortality analysis was an ITT (irrespective of crossover between  
3 treatments or failure of CABG patients to receive surgery). The paper does not report the search strategy used. The IPD meta analyses did not look at the longest follow-up  
4 of the VA trial comparing medical treatment to surgery in stable angina patients (22 years for VA study). Quality assessment of individual studies not reported\*. This meta  
5 analyses did not include all studies relevant to the question. We have separately assessed the quality of individual studies in the evidence review. Additional studies have  
6 been included in the study level meta analyses conducted by us. Note: One study (Norris 1981)[85] from this meta analyses was not included in our evidence review as the  
7 study did not meet our inclusion criteria (study population was recurrent MI).  
8 (b) None of the included studies reported of allocation concealment. Sub group analyses conducted for selected sub groups. If a study had no event in a given subgroup, it was  
9 omitted from the analysis for that sub group.  
10 (c) Studies included in this IPD - Norris 1981[85], Mathur 1977[86] (ECSS), Kloster 1979[74] (CASS).  
11

12 **Extension of survival (Yusuf IPD[65] meta analyses)**

13 Analysis of overall survival during the first 10 years after randomisation showed an improvement in survival with CABG surgery over medical treatment of 4.26 months with a  
14 1.96 SE of 2.35 months. The benefit seemed to increase with disease severity. The improvement in survival was greatest for patients with left main artery disease, intermediate  
15 for those with three vessel disease, and least for those with one vessel or two vessel disease (p=0.02 for trend) . Greater survival prolongation was also found among patients  
16 with abnormal exercise tests (p for interaction 0.71) and abnormal LV function (p<0.01) than in patients without these characteristics.

1 **Quality of Life data from studies:**

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3 **Alderman 1993[87] (CASS) – 5 year follow-up:**

4 In this RCT, Quality of Life was derived from a composite of subjective (questionnaire)  
5 and objective (exercise test) measures. Patients reported symptoms such as chest pain  
6 status, heart failure, activity limitation employment and recreational status, drug  
7 therapy, hospitalisation, smoking and supervised exercise program (i.e. whether the  
8 patient participated in such activities during the 2 months before follow-up). In  
9 addition blood pressure, cholesterol and weight was measure and patients took part  
10 in an exercise test.

11 *Results:* n=780 (n=390 in surgery and n=390 in CABG). At a mean of 5.5 years  
12 follow-up patients in the surgical group had significantly less chest pain, fewer activity  
13 limitations and required less therapy with nitrates and  $\beta$  -blockers. There was no  
14 significant difference in self reported symptoms of heart failure, employment and  
15 recreational status. The number of days in hospital was higher in the surgical  
16 compared to the medical group. All these results are graphically presented in the  
17 paper, but individual mean results are not given in the text. Treadmill exercise test  
18 results were significantly better for the surgical compared to the medical group (time  
19 to induced angina and ST segment depression). There were no significant differences  
20 in cholesterol levels, blood pressure, body weight and levels of smoking between the  
21 two groups. From these results the researchers deduced that CABG improves the  
22 quality of life as manifest by relief of chest pain, improvement in both subjective and  
23 objective measurements of functional status, and a diminished requirement for drug  
24 therapy.

25  
26 **Rogers 1990[81] (CASS) – 10 year follow-up:**

27 Same measures as described above.

28 *Results:* N=654 remaining at 10 year follow-up (CABG (n=333); Medical (n=324)).  
29 *Chest pain.* At 10 years the proportion of surgical patients who were asymptomatic  
30 had declined from medium term follow-up (to 47%) and the proportion of medical  
31 patients who were asymptomatic had increased (42%), which remains nonetheless a  
32 significant difference in favour of the surgical group. *Heart failure.* Absence of heart  
33 failure symptoms was reported by 72% in the medical and 75% in the surgical group  
34 (p=ns). *Activity limitations.* Proportion of patients without activity limitations at 10 year  
35 follow-up was not significant (34% vs. 28%). *Employment status.* 34% of the surgical  
36 group and 32% of the medical treatment group were employed after 10 years  
37 (p=ns). *Recreational status* did not differ after 10 years (proportion of patients who  
38 participate in moderate-strenuous exercise: 25% medical group and 26% surgical  
39 group. The authors concluded from this that the advantage reported in their Quality  
40 of Life assessment at short to medium follow-up length were much less apparent after  
41 a 10 year interval.

1 **11.2.3 Economic evidence**

2 One study[88] was found that included the relevant comparison. This is summarised in  
 3 the economic evidence profile below. See also Economics Evidence Tables in  
 4 Appendix G.

5 **Table 11.10: CABG vs. medical treatment - Economic study characteristics**

Study	Limitations	Applicability	Other Comments
Griffin 2007[88]	Potentially serious limitations (a)	Partial applicability (b)	Patients included in the analysis were those who had coronary angiography between April 1996 and April 1997 at three hospitals of one NHS trust in London and who were suitable for both CABG and PCI. Their suitability was assessed using the RAND appropriateness method. PCI was assessed as well but CABG was more cost-effective in this study.

- 6 e) *Not a randomised study; EQ-5D data were not collected at baseline and at one year and scores were*  
 7 *only predicted at these time points from other variables.*  
 8 f) *Criteria for assessment of the suitability for revascularisation could have changed since time of study.*  
 9 *PCI procedure could have been without stents.*

10 **Table 11.11: CABG vs. medical treatment - Economic summary of findings**

Study	Incremental cost per patient over 6 years (£)	Incremental QALYs per patient over 6 years	ICER (£/QALY)	Uncertainty
Griffin 2007[88]	7,169 (a, b)	0.3 (b, c)	18,603	For patients deemed appropriate for CABG only, ICER=£14,675/QALY At a threshold of £20,000/QALY all the strategies including PCI have a similar probability of being cost-effective.

- 12 (a) *2004 GBP. Costs included were cost of intervention, angiography, hospital stay, drugs, admissions for*  
 13 *chest pain, GP and outpatient visits, visits to the emergency department. Occurrence of admissions and*  
 14 *LOS obtained from the NHS-wide clearing service; data on drugs from hospital case notes, GP and*  
 15 *patients' questionnaires; unit costs from published studies and pricing lists for the UK.*  
 16 (b) *Discounted by 3.5%*  
 17 (c) *Based on EQ-5D data from participants in the study.*  
 18

19 **11.2.4 Evidence statements**

**Clinical**

**Multi -vessel disease- short term follow-up (1 year)**

**Soares 2006[67] (MASS-II):** Evidence from one RCT shows that there were significantly higher patients free of angina [RR 0.62 (0.5 to 0.76)] in CABG group compared to medical treatment group. There were significantly higher repeat revascularisations [RR 16 (2.14 to 119.52)] in the medical treatment group compared to CABG. There was no significant difference between medical treatment group and PCI for death [RR 0.38 (0.1 to 1.39)], MI [RR 2.5 (0.8 to 7.84)], stroke [RR 1 (0.2 to 4.9)]. There was no significant difference between medical treatment and CABG for

death in a sub group of patients with diabetes [RR 0.39 (0.07 to 2.07)] and no diabetes [RR 0.32 (0.07 to 1.52)]. [Follow-up 1 year]

**Multi -vessel disease- Medium term follow-up (2 to 4 years)**

**Read 1977[68] (VA); Varnauskas 1980[69] (ECSS):** Evidence from 2 RCTS's shows that there was no significant difference between medical treatment and CABG for deaths [RR 1.29 (0.96 to 1.74)]. [Follow-up 2 to 4 years]

**Varnauskas 1980[69] (ECSS):** Evidence from one RCT shows that there was significantly fewer death in the CABG compared to medical treatment. RR 2.85 (1.4 to 5.81) [follow-up 2 years]

**Guinn 1976[70] (VA):** Evidence from one RCTs to show that there was no significant difference between medical treatment and CABG for MI in the CABG [RR 2.05(0.76 to 5.54)]. [Follow-up 2.8 years]

**Guinn 1976[70] (VA); Varnauskas 1979[69] (ECSS):** Evidence from 2 RCTs shows that there were significantly more patients free of angina in the CABG group compared to medical treatment group. [RR 0.53 (0.47 to 0.60)]. [Follow-up 2 to 2.8 years]

**Varnauskas 1980[69] (ECSS):** Evidence from one RCT shows that there was no significant difference between medical treatment and CABG group for death in sub group 2 vessel disease RR 0.57 (0.21 to 1.54) .[Follow-up 2 years]

**Detre 1977[71] (VA); Varnauskas 1980[69] (ECSS):** Evidence from 2 RCTs shows that there were significantly fewer deaths in the CABG group compared to medical treatment [RR 1.57 (1.02 to 2.44)] in patients with sub group 3 vessel disease. [follow-up 2-4 years]]. But there was no significant difference between sub group of patients with 2 vessel or 3 vessel disease for death (p=0.07) at medium term follow-up (2- to 4 years).

**Guinn 1976[70]:** Evidence from one RCT shows that there was no significant difference between medical treatment and CABG for non protocol revascularisation. [RR 3.73 (0.43 to 32.4)]. [follow-up 2.8 years]

**Multi -vessel disease - Long term follow-up (>4 years)**

**Alderman 1990[72] (CASS); Frick 1985[73]; Kloster 1979[74]; Peduzzi 1998[75] (VA); Varnauskas 1988[76] (ECSS); Hueb 2010[77] (MASS –II):** Evidence from 6 RCTs shows that there was no significant difference between medical treatment and CABG for death [RR 1.08 (0.99 to 1.17)]. [Follow-up 5 to 22 years]

**Bhayana 1978[78] (VA); Varnauskas 1988[76] (ECSS):** Evidence from 2 RCTs shows that there was significantly fewer cardiac death in CABG compared to medical treatment [RR 1.44 (1.12 to 1.84)] [Follow-up 12 years]

**Fisher 1984[79] (CASS); Kloster 1979[74]; Peduzzi 1998[75] (VA); Hueb 2010[77] (MASS –II):** Evidence from 4 RCTs shows that there was no significant difference between medical treatment and CABG for MI [RR 0.94 (0.80 to 1.10)]. [Follow-up 5-22 years]

**Peduzzi 1992[80] (VA); Rogers 1990[81] (CASS); Varnauskas 1982[82] (ECSS); Hueb 2010[77] (MASS –II):** Evidence from 4 RCTs show that there were significantly more patients free of angina in the CABG group compared to medical treatment. [RR 0.73 (0.66 to 0.81)]. [Follow-up 5-15years]

**Peduzzi 1998[75] (VA); Rogers 1990[81] (CASS); Hueb 2010[77] (MASS –II):** Evidence from 3 RCTs shows that there were significantly more patients with non protocol revascularisation in the medical treatment group compared to CABG [RR 3.02 (2.56 to 3.55)]. [Follow-up 10-22 years]

**Hueb 2010[77] (MASS –II):** Evidence from 1 RCT shows that there was no significant difference between medical treatment and CABG for stroke [RR 0.82 (0.42 to 1.63)] [Follow-up 10 years]

**Alderman 1990[72] (CASS); Kloster 1979[74]; Varnauskas 1982[82] (ECSS):** Evidence from 3 RCTs' shows that there was significantly fewer deaths in the CABG group compared to medical treatment in a sub group of people with 2 vessel disease RR 1.64 (1.1 to 2.45) and 3 vessel disease RR 1.48 (1.07 to 2.06). [Follow-up 5-12 years] But there was no significant difference between sub groups 2 vessel and 3 vessel disease for death ( $p=0.70$ ) at long term follow-up (5-12 years).

**Alderman 1990[72] (CASS):** Evidence from one RCT shows that there was no significant difference between medical treatment and CABG for death in a sub group of people age >53 years. RR 1.18 (0.82 to 1.7). [Follow-up 10 years]

**Alderman 1990[72] (CASS):** Evidence from one RCT shows that there was no significant difference between medical treatment and CABG for death in a sub group of people age <47 years. RR 0.86 (0.46 to 1.60). [Follow-up 10 years]

**Alderman 1990[72] (CASS):** Evidence from one RCT shows that there was no significant difference between medical treatment and CABG for death in a sub group of people age 47-53 years. RR 1.54 (0.85 to 2.78). [Follow-up 10 years] But there was no significant difference between sub groups age <47 years, 47-53 years and >53 years for death ( $p= 0.41$ ) at long term follow-up

(10 years)

**Single vessel disease – Medium term follow-up (2- 4 years)**

**Hueb 1995[83] (MASS- I):** Evidence from one RCT shows that there were statistically significant higher no. of patients free of angina in the CABG group compared to medical treatment [RR 0.33 (0.23 to 0.46)]. There was no statistically significant difference medical and CABG for death [RR 0.32 (0.01 to 7.83)], stroke [0/72 in medical and 0/70 in CABG], MI [RR 1.94 (0.18 to 20.96)], and non protocol revascularisation [RR 14.59 (0.85 to 250.71)] [Follow-up 3 years]

**Single vessel disease - Long term follow-up (>4 years)**

**Alderman 1990[72] (CASS); Kloster 1979[74]; Hueb 1999[84] (MASS-I):** Evidence from 3 RCTs shows that there was no statistically significant difference between medical treatment and CABG for death [RR 1.41 (0.81 to 2.46)] [Follow-up 5-10 years]

**Hueb 1999[84] (MASS-I):** Evidence from one RCT shows that significantly higher no. of patients free of angina in the CABG group compared to medical treatment [RR 0.34 (0.22 to 0.54)]. There was significantly higher non protocol revascularisation in the medical treatment group compared to CABG group [RR 24.32 (1.47 to 402.97)]. There was no significant difference between medical treatment and CABG group for cardiac death [RR 0.97 (0.14 to 6.71)], MI [RR 0.97 (0.20 to 4.66)], stroke [RR 0.97 (0.06 to 15.24)] [Follow-up 5 years]

**Left main stem disease - Medium term follow-up (2 to 4 years)**

**Detre 1977[71] (VA); Varnauskas 1980[69] (ECSS):** Evidence from 2 RCTs shows that there was significantly fewer deaths in the CABG compared to medical treatment in patients with left main stem disease RR 4 (1.6 to 10.03).[follow-up 2-4 years]

**Left main stem disease- Long term follow-up (>4 years)**

**Alderman 1990[72] (CASS); Peduzzi 1998[75] (VA); Varnauskas 1982[82] (ECSS) :** Evidence from 3 RCTs shows that there was no significant difference between CABG and medical treatment for death in patients with left main stem disease [RR 1.18 (0.97 to 1.43)].[follow-up 10-22 years]

**Peduzzi 1998[75] (VA):** Evidence from one RCT shows that there was no significant difference medical treatment and CABG for MI [RR 0.85 (0.51 to 1.41)]. [follow-up 22 years]

**Left anterior descending artery - Long term follow-up (>4 years)**

**Alderman 1990[72] (CASS); Varnauskas 1988[76] (ECSS):**

Evidence from 2 RCTs shows that there was significantly fewer deaths in the CABG group compared to medical treatment in patients with involvement of left anterior descending artery RR 1.34 (1.09 to 1.66).[follow-up 10-12 years]

**Individual patient data meta-analyses [Medical vs. CABG ]**

**Yusuf 1994[65]:** Evidence from one IPD meta analyses shows that there were significantly fewer deaths in the CABG group compared to medical treatment [OR 0.83 (0.70 to 0.98)].[follow-up 10 years]

**Yusuf 1994[65]:** Evidence from one IPD meta analyses shows that there was no significant difference between medical treatment and CABG for death in sub group one vessel [OR 0.54 (0.22 to 1.33)] and 2 vessel [OR 0.84 (0.54 to 1.32)], significantly fewer deaths in CABG in patients with 3 vessel disease [OR 0.58 (0.42 to 0.80)], left main artery [OR 0.32 (0.15 to 0.70)] LAD disease [OR 0.50 (0.43 to 0.77)]. The benefits of surgery were similar among people with normal [OR 0.61 (0.46 to 0.81)] and abnormal LV function [OR 0.59 (0.39 to 0.91)] and all severity of angina classes [subgroup class 0, I, II-OR 0.63 (0.46 to 0.87)][sub group class III,IV - OR 0.57 (0.40 to 0.81)]. [Follow-up 5 years]

**Economic**

Medical treatment and CABG have the same probability of being cost-effective for patients suitable for both.

This evidence has potentially serious limitations and partial applicability.

1 **11.3 Medical interventions versus PCI**

2 **11.3.1 Clinical question**

3 What is the clinical and cost effectiveness of medical interventions versus PCI in  
4 people with stable angina?

5 **11.3.2 Clinical evidence**

6 The “Review Protocol” for this topic can be found in Appendix C, the “Search  
7 Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
8 E1, the “Clinical Evidence Tables” in Appendix E2, and the “Forest Plots” in Appendix  
9 F.

1 **Table 11.12: Multi-vessel disease- Short term follow-up (1 year) for stable angina**

Quality assessment							Summary of findings				Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		
							Multivessel disease - short term follow-up (1 year) Medical	PCI	Relative (95% CI)	Absolute	
<b>Death (follow-up 1 years)</b>											
Hueb 2004[66] (MASS-II)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	3/203 (1.5%)	9/205 (4.4%)	RR 0.34 (0.09 to 1.23)	29 fewer per 1000 (from 40 fewer to 10 more)	⊕⊕○○ LOW
<b>Q wave MI (follow-up 1 years)</b>											
Hueb 2004[66] (MASS-II)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	10/203 (4.9%)	16/205 (7.8%)	RR 0.63 (0.29 to 1.36)	29 fewer per 1000 (from 55 fewer to 28 more)	⊕⊕○○ LOW
<b>Stroke (follow-up 1 years)</b>											
Hueb 2004[66] (MASS-II)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	3/203 (1.5%)	2/205 (1%)	RR 1.51 (0.26 to 8.97)	5 more per 1000 (from 7 fewer to 78 more)	⊕⊕○○ LOW
<b>Non protocol revascularisation (follow-up 1 years) (d)</b>											
Hueb 2004[66] (MASS-II)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	16/203 (7.9%)	25/205 (12.2%)	RR 0.65 (0.36 to 1.17)	43 fewer per 1000 (from 78 fewer to 21 more)	⊕⊕○○ LOW
<b>Free of angina (follow-up 1 years)</b>											
Hueb 2004[66] (MASS-II)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (c)	none	74/203 (36.5%)	107/205 (52.2%)	RR 0.7 (0.56 to 0.87)	157 fewer per 1000 (from 68 fewer to 230 fewer)	⊕⊕○○ LOW
<b>Death- Sub group diabetes (follow-up 1 years)</b>											
Soares 2006[67] (MASS-II)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	2/75 (2.7%)	3/56 (5.4%)	RR 0.5 (0.09 to 2.88)	27 fewer per 1000 (from 49 fewer to 101 more)	⊕⊕○○ LOW
<b>Death- Subgroup no diabetes (follow-up 1 years)</b>											
Soares 2006[67] (MASS-II)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	2/128 (1.6%)	8/149 (5.4%)	RR 0.29 (0.06 to 1.35)	38 fewer per 1000 (from 50 fewer to 19 more)	⊕⊕○○ LOW

- 2 (a) Randomised. Allocation concealment unclear.
- 3 (b) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.
- 4 (c) 95% CI around the pooled estimate of effect includes appreciable benefit or appreciable harm.
- 5 (d) Medical treatment group – 12 non protocol CABG and 4 non protocol PCI; PCI group- 7 non protocol CABG and 18 non protocol PCI; CABG group-1 non protocol PCI.

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**Table 11.13: Multi-vessel disease- medium term follow-up (2 to 4 years) for stable angina**

Quality assessment							Summary of findings				
							No of patients		Effect		Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Multi vessel disease- medium term follow- up (2 to 4 years) Medical	PCI	Relative (95% CI)	Absolute	
<b>Death (follow-up 2.7 years)</b>											
Chamberlain 1997[89] (RITA-2)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	7/514 (1.4%)	11/504 (2.2%)	RR 0.62 (0.24 to 1.6)	8 fewer per 1000 (from 17 fewer to 13 more)	⊕⊕○○ LOW
<b>Cardiac death (follow-up 1.5-2.7 years)</b>											
Chamberlain 1997[89] (RITA-2); Pitt 1999[90] (AVERT)	randomised trials	serious (c)	no serious inconsistency	no serious indirectness	serious (b)	none	4/678 (0.6%)	6/681 (0.9%)	RR 0.67 (0.19 to 2.35)	3 fewer per 1000 (from 7 fewer to 12 more)	⊕⊕○○ LOW
<b>Non fatal MI (follow-up 1.5-2.7 years)</b>											
Chamberlain 1997[89] (RITA-2); Pitt 1999 (AVERT)	randomised trials	serious (c)	no serious inconsistency	no serious indirectness	serious (b)	none	14/678 (2.1%)	26/681 (3.8%)	RR 0.54 (0.28 to 1.02)	18 fewer per 1000 (from 27 fewer to 1 more)	⊕⊕○○ LOW
<b>Stroke (follow-up 1.5-2.7 years)</b>											
Chamberlain 1997[89] (RITA-2); Pitt 1999[90] (AVERT)	randomised trials	serious (c)	no serious inconsistency	no serious indirectness	serious (b)	none	6/678 (0.9%)	1/668 (0.1%)	RR 5.88 (0.71 to 48.69)	7 more per 1000 (from 0 fewer to 71 more)	⊕⊕○○ LOW
<b>Hospitalisation (for worsening of angina) no. of patients (follow-up 18 months)</b>											
Pitt 1999[90] (AVERT)	randomised trials	serious (d)	no serious inconsistency	no serious indirectness	serious (e)	none	11/164 (6.7%)	25/177 (14.1%)	RR 0.47 (0.24 to 0.93)	75 fewer per 1000 (from 10 fewer to 107 fewer)	⊕⊕○○ LOW
<b>Non protocol Revascularisation (follow-up 1.5-2.7 years)</b>											
Chamberlain 1997[89] (RITA-2); Pitt 1999[90] (AVERT)	randomised trials	serious (c)	serious (f)	no serious indirectness	serious (b)	none	151/678 (22.3%)	132/681 (19.4%)	RR 1.14 (0.93 to 1.4)	27 more per 1000 (from 14 fewer to 78 more)	⊕○○○ VERY LOW

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- (a) Strengths: multicentre (20 centres in UK and Ireland), stratified blocked randomisation. Sample size calculation reported. Intention to treat analysis reported. Loss to follow-up - 5.1% in PTCA and 3.3% in medical group (N=478 PTCA an n=497 at 2.7 yrs) Blind outcome assignment. Weakness: allocation concealment not reported.
- (b) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.
- (c) Blind outcome assignment in both studies. Both studies allocation concealment not reported. RITA-2 -stratified blocked randomisation. Sample size calculation reported. Intention to treat analysis reported. AVERT- No loss to follow-up.

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- 1 (d) *Strengths: open label randomised, multi centre, sample size calculation reported. Blind outcome assessment. No loss to follow-up. ITT reported. Limitations: allocation*  
2 *concealment not reported. This study is a 18 month follow-up of the AVERT trial.*  
3 (e) *95% CI around the pooled estimate of effect includes appreciable benefit or appreciable harm.*  
4 (f) *I<sup>2</sup>= 73%*

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1 **Table 11.14: Multi-vessel disease-long term follow-up (> 4 years follow-up) for stable angina**

Quality assessment							Summary of findings				Quality
							No of patients		Effect		
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Multivessel disease-long term follow-up (> 4 years follow-up) Medical	PCI	Relative (95% CI)	Absolute	
<b>Death (follow-up 2.7-10 years)</b>											
Boden 2007[91] (COURAGE) (I); Henderson 2003[92] (RITA-2); Hueb 2010[77] (MASS-II)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	201/1855 (10.8%)	177/188 (94.1%)	RR 1.14 (0.94 to 1.37)	132 more per 1000 (from 56 fewer to 348 more)	⊕⊕○○ LOW
<b>Cardiac death (follow-up 2.7-4.6 years)</b>											
Boden 2007[91] (COURAGE); Henderson 2003[92] (RITA-2)	randomised trials	serious (c)	serious (d)	no serious indirectness	serious (b)	none	47/1652 (2.8%)	36/1653 (2.2%)	RR 1.30 (0.85 to 2)	7 more per 1000 (from 3 fewer to 22 more)	⊕○○○ VERY LOW
<b>Non fatal MI (follow-up 2.7-10 years)</b>											
Boden 2007[91] (COURAGE) (I); Henderson 2003[92] (RITA-2); Hueb 2010[77] (MASS-II)	randomised trials	serious (a)	serious (e)	no serious indirectness	serious	none	193/1855 (10.4%)	202/1858 (10.9%)	RR 0.96 (0.80 to 1.16)	4 fewer per 1000 (from 22 fewer to 17 more)	⊕○○○ VERY LOW
<b>Non protocol Revascularisation (follow-up 2.7-10 years)</b>											
Boden 2007[91] (COURAGE) (I); Henderson 2003[92] (RITA-2); Hueb 2010[77] (MASS-II)	randomised trials	serious (a)	serious (f)	no serious indirectness	no serious imprecision	none	630/1855 (34%)	463/1858 (24.9%)	RR 1.36 (1.23 to 1.51)	90 more per 1000 (from 57 more to 127 more)	⊕⊕○○ LOW
<b>Stroke (follow-up 4.6-10 years)</b>											
Boden 2007[91] (COURAGE); Hueb 2010[77] (MASS-II)	randomised trials	serious (g)	no serious inconsistency	no serious indirectness	serious (b)	none	28/1341 (2.1%)	33/1354 (2.4%)	RR 0.86 (0.52 to 1.41)	3 fewer per 1000 (from 12 fewer to 10 more)	⊕⊕○○ LOW
<b>Free of angina (follow-up 4.6-10 years)</b>											
Boden 2007[91] (COURAGE); Folland 1997[93] (ACME); Hueb 2010[77] (MASS-II)	randomised trials	serious (a)	serious (h)	no serious indirectness	no serious imprecision	none	402/1391 (28.9%)	463/1405 (33%)	RR 0.88 (0.79 to 0.98)	40 fewer per 1000 (from 7 fewer to 69 fewer)	⊕⊕○○ LOW
<b>Death- sub group age &lt;65 yrs (follow-up 4.6 years)</b>											

Teo 2009[94] (COURAGE)	randomised trials	serious (i)	no serious inconsistency	no serious indirectness	serious (b)	none	41/693 (5.9%)	25/688 (3.6%)	RR 1.63 (1 to 2.65)	23 more per 1000 (from 0 more to 60 more)	⊕⊕○○ LOW
<b>MI - sub group age &lt;65 yrs (follow-up 4.6 years)</b>											
Teo 2009[94] (COURAGE)	randomised trials	serious (i)	no serious inconsistency	no serious indirectness	serious (b)	none	76/693 (11%)	83/688 (12.1%)	RR 0.91 (0.68 to 1.22)	11 fewer per 1000 (from 39 fewer to 27 more)	⊕⊕○○ LOW
<b>Free of angina- sub group age&lt;65 years (follow-up 4.6 years)</b>											
Teo 2009[94] (COURAGE)	randomised trials	serious (i)	no serious inconsistency	no serious indirectness	no serious imprecision	none	485/693 (70%)	481/688 (69.9%)	RR 1 (0.93 to 1.07)	0 fewer per 1000 (from 49 fewer to 49 more)	⊕⊕⊕○ MODERATE
<b>Death- sub group age &gt;65 yrs (follow-up 4.6 years)</b>											
Teo 2009[94] (COURAGE)	randomised trials	serious (j)	no serious inconsistency	no serious indirectness	serious (b)	none	54/444 (12.2%)	57/460 (12.4%)	RR 0.98 (0.69 to 1.39)	2 fewer per 1000 (from 38 fewer to 48 more)	⊕⊕○○ LOW
<b>MI- sub group age &gt;65 yrs (follow-up 4.6 years)</b>											
Teo 2009[94] (COURAGE)	randomised trials	serious (j)	no serious inconsistency	no serious indirectness	serious (b)	none	52/444 (11.7%)	60/460 (13%)	RR 0.9 (0.63 to 1.27)	13 fewer per 1000 (from 48 fewer to 35 more)	⊕⊕○○ LOW
<b>Free of angina- sub group age &gt;65 yrs (follow-up 4.6 years)</b>											
Teo 2009[94] (COURAGE)	randomised trials	serious (j)	no serious inconsistency	no serious indirectness	no serious imprecision	none	324/444 (73%)	368/460 (80%)	RR 0.91 (0.85 to 0.98)	72 fewer per 1000 (from 16 fewer to 120 fewer)	⊕⊕⊕○ MODERATE
<b>Death- sub group 2 vessel disease (follow-up 6 years)</b>											
Folland 1997[93] (ACME)	randomised trials	serious (k)	no serious inconsistency	no serious indirectness	serious (b)	none	10/50 (20%)	9/51 (17.6%)	RR 1.13 (0.5 to 2.55)	23 more per 1000 (from 88 fewer to 274 more)	⊕⊕○○ LOW
<b>Non fatal MI- sub group 2 vessel disease (follow-up 6 years)</b>											
Folland 1997[93] (ACME)	randomised trials	serious (k)	no serious inconsistency	no serious indirectness	serious (b)	none	7/50 (14%)	7/51 (13.7%)	RR 1.02 (0.39 to 2.7)	3 more per 1000 (from 84 fewer to 233 more)	⊕⊕○○ LOW

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- (a) Randomisation, sample size calculation, blind outcome assessment and, ITT reported in all 3 studies. Allocation concealment not reported in 3 studies.
- (b) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.
- (c) Randomisation, sample size calculation, blind outcome assessment and ITT reported in both the studies. Allocation concealment not reported in both the studies.

- 1 (d)  $I^2=62\%$   
 2 (e)  $I^2=69\%$   
 3 (f)  $I^2=83\%$   
 4 (g) Strengths: Randomisation reported in both studies. Allocation concealment unclear in both studies, ITT reported in both studies.  
 5 (h)  $I^2=62\%$   
 6 (i) Strengths: Randomisation method reported (permuted block design within strata -prior CABG/no prior CABG and by medical centre), sample size calculation reported,  
 7 Blind outcome assessment (clinical outcome adjudicated by an independent committee whose members were unaware of treatment assignments). 9% of patients were lost to  
 8 follow-up in the two groups (107 in the PCI group and 97 in the medical therapy group,  $p=0.51$ ). Intention to treat analysis reported.  
 9 (j) Strengths: Randomisation method reported (permuted block design within strata -prior CABG/no prior CABG and by medical centre), sample size calculation reported,  
 10 Blind outcome assessment (clinical outcome adjudicated by an independent committee whose members were unaware of treatment assignments). Loss to follow-up not  
 11 reported separately for subgroup age >65 years. Intention to treat analysis reported.  
 12 (k) Strengths: Randomised, baseline comparison made, intention to treat analysis used. Weakness: Randomisation method not clearly described. Allocation concealment not  
 13 reported.  
 14 (l) All patients received aspirin, and those who were undergoing PCI also received clopidogrel in accordance with treatment guidelines. Ant ischemic therapy included long  
 15 acting metoprolol, amlodipine, and isosorbide mononitrate, alone or in combination, together with simvastatin and either lisinopril or losartan for secondary prevention.  
 16

17 **Additional data:**

18 **(Multi vessel disease- Long term follow-up -RITA -2)**

19 Henderson 2003[92] (RITA-2): The prevalence of angina declined in both treatment groups during the first five years of follow-up, but this symptomatic improvement was much  
 20 more rapid in the PTCA group. At 3 months after randomisation, 19.4% and 35.9% of the PTCA and medical groups, respectively, had angina grade 2 or worse (difference  
 21 16.5%, 95% CI 11.0% to 21.9%). By 5 years follow-up, the prevalence of angina grade 2 or worse in the PTCA group remained steady at 15%, whereas in the medical  
 22 group the prevalence of angina was reduced to 21.4%. The 5 year treatment difference was thus much smaller, 6.4% in favour of PTCA (95% CI 1.5% to 11.3%,  $p=0.011$ ).  
 23 During the next 3 years, the prevalence of angina began to increase slightly in both treatment groups.  
 24 Sub group interaction- age <65 years and >65 years  
 25 There was no significant difference between sub groups age <65 years and >65 years for death ( $p=0.10$ ), MI ( $p=0.96$ ) and free of angina ( $p=0.06$ ).  
 26

27 **(Multi vessel disease- Long term follow-up (5 year follow-up)-MASS-II)**

28 Lopes 2008[95] (MASS-II):  $n=825$  ( $n=214$  single vessel disease,  $n=253$  two vessel disease,  $n=358$  three vessel disease)  
 29 Overall mortality was significantly higher in 3 vessel disease (17.8%) compared to 2 vessel disease (12.2%) and single vessel disease ( $n=6\%$ ) [ $p=0.001$ ]. Multivariate Cox  
 30 regression model (including variables such as age, hypertension, gender, hyperlipidemia, no. of coronary disease and treatment allocation) for mortality revealed a 3-fold  
 31 increased risk of mortality in 3 vessel disease comparing to single vessel disease [ $p=0.005$ , HR 3.14, 95% CI 1.4 to 97.0]. There was no significant difference between 2  
 32 vessel disease and single vessel disease for mortality [ $p=0.15$ , HR 1.89, 95% CI 0.75 to 4.56].  
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1 **Table 11.15: Single vessel disease - medium term follow-up (2 -4 years) for stable angina**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Single vessel disease - medium term follow-up (2 -4 years) Medical	PCI	Relative (95% CI)	Absolute	
<b>Death (follow-up 2-3 years)</b>											
Hartigan 1998[96] (ACME); Hueb 1995[83] (MASS-I)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	7/179 (3.9%)	6/177 (3.4%)	RR 1.14 (0.41 to 3.17)	5 more per 1000 (from 20 fewer to 74 more)	⊕⊕○○ LOW
<b>MI (follow-up 2-3 years)</b>											
Hartigan 1998[96] (ACME); Hueb 1995[83] (MASS-I)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	9/179 (5%)	12/177 (6.8%)	RR 0.74 (0.32 to 1.7)	18 fewer per 1000 (from 46 fewer to 47 more)	⊕⊕⊕○ MODERATE
<b>Hospitalisation (no. of patients) (follow-up 2-3 years)</b>											
Hartigan 1998[96] (ACME)	randomised trials	serious (c)	no serious inconsistency	no serious indirectness	serious <sup>2</sup>	none	69/107 (64.5%)	64/105 (61%)	RR 1.06 (0.86 to 1.3)	37 more per 1000 (from 85 fewer to 183 more)	⊕⊕○○ LOW
<b>Free of angina (follow-up 2-3 years)</b>											
Hartigan 1998[96] (ACME); Hueb 1995[83] (MASS-I)	randomised trials	serious (a)	serious (d)	no serious indirectness	no serious imprecision	none	73/179 (40.8%)	123/177 (69.5%)	RR 0.59 (0.48 to 0.72)	285 fewer per 1000 (from 195 fewer to 361 fewer)	⊕⊕○○ LOW
<b>Non protocol revascularisation (follow-up 2-3 years)</b>											
Hartigan 1998[96] (ACME); Hueb 1995[83] (MASS-I)	randomised trials	serious (a)	serious (e)	no serious indirectness	serious (f)	none	54/179 (30.2%)	76/177 (42.9%)	RR 0.7 (0.53 to 0.93)	129 fewer per 1000 (from 30 fewer to 202 fewer)	⊕○○○ VERY LOW
<b>Stroke (follow-up 3 years)</b>											
Hueb 1995[83] (MASS-I)	randomised trials	serious (g)	no serious inconsistency	no serious indirectness	no serious imprecision	none	0/72 (0%)	0/72 (0%)	not pooled	not pooled	⊕⊕⊕○ MODERATE

- 2 (a) Veteran affairs ACME study -Strengths: Randomised, baseline comparison made, intention to treat analysis used. Weakness: Randomisation method not clearly described.  
 3 Allocation concealment not reported. MASS-I: Strengths: Randomised, Baseline comparisons made. Number of patients lost to follow-up not reported. ITT reported.  
 4 Weakness: allocation concealment not reported. Blinding of outcome assessors not reported.  
 5 (b) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.  
 6 (c) Strengths: Randomised, baseline comparison made, intention to treat analysis used. Weakness: Randomisation method not clearly described. Allocation concealment not  
 7 reported.  
 8 (d)  $I^2 = 88\%$

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- (e)  $I^2 = 92\%$
- (f) 95% CI around the pooled estimate of effect includes appreciable benefit or appreciable harm.
- (g) Strengths: Randomised, Baseline comparisons made. Number of patients lost to follow-up not reported. ITT reported. Weakness: allocation concealment not reported. Blinding of outcome assessors not reported.

1 **Table 11.16: Single vessel disease - long term follow-up (>4 years) for stable angina**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Single vessel disease - long term follow-up (>4 years) Medical	PCI	Relative (95% CI)	Absolute	
<b>Death (follow-up 4.6-6 years)</b>											
Folland 1997[93] (ACME); Hueb 1995[83] (MASS-I)	randomised trials	serious (a)	no serious inconsistency (b)	no serious indirectness	serious (c)	none	22/184 (12%)	23/187 (12.3%)	RR 0.98 (0.57 to 1.68)	2 fewer per 1000 (from 53 fewer to 84 more)	⊕○○○ LOW
<b>Non fatal MI (follow-up 4.6-6 years)</b>											
Folland 1997[93] (ACME); Hueb 1995[83] (MASS-I)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (c)	none	11/184 (6%)	22/187 (11.8%)	RR 0.51 (0.26 to 1.02)	58 fewer per 1000 (from 87 fewer to 2 more)	⊕⊕○○ LOW
<b>Non protocol Revascularisation (follow-up 4.6-6 years)</b>											
Hueb 1995[83] (MASS-I)	randomised trials	serious (d)	no serious inconsistency	no serious indirectness	serious (c)	none	12/72 (16.7%)	29/72 (40.3%)	RR 0.41 (0.23 to 0.75)	238 fewer per 1000 (from 101 fewer to 310 fewer)	⊕○○○ VERY LOW
<b>cardiac death (follow-up 4.6-6 years)</b>											
Hueb 1995[83] (MASS-I)	randomised trials	serious (e)	no serious inconsistency	no serious indirectness	serious (c)	none	2/72 (2.8%)	4/72 (5.6%)	RR 0.5 (0.09 to 2.64)	28 fewer per 1000 (from 51 fewer to 91 more)	⊕⊕○○ LOW
<b>stroke (follow-up 5 years)</b>											
Hueb 1995[83] (MASS-I)	randomised trials	serious (e)	no serious inconsistency	no serious indirectness	serious (c)	none	1/72 (1.4%)	1/72 (1.4%)	RR 1 (0.06 to 15.68)	0 fewer per 1000 (from 13 fewer to 204 more)	⊕⊕○○ LOW
<b>Free of angina (follow-up 5 years)</b>											
Hueb 1995[83] (MASS-I)	randomised trials	serious (e)	no serious inconsistency	no serious indirectness	no serious imprecision	none	17/72 (23.6%)	44/72 (61.1%)	RR 0.39 (0.25 to 0.61)	373 fewer per 1000 (from 238 fewer to 458 fewer)	⊕⊕⊕○ MODERATE

- 2 (a) ACME study -Strengths: Randomised, baseline comparison made, intention to treat analysis used, no patients lost to follow up. Weakness: Randomisation method not clearly  
3 described. Allocation concealment not reported. MASS-I: Strengths: Randomised, Baseline comparisons made. Number of patients lost to follow-up not reported. ITT  
4 reported. Weaknesses: allocation concealment not reported. Blinding of outcome assessors not reported.  
5 (b)  $I^2 = 0\%$   
6 (c) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.  
7 (d) Strengths: Randomised, Baseline comparisons made. Number of patients lost to follow-up not reported. ITT reported. Weaknesses: allocation concealment not reported.  
8 Blinding of outcome assessors not reported.

1 (e) *Strengths: Randomised, Baseline comparisons made. Number of patients lost to follow-up not reported. ITT reported. Weaknesses: allocation concealment not reported.*  
2 *Blinding of outcome assessors not reported.*  
3

4 **Sub group interaction – single vessel and 2 vessel – Long term follow-up:**

5 *There was no significant difference between sub groups single and 2 vessel for death ( $p=0.61$ ) and MI ( $p=0.54$ ).*

1 **Quality of Life data from studies:**

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3 **Strauss 1995[97] (ACME):**

4 In ACME Quality of Life was assessed with the Psychologic General Well-Being Index  
5 (PGWB) developed to measure an individual's subjective sense of well-being or  
6 distress. It measures the patient's perception of his or her well-being in the month  
7 preceding assessment. Six categories of psychological well-being were assessed:  
8 anxiety, depressed mood, positive well-being, self-control, general health and  
9 vitality. The test consists of 22 questions, the responses to which are graded from 0  
10 (most negative) to 5 (most positive). There are three to five non-overlapping items or  
11 responses that form the subscales with which to measure the six states. The answers to  
12 the 22 questions are summed to yield an overall psychological well-being QOL score  
13 (maximum, 110).

14 **Results:** n=170 (n for each group not separately specified) with paired baseline and  
15 follow-up (6 months) data. At 6 month follow-up the mean change in Quality of Life  
16 score was significantly higher in the PCI group ( $+1.98 \pm 14.7$  Medical vs.  $+7.36$   
17  $\pm 15.6$ ;  $p < .02$ ). Within the subscales there was also a significant difference in  
18 perceived General Health which was significantly higher for the PCI group and all  
19 other subcomponents of the questionnaire showed the same trend (subscale mean  
20 change scores given in Figure, but not text).

21  
22 **Folland 1997[93] (ACME) – single vessel vs. double vessel disease:**

23 See above for details of measure.

24 **Results:** n=267 (n=35 PCI double-vessel, n=37 Medical double-vessel, n=95 PCI  
25 single-vessel; and n=100 Medical single-vessel). At 6 mean QOL scores improved for  
26 both treatment groups with double-vessel disease, but the difference between  
27 treatment groups was not significant ( $+4.4$  Medical vs.  $+1.3$  PCI,  $p = .32$ ). For patients  
28 with single-vessel disease there was significantly greater improvement in the PCI  
29 compared to the Medical group ( $+1.5$  Medical vs.  $+7.1$  PCI,  $p = .01$ ).

30  
31 **Pitt 1999[90] (AVERT trial):**

32 Quality of Life was assessed using the 36-item Medical Outcomes Study Short-Form  
33 General Health Survey (see below for details) at baseline, 6 and 18 months after  
34 randomization.

35 **Results:** n=341 (n=177 in PTCA and n=164 in medical treatment). Both treatment  
36 groups had a mean increase in the summary scores for physical and mental health at  
37 both 6-month and 18-month assessments, denoting an improvement in the quality of  
38 life from baseline. Mean increases in scores ranged from 2.9 to 6.3; the increases  
39 were slightly, but not significantly, larger in the angioplasty group. No further details  
40 were provided.

1

2 **Pocock 2000[98] (RITA-2):**

3 Patients assessed their Quality of Life using the SF-36 health survey, at baseline, 3  
4 months, 1 year and 3 years. The SF-36 comprises 36 items that can be combined into  
5 the following eight multi-item summary scores: physical functioning (10 items), vitality  
6 (4 items), bodily pain (2 items), mental health (5 items), social functioning (2 items) role  
7 limitation due to physical health (4 items) and due to emotional problems (3 items)  
8 and general health perceptions (5 items), plus one item assessing a change in health  
9 over the past year. Each summary score is obtained by simple unweighted summation  
10 of item scores and is then scaled from 0 to 100, with 0 and 100 indicating 'worst' and  
11 'best' possible health respectively.

12 **Results:** n=1018 (n=504 in PTCA and n=514 in medical treatment). Quality of life  
13 by SF-36 values (mean; SEM) reported in figures. Reported in text - The PTCA group  
14 showed highly significant superiority over the medical group in terms of physical  
15 functioning, vitality and general health at both 3 months and 1 year after  
16 randomisation. Mental health was also significantly better in the PTCA group at 3  
17 months and 1 year, although the magnitude of this difference was quite small. The  
18 slight superiority of the PTCA group in pain, social functioning and physical and  
19 emotional role functioning did not achieve such marked levels of statistical  
20 significance. None of the 8 SF-36 scores showed a significant treatment difference at  
21 3 years.

22 **Weintraub 2008[99] (COURAGE)**

23 **Measurement of health status:** Health status related to angina was assessed directly  
24 from patients at baseline; at 1, 3, 6 and 12 months; and at annual evaluations there  
25 after. Each assessment was performed with the use of the Seattle Angina  
26 Questionnaire, a 19 item questionnaire that quantifies physical limitations due to  
27 angina, any recent change in the severity of angina, the frequency of angina,  
28 satisfaction with treatment, and quality of life. Scores range from 0 to 100; higher  
29 scores indicate better health status.

30 **Measurement of general health status:** General health status was measured with the  
31 use of the RAND-36 health survey, which includes the following domains: physical  
32 functioning, role limitation due to physical problems, role limitation due to emotional  
33 problems, vitality, emotional well being social functioning, pain, and general health.  
34 Scores for each domain range from 0 to 100, with higher scores reflecting better  
35 health status. The RAND-36 health survey contains the same items as the Medical  
36 Outcomes Study 36-item Short Form General Health Survey (SF-36).

37 **Results:** N=2287 (n=1149 PCI and n=1138 in OMT). Patients were followed for a  
38 minimum of 30 months.

39 **Health status:** Baseline mean ( $\pm$  SD) Seattle Angina Questionnaire scores (which  
40 range from 0 to 100, with higher scores indicating better health status) were  $66 \pm 25$   
41 for physical limitations,  $54 \pm 32$  for angina stability,  $69 \pm 26$  for angina frequency,  
42  $87 \pm 16$  for treatment satisfaction, and  $51 \pm 25$  for quality of life. By 3 months, these  
43 scores had increased in the PCI group, as compared with medical therapy group, to  
44  $76 \pm 24$  versus  $72 \pm 23$  for physical limitation ( $p=0.004$ ),  $77 \pm 28$  versus  $73 \pm 27$  for

1 angina stability ( $p=0.002$ ),  $85\pm 22$  versus  $80\pm 23$  for angina frequency ( $p<0.001$ ),  
 2  $92\pm 12$  versus  $90\pm 14$  for treatment satisfaction ( $p<0.001$ ), and  $73\pm 22$  versus  
 3  $68\pm 23$  for quality of life ( $p<0.001$ ). In general, patients had an incremental benefit  
 4 from PCI for 6 months to 24 months; people with more severe angina had a greater  
 5 benefit from PCI.

6 **General Health status:** There were no significant differences at baseline between the  
 7 groups for any RAND-36 domain. There was improvement in all domains in both  
 8 groups between randomisation and follow-up at 1 to 3 months ( $p<0.001$  for all  
 9 comparisons). There was also an incremental advantage of PCI over medical therapy  
 10 at 3 months for the scores in five domains: physical functioning ( $69\pm 27$  vs.  
 11  $65\pm 26$ ,  $p<0.001$ ), role limitation-physical ( $60\pm 42$  vs.  $52\pm 43$ ,  $p<0.001$ ), vitality  
 12 ( $56\pm 23$  vs.  $53\pm 23$ ,  $p=0.008$ ), pain ( $72\pm 25$  vs.  $68\pm 26$ ,  $p=0.006$ ), and general health  
 13 ( $61\pm 21$  vs.  $58\pm 21$ ,  $p<0.001$ ). The benefit across domains was less consistent than seen  
 14 in the results for the Seattle Angina Questionnaire, with an advantage of PCI that was  
 15 noted in most but not all domains and that had a shorter duration. At 6 months, the  
 16 PCI group was more likely than the medical therapy group to have a clinically  
 17 significant improvement in physical functioning (50% vs. 43%) and role limitation-  
 18 physical (48% vs. 43%), but no advantage was observed at 12 months. There were  
 19 no significant subgroup interactions in the RAND-36 results.

### 20 11.3.3 Economic evidence

21 Two studies [100,101] were found that included the relevant comparison. These are  
 22 summarised in the economic evidence profile below. See also Economics Evidence  
 23 Tables in Appendix G.

24 **Table 11.17: PCI vs. medical treatment - Economic study characteristics**

Study	Limitations	Applicability	Other Comments
<b>Sculpher 2002</b> [100]	Minor limitations (a)	Partial applicability (b)	Based on the RITA-2 study [89] included in the clinical review. Patients had arteriographically proven coronary artery disease and were recruited from 20 centres in the UK and Ireland.
<b>Weintraub 2008</b> [101]	Minor limitations (c)	Partial applicability (d)	Based on the COURAGE trial [91]. Patients had stable coronary artery disease with >70% stenosis in at least one major epicardial coronary artery with objective evidence of myocardial ischemia or at least one coronary stenosis >80% and classic angina without provocative testing.

26 (a) No incremental analysis was conducted.

27 (b) Utility values were not estimated. Stents and other coronary interventional techniques were only used if  
 28 initial revascularisation with balloon angioplasty was unsatisfactory.

29 (c) Valuation of utilities not obtained from public but from patients. Effectiveness was estimated for the total  
 30 duration of the trial (4.6 years) while costs only for 3 years. These results were combined.

31 (d) Patients in the study were low risk. PCI group included angioplasty too. USA study.  
 32

1 **Table 11-18: PCI vs. medical treatment - Economic summary of findings**

Study	Incremental cost per patient over three years (£)	Incremental effectiveness	ICER (£/QALY)	Uncertainty
<b>Sculpher 2002</b> [100]	2,686 (a, b)	(c)	NA	Similar results when patients were stratified by CCS score, breathlessness, exercise time, and overall score. Similar results when no discount rate is applied, cost of visits for non-cardiac reasons is excluded, or when unit costs from the 5 hospitals are used separately.
<b>Weintraub 2008</b> [101]	6,174 (d, e)	0.05 QALYs (e, f)	125,759	Extrapolating beyond RCT follow-up: PCI is still significantly more costly and more effective (not sig). Use of drug-eluting stents: no revascularisation after PCI was assumed, additional cost of \$600 and clopidogrel for one year, PCI would not be cost-effective.  At a \$50k/QALY threshold PCI has a 25% probability of being cost-effective.

- 2 (a) 1999 GBP. Cost of cardiac procedures, in-hospital stay, subsequent procedures, GP and outpatient  
3 visits, antianginal and cardiac drugs.  
4 (b) Discounted by 6%  
5 (c) Number of deaths was higher in PTCA group (not statistically significant); number of deaths and MI was  
6 higher in PTCA group (statistically significant). More patients with grade 2 or worse angina in medical  
7 treatment group (statistically significant at 1 year, not statistically significant at 3 years).  
8 (d) 2008 GBP obtained by using the purchasing power parities and GDP deflator indexes  
9 (<http://eppi.ioe.ac.uk/costconversion/default.aspx>). Costs included were hospitalisation, PCI,  
10 medication, outpatient services.  
11 (e) Discounted by 3%  
12 (f) Utility values estimated with the standard gamble method from participants of the trial.  
13

14 **11.3.4 Evidence statements**

**Clinical Multi-vessel disease- short term follow-up (1 year) for stable angina**

**Hueb 2004**[66] (MASS-II): Evidence from one RCT shows that significantly higher number of patients were free of angina [RR 0.7 (0.56 to 0.87)] in the PCI group compared to medical treatment. There was no significant difference between medical treatment and PCI for death [RR 0.34 (0.09 to 1.23)], Q wave MI [RR 0.63 (0.29 to 1.36)], stroke [RR 1.51 (0.26 to 8.97)], non protocol revascularisation [RR 0.65 (0.36 to 1.17)].

**Soares 2006**[67] (MASS-II): Evidence from one RCT shows there was no significant difference between medical treatment and PCI for and death in a sub group of patients with diabetes [RR 0.5

(0.09 to 2.88)] and no diabetes [RR 0.29 (0.06 to 1.35)]. [Follow-up 1 year].

**Multi-vessel disease- medium term follow-up (2 to 4 years) for stable angina**

**Chamberlain 1997[89] (RITA-2):** Evidence from one RCT shows that there was no significant difference between medical treatment and PCI for death [RR 0.62 (0.24 to 1.6)]. [Follow-up 2.7 years]

**Chamberlain 1997[89] (RITA-2); Pitt 1999[90] (AVERT):** Evidence from 2 RCTs shows that there was no significant difference between medical treatment and PCI for cardiac death [RR 0.67 (0.19 to 2.35)], non fatal MI [RR 0.54 (0.28 to 1.02)], and stroke [RR 5.88 (0.71 to 48.69)]. [follow-up 1.5-2.7 years]

**Pitt 1999[90] (AVERT):** Evidence from one RCT shows that there were significantly fewer hospitalisations for worsening of angina in medical treatment compared to PCI [RR 0.47 (0.24 to 0.93)] [follow-up 18 months]

**Chamberlain 1997[89] (RITA-2); Pitt 1999[90] (AVERT):** Evidence from 2 RCTs shows that there was no significant difference medical treatment and PCI for non protocol revascularisation [RR 1.14 (0.93 to 1.4)]. [Follow-up 1.5-2.7 years]

**Multi vessel disease-long term follow-up (> 4 years follow-up) for stable angina**

**Boden 2007[91] (COURAGE); Henderson 2003[92] (RITA-2); Hueb 2010[77] (MASS-II):** Evidence from 3 RCTs shows that significantly higher number of patients were free of angina in the PCI compared to medical treatment [RR 0.88 (0.79 to 0.98)]. There was significantly higher non protocol revascularisation in medical treatment compared to PCI [RR 1.36 (1.23 to 1.51)]. There was no significant difference between medical treatment and PCI for death [RR 1.14 (0.94 to 1.37)], non fatal MI [RR 0.96 (0.80 to 1.16)] [follow-up 4.6-10 years]

**Boden 2007[91] (COURAGE); Henderson 2003[92] (RITA-2):** Evidence from 2 RCTs shows that there was no significant difference between medical treatment and PCI for cardiac death [RR 1.30 (0.85 to 2.00)], [follow-up 4.6-7 years]

**Boden 2007[91] (COURAGE); Hueb 2010[77] (MASS-II):** Evidence from 2 RCT shows that there was no significant difference medical treatment and PCI for stroke [RR 0.86 (0.52 to 1.41)] [follow-up 4.6 -10 years]

**Teo 2009[94] (COURAGE):** Evidence from one RCT shows that there was no significant difference medical treatment and PCI for

death [RR 0.98 (0.69 to 1.39)], MI [RR 0.9 (0.63 to 1.27)]. There were significantly more patients free of angina [RR 0.91 (0.85 to 0.98)] in PCI compared to medical treatment in patients aged >65 years. [Follow-up 4.6 years]

**Teo 2009[94] (COURAGE):** Evidence from one RCT shows that there was significantly higher death [RR 1.63 (1.00 to 2.65)] in medical treatment compared to PCI. There was no significant difference medical treatment and PCI for MI [RR 0.91 (0.68 to 1.22)] and free of angina [RR 1.00 (0.93 to 1.07)], in patients aged >65 years. [Follow-up 4.6 years] But there was no significant difference between sub groups age <65 years and >65 years for death (p=0.10), MI (p=0.96) and free of angina (p=0.06).

**Folland 1997[93] (ACME):** Evidence from one RCT shows that there was no significant difference between medical treatment and PCI for death [RR 1.13 (0.5 to 2.55)], non fatal MI [RR 1.02 (0.39 to 2.7)] in sub group of patients with 2 vessel disease. [Follow-up 6 years]

**Single vessel disease - medium term follow-up (2 -4 years) for stable angina**

**Hartigan 1998[96] (ACME); Hueb 1995[83] (MASS-I):** Evidence from 2 RCTs shows that there was statistically significant higher no. of patients free of angina [RR 0.59 (0.48 to 0.72)] in PCI compared to medical treatment; there was statistically significant higher non -protocol revascularisation the PCI group compared to medical treatment [RR 0.7 (0.53 to 0.93)]. There was no significant difference between medical treatment and PCI for death [RR 1.14 (0.41 to 3.17)], MI [RR 0.74 (0.32 to 1.7)], hospitalisation [RR 1.06 (0.86 to 1.3)] and stroke [PCI-0/72 and CABG-0/72]. [Follow-up 2-4 years]

**Single vessel disease - long term follow-up (>4 years) for stable angina**

**Folland 1997[93] (ACME); Hueb 1995[83] (MASS-I):** Evidence from 2 RCTs shows that there was no statistically significant difference between medical treatment and PCI for death [RR 0.98 (0.57 to 1.68)] non fatal MI [RR 0.51 (0.26 to 1.02)] [Follow-up 4.6 to 6 years]

**Hueb 1995[83] (MASS-I):** Evidence from one RCT shows that there was statistically significant higher no. of patients free of angina in the PCI group compared to medical treatment [RR 0.39 (0.25 to 0.61)]. There was no statistically significant difference between medical treatment and PCI for cardiac death [RR 0.5 (0.09 to 2.64)] non -protocol revascularisation [RR 0.41 (0.23 to 0.75)] and

stroke [RR 1 (0.06 to 15.68)]. [Follow-up 5 years]

**Economic**

Medical treatment is more cost-effective than early revascularisation with PCI in people with stable coronary artery disease. This evidence has minor limitations and partial applicability.

1 **11.4 Medical interventions versus PCI or CABG**

2 **11.4.1 Clinical question**

3 What is the clinical and cost effectiveness of medical interventions versus PCI or CABG  
4 in people with stable angina?

5 **11.4.2 Clinical evidence**

6 The “Review Protocol” for this topic can be found in Appendix C, the “Search  
7 Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
8 E1, the “Clinical Evidence Tables” in Appendix E2, and the “Forest Plots” in Appendix  
9 F.

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**Table 11.19: Multi-vessel disease- short term follow-up (1 year) for stable angina**

Quality assessment							Summary of findings				Quality
							No of patients		Effect		
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Multivessel disease- short term follow-up (1 year) Medical	PCI or CABG	Relative (95% CI)	Absolute	
<b>Death (follow-up 1 years)</b>											
Pfisterer 2003[102] (TIME)	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	12/148 (8.1%)	17/153 (11.1%)	RR 0.73 (0.36 to 1.47)	30 fewer per 1000 (from 71 fewer to 52 more)	⊕⊕⊕○ MODERATE
<b>MI (follow-up 1 years)</b>											
Pfisterer 2003[102] (TIME)	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	20/148 (13.5%)	14/153 (9.2%)	RR 1.48 (0.78 to 2.81)	44 more per 1000 (from 20 fewer to 167 more)	⊕⊕⊕○ MODERATE
<b>Non protocol revascularisation (follow-up 1 years)</b>											
Pfisterer 2003[102] (TIME)	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	71/148 (48%)	16/153 (10.5%)	RR 4.59 (2.8 to 7.51)	377 more per 1000 (from 189 more to 684 more)	⊕⊕⊕○ MODERATE

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(a) Strengths: randomised, low attrition bias (on-treatment analysis so no loss to follow up) Weaknesses: the potential for bias is substantial because both treatment groups contain failures of the other treatment. In addition the patient number is relatively low. No allocation concealment /No intention to treat analysis as it is an on treatment analysis.

1 **Table 11.20: Multi-vessel disease- medium term follow-up (2 to 4 years) for stable angina**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Multi vessel disease- medium term follow-up ( 2 to 4 years) Medical	PCI or CABG	Relative (95% CI)	Absolute	
<b>Death (follow-up 4 years)</b>											
Pfisterer 2004[103] (TIME)	randomised trial	Serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	31/139 (22.3%)	29/137 (21.2%)	RR 1.05 (0.67 to 1.65)	11 more per 1000 (from 70 fewer to 138 more)	⊕⊕⊕O MODERATE
<b>Non protocol revascularisation (follow-up 4 years)</b>											
Pfisterer 2004[103] (TIME)	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	4/139 (2.9%)	4/137 (2.9%)	RR 0.99 (0.25 to 3.86)	0 fewer per 1000 (from 22 fewer to 83 more)	⊕⊕⊕O MODERATE
<b>Non fatal MI (follow-up 4 years)</b>											
Pfisterer 2004[103] (TIME)	randomised trial	Serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	1/139 (0.7%)	6/137 (4.4%)	RR 0.16 (0.02 to 1.35)	37 fewer per 1000 (from 43 fewer to 15 more)	⊕⊕⊕O MODERATE

(a) Strengths: randomised, low attrition bias (on-treatment analysis so no loss to follow up) Weaknesses: the potential for bias is substantial because both treatment groups contain failures of the other treatment. In addition the patient number is relatively low. No allocation concealment /No intention to treat analysis as it is an on treatment analysis.

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1 **Table 11.21: Multi-vessel disease- short term follow-up (1 year) for stable angina- Angiography pre-randomisation**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Angiography pre randomisation- Multi vessel disease -short term follow-up (1 yr) Medical	PCI or CABG	Relative (95% CI)	Absolute	
<b>Death (follow-up 1 years)</b>											
Rogers 1995[104] (ACIP) (b)	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	8/183 (4.4%)	0/192 (0%)	RR 17.83 (1.04 to 306.73)	40 more per 1000 (from 10 more to 70 more)	⊕⊕⊕O MODERATE
<b>MI (follow-up 1 years)</b>											
Rogers 1995[104] (ACIP)	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	10/183 (5.5%)	5/192 (2.6%)	RR 2.1 (0.73 to 6.02)	29 more per 1000 (from 7 fewer to 131 more)	⊕⊕⊕O MODERATE
<b>Repeat revascularisation (follow-up 1 years)</b>											
Rogers 1995[104] (ACIP)	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	44/183 (24%)	18/192 (9.4%)	RR 2.56 (1.54 to 4.27)	147 more per 1000 (from 51 more to 307 more)	⊕⊕⊕O MODERATE

- (a) Strengths: randomised, baseline characteristics reported. Intention to treat analysis reported. At 1 year after entry, follow-up was 100% complete for death and 96% complete for other clinical events. Weaknesses: allocation concealment not reported. This is a 1 year follow-up of the ACIP study.
- (b) 3 arms to the study: 1) Pharmacologic therapy to suppress angina (angina guided therapy) 2) Pharmacologic therapy to suppress both angina and ambulatory ECG evidence of ischemia (ischemia guided strategy) 3) Revascularisation with either angioplasty or surgery. We have analysed data for only 2 arms – angina guided therapy vs. revascularisation.

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1 **Table 11.22: Multi-vessel disease- medium term follow-up (2-4 year) for stable angina- Angiography pre-randomisation**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Angiography pre randomisation- Multi vessel medium term follow-up (2 years) Medical	PCI or CABG	Relative (95% CI)	Absolute	
<b>Death (follow-up 2 years)</b>											
Davies 1997[105] (ACIP)	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	12/183 (6.6%)	2/192 (1%)	RR 6.3 (1.43 to 27.74)	53 more per 1000 (from 4 more to 267 more)	⊕⊕⊕O MODERATE
<b>Non protocol revascularisation (follow-up 2 years)</b>											
Davies 1997[105] (ACIP)	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	56/183 (30.6%)	25/192 (13%)	RR 2.35 (1.54 to 3.60)	175 more per 1000 (from 70 more to 338 more)	⊕⊕⊕O MODERATE

(a) Strengths: randomised, baseline characteristics reported. Intention to treat analysis reported. Weaknesses: allocation concealment not reported.

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1 **Table 11.23: Multi-vessel disease- Long term follow-up (>4 years) for stable angina**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Multi vessel disease- Long term follow-up (5 years) Medical	PCI or CABG	Relative (95% CI)	Absolute	
<b>Death (patients with type 2 diabetes) (follow-up 5 years)</b>											
Frye 2009[106] (BARI-2D) (b)	randomised trial	Serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	121/991 (12.2%)	112/953 (11.8%)	RR 1.04 (0.82 to 1.32)	5 more per 1000 (from 21 fewer to 38 more)	⊕⊕⊕○ MODERATE
<b>Death (in PCI stratum in BARI-2D) (follow-up 5 years)</b>											
Frye 2009[106] (BARI-2D) (b)	randomised trial	Serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	82/807 (10.2%)	86/798 (10.8%)	RR 0.94 (0.71 to 1.26)	6 fewer per 1000 (from 31 fewer to 28 more)	⊕⊕⊕○ MODERATE
<b>Death (in CABG stratum in BARI-2D) (follow-up 5 years)</b>											
Frye 2009[106] (BARI-2D) (b)	randomised trial	Serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	63/385 (16.4%)	51/378 (13.5%)	RR 1.21 (0.86 to 1.71)	28 more per 1000 (from 19 fewer to 96 more)	⊕⊕⊕○ MODERATE
<b>Freedom from CV events (death, MI or stroke) - PCI stratum (BARI-2D) (follow-up 5 years)</b>											
Frye 2009[106] (BARI-2D) (b)	randomised trial	Serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	637/807 (78.9%)	614/798 (76.9%)	RR 1.03 (0.97 to 1.08)	23 more per 1000 (from 23 fewer to 62 more)	⊕⊕⊕○ MODERATE
<b>Freedom from CV events (death, MI or stroke)- CABG stratum(BARI-2D) (follow-up 5 years)</b>											
Frye 2009[106] (BARI-2D) (b)	randomised trial	Serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	268/385 (69.6%)	293/378 (77.5%)	RR 0.9 (0.82 to 0.98)	78 fewer per 1000 (from 15 fewer to 140 fewer)	⊕⊕⊕○ MODERATE

(a) Strengths: Large scale randomised control trial (randomisation method not reported), intention to treat analysis, power calculation for 5 year follow-up reported, baseline comparisons were made Weaknesses: No allocation concealment reported, not all of the patients enrolled suffered from stable angina.

(b) All patients underwent clinically indicated coronary angiography before randomisation; most of them provided consent during screening before angiography but after meeting clinical eligibility requirements. Thus, the number of patients who were excluded for reasons unrelated to coronary anatomy was unavailable.

**Interaction between study group assignment in to PCI and CABG stratum in the BARI-2D trial**

At 5 years, the rates of death did not differ significantly between the revascularisation group and medical therapy group in either the CABG [RR 1.21 (0.86 to 1.71)] or the PCI stratum [RR 0.94 (0.71 to 1.26)]. The interaction between study group assignment and intended method of revascularisation was not significant for death (p=0.27). Patients in the CABG stratum who were assigned to the revascularisation group had significantly more patients were free from major cardiovascular events than did patients in the CABG stratum

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1 who were assigned to the medical therapy group [RR 0.90 (0.82 to 0.98)]. Freedom from major cardiovascular events among patients in the PCI stratum assigned to the  
2 revascularisation group did not differ significantly from those who were assigned to the medical therapy [RR (1.03 (0.97 to 1.08))].  
3 The interaction between study group assignment and intended method of revascularisation was significant for freedom from major cardiovascular events ( $p=0.01$ ), which indicated  
4 that the benefit associated with prompt coronary revascularisation, as compared with medical therapy, was significantly greater for patients selected for CABG than for patients  
5 selected for PCI.

1 **Quality of life data from studies:**

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3 **Medical vs. PCI or CABG**

4 **Pfisterer 2003[102] (TIME) – 6 months and 1 year follow-up**

5 Quality of Life was measured by items from three questionnaires that and 91% of  
6 surviving patients at 4 year follow-up provided data for this. The questionnaires were  
7 the short-form SF36 the Duke Activity Status Index (DASI) and the Rose questionnaire.

8 **Results:** N=282 (CABG or PCI n=140; Medical n=142). QOL increased in both  
9 groups. The Rose Score and the General Health component of SF36 showing  
10 significantly larger improvements for the revascularization group compared to the  
11 medical group at the time of the first follow-up (6 months). However, improvements  
12 were no longer significant between the 2 treatment groups after 1 year.  
13 Improvements in the DASI were not significantly different between groups at 6 or 12  
14 months follow-up.

15  
16 **Pfisterer 2004[103] (TIME) – 4 year follow-up:**

17 See above, instead of SF36 they reported SF12 results.

18 **Results:** N=282 (CABG or PCI n=140; Medical n=142). After 4 years cores from the  
19 Rose, SF12 physical component and DASI continued to be significantly improved  
20 compared to baseline. However, none of the group differences were significant. The  
21 SF12 mental-component summary scores did not change significantly in either  
22 treatment group ( $p=.29$ ) compared to baseline and remained constant throughout the  
23 entire study period.

24  
25 **Medical vs. PCI vs. CABG**

26 **Favarato 2007[107] (MASS-II)**

27 In the MASS II study Quality of Life was assessed using short form 36 (SF-36) at 1  
28 year. The SF-36 comprises 36 items that can be combined in to the following 8 multi-  
29 item summary scores: physical functioning (10 items), vitality (four items), bodily pain  
30 (2 items), mental health (5 items), social functioning (2 items), role limitations due to  
31 physical health (4 items) and due to emotional problems (3 items) and general health  
32 perceptions (5 items), plus one item assessing a change in health over the past year.  
33 Each summary score is obtained by simple unweighted summation of item scores and is  
34 scaled from 0 to 100, with 0 and 100 indicating 'worst' and 'best' possible health,  
35 respectively (higher scores indicate better perceived health).

36 **Results:** N=522 (n=17 medical treatment, n=180 PCI, n=175 CABG). The SF-36  
37 mean scores for CABG, PCI and Medical treatment respectively were: Role physical  
38 (48.37 vs. 50 vs. 40.26); role emotional (66.08 vs. 63.48 vs. 62.63); mental health  
39 (70.69 vs. 70.43 vs. 68.13); vitality (71.33 vs. 67.37 vs. 61.59); physical functioning  
40 (71.51 vs. 68.29 vs. 62.63); bodily pain (72.24 vs. 70.10 vs. 64.92); general health

(76.59 vs. 71.32 vs. 69.58); social functioning (81.89 vs. 81.82 vs. 77.05). None of the eight SF-36 scores showed a significant treatment difference at 12 months between PCI and CABG. However, the CABG group better than the medical group in terms of vitality ( $p < 0.001$ ), physical functioning ( $p < 0.001$ ) and general health ( $p < 0.001$ ) at 12 months. The PCI group showed significantly superiority over the medical group just in terms of vitality and functioning at 12 months ( $p < 0.001$ ). All these treatment comparisons were done using the analysis of co-variance, adjusting for the patient's baseline scores.

### 11.4.3 Economic evidence

One study[108] was found that included the relevant comparison in people with type 2 diabetes mellitus. This is summarised in the economic evidence profile below. See also Economics Evidence Tables in Appendix G.

**Table 11.24: CABG and PCI vs. medical treatment - Economic study characteristics**

Study	Limitations	Applicability	Other Comments
Hlatky 2009[108]	Potentially serious limitations (a)	Partial applicability (b)	Based on the BARI 2D trial. Patients had type 2 diabetes mellitus and stable, angiographically documented coronary disease.

- (a) Not clear how utility estimates were used to calculate results in the study. In the clinical paper the probability of cardiovascular events was lower in the CABG stratum and this was inconsistent with the QALYs calculation. QALYs were not adjusted by baseline values.
- (b) USA study.

**Table 11.25: CABG and PCI vs. medical treatment - Economic summary of findings**

Study	Incremental cost per patient over 4 years	Incremental QALYs per patient over 4 years	ICER (£/QALY)	Uncertainty
Hlatky 2009[108]	Medical treatment costs saving (a)	Medical treatment more effective (b)	Medical treatment dominant	Medical therapy was not dominant but still cost-effective when: <ul style="list-style-type: none"> <li>- results were extrapolated to lifetime assuming costs after 4 years are the same in the 2 groups</li> <li>- QALYs were adjusted by baseline values</li> <li>- a reduced survival after MI (2 and 3 years) and after non-fatal stroke (3 years) was assumed.</li> </ul>

- (a) Costs included were hospitalisation, outpatient visits, nursing home/rehab, medications, test and procedure. Hospital costs calculated using a ratio of cost to charges. Discounted by 3%.
- (b) Total QALYs at 4 years were higher in the medical treatment group compared to both the PCI and CABG strata.

1 11.4.4 Evidence statements

**Clinical** **Multi-vessel disease- short term follow-up (1 year) for stable angina**

**Pfisterer 2003[102] (TIME):** Evidence from one RCT shows that there was no significant difference medical treatment or revascularisation (PCI or CABG) for death [RR 0.73 (0.36 to 1.47)], MI [RR 1.48 (0.78 to 2.81)]. There was significantly higher non protocol revascularisation [RR 4.59 (2.80 to 7.51)] in the medical treatment compared to revascularisation (PCI or CABG). [Follow-up 1 year]

**Angiography pre-randomisation – Multi- vessel disease- short term follow-up (1 year) for stable angina**

**Rogers 1995[104] (ACIP):** Evidence from one RCT shows that there were significantly higher deaths in medical treatment compared to revascularisation (medical or PCI), there no significant difference medical treatment or revascularisation (PCI or CABG) for MI [RR 2.10 (0.73 to 6.02)], MI [RR 1.64 (0.95 to 2.84)] .There was significantly higher non protocol revascularisation [RR 2.56 (1.54 to 4.27)] in the medical treatment group compared to revascularisation (PCI or CABG).[follow-up 1 year]

**Multi-vessel disease- medium term follow-up ( 2 to 4 years) for stable angina**

**Pfisterer 2004[103] (TIME):** Evidence from 1 RCT shows that there was no significant difference between medical treatment and revascularisation (PCI or CABG) for death [RR 1.05 (0.67 to 1.65)], non fatal MI, [RR 0.16 (0.02 to 1.35)] . There were significantly higher non protocol revascularisations [RR 0.99 (0.25 to 3.86)] in medical treatment compared to revascularisation (PCI or CABG). [Follow-up 4 years]

**Angiography pre-randomisation – Multi-vessel disease- medium term follow-up ( 2 to 4 years) for stable angina**

**Davies 1997[105] (ACIP):** Evidence from one RCT shows that there was no significant difference between medical treatment and revascularisation (PCI or CABG) for death [RR 6.30 (1.43 to 27.74)]. There were significantly higher non protocol revascularisations [RR 2.35 (1.54 to 3.60)] in the medical treatment group compared to revascularisation (PCI or CABG). [Follow-up 2 years]

**Multi-vessel disease - Long term follow-up (5 years) for stable angina**

**Frye 2009[106] (BARI-2D):** Evidence from one RCT in patients with type 2 diabetes shows that there was no significant difference

between medical treatment and revascularisation (PCI or CABG) for death [RR 1.04 (0.82 to 1.32)] [Follow-up 5.3 years]

**Frye 2009[106] (BARI-2D):** Evidence from one RCT in patients with type 2 diabetes shows that the rates of death did not differ significantly between the revascularisation group and medical treatment group in either the CABG [RR 1.21 (0.86 to 1.71)] or the PCI stratum [RR 0.94 (0.71 to 1.26)]. The interaction between study group assignment and intended method of revascularisation was not significant for death (p=0.27). [Follow-up 5.3 years]

**Frye 2009[106] (BARI-2D):** Evidence from one RCT in patients with type 2 diabetes shows that patients in the CABG stratum who were assigned to the revascularisation group had significantly more patients were free from major cardiovascular events than did patients in the CABG stratum who were assigned to the medical therapy group [RR 0.90 (0.82 to 0.98)]. Freedom from major cardiovascular events among patients in the PCI stratum assigned to the revascularisation group did not differ significantly from those who were assigned to the medical therapy [RR (1.03 (0.97 to 1.08))]. The interaction between study group assignment was and intended method of revascularisation significant for freedom from major cardiovascular events (p=0.01). [Follow-up 5.3 years]

**Economic**

Medical treatment is more cost-effective than early revascularisation with either CABG or PCI in people with type 2 diabetes mellitus and stable coronary artery disease. This evidence has potentially serious limitations and partial applicability.

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2 **11.5 Recommendations and link to evidence (medical vs. recascularisation)**

Recommendation	<p><b>Do not routinely offer percutaneous coronary intervention (PCI) or coronary artery bypass grafting (CABG) to people whose symptoms are controlled with drug treatment.</b></p> <p><b>Consider revascularisation (PCI or CABG) for people whose symptoms are not controlled with drug treatment.</b></p>
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Relative values of different outcomes

Outcomes of interest included long-term mortality (total and cardiovascular), rates of major adverse cardiovascular events (myocardial infarction, stroke, myocardial revascularisation), measures of symptom severity (frequency of angina, exercise test outcomes), and quality of life.

Trade off between clinical benefits and

**Medical treatment versus CABG**

harms

The randomised trials of initial treatment strategies of coronary artery bypass surgery versus continued medical therapy included in the evidence review were mainly conducted in the 1970's. The individual patient data meta-analyses of the trials suggests that coronary artery bypass surgery is associated with a survival advantage that persists for about ten years and translates into an extension of life of around four months (95% CI 1.91-6.61). This benefit is greater for people with three-vessel or left main stem disease (extensions of survival of 5.7 and 19.3 months, respectively) than for people with less extensive disease (one or two vessel disease). Moreover, a risk model incorporating multiple clinical and angiographic predictor variables suggests that absolute survival benefit is greatest in patients at highest baseline risk[65]. The trials also confirm that coronary artery bypass surgery is associated with a greater improvement in symptoms of angina and quality of life than continued medical therapy, and this benefit persists for up to ten years (CASS)[72,79,81]. These trials of coronary surgery have several important limitations, which restrict their relevance to current clinical practice:

The trials recruited a highly selected group of patients who were considered suitable for management with either initial treatment strategy. Moreover, the individual trials have limited statistical power and the meta-analysis of the trials included only 2649 patients, of whom only 150 patients had left main stem disease.

Interpretation of the trial results is confounded by relatively high rates of myocardial revascularisation among patients initially assigned to a strategy of continued medical therapy. In the individual patient data meta-analysis 25% and 41% of medical patients underwent coronary surgery by five and ten years, respectively. This high crossover rate may dilute any long term differences between the two treatment strategies and lead to an underestimation of the true treatment effect.

The profile of patients undergoing coronary artery bypass surgery has changed over time. In the individual patient data meta-analysis almost all were male (96.8%), the majority were aged between 40 and 60 years (mean age 50.8 years), and the mean left ventricular ejection fraction was 59.4[65]. It is likely that most of the patients enrolled in the trials would therefore have been good candidates for surgical revascularisation. By contrast, in contemporary surgical practice coronary artery bypass operations are often

carried out in elderly people with extensive coronary artery disease, impaired left ventricular function, and multiple co-morbidities.

Since the early trials of CABG were conducted there have been major advances in the surgical treatment of people with angina. For example, in the individual patient data meta-analysis internal mammary artery grafts were used in only 9.9% of patients assigned to coronary bypass surgery, but internal mammary artery grafts are associated with improved long-term outcome and are used routinely in contemporary cardiac surgical practice[109,110].

Most of the trials were conducted before the widespread introduction of contemporary secondary prevention measures, which improve long-term outcome in people with coronary artery disease. In the individual patient data meta-analysis only 18.8% of the medical patients were taking antiplatelet agents, and statins, angiotensin-converting enzyme inhibitors, and thienopyridines were not used. It is noteworthy that treatment with a statin alone can result in a reduction in coronary heart disease mortality similar to that associated with coronary bypass surgery[111]. In a large meta-analysis a reduction in mean LDL cholesterol of 1.09mmol/L resulted in a 12% reduction in all-cause mortality and a 19% reduction in coronary heart disease mortality[112]. In the individual patient data meta-analysis coronary artery surgery was associated with a 39% relative risk reduction in mortality at five years, and a 17% reduction at ten years. Whether bypass surgery confers incremental prognostic benefit in people with three vessel or left main stem disease who are also treated with contemporary secondary prevention therapies (antiplatelet agents, statins, renin-angiotensin system inhibitors) is unknown.

### **Medical treatment versus PCI**

Randomised trials of initial treatment strategies of percutaneous coronary intervention versus continued medical therapy recruited patients who were considered suitable for either treatment strategy. Overall we found no evidence of a beneficial effect of percutaneous coronary intervention on medium or long-term mortality. Percutaneous coronary intervention was associated with better symptom relief than continued medical therapy, but this treatment difference attenuated over several years, partly because many patients assigned to medical

therapy subsequently underwent myocardial revascularisation. Several of the trials also reported greater improvements in quality of life scores among coronary intervention patients, but this was not confirmed in all trials (AVERT)[90] or sustained long-term (RITA-2[89,92], COURAGE[91,94]). Treatment effects were consistent among people with single vessel and multi-vessel disease.

Several issues limit the relevance of these trials to contemporary practice:

The trials recruited a highly selected group of patients and it is likely that high risk patients were systematically excluded. For example, in RITA-2[89,92], participating hospitals carried out over 70 000 coronary arteriograms during the recruitment period, but only identified 2750 eligible patients of whom 1018 patients were randomized (RITA-2)[89,92]. In COURAGE[91,94] 35,539 patients were screened of whom 3071 met the eligibility criteria and 2287 were subsequently randomised (6.4% of the screened population). COURAGE [91,94] excluded high risk patients with severe (CCS class IV) angina, marked ischaemia on an exercise test, or impaired left ventricular function.

Several of the trials (RITA-2[89,92], ACME [93,96], AVERT [90], MASS-I[83,84]) recruited patients before coronary artery stents were available. Only two trials compared an initial treatment strategy of percutaneous coronary intervention using bare metal stents with an initial treatment strategy of continued medical therapy (MASS-II[66,67,77], COURAGE[91,94]). We found no trials of percutaneous coronary intervention with drug-eluting stents versus optimal medical therapy.

In several of the trials the use of statins and ACE inhibitors was suboptimal by contemporary standards. The COURAGE[91,94] trial is the largest trial to compare percutaneous coronary intervention and 'optimal' medical therapy (including anti-platelet therapy, anti-ischaemic therapy, renin-angiotensin system inhibition, and lipid-lowering therapy) with optimal medical therapy alone, but with only 413 end point events the trial has limited statistical power for the primary endpoint (death or non-fatal myocardial infarction). Moreover, after a mean follow-up of 4.6-years vital status was unknown in 8.4% of the patients in COURAGE [91,94].

All of the trials reported high rates of revascularisation

procedures among patients assigned to medical therapy, which may have reduced any real differences between the two treatment strategies [non-protocol revascularisation rates - 630/1855 (34%) in medical group and 463/1858 (24.9%) in PCI group at long term follow-up].

The spectrum of patients considered suitable for PCI has changed over time and with evolving techniques and increasing operator experience a wider range of people with a more complex pattern of coronary artery disease are now considered eligible for percutaneous treatment.

### **Medical treatment versus myocardial revascularisation**

These trials compared initial treatment strategies of continued medical therapy with myocardial revascularisation (either PCI or CABG) in patients considered suitable for either treatment strategy.

The ACIP[104,105] trial reported lower short- and medium-term mortality in patients assigned to a revascularisation strategy, but there was no difference in mortality between the treatment groups in TIME[102,103] or BARI-2D[106]. In BARI-2D[106], among patients prospectively stratified to revascularisation by CABG (CABG stratum) there was a significant difference in major cardiovascular events (death, myocardial infarction, or stroke) between surgical and medical treatment groups. This difference was driven mainly by a difference in the rate of myocardial infarction but there was no difference in five year survival between patients randomised to revascularisation and medical therapy [113].

Interpretation of these trials is complicated by several limitations:

ACIP [104,105] recruited patients before the introduction of coronary artery stents and amongst 192 patients assigned to revascularisation 102 patients were selected for coronary balloon angioplasty and 90 patients for coronary bypass surgery. Aspirin was prescribed to 89% of patients but statins were not used.

The TIME [102,103] trial did not report the use of stents among patients who underwent percutaneous coronary intervention. Lipid-lowering therapy was used in only

23% of patients.

In BARI-2D [106] most patients were treated with statins and renin-angiotensin system inhibitors. Among 1176 patients assigned to the revascularization group, 765 underwent PCI, of whom 34.7% received a drug-eluting stent and 56.0% received a bare metal stent (9.3% did not receive a stent). In BARI-2D[106] 42.1% of patients in the medical group underwent myocardial revascularization during follow-up.

The GDG concluded that there is no evidence that myocardial revascularisation confers incremental prognostic benefit in people with stable angina who are also treated with contemporary medical therapy (antiplatelet agents, statins, and renin-angiotensin system inhibitors). All people with angina should be offered appropriate medical therapy and only considered for invasive investigation and myocardial revascularisation if anginal symptoms are not optimally controlled by anti-anginal medication.

Economic considerations

Medical treatment is more cost-effective than early revascularisation with either CABG or PCI in people with stable coronary artery disease including people with type 2 diabetes mellitus.

Quality of evidence

See discussion under balance of benefits and harms above.

The economic evidence regarding people with stable coronary artery disease has overall minor limitations but partial applicability. The economic evidence regarding people with type 2 diabetes and stable coronary artery disease has potentially serious limitations (unclear QALY calculations) and partial applicability (USA study).

## 1 11.6 Recommendations and link to evidence (Subgroup populations)

<p>Recommendation</p>	<p><b>Do not exclude people with stable angina from treatment based on their age alone.</b></p> <p><b>Do not investigate or treat symptoms of stable angina differently in men and women or in different ethnic groups.</b></p>
<p>Other considerations</p>	<p>Elderly people commonly present with symptoms of cardiovascular disease. The very old (&gt; 80 or 85 years</p>

depending on definition used) and those with co-morbidity are commonly not included in trials and the GDG considered that this was an area where consensus recommendation was required. The GDG considered that there was no clear evidence that age influenced response to treatment although greater age can be associated with frailty and co-morbidity. The absolute risk for the elderly will however be greater. The GDG agreed that age alone should not preclude consideration for medical or revascularisation treatment.

Review protocols included women and people belonging to South Asian ethnic group. No evidence was found to indicate that investigation or treatment should differ for these people.

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## 2 **12 Revascularisation**

### 3 **12.1 Introduction**

4 People with stable angina may be considered for myocardial revascularisation with  
5 percutaneous coronary intervention or coronary artery bypass surgery. The choice  
6 between the two revascularisation procedures is influenced by the result of coronary  
7 arteriography. Some patients are not angiographically suitable for percutaneous or  
8 surgical revascularisation but many patients are technically suitable for either  
9 revascularisation technique. Over the last three decades randomised controlled trials  
10 have compared strategies of percutaneous coronary (balloon) angioplasty (1980s),  
11 percutaneous coronary intervention using bare metal stents (1990s), and percutaneous  
12 coronary intervention using drug-eluting stents (2000s) with coronary artery bypass  
13 surgery in patients who are suitable for either method of revascularisation.

14 The following key clinical question is addressed in this chapter: In adults with stable  
15 angina, what is the clinical/cost effectiveness of revascularisation techniques to  
16 alleviate angina symptoms and to improve long term outcomes?

17 The evidence review for the determination of the clinical effectiveness of PCI vs.  
18 CABG included a total of 42 papers.

19 Twenty seven papers included patients with multi- vessel disease, 10 papers focused  
20 on single vessel disease, two paper focused on patients with left main coronary  
21 disease , two papers included patients with left main coronary artery or three vessel  
22 disease and one included paper was an IPD meta-analysis of trials comparing PCI  
23 and CABG.

24 The included IPD [114] included 10 trials with data on 7812 patients with a median  
25 follow-up of 5.9 years. PCI with balloon angioplasty was used in 6 trials and bare  
26 metal stents in 4 trials. Of the 10 trials, 3 trials (BARI [115], ERACI-II[116,117] and  
27 Toulouse[118]) were not included in the study level meta-analyses as they did not  
28 meet the inclusion criteria for the minimum number of stable angina patients. The  
29 results of the IPD meta- analyses should be considered taking into account the inclusion  
30 of unstable angina population in the results.

31 The main outcomes analysed were:

- 32
- Death (all causes)

- 1           • Cardiac death
- 2           • Non fatal MI
- 3           • Stroke
- 4           • Repeat revascularisation (PCI and/or CABG)
- 5           • Free of angina

6           Outcomes were also analysed separately for the sub-groups: Diabetes (yes and no),  
7           age (>65 and <65 years), and number of vessels.

8           The results of the review have been analysed based on the involvement of vessels  
9           (Multi vessel disease, single vessel disease, Left main coronary disease and Left main  
10          coronary artery or 3 vessel disease) and follow-up periods (Immediate follow-up (in-  
11          hospital), short term-follow-up (1 yr), medium term follow-up (2-4 yrs) and long term  
12          follow-up (>5 yrs).

13

14          In this guideline, the classification 'Multi-vessel disease' includes studies with patients  
15          having:

- 16           • Multi-vessel disease only (2 vessel disease, 3 vessel disease)
- 17           • Multi-vessel disease along with single vessel disease

18          This was because most of the studies on revascularisation for stable angina did not  
19          report data separately for multi vessel disease and single vessel disease  
20          separately. Sub group analysis was conducted separately for 2 vessel and 3 vessel  
21          diseases if the results were reported separately in the studies.

22

23          The main results of the review are presented as follows:

24          **A. Multi-vessel disease**

- 25           • Multi-vessel disease - Immediate follow-up for Stable angina
- 26           • Multi -vessel disease -short term follow-up (1 year) for Stable angina
- 27           • Multi-vessel disease - medium term follow-up (2 to 4 years) for Stable angina
- 28           • Multi-vessel disease - Long term follow-up (> 5 years) for Stable angina

29          **B. Single vessel disease**

- 30           • Single vessel disease - short term follow-up (1 year) for Stable angina

- 1           • Single vessel disease- medium term follow-up (2 to 4 years) for Stable angina
- 2           • Single vessel disease- Long term follow-up (>5 years) for Stable angina
- 3           **C. Left main coronary disease**
- 4           ➤ Left main coronary disease - short term follow-up (1 year) for Stable angina
- 5           **D. Left main coronary artery or 3 vessel disease**
- 6           ➤ Left main coronary artery or 3 vessel disease - short term follow-up (1 year) for
- 7           Stable angina
- 8           **E. IPD meta-analyses (Multi-vessel disease- Immediate, short and Long term**
- 9           **follow-up)**
- 10

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2 **12.2 Multi-vessel disease**

3 **12.2.1 Clinical evidence**

4 The "Review Protocol" for this topic can be found in Appendix C, the "Search  
5 Strategies" in Appendix D, the "List of Included and Excluded Studies" in Appendix  
6 E1, the "Clinical Evidence Tables" in Appendix E2, and the "Forest Plots" in Appendix  
7 F.

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1 **Table 12.1: Multi -vessel disease- Immediate follow-up for Stable angina**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Multi vessel disease-Immediate follow-up PCI	CABG	Relative (95% CI)	Absolute	
<b>Stroke</b>											
Eefting 2003[119] Hamm 1994[120] (GABI) Hampton 1993[121] (RITA) King 1994[122] (EAST) Zhang 2006[123] (SOS)	randomised trial	serious (a)	no serious inconsistency (b)	no serious indirectness	serious (c)	none	5/1509 (0.3%)	15/1495 (1%)	RR 0.35 (0.13 to 0.92)	6 fewer per 1000 (from 1 fewer to 9 fewer)	⊕⊕○○ LOW

- 2 (a) All studies randomised, 3 out of 5 studies reported allocation concealment, 4 out of studies blind outcome assessment, ITT reported in all studies.
- 3 (b) No heterogeneity.
- 4 (c) 95% CI around the pooled estimate of effect includes appreciable benefit or appreciable harm.

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**Table 12.2: Multi-vessel disease-short term follow-up (1 year) for Stable angina**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Multi vessel disease-short term follow-up (1 yr)	control	Relative (95% CI)	Absolute	
<b>Death (all causes) (follow-up 1 years)</b>											
Eefting 2003[119]; Hamm 1994[120] (GABI); Rickards 1995 (CABRI); Serruys 2001[124] (ARTS); Sigwart 2002[125] (SoS); Hueb 2004[66] (MASS-II)	randomised trials	serious (a)	serious (b)	no serious indirectness	serious (c)	none	61/2127 (2.9%)	56/2102 (2.7%)	RR 1.06 (0.75 to 1.52)	2 more per 1000 (from 7 fewer to 14 more)	⊕○○○ VERY LOW
<b>Cardiac mortality (follow-up 1 years)</b>											
Eefting 2003[119]	randomised trials	no serious limitations (d)	no serious inconsistency	no serious indirectness	serious (c)	none	0/138 (0%)	2/142 (1.4%)	RR 0.21 (0.01 to 4.25)	11 fewer per 1000 (from 14 fewer to 46 more)	⊕⊕⊕○ MODERATE
<b>Non fatal MI (follow-up 1 years)</b>											
Eefting 2003[119]; Hamm 1994[120] (GABI); Rickards 1995[126] (CABRI); Serruys 2001[124] (ARTS); Sigwart 2002[125] (SoS); Hueb 2004[66] (MASS-II)	randomised trials	serious (a)	serious (e)	no serious indirectness	serious (c)	None	114/2127 (5.4%)	105/2102 (5%)	RR 1.07 (0.83 to 1.39)	3 more per 1000 (from 8 fewer to 19 more)	⊕○○○ VERY LOW
<b>Repeat revascularisation (follow-up 1 years)</b>											
Eefting 2003[119]; Hamm 1994[120] (GABI); Rickards 1995[126] (CABRI); Serruys 2001[124] (ARTS); Sigwart 2002[125]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	538/2127 (25.3%)	93/2102 (4.4%)	RR 5.64 (4.57 to 6.97)	205 more per 1000 (from 158 more to 264 more)	⊕⊕⊕○ MODERATE

(SoS); Hueb 2004[66] (MASS-II)												
<b>Free of angina (follow-up 1 years)</b>												
Eefting 2003[119]; Hamm 1994[120] (GABI); Rickards 1995[126] (CABRI); Serruys 2001[124] (ARTS); Sigwart 2002[125] (SoS); Hueb 2004[66] (MASS-II)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	1435/2110 (68%)	1620/2095 (77.3%)	RR 0.88 (0.85 to 0.91)	93 fewer per 1000 (from 70 fewer to 116 fewer)	⊕⊕⊕⊕	MODERATE
<b>Stroke (follow-up 1 years)</b>												
Eefting 2003[119]; Serruys 2001[124] (ARTS); Sigwart 2002[125] (SoS); Hueb 2004[66] (MASS-II)	randomised trials	no serious limitations (f)	no serious inconsistency	no serious indirectness	serious (c)	None	19/1431 (1.3%)	24/1450 (1.7%)	RR 0.80 (0.44 to 1.45)	3 fewer per 1000 (from 9 fewer to 7 more)	⊕⊕⊕⊕	MODERATE
<b>Subgroup-diabetes- Death (all causes) (follow-up 1 years)</b>												
Abizaid 2001[127] (ARTS); Kapur 2010[128] (CARDia trial); Hueb 2004[66] (MASS-II)	randomised trials	no serious limitations (g)	no serious inconsistency	no serious indirectness	serious (c)	None	18/422 (4.3%)	15/403 (3.7%)	RR 1.15 (0.58 to 2.25)	6 more per 1000 (from 16 fewer to 47 more)	⊕⊕⊕⊕	MODERATE
<b>Subgroup diabetes-MI (follow-up 1 years)</b>												
Abizaid 2001[127] (ARTS); Kapur 2010[128] (CARDia trial)	randomised trials	no serious limitations (g)	no serious inconsistency	no serious indirectness	serious (h)	none	32/366 (8.7%)	17/344 (4.9%)	RR 1.79 (1.01 to 3.17)	39 more per 1000 (from 0 more to 107 more)	⊕⊕⊕⊕	MODERATE
<b>Subgroup diabetes- Repeat revascularisation (follow-up 1 years)</b>												
Abizaid 2001[127] (ARTS); Kapur 2010[128] (CARDia trial)	randomised trials	no serious limitations (g)	no serious inconsistency	no serious indirectness	no serious imprecision	None	55/366 (15%)	8/344 (2.3%)	RR 6.36 (3.07 to 13.16)	125 more per 1000 (from 48 more to 283 more)	⊕⊕⊕⊕	HIGH
<b>Sub group diabetes- Non fatal stroke (follow-up 1 years)</b>												
Kapur 2010[128] (CARDia trial)	randomised trials	no serious limitations (i)	no serious inconsistency	no serious indirectness	serious (c)	None	1/254 (0.4%)	7/248 (2.8%)	RR 0.14 (0.02 to 1.13)	24 fewer per 1000 (from 28 fewer to 4 more)	⊕⊕⊕⊕	MODERATE

<b>Subgroup age&gt;65 yrs- Death (all causes) (follow-up 1 years)</b>											
Zhang 2006[123] (SOS)	randomised trials	no serious limitations (j)	no serious inconsistency	no serious indirectness	serious (c)	None	4/190 (2.1%)	1/205 (0.5%)	RR 4.32 (0.49 to 38.27)	16 more per 1000 (from 2 fewer to 182 more)	⊕⊕⊕○ MODERATE
<b>subgroup age&gt;65 yrs-MI (follow-up 1 years)</b>											
Zhang 2006[123] (SOS)	randomised trials	no serious limitations (j)	no serious inconsistency	no serious indirectness	serious (c)	None	13/190 (6.8%)	17/205 (8.3%)	RR 0.83 (0.41 to 1.65)	14 fewer per 1000 (from 49 fewer to 54 more)	⊕⊕⊕○ MODERATE
<b>Subgroup age&gt;65 yrs- stroke (follow-up 1 years)</b>											
Zhang 2006[123] (SOS)	randomised trials	no serious limitations (j)	no serious inconsistency	no serious indirectness	serious (c)	None	5/190 (2.6%)	5/205 (2.4%)	RR 1.08 (0.32 to 3.67)	2 more per 1000 (from 17 fewer to 65 more)	⊕⊕⊕○ MODERATE
<b>subgroup age&gt;65 yrs- repeat revascularisation (follow-up 1 years)</b>											
Zhang 2006[123] (SOS)	randomised trials	no serious limitations (j)	no serious inconsistency	no serious indirectness	no serious imprecision	None	37/190 (19.5%)	7/205 (3.4%)	RR 5.7 (2.61 to 12.48)	160 more per 1000 (from 55 more to 392 more)	⊕⊕⊕⊕ HIGH
<b>Sub group age &lt;65 yrs- Death (follow-up 1 years)</b>											
Zhang 2006[123] (SOS)	randomised trials	no serious limitations (j)	no serious inconsistency	no serious indirectness	serious (c)	None	8/298 (2.7%)	3/295 (1%)	RR 2.64 (0.71 to 9.85)	17 more per 1000 (from 3 fewer to 90 more)	⊕⊕⊕○ MODERATE
<b>Sub group age &lt;65 yrs-MI (follow-up 1 years)</b>											
Zhang 2006[123] (SOS)	randomised trials	no serious limitations (j)	no serious inconsistency	no serious indirectness	serious (c)	None	8/298 (2.7%)	17/295 (5.8%)	RR 0.47 (0.2 to 1.06)	31 fewer per 1000 (from 46 fewer to 3 more)	⊕⊕⊕○ MODERATE
<b>Sub group age&lt;65 yrs- Stroke (follow-up 1 years)</b>											
Zhang 2006[123] (SOS)	randomised trials	no serious limitations (j)	no serious inconsistency	no serious indirectness	serious (c)	None	2/298 (0.7%)	3/295 (1%)	RR 0.66 (0.11 to	3 fewer per 1000 (from	⊕⊕⊕○ MODERATE

									3.92)	9 fewer to 30 more)	
<b>Sub group age&lt;65 yrs- Repeat revascularisation (follow-up 1 years)</b>											
Zhang 2006[123] (SOS)	randomised trials	no serious limitations (j)	no serious inconsistency	no serious indirectness	no serious imprecision	None	48/298 (16.1%)	14/295 (4.7%)	RR 3.39 (1.91 to 6.02)	113 more per 1000 (from 43 more to 238 more)	⊕⊕⊕⊕ HIGH

- 1 (a) All studies randomised, ITT reported in all studies, 4 out of 6 studies reported allocation concealment, all studies reported of blind outcome assessment.
- 2 (b) I<sup>2</sup>=47%.
- 3 (c) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.
- 4 (d) Method of Randomisation and allocation concealment reported. No loss to follow up. Analysis was conducted on an intent-to-treat basis. An independent committee blinded to the treatment allocation evaluated all events. Risk of bias was low
- 5 (e) I<sup>2</sup>=65%. Considerable heterogeneity
- 6 (f) Randomised, allocation concealment, blind outcome assessment and ITT reported in all studies.
- 7 (g) Randomisation and ITT reported in all studies. , allocation concealment, blind outcome assessment in 2 of 3 studies
- 8 (h) 95% CI around the pooled estimate of effect includes appreciable benefit or appreciable harm.
- 9 (i) Strengths: Randomisation undertaken either by a local secure computer-based system or telephone contact with the coordinating centre stratifying for urgency of intervention, sex, and number of diseased vessels. Allocation concealment reported. Sample size calculation reported. Blind outcome assessors. ITT used. Weakness: None
- 10 (j) Multi centre, Randomisation method reported, allocation concealment reported, sample size calculation reported, baseline comparisons made, Numbers lost to follow
- 11 reported (1 year- 8/488 (1.6%) in PCI and 13/500 (2.6%) in CABG) (not reported separately for >65 yrs of age), Intention to treat analysis reported. Blind outcome
- 12 assessment (A clinical events committee, consisting of study interventionists and surgeons, adjudicated all outcome measures. The members of the clinical events committee
- 13 did not adjudicate patients treated at their own centres and were blinded to the randomisation allocation and of the identities of patients and centres). Not reported if blind
- 14 outcome assessment for quality of life. Patients aware of treatment allocation. \* This study reports 1 year follow-up of the SOS trial reporting outcomes in the subgroup of
- 15 people aged ≥ 65 years.

**Sub group interaction:**

Age >65 yrs and Age <65 yrs: There was no significant difference between sub group of patients with age >65 yrs and age <65 yrs for death (p=0.70), MI (p=0.12), repeat revascularisation (p=0.29) at short term follow-up

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**Table 12.3: Multi-vessel disease-medium term follow-up (2 to 4 yrs) for Stable angina**

Quality assessment							Summary of findings				
							No of patients		Effect		Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Multi vessel disease-medium term follow-up (>1-4 yrs)	control	Relative (95% CI)	Absolute	
<b>Death (all causes) (follow-up 2-4 years)</b>											
Hampton 1993[121] (RITA); King 1994[122] (EAST); Legrand 2004[129] (ARTS); Martuscelli 2008[130] (CABRI); Sigwart 2002[125] (SOS)	randomised trials	no serious limitations (a)	serious (b)	no serious indirectness	serious (c)	none	89/1916 (4.6%)	71/1903 (3.7%)	RR 1.23 (0.91 to 1.67)	9 more per 1000 (from 3 fewer to 25 more)	⊕⊕○○ LOW
<b>Cardiac mortality (follow-up 2-4 years)</b>											
Hampton 1993[121] (RITA); Sigwart 2002[125] (SoS)	randomised trials	no serious limitations (d)	no serious inconsistency	no serious indirectness	serious (c)	none	13/998 (1.3%)	8/1001 (0.8%)	RR 1.64 (0.68 to 3.92)	5 more per 1000 (from 3 fewer to 23 more)	⊕⊕⊕○ MODERATE
<b>Non fatal MI (follow-up 2-4 years)</b>											
Hampton 1993[121] (RITA); King 1994[122] (EAST); Legrand 2004[129] (ARTS); Martuscelli 2008[130] (CABRI)	randomised trials	no serious limitations (e)	serious (f)	no serious indirectness	serious (c)	none	115/1428 (8.1%)	101/1403 (7.2%)	RR 1.12 (0.87 to 1.45)	9 more per 1000 (from 9 fewer to 32 more)	⊕⊕○○ LOW
<b>Repeat revascularisation (follow-up 2-4 years)</b>											
Hampton 1993[121] (RITA); King 1994[122] (EAST); Legrand 2004[129] (ARTS); Sigwart 2002[125] (SoS)	randomised trials	no serious limitations (e)	serious (g)	no serious indirectness	no serious imprecision	none	590/1796 (32.9%)	121/1800 (6.7%)	RR 4.87 (4.06 to 5.85)	260 more per 1000 (from 206 more to 326 more)	⊕⊕⊕○ MODERATE
<b>Free of angina (follow-up 2-4 years)</b>											
Unger 2003[131] (ARTS)	randomised trials	no serious limitations (h)	no serious inconsistency	no serious indirectness	no serious imprecision	none	478/600 (79.7%)	527/605 (87.1%)	RR 0.91 (0.87 to 0.96)	78 fewer per 1000 (from 35 fewer to 113 fewer)	⊕⊕⊕⊕ HIGH
<b>Stroke (follow-up 2-4 years)</b>											
Legrand 2004[129] (ARTS)	randomised trials	no serious limitations (h)	no serious inconsistency	no serious indirectness	serious (c)	none	20/600 (3.3%)	20/605 (3.3%)	RR 1.01 (0.55 to 1.85)	0 more per 1000 (from 15 fewer to 28 more)	⊕⊕⊕○ MODERATE

<b>Sub group diabetes- Mortality (follow-up 2-4 years)</b>											
Booth 2008[132] (SoS); Kurbaan 2001[133] (CABRI); Legrand 2004[129] (ARTS)	randomised trials	no serious limitations (i)	no serious inconsistency	no serious indirectness	serious (c)	none	25/242 (10.3%)	13/233 (5.6%)	RR 1.87 (0.99 to 3.5)	49 more per 1000 (from 1 fewer to 139 more)	⊕⊕⊕O MODERATE
<b>Sub group diabetes- MI (follow-up 3 years)</b>											
Legrand 2004[129] (ARTS)	randomised trials	no serious limitations (j)	no serious inconsistency	no serious indirectness	serious (c)	none	11/112 (9.8%)	6/96 (6.3%)	RR 1.57 (0.6 to 4.09)	36 more per 1000 (from 25 fewer to 193 more)	⊕⊕⊕O MODERATE
<b>Sub group diabetes- Repeat revascularisation (follow-up 2-4 years)</b>											
Booth 2008[132] (SoS) Legrand 2004[129] (ARTS)	randomised trials	no serious limitations (k)	no serious inconsistency	no serious indirectness	no serious imprecision	none	63/180 (35%)	12/170 (7.1%)	RR 4.84 (2.71 to 8.64)	271 more per 1000 (from 121 more to 539 more)	⊕⊕⊕⊕ HIGH
<b>Sub group- Left Anterior descending coronary artery proximally- Death (follow-up 3 years)</b>											
Aoki 2004[134] (ARTS)	randomised trials	no serious limitations (l)	no serious inconsistency	no serious indirectness	serious (c)	none	11/246 (4.5%)	11/253 (4.3%)	RR 1.03 (0.45 to 2.33)	1 more per 1000 (from 24 fewer to 58 more)	⊕⊕⊕O MODERATE
<b>Sub group LAD artery- Stroke (follow-up 3 years)</b>											
Aoki 2004[134] (ARTS)	randomised trials	no serious limitations (l)	no serious inconsistency	no serious indirectness	serious (c)	none	5/246 (2%)	7/253 (2.8%)	RR 0.73 (0.24 to 2.28)	7 fewer per 1000 (from 21 fewer to 35 more)	⊕⊕⊕O MODERATE
<b>Sub group LAD artery- MI (follow-up 3 years)</b>											
Aoki 2004[134] (ARTS)	randomised trials	no serious limitations (l)	no serious inconsistency	no serious indirectness	serious (c)	none	17/246 (6.9%)	16/253 (6.3%)	RR 1.09 (0.56 to 2.11)	6 more per 1000 (from 28 fewer to 70 more)	⊕⊕⊕O MODERATE
<b>Sub group LAD artery- Repeat revascularisation (follow-up 3 years)</b>											
Aoki 2004[134] (ARTS)	randomised trials	no serious limitations (l)	no serious inconsistency	no serious indirectness	no serious imprecision	none	54/246 (22%)	12/253 (4.7%)	RR 4.63 (2.54 to 8.44)	172 more per 1000 (from 73 more to 353 more)	⊕⊕⊕⊕ HIGH

- 1 (a) Randomisation, allocation concealment, and ITT reported in all studies. Blind outcome assessment in 4 out of 5 studies
- 2 (b)  $I^2=60\%$ . Substantial heterogeneity
- 3 (c) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.
- 4 (d) Randomisation, allocation concealment, blind outcome assessment and ITT reported in both studies.
- 5 (e) Randomisation, allocation concealment, and ITT reported in all studies.

- 1 (f)  $I^2=42\%$ . Moderate heterogeneity  
2 (g)  $I^2=77\%$ . High heterogeneity  
3 (h) Multi centre, randomised, allocation concealment reported, sample size calculation reported, baseline comparisons made. Nos. lost to follow-up reported). Intention to treat  
4 analysis reported. Clinical events adjudicated by an independent committee. No risk of bias.  
5 (i) Randomisation, allocation concealment, ITT and blind outcome assessment reported in all studies.  
6 (j) Multi centre, randomised, allocation concealment reported, sample size calculation reported, baseline comparisons made. Nos. lost to follow-up reported (0.4%;  
7 6/1205\*\*). Intention to treat analysis reported. Clinical events adjudicated by an independent committee. No risk of bias. \* This study is a 3 year follow-up of the ARTS  
8 trial. \*\* 1 patient was lost to follow-up, 3 were alive but withdrew their consent from further participation in the trial, and 2 patients were never treated by either modality.  
9 (k) Randomisation, allocation concealment, ITT and blind outcome assessment reported in both studies.  
10 (l) Multi centre, randomised, allocation concealment reported, sample size calculation reported, baseline comparisons made. Nos. lost to follow-up reported (1.2%; 3/243 in  
11 stenting and 3.1%; 8/253 in CABG). Intention to treat analysis reported. Clinical events adjudicated by an independent committee. No risk of bias. \* This study is a sub-  
12 analysis of the ARTS trial comparing 3 year outcomes after stenting vs. CABG in patients with multi vessel disease involving the proximal left anterior descending artery.

1 **Additional data for Multi-vessel disease-medium term follow-up (2 to 4 yrs)**

2

3 **Martuscelli 2008[130] (CABRI) (Follow-up 30 months)**

4 No. of participants: n= 223 (CABG (n=103); PTCA (n=120))

5 At 30 months, of the patients initially randomised to PTCA, required a significantly  
6 higher second revascularisation (46.7% (n=56) vs. 5.8% (n=6); p<0.01) and a third  
7 revascularisation (10% (n=12) vs. 1 (1%); p<0.05).

8

9 **Hampton 1993[121] (RITA) (2.5 Years)**

10 No. of participants: n=1011 (n=501 in the CABG and n=510 in the PTCA)

11 There was striking improvement in reported angina in both the treatment groups at all  
12 follow-ups (1 month, 6 months, 1 and 2 years). However, at every point there was a  
13 significant excess of patients with angina in the PTCA group. At 6 months 11% of  
14 CABG patients had anginal symptoms compared with 31.6% of PTCA patients  
15 (RR=0.35, 95% CI 0.26-0.47; p<0.001). Two years after randomisation the  
16 prevalence of angina in the CABG group had increased to 21.5% but this was still  
17 significantly less than the 31.3% for PTCA patients (p=0.007).

18

19 **Legrand 2004[129] (ARTS) (3 YEARS)**

20 No. of participants: n=1205 (n=600 in stent and n=605 in CABG)

21 After 3 years patients in the surgery group had significantly less angina (12.8% in  
22 surgery vs. 18.4% in the stenting group, p=0.011)

23

24 **King 1994[122] (EAST) (3 years)**

25 No. of participants: n=392 (n=194 in the CABG group and n=198 in the PTCA  
26 group)

27 Angina was more prevalent in the PTCA group at 3 years, with 20% of the patients  
28 having CCS class II, III, or IV angina, as compared with 12% of patients in the CABG  
29 group (p=0.039).

1 **Table 12.4: Multi-vessel disease- Long term follow-up (> 5 years) for Stable angina**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Multi vessel disease- Long term follow-up (> 5 yrs)	control	Relative (95% CI)	Absolute	
<b>Death (all causes) (follow-up 5-13 years)</b>											
Buszman 2009[135] (SoS); Henderson 1998[136] (RITA); Kaehler[137] (GABI 2005); Serruys 2005[138] (ARTS); Hueb 2010[77] (MASS-11)	randomised trials	serious limitations (a)	no serious inconsistency	no serious indirectness	No serious imprecision	none	169/1296 (13%)	166/1297 (12.8%)	RR 1.01 (0.83 to 1.23)	5 more per 1000 (from 22 fewer to 29 more)	⊕⊕⊕○ MODERATE
<b>Cardiac mortality (follow-up 5-13 years)</b>											
Booth 2008[132] (SoS); Henderson 1998[136] (RITA); Kaehler 2005[137] (GABI)	randomised trials	no serious limitations (c)	no serious inconsistency	no serious indirectness	serious (b)	none	47/929 (5.1%)	38/939 (4%)	RR 1.24 (0.82 to 1.87)	10 more per 1000 (from 7 fewer to 35 more)	⊕⊕⊕○ MODERATE
<b>Non fatal MI (follow-up 5-10 years)</b>											
Henderson 1998[136] (RITA); Serruys 2005[138] (ARTS); Hueb 2010[77] (MASS-11)	randomised trials	serious limitations (d)	no serious inconsistency	no serious indirectness	serious (b)	none	102/1082 (9.4%)	80/1087 (7.4%)	RR 1.28 (0.97 to 1.69)	21 more per 1000 (from 2 fewer to 51 more)	⊕⊕⊕○ LOW
<b>Repeat revascularisation (follow-up 5-13 years)</b>											
Buszman 2009[135] (SoS); Henderson 1998[136] (RITA); Kaehler 2005[137] (GABI); King 2000[139] (EAST); Serruys 2005[138] (ARTS); Hueb 2010[77] (MASS-11)	randomised trials	no serious limitations (e)	serious (f)	no serious indirectness	no serious imprecision	none	671/1494 (44.9%)	251/1491 (16.8%)	RR 2.65 (2.35 to 2.98)	278 more per 1000 (from 227 more to 333 more)	⊕⊕⊕○ MODERATE
<b>Stroke (follow-up 5-10 years)</b>											
Serruys 2005[138] (ARTS); Hueb 2010 (MASS-11)	randomised trials	no serious limitations (g)	no serious inconsistency	no serious indirectness	serious (b)	none	34/805 (4.2%)	38/808 (4.7%)	RR 0.90 (0.57 to 1.41)	5 fewer per 1000 (from 20 fewer to 19 more)	⊕⊕⊕○ MODERATE
<b>Sub group diabetes - Death (all causes) (follow-up 05-10 years)</b>											
Booth 2008[132] (SoS); Henderson 1998[136] (RITA); Serruys 2005[138] (ARTS)	randomised trials	no serious limitations (h)	serious (i)	no serious indirectness	serious (b)	none	29/209 (13.9%)	20/203 (9.9%)	RR 1.43 (0.83 to 2.47)	42 more per 1000 (from 17 fewer to 145 more)	⊕⊕⊕○ LOW

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										more)	
<b>Sub group diabetes- Repeat revascularisation (follow-up 5 years)</b>											
Serruys 2005[138] (ARTS)	randomised trials	no serious limitations (g)	no serious inconsistency	no serious indirectness	no serious imprecision	none	48/112 (42.9%)	10/96 (10.4%)	RR 4.11 (2.2 to 7.68)	324 more per 1000 (from 125 more to 696 more)	⊕⊕⊕⊕ HIGH
<b>Sub group diabetes- stroke (follow-up 5 years)</b>											
Serruys 2005[138] (ARTS)	randomised trials	no serious limitations (g)	no serious inconsistency	no serious indirectness	no serious imprecision	none	7/112 (6.3%)	7/96 (7.3%)	RR 0.86 (0.31 to 2.36)	10 fewer per 1000 (from 50 fewer to 99 more)	⊕⊕⊕⊕ HIGH
<b>Sub group diabetes- MI (follow-up 5 years)</b>											
Serruys 2005[138] (ARTS)	randomised trials	no serious limitations (g)	no serious inconsistency	no serious indirectness	serious (b)	none	12/112 (10.7%)	7/96 (7.3%)	RR 1.47 (0.6 to 3.58)	34 more per 1000 (from 29 fewer to 188 more)	⊕⊕⊕⊕ MODERATE
<b>Free of angina (follow-up 5-10 years)</b>											
Serruys 2005[138] (ARTS); Hueb 2010 (MASS-II)	randomised trials	no serious limitations (g)	no serious inconsistency	no serious indirectness	no serious imprecision	none	587/805 (72.9%)	641/808 (79.3%)	RR 0.92 (0.87 to 0.97)	68 fewer per 1000 (from 25 fewer to 110 fewer)	⊕⊕⊕⊕ MODERATE
<b>Sub group-no diabetes -Death (all causes) (follow-up 5-10 years)</b>											
Booth 2008[132] (SoS); Serruys 2005[138] (ARTS)	randomised trials	no serious limitations (j)	serious (k)	no serious indirectness	serious (b)	none	74/908 (8.1%)	68/935 (7.3%)	RR 1.12 (0.82 to 1.54)	9 more per 1000 (from 13 fewer to 39 more)	⊕⊕⊕⊕ LOW
<b>Sub group 2 vessel- Death (follow-up 10 years)</b>											
Booth 2008[132] (SoS)	randomised trials	no serious limitations (l)	no serious inconsistency	no serious indirectness	serious (b)	none	31/305 (10.2%)	16/264 (6.1%)	RR 1.68 (0.94 to 3)	41 more per 1000 (from 4 fewer to 121 more)	⊕⊕⊕⊕ MODERATE
<b>Sub group 3 vessel- Death (follow-up 10 years)</b>											
Booth 2008[132] (SoS)	randomised trials	no serious limitations (l)	no serious inconsistency	no serious indirectness	serious (b)	none	22/183 (12%)	18/236 (7.6%)	RR 1.58 (0.87 to 2.85)	44 more per 1000 (from 10 fewer to 141 more)	⊕⊕⊕⊕ MODERATE
<b>Sub group no diabetes- stroke (follow-up 5 years)</b>											
Serruys 2005[138] (ARTS)	randomised trials	no serious limitations (g)	no serious inconsistency	no serious indirectness	serious (b)	none	16/488 (3.3%)	14/509 (2.8%)	RR 1.19 (0.59 to 2.42)	5 more per 1000 (from 11 fewer to 39 more)	⊕⊕⊕⊕ MODERATE

											more)	
<b>Sub group no diabetes- MI (follow-up 5 years)</b>												
Serruys 2005[138] (ARTS)	randomised trials	no serious limitations (l)	no serious inconsistency	no serious indirectness	serious (b)	none	38/488 (7.8%)	31/509 (6.1%)	RR 1.28 (0.81 to 2.02)	17 more per 1000 (from 12 fewer to 62 more)	⊕⊕⊕○	MODERATE
<b>Sub group no diabetes- Repeat revascularisation (follow-up 5 years)</b>												
Serruys 2005[138] (ARTS)	randomised trials	no serious limitations (l)	no serious inconsistency	no serious indirectness	no serious imprecision	none	134/488 (27.5%)	43/509 (8.4%)	RR 3.25 (2.36 to 4.48)	190 more per 1000 (from 115 more to 294 more)	⊕⊕⊕⊕	HIGH

- 1 (a) Randomisation reported in all studies. Allocation concealment, ITT reported in 4/5 studies. Blind outcome assessment reported in 3 out of 5 studies.
- 2 (b) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.
- 3 (c) Randomisation, allocation concealment, ITT and blind outcome assessment reported in 2 out of 3 studies.
- 4 (d) Randomisation, ITT and reported in all 3 studies. Allocation concealment and blind outcome assessment reported in 2 out of 3 studies.
- 5 (e) Randomisation, allocation concealment, ITT and blind outcome assessment reported in 4 out of 5 studies.
- 6 (f) I<sup>2</sup>=95%. High heterogeneity
- 7 (g) Both randomised, allocation concealment reported in 1 out of 2 studies
- 8 (h) Randomisation, allocation concealment, ITT and blind outcome assessment reported in 2 out of 3 studies.
- 9 (i) I<sup>2</sup>=71%. High heterogeneity
- 10 (j) Randomisation, allocation concealment, ITT and blind outcome assessment reported in both studies.
- 11 (k) I<sup>2</sup>=42 %. Moderate heterogeneity
- 12 (l) Multi centre, randomisation method reported, allocation concealment reported, sample size calculation reported, baseline comparisons made, Numbers lost to follow
- 13 reported (5 years- (1.8%) 9/479 in PCI and (3%) 15/500 in CABG), Intention to treat analysis reported. Blind outcome assessment (A clinical events committee,
- 14 consisting of study interventionists and surgeons, adjudicated all outcome measures. The members of the clinical events committee did not adjudicate patients treated at their
- 15 own centres and were blinded to the randomisation allocation and of the identities of patients and centres). Patients aware of treatment allocation.
- 16
- 17 **Sub group interaction:**
- 18 Diabetes and no diabetes: There was no significant difference between diabetes and no diabetes sub group of patients for death (p=0.45), MI (p=0.79) and repeat
- 19 revascularisation (p=0.51) at long term follow-up.
- 20
- 21 Single, vessel, 2 vessel and 3 vessel disease: There was no significant difference between single, 2 vessel and 3 vessel disease for death all causes (p=0.17) at long term follow-up.

1           **Additional data for Multi-vessel disease- Long term follow-up > 5 years)**

2           **Buszman 2009[135] (SoS) (10 years)**

3           No. of participants: N=100 (PCI (n=50); CABG (n=50)

4           At 10 years, there was significant improvement of anginal symptoms in both groups.  
5           Improvement in anginal symptoms was reported in 88.9% PCI patients and 84.38%  
6           CABG patients; p=ns.

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8           **Quality Of Life data for Multi-vessel disease:**

9           **Pocock1996[4]**

10          One RCT[4] assessed quality of life by a Self reported health status (Nottingham  
11          Health Profile (NHP) which consisted of 2 parts. Part 1 included 38 statements  
12          describing levels of physical, social or emotional distress which are grouped in to 6  
13          dimensions: energy (3 statements), pain (8), emotional reactions (9), sleep (5), social  
14          isolation (5), and physical mobility (8). Scores were calculated for each of the 6  
15          dimensions by summing the number of positive (yes) responses: the higher the score,  
16          the greater the impairment of health. Part 2 of the NHP assessed whether an  
17          individual's health is causing problems with seven aspects of daily life: work, tasks  
18          around the home, social life, home relationships, sex life hobbies and interests and  
19          holidays. For both parts 1 and 2, NHP weighted mean scores were compared with  
20          population norms of the same age and sex derived from a general community survey.

21          **Results:** n=1011 (n=501 in the CABG and n=510 in the PCI). For both PCI and  
22          CABG groups there were marked improvements from baseline in all domains: energy,  
23          pain, emotional reactions, sleep, social isolation and physical mobility. There was no  
24          significant difference between the groups for individual domains. When all items  
25          were combined, the treatment difference at 2 years was 0.79 item (p=0.10) in  
26          favour of the CABG group.

27

28          **Eefting 2003[119]**

29          In one RCT[119] quality-of-life was assessed by the Short Form-36 generic instrument:  
30          scores ranged from 0 (worst) to 100 (best imaginable health status). The following  
31          domains were assessed: Physical functioning, role physical, role emotional, pain,  
32          vitality, general health perception, general mental health.

33          **Results:** n=280 (n=138 PCI and n=142 CABG). At 12 months there was no  
34          significant difference between PCI and CABG groups for any of the domains except  
35          for General Health Perception which was significantly higher in the CABG group  
36          (61.6 vs. 66.9; p=0.03).

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1        **Zhang 2003[140] (SOS)**

2        In one RCT[140] cardiac related health status was assessed with the Seattle Angina  
3        Questionnaire (SAQ), a 19 item self-administered questionnaire that measures 5  
4        domains of CAD related health status: physical limitation, angina stability, angina  
5        frequency, treatment satisfaction, and disease perception/quality of life. Scores  
6        range from 0 to 100 for each domain, with higher scores indicating better functioning.  
7        Each domain measures a unique dimension of CAD, and no summary score is  
8        available.

9        **Results:** At 1 year physical limitation, angina frequency, and quality of life improved  
10       from baseline within each treatment group. However, the greatest overall changes  
11       from baseline, as well as the greatest influence of CABG vs. PCI were seen for the  
12       angina frequency domain (PCI (n=476) vs. CABG (n=496) Physical limitation:  
13       75.2±21.3 vs. 76.6±20.7, p=0.36; Angina frequency: 86.9±19.8 vs. 89.6±18.2,  
14       p=0.03; Treatment satisfaction: 91.2±13.1 vs. 90.0±16.0, p=0.73; Quality of life:  
15       69.8±23.0 vs. 71.5±21.4, p=0.41).

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17       **Legrand 2004[129] (ARTS)**

18       One RCT[129] assessed quality of life by EQ-5D questionnaire.

19       Higher scores on the EQ-5D summary indicate a good quality of life; whereas low  
20       scores on the 5 items of EQ-5D domain reflect a favourable assessment of each  
21       component. The following domains were assessed: Mobility, Self-care, Usual activity,  
22       Pain or discomfort, Anxiety or depression

23       **Results:** n=1205 (n=600 PCI and n=605 in CABG). EQ-5D was assessed at 1 and 3  
24       years. At one year there was significant difference in scores between PCI and CABG,  
25       with benefit observed after CABG in specific domains such as 'mobility' (1.4±2.8 vs.  
26       1.1 ±2.8; p=0.05), 'usual activity' (1.0 ±1.9 vs. 0.8±1.8; p=0.01) and 'anxiety or  
27       depression' (2.5±4.5 vs. 2.0±4.1; p=0.04). At 3 years, there were no significant  
28       differences in quality of life between PCI and CABG (EQ-5D summary: PCI vs. CABG:  
29       85±17 vs. 86±17, p=0.74; EQ-5D domain: Mobility: 1.7±3.0 vs. 1.5±2.9, p=0.46;  
30       Self-care: 0.6±2.5 vs. 0.5±2.3, p=0.87; Usual activity: 1.0±1.9 vs. 0.8±1.7, p=0.09;  
31       Pain or discomfort: 4.9±6.9 vs. 5.2±7.7, p=0.78; Anxiety or depression: 2.4±4.8 vs.  
32       2.2±4.4, p=0.77). More specifically the benefits from CABG seen at one year had  
33       disappeared by 3 years

34       **12.2.2       Economic evidence**

35       Eleven studies [88,119,123,127,129,136,141-145] were found that included the  
36       relevant comparison. These are summarised in the Economics Evidence Tables in  
37       Appendix G. However, none of the studies fully met our quality and applicability  
38       criteria. It was thus decided to build an original economic model to compare PCI and  
39       CABG, which is reported in the economic profile tables below. Please see cost-  
40       effectiveness analysis in Appendix H for further details.

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**Table 12.5: CABG vs. PCI - Economic study characteristics**

Study	Limitations	Applicability	Other Comments
<b>NGGC model</b> (Appendix H)	Minor limitations (a)	Direct applicability	Based on the systematic review (see 12.2.1) including only studies where PCI was with stents. Patients had multi vessel disease and were suitable for both PCI and CABG.

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(e) Based on clinical data up to 10 years (limited time horizon).

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**Table 12.6: CABG vs. PCI - Economic summary of findings**

Study	Incremental cost per patient over ten years (£)	Incremental effectiveness (QALYs)	ICER (£/QALY)	Uncertainty
<b>NGGC model</b> (Appendix H)	2,427 (a, b, c)	0.0694 (b, c)	34,971 (c)	95% CI: CABG dominant – PCI dominant. At a willingness to pay of £20,000/QALY, PCI has 63% of probability of being cost-effective, while CABG has 37% of probability. If more than 85% of the repeat procedures are CABG, PCI is no longer cost-effective.

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(a) Cost of initial procedures, further revascularisations, further investigations, medications, treatment of myocardial infarctions.

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(b) Discounted by 3.5%.

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(c) Results of probabilistic analysis.

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Patients in the model had multi-vessel disease; in single vessel disease the repeat revascularisation rate is generally lower compared to multi-vessel disease and PCI is likely to be an even more cost-effective option for this group of patients.

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The other studies considered for inclusion[88,119,123,127,129,136,141-145] (see economic evidence tables in Appendix G) consistently reported higher cost of CABG compared to PCI. The difference in costs tends to decrease when a longer follow-up time was considered (e.g. in the ARTS study [129], RITA trial [136]). Of the other three cost-utility analyses[88,119,145], two[119,145] showed that CABG was not cost-effective but their analysis was limited to a one-year time horizon. The other analysis[88] concluded that CABG was cost-effective in patients suitable for both procedures; however this study was based on non-randomised data and probably most of the PCI procedures were without stents.

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**12.2.3 Evidence statements**

**Clinical Multi -vessel disease (Immediate follow-up)**

**Eefting 2003[119]; Hamm 1994[120] (GABI); Hampton 1993[121] (RITA); King 1994[122] (EAST); Zhang 2006[123]**

**(SoS):** Evidence from 5 RCTs shows that there was significantly higher stroke in CABG patients compared to PCI [RR 0.35 (0.13 to 0.92)] at immediate follow-up (in-hospital event). [Immediate follow-up].

#### **Multi-vessel disease (Short term follow-up – 1 year)**

**Eefting 2003[119]; Hamm 1994[120] (GABI); Rickards 1995[126] (CABRI); Serruys 2001[124] (ARTS); Sigwart 2002[125] (SoS); Hueb 2004[66] (MASS-II):** Evidence from 6 RCTs shows that there were significantly higher repeat revascularisations in the PCI group compared to CABG [RR 5.64 (4.57 to 6.97)] . There were significantly more patients in CABG were free of angina compared to PCI [RR 0.88 (0.85 to 0.91)]. There was no significant difference between PCI and CABG for death (all causes) [RR 1.06 (0.75 to 1.52) and non fatal MI [RR 1.07 (0.83 to 1.39)] [1 year follow-up]

**Eefting 2003[119]:** Evidence from one RCT shows that there was no significant difference between PCI and CABG for cardiac mortality [RR 0.21 (0.01 to 4.25)]. [1 year follow-up].

**Eefting 2003[119]; Serruys 2001[124] (ARTS); Sigwart 2002[125] (SoS); Hueb 2004[66] (MASS-II):** Evidence from 4 RCTs shows that there was no significant difference between PCI and CABG for stroke. [RR 0.80 (0.44 to 1.45)]. [1 year follow-up].

**Abizaid 2001[127] (ARTS); Kapur 2010[128] (CARDia):** Evidence from 2 RCT shows that there was significantly higher repeat revascularisation [RR 6.36 (3.07 to 13.16)] and there was no significant difference between PCI and CABG for MI [RR 1.79 (1.01 to 3.17)] in a sub group of people with diabetes. [1 year follow-up]

**Abizaid 2001 (ARTS); Kapur 2009[128] (CARDia ); Hueb 2004[66] (MASS-II):** Evidence from 3RCT shows that there was no significant difference between PCI and CABG for death [RR 1.15 (0.58 to 2.25)] in a sub group of people with diabetes. [1 year follow-up]

**Kapur 2010[128] (CARDia ):** Evidence from one RCT shows that there was no significant difference between PCI and CABG for stroke [RR 0.14 (0.02 to 1.13)] in a sub group of patients with diabetes [1 year follow-up]

**Zhang 2006[123] (SoS):** Evidence from one RCT shows that there was significantly higher repeat revascularisation [RR 5.7 (2.61 to 12.48)] in the PCI compared to CABG and there was no significant difference between PCI and CABG for death all causes [RR 4.32 (0.49 to 38.27)], MI [RR 0.83 (0.41 to 1.65)], stroke [RR 1.08 (0.32 to 3.67)], in a sub group of people aged > 65 years. [1 year follow-up].

**Zhang 2006[123] (SoS):** Evidence from one RCT shows that there was significantly higher repeat revascularisation [RR 3.39 (1.91 to 6.02)] in the PCI compared to CABG and there was no significant difference between PCI and CABG for death all causes [RR 2.64 (0.71 to 9.85)], MI [RR 0.47 (0.20 to 1.06)], stroke [RR 0.66 (0.11 to 3.92)], in a sub group of people aged < 65 years. [1 year follow-up].

Sub group interaction: There was no significant difference between sub group of patients with age >65 yrs and age <65 yrs for death ( $p=0.70$ ), MI ( $p=0.12$ ), repeat revascularisation ( $p=0.29$ ) at short term follow-up

#### **Multi -vessel disease (Medium term follow-up - >1 to 4 years)**

**Hampton 1993[121] (RITA); King 1994[122] (EAST); Legrand[129] 2004 (ARTS); Martuscelli 2008[130] (CABRI); Sigwart 2002[125] (SoS):** Evidence from 5 RCTs shows that there was no significant difference between PCI and CABG for death (all causes) [RR 1.23 (0.91 to 1.67)]. [2 – 4 years follow-up]

**Hampton 1993[121] (RITA); Sigwart 2002[125] (SOS):** Evidence from 2 RCTs shows that there was no significant difference between PCI and CABG for cardiac mortality [RR 1.64 (0.68 to 3.92)] [2– 4 years follow-up]

**Hampton 1993[121] (RITA); King 1994[122] (EAST); Legrand 2004 (ARTS); Sigwart 2002[125] (SOS):** Evidence from 4 RCTs shows that there was no significant difference between PCI and CABG for non fatal MI [RR 1.12 (0.87 to 1.45)] [2 – 4 years follow-up]

**Hampton 1993[121] (RITA); King 1994[122] (EAST); Legrand 2004 (ARTS); Sigwart 2002[125] (SOS):** Evidence from 4 RCTs shows that there was significantly higher repeat revascularisation in the PCI group compared to CABG [RR 4.87 (4.06 to 5.85)]. [2– 4 years follow-up]

**Unger 2003[131] (ARTS):** Evidence from one RCTs shows that there was significantly higher patients free of angina in the CABG group compared to PCI [RR 0.91 (0.87 to 0.96)]. [2 years follow-up].

**Legrand 2004[129] (ARTS):** Evidence from one RCT shows that there was no significant difference between PCI and CABG for stroke [RR 1.01 (0.55 to 1.85)] for the entire group; and MI in a sub group of patients with diabetes [RR 1.57 (0.60 to 4.09)] [2 – 4 years follow-up]

**Booth 2008[132] (SOS); Kurbaan 2001[133] (CABRI); Legrand 2004[129] (ARTS):** Evidence from 3 RCTs shows that there was no significant difference in mortality [RR 1.87 (0.99 to 3.50)] in the PCI group compared to CABG in a sub group of patients with diabetes [2– 4 years follow-up].

**Booth 2008[132] (SoS); Legrand 2004[129] (ARTS):** Evidence from 2 RCTs shows that there was significantly higher repeat revascularisation in the PCI group compared to CABG [RR 4.84 (2.71 to 8.64)] in a sub group of patients with diabetes [2 – 4 years follow-up]

**Aoki 2004[134] (ARTS):** Evidence from one RCTs shows there was significantly higher repeat revascularisation in the PCI group compared to CABG [RR 4.63 (2.54 to 8.44)], there was no significant difference between PCI and CABG for death all causes [RR 1.03 (0.45 to 2.33)], stroke [RR 0.73 (0.24 to 2.28)], MI [RR

1.09 (0.56 to 2.11)], in a sub group of patients with involvement of the left Anterior descending coronary artery proximally. [3 years follow-up]

**Multi -vessel disease (Long term follow-up > 5 years)**

**Buszman 2009[135] (SOS); Henderson 1998[136] (RITA); Kaehler 2005[137] (GABI); Serruys 2005[138] (ARTS); Hueb 2010[77] (MASS-II):** Evidence from 5 RCTs shows that there was no significant difference between PCI and CABG for death (all causes) [RR 1.01 (0.83 to 1.23)] [5-13 years follow-up]

**Booth 2008[132] (SOS); Henderson 1998[136] (RITA); Kaehler 2005[137] (GABI):** Evidence from 3 RCTs shows that there was no significant difference between PCI and CABG for cardiac mortality [RR 1.24 (0.82 to 1.87)] [5-13 years follow-up]

**Serruys 2005[138] (ARTS); Hueb 2010[77] (MASS-II):** Evidence from 2 RCTs shows that there was no significant difference between PCI and CABG for stroke [RR 0.90 (0.57 to 1.41)] [5-10 years follow-up]

**Henderson 1998[136] (RITA); Serruys 2005[138] (ARTS); Hueb 2010[77] (MASS-II):** Evidence from 3 RCTs shows that there was no significant difference in non fatal MI in the PCI group compared to CABG [RR 1.28 (0.97 to 1.69)] [5-10 years follow-up]

**Buszman 2009[135] (SoS); Henderson 1998[136] (RITA); Kaehler 2005[137] (GABI); King 2000[139] (EAST); Serruys 2005[138] (ARTS); Hueb 2010[77] (MASS-II):** Evidence from 6 RCTs shows that there was significantly higher repeat revascularisation in the PCI group compared to CABG [RR 2.65 (2.35 to 2.98)] [5-13 years follow-up]

**Serruys 2005[138] (ARTS); Hueb 2010[77] (MASS-II):** Evidence from 2 RCTs shows that there was significantly more patients free of angina in CABG compared to PCI [RR 0.92 (0.87 to 0.97)] [5 yrs -10 yrs follow-up].

**Serruys 2005[138] (ARTS):** Evidence from one RCTs shows that there was no significant difference between PCI and CABG for, MI [RR 1.47 (0.6 to 3.58)], and stroke [RR 0.86 (0.31 to 2.36)] in a sub group of patients with diabetes. However there were significantly higher repeat revascularisations [RR 4.11 (2.2 to 7.68)] in the PCI group compared to CABG in a subgroup of patients with diabetes [5 years follow-up].

**Booth 2008[132] (SOS); Henderson 1998[136] (RITA); Serruys 2005[138] (ARTS):** Evidence from 3 RCTs shows that there was no significant difference between PCI and CABG for death [RR 1.43 (0.83 to 2.47)] in a subgroup of patients with diabetes [5- 10 years follow-up].

**Booth 2008[132] (SoS); Serruys 2005[138] (ARTS):** Evidence from 2 RCTs' shows that there was no significant difference between PCI and CABG for death [RR 1.12 (0.82 to 1.54)] in sub group of patients with no diabetes [5 years follow-up].

**Serruys 2005[138] (ARTS):** Evidence from one RCT shows that there was no significant difference between PCI and CABG for stroke [RR 1.19 (0.59 to 2.42)], MI [RR 1.28 (0.81 to 2.02)], and there were significantly more patients with repeat revascularisation [RR 3.25 (2.36 to 4.48)] in PCI compared to CABG in a sub group of patients with no diabetes. [5 years follow-up].

Sub group interaction: There was no significant difference between diabetes and no diabetes sub group of patients for death ( $p=0.45$ ), MI ( $p=0.79$ ) and repeat revascularisation ( $p=0.51$ ) at long term follow-up.

**Booth 2008[132] (SoS):** Evidence from one RCT shows that there was no significant difference between PCI and CABG for death in sub group 2 vessel disease [RR 1.68 (0.94 to 3.00)] and sub group 3 vessel disease [RR 1.58 (0.87 to 2.85)] [5 yrs follow-up].

Sub group interaction: There was no significant difference between single, 2 vessel and 3 vessel disease for death all causes ( $p=0.17$ ) at long term follow-up.

#### **Economic**

In people with multi vessel disease who are suitable for both CABG and PCI, PCI is more cost-effective. In people with single vessel disease PCI is likely to be even more cost-effective. This evidence

has minor limitations and direct applicability but there is some uncertainty around this conclusion.

1 **12.3 Single vessel disease**

2 **12.3.1 Clinical evidence**

3 The "Review Protocol" for this topic can be found in Appendix C, the "Search  
4 Strategies" in Appendix D, the "List of Included and Excluded Studies" in Appendix  
5 E1, the "Clinical Evidence Tables" in Appendix E2, and the "Forest Plots" in Appendix  
6 F.

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1 **Table 12.7: Single vessel disease - short term follow-up (1 year) for Stable angina**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Single vessel disease - short term follow-up (1 yr)	control	Relative (95% CI)	Absolute	
<b>Death (all causes) (follow-up 1 years)</b>											
Cisowski 2002[146]	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	serious (b)	none	1/50 (2%)	0/50 (0%)	RR 3 (0.13 to 71.92)	20 more per 1000 (from 30 fewer to 70 more)	⊕⊕⊕O MODERATE
<b>MI (follow-up 1 years)</b>											
Cisowski 2002[146]	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	0/50 (0%)	0/50 (0%)	not pooled	not pooled	⊕⊕⊕⊕ HIGH
<b>Free of angina (follow-up 1 years)</b>											
Cisowski 2002[146]	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	serious (b)	none	21/50 (42%)	24/50 (48%)	RR 0.88 (0.57 to 1.35)	58 fewer per 1000 (from 206 fewer to 168 more)	⊕⊕⊕O MODERATE

(a) Randomised, comparable at baseline, blind outcome assessment. Randomisation and allocation concealment methods not reported, high attrition: at 1 yr follow-up: 44% in PCI; 52% in E-ACAB)

(b) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.

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**Table 12.8: Single vessel disease- medium term follow-up (2 to 4 years) for Stable angina**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Single vessel disease- medium term follow-up (>1-4 yrs)	control	Relative (95% CI)	Absolute	
<b>Death (all causes) (follow-up 2. years)</b>											
Drenth 2004[147]; Goy 2000[148] (SIMA); Hueb 1995[83] (MASS-1)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	2/185 (1.1%)	6/180 (3.3%)	RR 0.37 (0.09 to 1.60)	21 fewer per 1000 (from 30 fewer to 20 more)	⊕⊕○○ LOW
<b>Cardiac death (follow-up 2-2.5 years)</b>											
Drenth 2004[147]; Goy 1994[149]; Goy 2000[148] (SIMA)	randomised trials	serious (e)	no serious inconsistency	no serious indirectness	serious (b)	none	1/181 (0.6%)	4/176 (2.3%)	RR 0.39 (0.08 to 2)	14 fewer per 1000 (from 21 fewer to 23 more)	⊕⊕○○ LOW
<b>MI (follow-up 2-2.5 years)</b>											
Drenth 2004[147]; Goy 1994[149]; Goy 2000[148] (SIMA); Hueb 1995[83] (MASS-1)	randomised trials	serious (c)	no serious inconsistency	no serious indirectness	serious (d)	none	18/253 (7.1%)	6/246 (2.4%)	RR 2.92 (1.18 to 7.21)	47 more per 1000 (from 4 more to 151 more)	⊕⊕○○ LOW
<b>Repeat revascularisation (follow-up 2-2.5 years)</b>											
Drenth 2004[147]; Goy 1994[149]; Goy 2000[148] (SIMA); Hueb 1995[83] (MASS-1)	randomised trials	serious (c)	no serious inconsistency	no serious indirectness	no serious imprecision	none	67/253 (26.5%)	4/246 (1.6%)	RR 13.27 (5.41 to 32.51)	200 more per 1000 (from 72 more to 512 more)	⊕⊕⊕○ MODERATE
<b>Free of angina</b>											
Drenth 2004[147]; Goy 1994[149]; Hueb 1995[83] (MASS-1)	randomised trials	serious (e)	no serious inconsistency	no serious indirectness	serious (d)	none	144/191 (75.4%)	168/184 (91.3%)	RR 0.83 (0.75 to 0.91)	155 fewer per 1000 (from 82 fewer to 228 fewer)	⊕⊕○○ LOW
<b>Stroke (follow-up 2 years)</b>											
Drenth 2004[147]; Goy 2000[148] (SIMA)	randomised trials	serious (f)	no serious inconsistency	no serious indirectness	serious (b)	none	2/113 (1.8%)	0/110 (0%)	RR 5 (0.25 to 101.63)	20 more per 1000 (from 20 fewer to 50 more)	⊕⊕○○ LOW

4 (a) Randomisation, ITT reported in both studies. Allocation concealment not reported in 2 out of 3 studies.  
 5 (b) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.  
 6 (c) Randomisation, ITT reported in all 4 studies. Allocation concealment not reported in all 4 studies.

- 1 (d) *95% CI around the pooled estimate of effect includes appreciable benefit or appreciable harm.*
- 2 (e) *Randomisation, ITT reported all 3 studies. Allocation concealment not reported in all 3 studies.*
- 3 (f) *Randomisation, ITT reported both studies. Allocation concealment not reported in all both studies.*
- 4
- 5

1 **Table 12.9: Single vessel disease- Long term follow-up (>5 years) for Stable angina**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Single vessel disease- Long term follow-up (>5 yrs)	control	Relative (95% CI)	Absolute	
<b>Death (all causes) (follow-up 5-10 years)</b>											
Goy 2008[150] (SIMA); Henderson 1998[136] (RITA); Hueb 1999[84] (MASS-I)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	28/367 (7.6%)	27/351 (7.7%)	RR 0.99 (0.60 to 1.65)	1 fewer per 1000 (from 31 fewer to 50 more)	⊕⊕○○ LOW
<b>Cardiac death (follow-up 10 years)</b>											
Goy 2008[150] (SIMA); Hueb 1999[84] (MASS-I)	randomised trials	serious (c)	no serious inconsistency	no serious indirectness	serious (b)	none	6/134 (4.5%)	3/129 (2.3%)	RR 1.93 (0.49 to 7.55)	22 more per 1000 (from 12 fewer to 152 more)	⊕⊕○○ LOW
<b>MI (follow-up 5-10 years)</b>											
Goy 2008[150] (SIMA); Henderson 1998[136] (RITA); Hueb 1999[84] (MASS-I)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	38/367 (10.4%)	23/351 (6.6%)	RR 1.58 (0.96 to 2.59)	38 more per 1000 (from 3 fewer to 104 more)	⊕⊕○○ LOW
<b>Repeat revascularisation (follow-up 5-10 years)</b>											
Goy 2008[150] (SIMA); Henderson 1998[136] (RITA); Hueb 1999[84] (MASS-I)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	156/367 (42.5%)	32/351 (9.1%)	RR 4.60 (3.25 to 6.50)	328 more per 1000 (from 205 more to 501 more)	⊕⊕⊕○ MODERATE
<b>Free of angina (follow-up 5 years)</b>											
Hueb 1999[84] (MASS-I);	randomised trials	serious (d)	no serious inconsistency	no serious indirectness	serious (b)	none	44/72 (61.1%)	48/70 (68.6%)	RR 0.89 (0.7 to 1.14)	75 fewer per 1000 (from 206 fewer to 96 more)	⊕⊕○○ LOW

- 2 (a) Allocation concealment, method of randomisation and blinding of outcome assessors reported in 1 out of 3 studies. IIT reported in all studies.
- 3 (b) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.
- 4 (c) Randomised. Intention to treat analysis reported in both, Allocation concealment and blinding of outcome assessors not reported in both the studies.
- 5 (d) Randomised. Allocation concealment not reported. Blinding of outcome assessors not reported. ITT reported.
- 6

1 **Additional data for Single vessel disease - short term follow-up (1 year)**

2 **Goy 2000[148] (SIMA) (Follow-up 1 year)**

3 No. of participants: N=123 (CABG (n=60); Stent (n=63))

4 At 1 year follow-up, 56 patients (95%) in the CABG group and 56 (91%) in the stent  
5 group were in CCS class 0 or 1 (p=0.90). Only 3 patients in the CABG group were in  
6 class III or IV compared with 6 patients in the stent group (p=0.08). The functional  
7 class showed no significant differences between the 2 groups

8

9 **Additional data – Hampton 1993[121] (RITA trial) – Medium term follow-up (2.5**  
10 **years)**

11 **Sub group interaction for single vessel and multi vessel disease:**

12 At 2.5 years the risk of death or infarction appeared unrelated to the number of  
13 treatment vessels at randomisation, there being 40 primary endpoints in the 456  
14 single vessel disease patients (16 CABG, 24 PTCA) and 53 primary endpoints in the  
15 555 multi vessel patients (27 CABG, 26 PTCA). The relative risk single: multi vessel is  
16 0.91 (95% CI 0.60-1.40, p=0.66). There is no evidence that any treatment difference  
17 depends on the number of disease vessels (interaction test p=0.35).

18

19 **Additional data for Single vessel disease - Long term follow-up - >5 years)**

20 **Goy 2008[150] (SIMA) (Follow-up 10 years)**

21 No. of participants: n=62 in PCI and n=59 CABG

22 At 10 years, most of the patients in both groups were asymptomatic (93%) or  
23 suffered mild angina. Angina functional class showed no significant differences  
24 between the PTCA and CABG. (No further details reported)

25

26 **Quality Of Life data for Single vessel disease:**

27

28 **Drenth 2004[147]:**

29 In this RCT assessments of Functional Health Status (FHS) were performed with SF-36  
30 questionnaire. SF-36 comprises 36 items covering the above 8 domains. These items  
31 were scored on a 0 to 100 range. Next, the items in the same domain were  
32 averaged together to create domain scores. For each domain, a high score indicates  
33 a more favourable health status (i.e., better physical functioning, less emotional  
34 problems, less pain and so forth).

1       **Results:** n=102 (n=51 in surgery and n=51 in PTCA). Both angioplasty and surgery  
2       resulted in good FHS in patients treated for an isolated high grade narrowing of the  
3       proximal LAD artery at 4 year follow-up. FHS did not differ between angioplasty  
4       and surgery in all domains. (Angioplasty vs. surgery: physical functioning: 77 vs. 81,  
5       p=0.48; Social functioning: 87 vs. 87, p=0.89; Role-physical: 76 vs. 78, p=0.81;  
6       Role-emotional: 87 vs. 85, p=0.98; Mental health: 82 vs. 81, p=0.86; Vitality: 70 vs.  
7       70, p=0.96; Bodily pain: 90 vs.88, p=0.97; General health perception: 69 vs. 70,  
8       p=0.78).

9

#### 10       **Goy 2000[148] (SIMA):**

11       In this RCT Quality of life was assessed with SF-36 and the Seattle questionnaire  
12       between 9-15 months.

13       **Results:** N=123 (CABG (n=60); PCI (n=63)). The quality of life questionnaires did not  
14       show significant differences between PCI and CABG. Only perception of the disease  
15       was more marked (but not significantly) after surgery.

16

#### 17       **12.3.2       Economic evidence**

18       No economic studies were identified specifically on this population. However the  
19       results of our economic model (see Appendix H and section 12.2.2) are likely to be  
20       applicable to people with single vessel disease.

21

#### 22       **12.3.3       Evidence statements**

##### **Clinical                Single vessel disease (Short term follow-up – 1 year)**

**Cisowski 2002[146]:** Evidence from one RCT shows that there was no significant difference between PCI and CABG for death (all causes) [RR 3 (0.13 to 71.92)], MI (not pooled- 0/50 in both groups) and free of angina [RR 0.88 (0.57 to 1.35)]. [1 year follow-up].

##### **Single vessel disease (Medium term follow-up - 2 to 4 years)**

**Drenth 2004[147]; Goy 2000[148] (SIMA); Hueb 1995[83] (MASS-I):** Evidence from 3 RCTS shows that there was no significant difference between PCI and CABG for death (all causes) [RR 0.37 (0.09 to 1.60) [2 -4 years follow-up].

**Drenth 2004[147]; Goy 1994[149]; Goy 2000[148] (SIMA):** Evidence from 3 RCTs shows that there was no significant difference between PCI and CABG for cardiac death [RR 0.39 (0.08 to 2)]. [2-4 years follow-up].

**Drenth 2004[147]; Goy 1994[149]; Goy 2000[148] (SIMA); Hueb 1995[83] (MASS-I):** Evidence from 4 RCTs shows that there were significantly more patients with MI [RR 2.92 (1.18 to 7.21)] in PCI compared to CABG, significantly higher repeat revascularisation [RR 13.27 (5.41 to 32.51)] in PCI compared to CABG [2-4 years follow-up].

**Drenth 2004[147]; Goy 1994[149]; Hueb 1995[83] (MASS-I):** Evidence from 3 RCTs shows that there was significantly more patients free of angina in the CABG group compared to PCI [RR 0.83 (0.75 to 0.91)] [2-4 years follow-up].

**Drenth 2004[147]; Goy 2000[148] (SIMA):** Evidence from 2 RCTs shows that there was no significant difference between PCI and CABG for stroke [RR 5.00 (0.25 to 101.63)] [2-4 years follow-up].

#### **Single vessel disease (Long term follow-up >5 years)**

**Goy 2008[148] (SIMA); Henderson 1998[136] (RITA); Hueb 1999[83] (MASS-I):** Evidence from 3 RCTs shows that there was significantly higher repeat revascularisation [RR 4.60 (3.25 to 6.50)] in the PCI group compared to CABG, and there was no significant difference between PCI and CABG for death (all causes) [RR 0.99 (0.60 to 1.65)] and MI [RR 1.58 (0.96 to 2.59)] [5- 10 years follow-up].

**Goy 2008[148] (SIMA); Hueb 1999[83] (MASS-I):** Evidence from 2 RCTs shows that there was no significant difference between PCI and CABG for cardiac death [RR 1.93 (0.49 to 7.55)] [10 years follow-up].

**Hueb 1999[83] (MASS-I):** Evidence from one RCT shows that there was no significant difference between PCI and CABG for free of angina [RR 0.89 (0.7 to 1.14)] [5 years follow-up].

**Economic** No economic studies were identified specifically on this population. The results of the economic model on people with multi-vessel disease are likely to be applicable to people with single vessel disease. Therefore PCI is more cost-effective than CABG in people eligible for both procedures. This evidence has minor limitations and direct applicability but there is some uncertainty around this conclusion.

1 **12.4 Left main coronary disease**

2 **12.4.1 Clinical evidence**

3 The “Review Protocol” for this topic can be found in Appendix C, the “Search  
4 Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
5 E1, the “Clinical Evidence Tables” in Appendix E2, and the “Forest Plots” in Appendix  
6 F.

7 .

1 **Table 12.10: Left main coronary disease - short term follow-up (1 year) for Stable angina**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Left main coronary disease - Short term follow-up (1 yr)	control	Relative (95% CI)	Absolute	
<b>Death (follow-up 1 years)</b>											
Buszman 2008[151] (LEMANS); Morice 2010[152] (SYNTAX)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	16/409 (3.9%)	19/401 (4.7%)	RR 0.83 (0.43 to 1.59)	8 fewer per 1000 (from 27 fewer to 28 more)	⊕⊕○○ LOW
<b>non fatal MI (follow-up 1 years)</b>											
Buszman 2008[151] (LEMANS); Morice 2010[152] (SYNTAX)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	16/409 (3.9%)	17/401 (4.2%)	RR 0.92 (0.47 to 1.8)	3 fewer per 1000 (from 22 fewer to 34 more)	⊕⊕○○ LOW
<b>Stroke (follow-up 1 years)</b>											
Buszman 2008[151] (LEMANS); Morice 2010[152] (SYNTAX)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	1/409 (0.2%)	11/401 (2.7%)	RR 0.13 (0.02 to 0.7)	24 fewer per 1000 (from 8 fewer to 27 fewer)	⊕⊕⊕○ MODERATE
<b>Repeat revascularisation (follow-up 1 years)</b>											
Buszman 2008[151] (LEMANS); Morice 2010[152] (SYNTAX)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	58/409 (14.2%)	28/401 (7%)	RR 2.04 (1.33 to 3.13)	73 more per 1000 (from 23 more to 149 more)	⊕⊕⊕○ MODERATE
<b>Cardiac death (follow-up 1 years)</b>											
Morice 2010[152] (SYNTAX)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	14/357 (3.9%)	8/348 (2.3%)	RR 1.71 (0.72 to 4.02)	16 more per 1000 (from 6 fewer to 69 more)	⊕⊕○○ LOW

- 2 (a) Buszman 2008[151] (LEMANS) Randomised, baseline comparisons made, blind outcome assessment for some outcomes (all clinical outcomes were analysed by the Clinical  
3 Event Committee. Echocardiographic and stress test recordings were read centrally by a group of independent investigators unaware of treatment assignment). Intention to  
4 treat analysis reported. Allocation concealment not reported, nos. lost to follow-up not reported, small sample size. \*This study reports 1 year follow-up results of the  
5 LEMANS (study of unprotected Left main stenting versus bypass surgery) study. Morice 2010[152] (SYNTAX) Strengths - Randomised, allocation concealment  
6 reported. n=12 withdrew consent in CABG group (N=336, 96.6% follow-up at 12 months) and n=1 lost to follow-up and n=1 discontinued treatment in PCI group  
7 (n=355, 99.4% follow-up at 12 months). Baseline comparisons made. ITT not reported. \*This study presents the outcomes in the pre-specified subgroup of patients  
8 (n=705) with LM disease in the SYNTAX trial.  
9 (b) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.

1        **Additional data (for Left main coronary disease - short term follow-up 1 year)**

2        **Buszman 2008[151] (LEMANS) (Follow-up 1 year)**

3        No. of participants: (n=52 in PCI and n=53 in CABG)

4        Patients after PCI had more angina after 6 months (p=0.01) but had similar angina  
5        status to CABG patients after 12 months (p=0.11).

6        **12.4.2        Economic evidence**

7        No economic studies were identified specifically on this population.

8

9        **12.4.3        Evidence statements**

**Clinical**

**Left main coronary artery stenosis (Short term follow – 1 year)**

**Buszman 2008[151] (LEMANS); Morice 2010[152] (SYNTAX):** Evidence from 2 RCTs shows that there was statistically significant higher stroke in the CABG group compared to PCI [RR 0.13 (0.02 to 0.7)]. There were statistically significant higher repeat revascularisations in the PCI group compared to CABG [RR 2.04 (1.33 to 3.13)]. There was no statistically significant difference between PCI and CABG for death [RR 0.83 (0.43 to 1.59)] and non fatal MI [RR 0.92 (0.47 to 1.8)]. [Follow-up 1 year]

**Morice 2010[152] (SYNTAX):** Evidence from 1 RCT shows that there was no statistically significant difference between PCI and CABG for cardiac death [RR 1.71 (0.72 to 4.02)] [Follow-up 1 year]

**Economic**

No economic studies were identified specifically on this population.

10

11        **12.5 Left main coronary artery or 3 vessel disease**

12        **12.5.1        Clinical evidence**

13        The “Review Protocol” for this topic can be found in Appendix C, the “Search  
14        Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
15        E1, the “Clinical Evidence Tables” in Appendix E2, and the “Forest Plots” in Appendix  
16        F.

17

1 **Table 12.11: Left main coronary artery or 3 vessel disease short term follow-up (1 year) for Stable angina**

Quality assessment							Summary of findings				
							No of patients		Effect		Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Left main coronary artery or 3 vessel disease short term follow-up (1yr)	control	Relative (95% CI)	Absolute	
<b>Death (all causes) (follow-up 1 years)</b>											
Serruys 2009[153] (SYNTAX) (d)	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	serious (b)	none	39/891 (4.4%)	30/849 (3.5%)	RR 1.24 (0.78 to 1.98)	8 more per 1000 (from 8 fewer to 35 more)	⊕⊕⊕○ MODERATE
<b>cardiac mortality (follow-up 1 years)</b>											
Serruys 2009[153] (SYNTAX)	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	serious (b)	none	33/891 (3.7%)	18/849 (2.1%)	RR 1.75 (0.99 to 3.08)	16 more per 1000 (from 0 fewer to 44 more)	⊕⊕⊕○ MODERATE
<b>Stroke (follow-up 1 years)</b>											
Serruys 2009[153] (SYNTAX)	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	5/891 (0.6%)	19/849 (2.2%)	RR 0.25 (0.09 to 0.67)	17 fewer per 1000 (from 7 fewer to 20 fewer)	⊕⊕⊕⊕ HIGH
<b>MI (follow-up 1 years)</b>											
Serruys 2009[153] (SYNTAX)	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	serious (b)	none	43/891 (4.8%)	28/849 (3.3%)	RR 1.46 (0.92 to 2.33)	15 more per 1000 (from 3 fewer to 44 more)	⊕⊕⊕○ MODERATE
<b>Repeat revascularisation (follow-up 1 years)</b>											
Serruys 2009[153] (SYNTAX)	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	120/891 (13.5%)	50/849 (5.9%)	RR 2.29 (1.67 to 3.14)	76 more per 1000 (from 39 more to 126 more)	⊕⊕⊕⊕ HIGH
<b>Sub group diabetes (Death) (follow-up 1 years)</b>											
Banning 2010[154] (SYNTAX)	randomised trials	no serious limitations (c)	no serious inconsistency	no serious indirectness	serious (b)	none	19/227 (8.4%)	13/204 (6.4%)	RR 1.31 (0.67 to 2.59)	20 more per 1000 (from 21 fewer to 101 more)	⊕⊕⊕○ MODERATE
<b>Sub group diabetes (cardiac death) (follow-up 1 years)</b>											
Banning 2010[154] (SYNTAX)	randomised trials	no serious limitations (c)	no serious inconsistency	no serious indirectness	serious (b)	none	16/227 (7%)	8/204 (3.9%)	RR 1.8 (0.79 to 4.11)	31 more per 1000 (from 8 fewer to 122 more)	⊕⊕⊕○ MODERATE
<b>Sub group diabetes (stroke) (follow-up 1 years)</b>											
Banning 2010[154] (SYNTAX)	randomised trials	no serious limitations (c)	no serious inconsistency	no serious indirectness	serious (b)	none	2/227 (0.9%)	5/204 (2.5%)	RR 0.36 (0.07 to 1.83)	16 fewer per 1000 (from 23 fewer to 20 more)	⊕⊕⊕○ MODERATE

Sub group diabetes (MI) (follow-up 1 years)											
Banning 2010[154] (SYNTAX)	randomised trials	no serious limitations (c)	no serious inconsistency	no serious indirectness	serious (b)	none	11/227 (4.8%)	9/204 (4.4%)	RR 1.1 (0.46 to 2.6)	4 more per 1000 (from 24 fewer to 71 more)	⊕⊕⊕○ MODERATE
Sub group diabetes (Repeat revascularisation) (follow-up 1 years)											
Banning 2010[154] (SYNTAX)	randomised trials	no serious limitations (c)	no serious inconsistency	no serious indirectness	no serious imprecision	none	46/227 (20.3%)	13/204 (6.4%)	RR 3.18 (1.77 to 5.71)	139 more per 1000 (from 49 more to 300 more)	⊕⊕⊕⊕ HIGH
Sub group no diabetes (Death) (follow-up 1 years)											
Banning 2010[154] (SYNTAX)	randomised trials	no serious limitations (c)	no serious inconsistency	no serious indirectness	serious (b)	none	20/664 (3%)	17/645 (2.6%)	RR 1.14 (0.6 to 2.16)	4 more per 1000 (from 11 fewer to 31 more)	⊕⊕⊕○ MODERATE
Sub group no diabetes (no cardiac death) (follow-up 1 years)											
Banning 2010[154] (SYNTAX)	randomised trials	no serious limitations (c)	no serious inconsistency	no serious indirectness	serious (b)	none	17/664 (2.6%)	10/645 (1.6%)	RR 1.65 (0.76 to 3.58)	10 more per 1000 (from 4 fewer to 40 more)	⊕⊕⊕○ MODERATE
Sub group no diabetes (stroke) (follow-up 1 years)											
Banning 2010[154] (SYNTAX)	randomised trials	no serious limitations (c)	no serious inconsistency	no serious indirectness	no serious imprecision	none	3/664 (0.5%)	14/645 (2.2%)	RR 0.21 (0.06 to 0.72)	17 fewer per 1000 (from 6 fewer to 20 fewer)	⊕⊕⊕⊕ HIGH
Sub group no diabetes (MI) (follow-up 1 years)											
Banning 2010[154] (SYNTAX)	randomised trials	no serious limitations (c)	no serious inconsistency	no serious indirectness	serious (b)	none	32/664 (4.8%)	19/645 (2.9%)	RR 1.64 (0.94 to 2.86)	19 more per 1000 (from 2 fewer to 55 more)	⊕⊕⊕○ MODERATE
Sub group no diabetes (Repeat revasc) (follow-up 1 years)											
Banning 2010[154] (SYNTAX)	randomised trials	no serious limitations (c)	no serious inconsistency	no serious indirectness	no serious imprecision	none	74/664 (11.1%)	37/645 (5.7%)	RR 1.94 (1.33 to 2.84)	54 more per 1000 (from 19 more to 106 more)	⊕⊕⊕⊕ HIGH

- 1 a) Randomised, allocation concealment reported, baseline comparisons made, nos. lost to follow-up reported ((5.4% in CABG and 1.3% in PCI group), Intention to treat
- 2 analysis reported. Blind outcome assessment (adjudicated by an independent Clinical Events Committee).Patients aware of the intervention allocated.
- 3 b) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.
- 4 c) Strengths- Randomised, ITT used, one year MACCE was evaluated in 849 (94.6%) CABG patients (645 non diabetic and 204 medically treated diabetes) and 891
- 5 (98.7%) PES patients (664 non diabetic and 227 with medically treated diabetes). Allocation concealment reported. Baseline comparisons made. This is a sub group
- 6 analysis of the SYNTAX trial.
- 7 d) Authors note:
- 8 ○ Most cases of stent thrombosis occurred within 30 days after the procedure, and the 12 month rate of stent thrombosis in the PCI group was similar to the rate of
- 9 symptomatic graft occlusion in the CABG group. Stent thrombosis often has more serious consequences for patients (rate of death, approximately 30%, rate of
- 10 MI approximately 60%) than does graft occlusion, which often results only in angina leading to revascularisation.

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- *The use of antiplatelet medication was high among patients in the PCI group (with 71.1% receiving a thienopyridine at 12 months). The authors report that the low rate of stroke among patients with PCI may have resulted from the use of highly effective dual antiplatelet therapy which prevents thrombo embolic events.*
- *More patients in the CABG group than in PCI declined to participate after proving consent; this imbalance was due to the greater invasiveness of CABG.*



the CABG group compared to PCI. There was no significant difference between PCI and CABG for death (all causes) [RR 1.24 (0.78 to 1.98)], cardiac mortality [RR 1.75 (0.99 to 3.08)], and MI [RR 1.46 (0.92 to 2.33)]. [1 year follow-up]

**Banning 2010[154] (SYNTAX):** Evidence from one RCT shows that there was no significant difference between PCI with PES and CABG for death [RR 1.31 (0.67 to 2.59)], cardiac death [RR 1.8 (0.79 to 4.11)], stroke [RR 0.36 (0.07 to 1.83)], MI [RR 1.1 (0.46 to 2.6)]. There were significantly higher repeat revascularisations [RR 3.18 (1.77 to 5.71)] in the PCI group compared to CABG in a sub group of patients with diabetes. [1 year follow-up].

**Banning 2010[154] (SYNTAX):** Evidence from one RCT shows that there was no significant difference between PCI with PES and CABG for death [RR 1.14 (0.6 to 2.16)], cardiac death [RR 1.65 (0.76 to 3.58)], MI [RR 1.64 (0.94 to 2.86)]. There was significantly higher stroke\* [RR 0.21 (0.06 to 0.72) ] in the CABG group and higher repeat revascularisation [RR 1.94 (1.33 to 2.84) ] in the PCI group compared to CABG in the sub group of patients with no diabetes. [1 year follow-up]. \*Authors report that this value did not reach statistical significance in diabetes patients, possibly because of the small size in the diabetic group.

**Economic** No economic studies were identified specifically on this population.

1 **12.6 IPD Meta-analyses (Multi vessel disease- immediate, short and long**  
2 **term follow-up)**

3 **12.6.1 Clinical evidence**

4 The “Review Protocol” for this topic can be found in Appendix C, the “Search  
5 Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
6 E1, the “Clinical Evidence Tables” in Appendix E2, and the “Forest Plots” in Appendix  
7 F.

1 **Table 12.12: IPD meta-analyses (Multi -vessel disease- Immediate, short and Long term follow-up)**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							IPD meta analyses (PCI)	CABG	Relative (95% CI)	Absolute	
<b>Death (follow-up median 5.9 years)</b>											
Hlatky 2009[114]	randomised trial	no serious limitations (a)	no serious inconsistency	serious (b)	no serious imprecision	none	628/3923 (16%)	575/3889 (14.8%)	HR 0.91 (0.82 to 1.02)	12 fewer per 1000 (from 25 fewer to 3 more)	⊕⊕⊕○ MODERATE
<b>Death - Age &lt;55 years (follow-up 5.9 years)</b>											
Hlatky 2009[114]	randomised trial	no serious limitations (a)	no serious inconsistency	serious (b)	no serious imprecision	none	88/1122 (7.8%)	107/1063 (10.1%)	HR 1.25 (0.94 to 1.66)	24 more per 1000 (from 6 fewer to 61 more)	⊕⊕⊕○ MODERATE
<b>Death- age 55-64 years (follow-up median 5.9 years)</b>											
Hlatky 2009[114]	randomised trial	no serious limitations (a)	no serious inconsistency	serious (b)	no serious imprecision	none	220/1456 (15.1%)	201/1477 (13.6%)	HR 0.90 (0.75 to 1.09)	13 fewer per 1000 (from 32 fewer to 11 more)	⊕⊕⊕○ MODERATE
<b>Death-&gt;65 years (follow-up median 5.9 years)</b>											
Hlatky 2009[114]	randomised trial	no serious limitations (a)	no serious inconsistency	serious (b)	no serious imprecision	none	319/1341 (23.8%)	267/1347 (19.8%)	HR 0.82 (0.7 to 0.97)	32 fewer per 1000 (from 5 fewer to 55 fewer)	⊕⊕⊕○ MODERATE
<b>Death- women (follow-up median 5.9 years)</b>											
Hlatky 2009[114]	randomised trial	no serious limitations (a)	no serious inconsistency	serious (b)	no serious imprecision	none	164/992 (16.5%)	162/909 (17.8%)	HR 1.02 (0.82 to 1.27)	3 more per 1000 (from - 30 fewer to 42 more)	⊕⊕⊕○ MODERATE
<b>Death- men (follow-up median 5.9 years)</b>											
Hlatky 2009[114]	randomised trial	no serious limitations (a)	no serious inconsistency	serious (b)	no serious imprecision	none	464/3001 (15.5%)	413/2980 (13.9%)	HR 0.88 (0.77 to 1)	16 fewer per 1000 (from 30 fewer to 0 more)	⊕⊕⊕○ MODERATE
<b>Death- No diabetes (follow-up median 5.9 years)</b>											
Hlatky 2009[114]	randomised trial	no serious limitations (a)	no serious inconsistency	serious (b)	no serious imprecision	none	448/3298 (13.6%)	432/3263 (13.2%)	HR 0.98 (0.86 to 1.12)	2 fewer per 1000 (from 17 fewer to 15 more)	⊕⊕⊕○ MODERATE
<b>Death- Diabetes (follow-up median 5.9 years)</b>											
Hlatky 2009[114]	randomised trial	no serious limitations (a)	no serious inconsistency	serious (b)	no serious imprecision	none	179/618 (29%)	143/615 (23.3%)	HR 0.70 (0.56 to 0.87)	64 fewer per 1000 (from 27 fewer to 95 fewer)	⊕⊕⊕○ MODERATE
<b>Death- stable symptoms (follow-up median 5.9 years)</b>											
Hlatky 2009[114]	randomised trial	no serious limitations (a)	no serious inconsistency	serious (b)	no serious imprecision	none	256/1900 (13.5%)	205/1840 (11.1%)	HR 0.83 (0.69 to 0.99)	18 fewer per 1000 (from 1 fewer to 33 fewer)	⊕⊕⊕○ MODERATE
<b>Death- unstable symptoms (follow-up median 5.9 years)</b>											
Hlatky 2009[114]	randomised trial	no serious limitations (a)	no serious inconsistency	serious (b)	no serious imprecision	none	266/1306 (20.4%)	262/1347 (19.5%)	HR 0.95 (0.8 to 1.12)	9 fewer per 1000 (from 36 fewer to 21 more)	⊕⊕⊕○ MODERATE
<b>Death- Normal LV function (follow-up median 5.9 years)</b>											
Hlatky	randomised	no serious	no serious	serious (b)	no serious	none	398/2791	375/2789	HR 0.92 (0.8	10 fewer per 1000 (from	⊕⊕⊕○

2009[114]	trial	limitations (a)	inconsistency		imprecision		(14.3%)	(13.4%)	to 1.06)	25 fewer to 7 more)	MODERATE
<b>Death- abnormal LV function (follow-up median 5.9 years)</b>											
Hlatky 2009[114]	randomised trial	no serious limitations (a)	no serious inconsistency	serious (b)	no serious imprecision	none	151/615 (24.6%)	126/551 (22.9%)	HR 0.93 (0.73 to 1.18)	14 fewer per 1000 (from 56 fewer to 35 more)	⊕⊕⊕O MODERATE
<b>Death- less than 3 diseased vessels (follow-up median 5.9 years)</b>											
Hlatky 2009[114]	randomised trial	no serious limitations (a)	no serious inconsistency	serious (b)	no serious imprecision	none	371/2523 (14.7%)	325/2386 (13.6%)	HR 0.91 (0.78 to 1.06)	11 fewer per 1000 (from 28 fewer to 8 more)	⊕⊕⊕O MODERATE
<b>Death- 3 vessel disease (follow-up median 5.9 years)</b>											
Hlatky 2009[114]	randomised trial	no serious limitations (a)	no serious inconsistency	serious (b)	no serious imprecision	none	253/1376 (18.4%)	248/1477 (16.8%)	HR 0.91 (0.77 to 1.09)	14 fewer per 1000 (from 36 fewer to 14 more)	⊕⊕⊕O MODERATE
<b>Death- No proximal LAD (follow-up median 5.9 years)</b>											
Hlatky 2009[114]	randomised trial	no serious limitations (a)	no serious inconsistency	serious (b)	no serious imprecision	none	310/1636 (18.9%)	278/1567 (17.7%)	HR 0.92 (0.79 to 1.09)	13 fewer per 1000 (from 34 fewer to 14 more)	⊕⊕⊕O MODERATE
<b>Death- Proximal LAD (follow-up median 5.9 years)</b>											
Hlatky 2009[114]	randomised trial	no serious limitations (a)	no serious inconsistency	serious (b)	no serious imprecision	none	268/1684 (15.9%)	249/1707 (14.6%)	HR 0.90 (0.75 to 1.07)	14 fewer per 1000 (from 34 fewer to 9 more)	⊕⊕⊕O MODERATE
<b>Death- balloon angioplasty trials (follow-up median 5.9 years)</b>											
Hlatky 2009[114]	randomised trial	no serious limitations (a)	no serious inconsistency	serious (b)	no serious imprecision	none	481/2405 (20%)	436/2356 (18.5%)	HR 0.91 (0.8 to 1.03)	15 fewer per 1000 (from 34 fewer to 5 more)	⊕⊕⊕O MODERATE
<b>Death- BMS trials (follow-up median 5.9 years)</b>											
Hlatky 2009[114]	randomised trial	no serious limitations (a)	no serious inconsistency	serious (b)	no serious imprecision	none	147/1518 (9.7%)	139/1533 (9.1%)	HR 0.94 (0.74 to 1.18)	5 fewer per 1000 (from 23 fewer to 15 more)	⊕⊕⊕O MODERATE
<b>Frequency of angina (Follow-up 1 year)</b>											
Hlatky 2009[114]	randomised trial	no serious limitations (a)	no serious inconsistency	serious (b)	no serious imprecision	none	856/3240 (26.4%)	439/3228 (13.6%)	RR 1.94 (1.75 to 2.16)	128 more per 1000 (from 102 more to 158 more)	⊕⊕⊕O MODERATE
<b>Stroke (Follow-up 90 days)</b>											
Hlatky 2009[114]	randomised trial	no serious limitations (a)	no serious inconsistency	serious (c)	no serious imprecision	none	12/2269 (0.5%)	26/2268 (1.1%)	RR 0.46 (0.23 to 0.91)	6 fewer per 1000 (from 1 fewer to 8 fewer)	⊕⊕⊕O MODERATE

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- (a) This is an IPD (Individual patient data) meta-analyses. Review addresses an appropriate and clearly focused question. The review included only RCTs which was relevant to the review question. There was adequate description of the methodology used in the meta-analysis. The papers report the search strategy used in detail. The authors report that all the included trials were reviewed and approved by ethics committees. All analyses followed the Intention to treat principle. This IPD meta analyses included 10 trials. Note: The IPD included 3 trials which were not included in the study level meta-analyses 1) BARI[115] -<30% with stable angina, 2) ERACI-II[116,117] - 92% unstable angina and 3) Toulouse[118]) - Study reports- Few patients presented with stable angina, whereas the majority complained of unstable angina or recent MI
- (b) 4 studies from the IPD meta-analyses did not have sufficient stable angina population (BARI[115], ERACI-II[116,117], Toulouse[118]).
- (c) Stroke data available from 7 trials

**Sub group interaction:**

## DRAFT FOR CONSULTATION

1 *In patients with diabetes (CABG, n=615; PCI, n=618), mortality was substantially lower in the CABG group than in the PCI group (HR 0.70, 0.56-0.87); however, mortality was*  
2 *similar between groups in patients without diabetes (HR 0.98, 0.86-1.12; p=0.014 for interaction). Patient age modified the effect of treatment on mortality, with hazard ratios of*  
3 *1.25 (0.94-1.66) in patients younger than 55 years, 0.90 (0.75-1.09) in patients aged 55-64 years, and 0.82 (0.70-0.97) in patients 65 years and older (p=0.002 for*  
4 *interaction). Treatment effect was not modified by the number of diseased vessels (p=0.98 for interaction), gender (p=0.25 for interaction), stable/unstable symptoms (p=0.30 for*  
5 *interaction), LV function (p=0.87 for interaction), involvement of proximal LAD (p=0.77 for interaction), and angioplasty/bare metal stents (p=0.19 for interaction).*

1 **12.6.2 Economic evidence**

2 See 12.2.212.2.2.

3 **12.6.3 Evidence statements**

<b>Clinical</b>	<p><b><u>IPD meta analyses [ Multi vessel disease -Immediate, short and Long term follow-up]</u></b></p> <p><b>Hlatky 2009[114]:</b> Evidence from one IPD meta-analyses shows that at 90 days stroke was significantly higher in the CABG group compared to PCI [RR 0.46 (0.23 to 0.91)]. [90 days follow-up].</p> <p><b>Hlatky 2009[114]:</b> Evidence from one IPD meta-analyses shows that at 1 year angina was significantly less frequent in the CABG group compared to PCI [RR 1.94 (1.75 to 2.16)] [1 year] follow-up.</p> <p><b>Hlatky 2009[114]:</b> Evidence from one IPD meta analyses shows that there was no significant difference between PCI and CABG for death [HR 0.91, 95% CI 0.82 to 1.02]]. There was significantly higher mortality in PCI compared to CABG in patients with diabetes [HR 0.70, 0.56 to 0.87]], however mortality was similar between PCI and CABG groups for patients with no diabetes [HR 0.98, 0.86 to 1.12; p=0.014 for interaction]]. There was no significant difference in mortality between PCI and CABG in patients younger than 55 years [HR 1.25, 0.94 to 1.66]] and in patients aged 55-64 years [HR 0.90 (0.75 to 1.09)], however mortality was significantly lower in CABG compared to PCI in patients 65 years and older [HR 0.82 (0.70 to 0.97) p=0.002 for interaction]]. There was no significant difference in mortality between PCI and CABG groups when assessed by bare metal stents [HR 0.94 (0.74 to 1.18)] or balloon angioplasty [HR 0.91 (0.80 to 1.03)] (p=0.19 for interaction). There was no significant difference in mortality between PCI and CABG in patients less than 3 diseased vessels [HR 0.91 (0.78 to 1.06)] or 3 vessel disease [HR 0.91 (0.77 to 1.09)] (p=0.98 for interaction). There was no significant difference in mortality between PCI and CABG in patients with no proximal LAD [HR 0.92 (0.79 to 1.09)] or with proximal LAD [HR 0.90 (0.75 to 1.07)] (p=0.77 for interaction) [median 5.9 years follow-up].</p>
<b>Economic</b>	<p>In people with multi vessel disease who are suitable for both CABG and PCI, PCI is more cost-effective. This evidence has minor limitations and direct applicability but there is some uncertainty around this conclusion.</p>

1 **12.7 Recommendations and link to evidence**

<b>Recommendation</b>	<p><b>Consider the relative risks and benefits of PCI and CABG using a systematic approach to assess the severity and complexity of the person’s coronary disease, in addition to other relevant clinical factors and comorbidities.</b></p> <p><b>Consider PCI in preference to CABG for people who have single-vessel disease or multi-vessel disease, including left main stem disease, and who have continuing symptoms despite optimal medical treatment and the anatomy is suitable for PCI.</b></p> <p><b>Consider CABG for people with single-vessel disease or multi-vessel disease, including left main stem disease, and continuing symptoms despite optimal medical treatment if the anatomy is unsuitable for PCI.</b></p> <p><b>Consider CABG in preference to PCI for people with multi-vessel disease who have continuing symptoms despite optimal medical treatment and who:</b></p> <ul style="list-style-type: none"> <li>• <b>are over 65 years and/or</b></li> <li>• <b>have diabetes.</b></li> </ul>
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**Relative values of different outcomes**

Outcomes of interest included long-term mortality (total and cardiovascular), rates of major adverse cardiovascular events (myocardial infarction, stroke, myocardial revascularisation), measures of symptom severity (frequency of angina, exercise test outcomes), and quality of life.

**Trade off between clinical benefits and harms**

The trials of myocardial revascularisation in this review compared an initial treatment strategy of coronary artery bypass surgery with initial strategies of coronary balloon angioplasty, or percutaneous coronary intervention using either bare-metal or drug-eluting coronary stents. The trials recruited highly selected patients who were considered suitable for either revascularisation strategies and the trial results do not apply to all patients being considered for myocardial revascularisation procedures in contemporary practice.

**Mortality**

None of the individual trials of coronary artery bypass surgery versus percutaneous coronary intervention has sufficient statistical power to reliably detect potentially important differences in long-term mortality between the two treatment strategies. Our analysis of pooled data from the trials provides evidence that mortality in the medium to long term is comparable between the two treatment groups.

The individual patient data meta-analysis combines data from all larger trials of bypass surgery versus percutaneous coronary intervention. Overall the individual patient data meta-analysis reported no significant difference in mortality between the two treatment strategies. Subgroup analyses demonstrated a significant interaction between age and treatment effect, suggesting that CABG may confer a prognostic advantage in older patients (aged over 65 years). In addition there was a significant interaction between diabetes and treatment effect suggesting that coronary bypass surgery may additionally confer prognostic advantage in people with diabetes.

### **Stroke**

Several trials reported on short term risk of stroke. In our analysis there was an excess risk of stroke in the coronary bypass surgery group (1.0% versus 0.3%) and this was confirmed in the individual patient data meta-analysis (1.1% versus 0.5% at 90 days). The GDG were concerned that the clinical significance of stroke (disabling versus non-disabling) is not reported consistently in the trials, and the difference in stroke risk may be partly due to bias resulting from different protocols for detection and diagnosis of stroke in the two treatment groups. There is no evidence of a difference in stroke risk between the treatment groups beyond the early follow-up phase.

### **Repeat revascularisation**

The trials consistently reported higher rates of repeat (non-protocol) revascularisation in the percutaneous coronary intervention group than in the surgery group. Revascularisation rates among patients assigned to percutaneous coronary intervention were higher in the early balloon angioplasty trials than in the later bare metal or drug-eluting stent trials (Figure 12.1- figure prepared for GDG)).

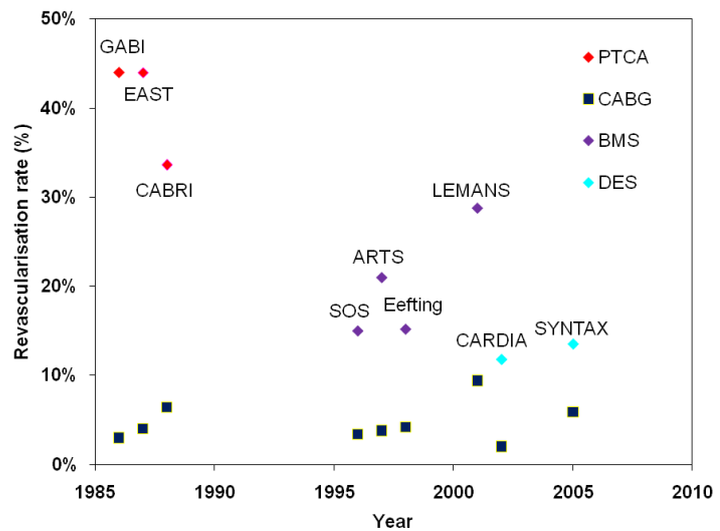


Figure 12.1: revascularisation rates at one year follow-up in trials of percutaneous coronary intervention versus coronary artery bypass surgery. The trials are plotted against the year in which trial recruitment started. For each trial the squares show the revascularisation rate for patients in the surgical group and the diamonds show the rate for patients in the percutaneous coronary intervention group.

### Angina

The trials and the individual patient data meta-analysis provide evidence that initial strategies of coronary artery bypass surgery and percutaneous coronary intervention are effective treatments for angina in the medium and long-term. Nevertheless, freedom from angina was consistently higher among patients treated by coronary artery surgery than by percutaneous coronary intervention, both in trials of balloon angioplasty and in trials that used bare metal stents. The magnitude of the difference in angina prevalence between the two treatment strategies is small but was still evident in the ARTS trial after five years. We found no information from randomised trials about the effect of percutaneous coronary intervention with drug-eluting stents on the prevalence of angina.

The results of the trials of PCI versus CABG are consistent across subgroups with single and multi-vessel disease.

### Limitations

The patients in the trials of percutaneous coronary intervention versus coronary artery bypass surgery were highly selected and considered angiographically suitable for either revascularisation procedure. For example in RITA-1 22800 patients with a clinical indication for myocardial revascularisation were considered for the trial, 4800 were eligible for the trial, but only 1011 were randomised. Also, eligibility for percutaneous treatment of coronary artery

disease has evolved over time as improvement in technique and equipment have allowed treatment of more complex patterns of disease. The trial results may therefore not be generalisable to the wider population of people with stable angina and require cautious interpretation.

The IPD analysis included the BARI and ERACI trials, but these trials were excluded from our analysis because they enrolled a high proportion of patients with acute coronary syndrome. People with acute coronary syndrome are at higher risk of adverse cardiovascular events than people with stable angina and this may influence the relative effects of CABG and PCI on outcome.

Interpretation of the trials of CABG versus PCI is confounded by changes in surgical and interventional technique over time. In particular the introduction of bare-metal and drug-eluting stents has improved the acute results of PCI and reduces the subsequent risk of restenosis and repeat revascularisation procedures[62,64,155]. The IPD meta-analysis[114]) included patients from the balloon angioplasty era and is therefore only partially applicable to current practice. On the other hand, inclusion of trials of balloon angioplasty allows analysis of longer term follow-up data, which is not currently available for trials of bare metal or drug-eluting stents.

The IPD meta-analysis reported an interaction between treatment effect and diabetes, with a survival advantage from CABG in people with diabetes. However, recent trials that used bare metal or drug-eluting stents have not demonstrated a survival advantage of surgical revascularisation over a PCI-based strategy (SYNTAX, CARDIA, LEMANS, ARTS, SOS), either in the entire trial populations or in the diabetic subgroups. SYNTAX and CARDIA used first-generation drug-eluting stents, but recent trials have shown that second generation drug-eluting stents are associated with superior clinical outcomes including reduced risks of stent thrombosis and requirement for repeat revascularisation[156,157]. The GDG concluded that the relative effects of PCI with drug-eluting stents and coronary artery bypass surgery on mortality in people with diabetes is uncertain and requires further investigation.

## **Conclusions**

The GDG concluded that there is no definitive evidence that one revascularisation strategy confers a prognostic advantage over the other strategy in contemporary clinical practice.

The trials provide evidence that both revascularisation strategies relieve angina but coronary artery bypass surgery

provides superior relief of angina in the medium term when compared with balloon angioplasty or percutaneous coronary intervention with bare metal stents.

The choice of revascularisation strategy will depend on many factors including angiographic suitability, patient choice, age, and the presence of diabetes and other comorbidities.

<b>Economic considerations</b>	PCI is more cost-effective than CABG in people with multi vessel disease eligible for both procedures. There is however some uncertainty around this conclusion. CABG could still be more cost-effective in high risk patients.
<b>Quality of evidence</b>	<p>We found significant heterogeneity between the trials included in this review, probably partly related to differences in inclusion criteria and to different revascularisation techniques.</p> <p>The economic evidence has minor limitations and direct applicability.</p>
<b>Other considerations</b>	The economic evidence is based on an analysis of data from trials that recruited patients with multivessel disease, as we found only limited data from trials of PCI versus CABG in patients with single vessel disease. Patients undergoing PCI for single vessel disease generally require fewer stents than patients with multivessel disease and are therefore likely to incur lower costs. Patients undergoing CABG for single vessel disease may also incur lower costs than patients with multivessel disease, but there is no consistent evidence that the clinical results of the two revascularisation strategies differ between subgroups with single and multivessel disease. The GDG therefore considered that PCI is likely to be a cost-effective strategy in patients with single vessel disease.

<b>Recommendation</b>	<p><b>Consider the risks and benefits of continuing drug treatment or performing revascularisation (PCI or CABG) after coronary angiography.</b></p> <p><b>Consider whether the decision to continue drug treatment or perform revascularisation (PCI or CABG) needs to be discussed by a multidisciplinary team. The team should include an interventional cardiologist and a cardiac surgeon.</b></p> <p><b>Consider the relative risks and benefits of PCI and CABG using a systematic approach to assess the severity and complexity of the person's coronary disease, in addition to other relevant clinical factors and comorbidities.</b></p>
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Quality of evidence	No evidence was reviewed for this recommendation.
Other considerations	The GDG considered that review of treatment options for people with stable angina within a multidisciplinary team meeting that includes a cardiac surgeon and an interventional cardiologist can be helpful. They did not think this is required for all patients but that the balance of risks and benefits in individual patients can be finely balanced and may best be made by review and discussion with professionals from different disciplines.

The GDG considered that the review of treatment options should be approached systematically, taking account of the severity and complexity of the patient's coronary artery disease and any other relevant clinical factors and comorbidities. The GDG were aware that tools have been developed to support this process and scores that predict risk of revascularisation procedures are in clinical use (e.g. EUROSCORE [[www.euroscore.org](http://www.euroscore.org)]). The SYNTAX score was developed to risk stratify participants in the SYNTAX clinical trial and in subgroup analyses high SYNTAX scores were associated with better one year outcome among patients assigned to CABG than among patients assigned to PCI. [153] Longer term follow-up data from the SYNTAX trial, and validation of the SYNTAX score in larger patient populations are not available. In the interim, the GDG considered that there is insufficient evidence to recommend the routine use of any particular score or method to decide on appropriate intervention.

<b>Recommendation</b>	<p><b>Ensure people with stable angina receive balanced information and have the opportunity to discuss the benefits, limitations and risks of continuing drug treatment, PCI and CABG to help them make an informed decision about their treatment.</b></p> <p><b>Explain to the person that:</b></p> <ul style="list-style-type: none"> <li>• <b>The purpose of revascularisation is to improve the symptoms of stable angina.</b></li> <li>• <b>PCI and CABG are effective in relieving symptoms.</b></li> <li>• <b>CABG is slightly more effective than PCI in relieving symptoms of stable angina in the longer term.</b></li> <li>• <b>Repeat revascularisation may be necessary after either PCI or CABG and the rate is higher after PCI than CABG.</b></li> </ul>
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	<ul style="list-style-type: none"> <li>● <b>Stroke is uncommon after either PCI or CABG, and the incidence is similar between the two procedures.</b></li> </ul> <p><b>Inform the person about the practical aspects of PCI and CABG. Include information about:</b></p> <ul style="list-style-type: none"> <li>● <b>vein and/or artery harvesting</b></li> <li>● <b>likely length of hospital stay</b></li> <li>● <b>recovery time</b></li> <li>● <b>drug treatment after the procedure.</b></li> </ul>
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Quality of evidence	No evidence was specifically reviewed for these recommendations.
Other considerations	<p>These recommendations were informed by the evidence from the reviews on medical versus revascularization treatment and PCI versus CABG and by the professional opinion and views of the GDG</p> <p>The GDG considered it important that patients are given full information about the relative benefits and risks of continuing medical therapy or undergoing revascularisation. The areas of information listed by the GDG is not exhaustive but included the areas they considered should be included in informing patients.</p>

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2 **12.8 Research recommendation A**

3 The GDG recommended the following research question:

4 ➤ **Research question:** Do people with stable angina and evidence of reversible  
 5 ischaemia on non-invasive functional testing who are on optimal drug treatment  
 6 benefit from routine coronary angiography with a view to revascularisation?

7 ➤ **Why this is important:** Revascularisation has traditionally been offered to people  
 8 with stable angina who have evidence of reversible ischaemia on non-invasive  
 9 functional testing. Recent trials in people with stable angina (COURAGE, BARI-2D,  
 10 MASS II) have not shown survival benefit from revascularisation compared with drug  
 11 treatment. In the nuclear substudy of COURAGE (n = 314), PCI was shown to be  
 12 more effective in treating ischaemia than optimal drug treatment, and in  
 13 multivariate analyses reduction of ischaemia was associated with greater event-free  
 14 survival. It is unclear, however, whether people on optimal drug treatment who have  
 15 evidence of inducible ischaemia on non-invasive functional testing should routinely

1 have coronary angiography and revascularisation. This question is particularly  
2 relevant for people who have responded adequately (say Canadian  
3 Cardiovascular Class 1 or 2) to optimal drug treatment and in whom, based on  
4 symptoms alone, revascularisation is not indicated. To answer this question we  
5 recommend a randomised trial of interventional management versus continued drug  
6 treatment in people with stable angina and myocardial ischaemia on non-invasive  
7 functional testing, with all-cause mortality and cardiovascular mortality as the  
8 primary endpoints.

## 10 **12.9 Research recommendation B**

11 The GDG recommended the following research question:

- 12 ➤ **Research question:** In people with stable angina and multivessel disease (including  
13 left main stem [LMS] disease) whose symptoms are controlled on optimal drug  
14 treatment, would an initial treatment strategy of revascularisation be clinically and  
15 cost effective compared with continued drug treatment?
  
- 16 ➤ **Why this is important:** Research is needed to determine whether early investigation  
17 and revascularisation can improve longer term survival. People with stable angina  
18 may be disadvantaged if they do not have tests to identify whether they have a  
19 higher risk profile for early cardiac death, which could be reduced by  
20 revascularisation. This disadvantage could be magnified when people who are  
21 deemed to fall into very high risk groups (for example, LMS stenosis > 50% in the  
22 MASS II trial) are excluded from randomised trials, resulting in the benefits of  
23 revascularisation being underestimated. We propose a randomised trial comparing  
24 an initial strategy of revascularisation (PCI or CABG) with an initial strategy of  
25 continued drug treatment in people with multivessel disease (including LMS disease)  
26 in whom revascularisation is not needed for symptom relief. The trial should use  
27 drug-eluting stents and wider inclusion criteria than BARI-2D and COURAGE.

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## 2 **13 Secondary prevention**

### 3 **13.1 Introduction**

4 The aim of treatment for people with stable angina is to reduce symptoms suffered  
5 by patients and also to improve long term outcomes. Secondary prevention measures  
6 are important to reduce the progression of cardiovascular disease and are of  
7 established benefit for patients in certain circumstances e.g. post myocardial  
8 infarction. NICE have published a guideline NICE Clinical Guideline 67 Lipid  
9 modification which recommends statins for all patients with evidence of cardiovascular  
10 disease. This review therefore examined the evidence for use of aspirin and ace  
11 inhibitors in people with stable angina.

12

1 **13.2 Aspirin**

2 Aspirin is an anti-platelet agent. Anti-platelet agents decrease platelet aggregation  
3 and may inhibit thrombus formation. Clopidogrel and dypiridamole do not have  
4 licences for use in stable angina.

5 **13.2.1 Clinical question**

6 What is the clinical effectiveness of aspirin to improve long term outcomes in people  
7 with stable angina?

8

9 **13.2.2 Clinical evidence**

10 The “Review Protocol” for this topic can be found in Appendix C, the “Search  
11 Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
12 E1, the “Clinical Evidence Tables” in Appendix E2, and the “Forest Plots” in Appendix  
13 F.

1 **Table 13.1: Aspirin vs. placebo for stable angina**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Aspirin	Placebo	Relative (95% CI)	Absolute	
<b>Non fatal MI (follow-up 50-60 months)</b>											
Juul-Moller 1992[158]; Ridker 1991[159]	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	14/1187 (1.2%)	94/1181 (8%)	RR 0.14 (0.08 to 0.25)	69 fewer per 1000 (from 60 fewer to 74 fewer)	⊕⊕⊕O MODERATE
<b>Fatal MI (follow-up 50-60 months)</b>											
Juul-Moller 1992[158]; Ridker 1991[159]	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	serious imprecision (g)	None	15/1187 (1.3%)	19/1181 (1.6%)	RR 0.79 (0.41 to 1.53)	3 fewer per 1000 (from 9 fewer to 8 more)	⊕⊕⊕O LOW
<b>Cardiovascular death (follow-up 60.2 months)</b>											
Ridker 1991[159] (d)	randomised trial	serious (b)	no serious inconsistency	no serious indirectness	serious imprecision (g)	None	6/178 (3.4%)	7/155 (4.5%)	RR 0.75 (0.26 to 2.17)	11 fewer per 1000 (from 33 fewer to 53 more)	⊕⊕⊕O LOW
<b>Sudden death (follow-up median 50 months)</b>											
Juul-Moller 1992[158]; (e)	randomised trial	serious (c)	no serious inconsistency	no serious indirectness	serious imprecision (g)	None	19/1009 (1.9%)	31/1026 (3%)	RR 0.62 (0.35 to 1.1)	11 fewer per 1000 (from 20 fewer to 3 more)	⊕⊕⊕O LOW
<b>Vascular events (follow-up median 50 months) (f)</b>											
Juul-Moller 1992[158];	randomised trial	serious (c)	no serious inconsistency	no serious indirectness	serious imprecision (h)	None	108/1009 (10.7%)	161/1026 (15.7%)	RR 0.68 (0.54 to 0.86)	50 fewer per 1000 (from 22 fewer to 72 fewer)	⊕⊕⊕O LOW
<b>Vascular deaths (follow-up median 50 months)</b>											
Juul-Moller 1992[158];	randomised trial	serious (c)	no serious inconsistency	no serious indirectness	serious imprecision (g)	None	51/1009 (5.1%)	70/1026 (6.8%)	RR 0.74 (0.52 to 1.05)	18 fewer per 1000 (from 33 fewer to 3 more)	⊕⊕⊕O LOW
<b>All cause mortality (follow-up median 50 months)</b>											
Juul-Moller 1992[158];	randomised trial	serious (c)	no serious inconsistency	no serious indirectness	serious imprecision (g)	None	82/1009 (8.1%)	106/1026 (10.3%)	RR 0.79 (0.6 to 1.04)	22 fewer per 1000 (from 41 fewer to 4 more)	⊕⊕⊕O LOW
<b>Haemorrhagic adverse events (follow-up median 50 months)</b>											
Juul-Moller 1992[158];	randomised trial	serious (c)	no serious inconsistency	no serious indirectness	serious imprecision (g)	None	27/1009 (2.7%)	16/1026 (1.6%)	RR 1.72 (0.93 to 3.17)	12 more per 1000 (from 1 fewer to 35 more)	⊕⊕⊕O LOW
<b>Non haemorrhagic adverse events (follow-up median 50 months)</b>											

Juul-Moller 1992[158];	randomised trial	serious (c)	no serious inconsistency	no serious indirectness	serious imprecision (g)	None	174/1009 (17.2%)	168/1026 (16.4%)	RR 1.05 (0.87 to 1.28)	8 more per 1000 (from 21 fewer to 46 more)	⊕⊕⊕○ LOW
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- 1 (a) *Juul-Moller 1992[158]: Multicentre Randomised, double blind, low drop out rate (0.5% drop out after 50 months), sample size calculation reported, baseline comparisons*
- 2 *made, Allocation concealment not reported, Intention to treat analysis not reported. Ridker 1991 Juul-Moller 1992[158]: Randomised, double blind, baseline comparisons*
- 3 *made, Intention to treat analyses used. Allocation concealment not reported.*
- 4 (b) *Randomised, double blind, baseline comparisons made, Intention to treat analyses used. Allocation concealment not reported.*
- 5 (c) *Multicentre randomised, double blind, low drop out rate (0.5% drop out after 50 months), sample size calculation reported, baseline comparisons made. Allocation*
- 6 *concealment not reported, Intention to treat analysis not reported.*
- 7 (d) *Drug dosage: Alternate day aspirin therapy (325 mg)*
- 8 (e) *Drug dosage: Aspirin 75 mg daily. All patients were treated with Sotalol, median dose was 160 (40-480 mg) daily.*
- 9 (f) *Vascular events (first occurrence of MI, stroke or vascular death)*
- 10 (g) *95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.*
- 11 (h) *95% CI around the pooled estimate of effect includes appreciable benefit or appreciable harm.*

1

2 **13.2.3 Economic evidence**

3 No economic studies were identified on this question. We calculated the daily and  
4 annual cost of aspirin based on the unit cost reported in the BNF59[18].

5 **Table 13.2: Drug cost - aspirin**

	Cost per day (£)	Cost per year (£)
Aspirin 75 mg, 1/day	0.035	12.8

6

7 The costs of adverse effects and events further down the line were not estimated.

8 **13.2.4 Evidence statements**

**Clinical**

**Aspirin vs. placebo**

**Juul-Moller 1992[158]; Ridker 1991[159]:** Evidence from 2 RCTs shows that there were significantly fewer patients with non fatal MI in the aspirin group compared to placebo. [RR 0.14 (0.08 to 0.25)]. (Follow-up 50-60 months)

**Juul-Moller 1992[158]; Ridker 1991[159]:** Evidence from 2 RCTs shows that there was no significant difference between aspirin and placebo for fatal [MI RR 0.79 (0.41 to 1.53)]. (Follow-up 50-60 months)

**Ridker 1991[159]:** Evidence from one RCT shows that there was no significant difference between aspirin and placebo for cardiovascular death [RR 0.75 (0.26 to 2.17)].[follow-up 60.2 months)

**Juul-Moller 1992[158]:** Evidence from one RCT shows that there were significantly fewer vascular events (first occurrence of MI, stroke or vascular death) in the aspirin group compared to placebo [RR 0.68 (0.54 to 0.86)]. (Follow-up median 50 months)

**Juul-Moller 1992[158]:** Evidence from one RCT shows that there was no significant difference between aspirin and placebo for sudden death [RR 0.62 (0.35 to 1.1)], vascular deaths (i.e, fatal vascular events) [RR 0.74 (0.52 to 1.05)] and all cause mortality [RR 0.79 (0.6 to 1.04)]. (Follow-up median 50 months)

**Juul-Moller 1992[158]:** Evidence from one RCT shows that there was no significant difference between aspirin and placebo for haemorrhagic adverse events [RR 1.72 (0.93 to 3.17)] and non-haemorrhagic adverse events [RR 1.05 (0.87 to 1.28)]. (Follow-up median 50 months)

**Economic**

No economic evidence was found on this question. A simple cost

analysis showed low drug costs of aspirin.

1 13.2.5 Recommendations and link to evidence

<b>Recommendation</b>	<b>Consider aspirin 75 mg daily for people with stable angina, taking into account the risk of bleeding and comorbidities.</b>
<b>Relative values of different outcomes</b>	The GDG were interested in a reduction in morbidity and mortality associated with use of aspirin for secondary prevention.
<b>Trade off between clinical benefits and harms</b>	<p>Aspirin use was associated with statistically significant reduction of non fatal MI and vascular events. All cause mortality and vascular deaths were not statistically significant but the GDG was impressed by a clinically significant risk reduction which approached statistical significance.</p> <p>There was a trend towards increased bleeding risk associated with the use of aspirin. The GDG were aware of recent debates concerning the use of aspirin for primary prevention and considered it likely that within the population of people with stable angina some are at higher risk of future cardiovascular events than others. For those at lowest risk the harms from aspirin might outweigh the benefits but there is currently no way of risk stratifying people with stable angina</p>
<b>Economic considerations</b>	The small drug cost of treatment with aspirin is likely to be offset by the improvement in clinical outcomes.
<b>Quality of evidence</b>	The quality for outcomes was low using GRADE methodology and the lack of precision contributed to this. The GDG however considered that the quality of the evidence was adequate to make a recommendation and consistent with what is known about use if aspirin across primary and secondary prevention.
<b>Other considerations</b>	The GDG agreed that aspirin should be considered for people with stable angina but did not think it should be offered to all patients. Healthcare professionals should take into consideration bleeding risk and co-morbidities when considering prescription of aspirin.

2

1 **13.3 ACE Inhibitors**

2 ACE inhibitors (angiotensin converting enzyme inhibitors) block the conversion of  
3 angiotensin 1 to angiotensin 11. They therefore lower arteriolar resistance and  
4 increase venous capacity; increase cardiac output and lower renovascular resistance.  
5 They are used to treat raised blood pressure but have been shown also to be  
6 beneficial for people with conditions such as heart failure.

7 **13.3.1 Clinical question**

8 What is the clinical /cost effectiveness of ACE inhibitors /ARB's for the management  
9 of angina?

10 **13.3.2 Clinical evidence**

11 The "Review Protocol" for this topic can be found in Appendix C, the "Search  
12 Strategies" in Appendix D, the "List of Included and Excluded Studies" in Appendix  
13 E1, the "Clinical Evidence Tables" in Appendix E2, and the "Forest Plots" in Appendix  
14 F.

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1 **Table 13.3: ACE inhibitors +background medication vs. placebo +background medication**

Quality assessment							Summary of findings				
							No of patients		Effect		Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	ACE +background medication	Placebo +background medication	Relative (95% CI)	Absolute	
<b>Combined (death from CV causes or non fatal MI) (follow-up mean 4.8 years)</b>											
Braunwald 2004[160] (e,g,h)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	344/4158 (8.3%)	352/4132 (8.5%)	RR 0.97 (0.84 to 1.12)	3 fewer per 1000 (from 14 fewer to 10 more)	⊕⊕⊕O MODERATE
<b>Death from cardio vascular causes (follow-up 3- 4.8 years)</b>											
Braunwald 2004[160]; Pitt 2001[161] (f,i)	randomised trials	serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	none	159/5036 (3.2%)	166/5004 (3.3%)	RR 0.95 (0.77 to 1.18)	2 fewer per 1000 (from 8 fewer to 6 more)	⊕⊕⊕O MODERATE
<b>Non fatal MI (follow-up 3- 4.8 years)</b>											
Braunwald 2004[160]; Pitt 2001[161]	randomised trials	serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	none	258/5036 (5.1%)	260/5004 (5.2%)	RR 0.99 (0.83 to 1.17)	1 fewer per 1000 (from 9 fewer to 9 more)	⊕⊕⊕O MODERATE
<b>Death from non cardiovascular or unknown causes (follow-up 3- 4.8 years)</b>											
Braunwald 2004[160]; Pitt 2001[161]	randomised trials	serious (b)	no serious inconsistency	no serious indirectness	serious (c)	none	167/5036 (3.3%)	195/5004 (3.9%)	RR 0.85 (0.69 to 1.04)	6 fewer per 1000 (from 12 fewer to 2 more)	⊕⊕OO LOW
<b>Hospitalised with unstable angina (follow-up 36 months)</b>											
Pitt 2001[161]	randomised trials	serious (d)	no serious inconsistency	no serious indirectness	serious (c)	none	52/878 (5.9%)	45/872 (5.2%)	RR 1.15 (0.78 to 1.69)	8 more per 1000 (from 11 fewer to 36 more)	⊕⊕OO LOW
<b>All causes death (follow-up 36 months)</b>											
Pitt 2001[161]	randomised trials	serious (d)	no serious inconsistency	no serious indirectness	serious (c)	none	27/878 (3.1%)	27/872 (3.1%)	RR 0.99 (0.59 to 1.68)	0 fewer per 1000 (from 13 fewer to 21 more)	⊕⊕OO LOW
<b>Hospitalisation due to CHF (follow-up mean 4.8 years)</b>											
Braunwald 2004[160]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (c)	none	105/4158 (2.5%)	134/4132 (3.2%)	RR 0.78 (0.61 to 1)	7 fewer per 1000 (from 13 fewer to 0 more)	⊕⊕OO LOW
<b>Death from CHF (follow-up mean 4.8 years)</b>											
Braunwald 2004[160]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (c)	none	15/4158 (0.4%)	25/4132 (0.6%)	RR 0.6 (0.31 to 1.13)	2 fewer per 1000 (from 4 fewer to 1 more)	⊕⊕OO LOW

- 1 (a) Block randomisation, double blind, sample size calculation reported, large sample (n=8290), Loss to follow-up (1.6% (68) in the placebo group and 1.6% (66) in the  
2 Trandolapril group) and intention to treat analysis used. Allocation concealment not reported.
- 3 (b) Both studies randomised, double blind and ITT used in both studies. Allocation concealment not used in both studies.
- 4 (c) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm
- 5 (d) Randomised, double blind, baseline comparisons made, sample size calculation reported, four patients lost to follow-up at 3 years, intention to treat analysis used.  
6 Allocation concealment not reported.
- 7 (e) Drug used: Trandolapril 2 mg per day
- 8 (f) Pitt 2001[161]: Drugs used: Quinapril 20 mg once daily.
- 9 (g) Ejection fraction >40% and <50% [% of patients]: 15% in both Trandolapril, and placebo groups.
- 10 (h) Background medications: 60% of patients on BB, 36% on CCB.
- 11 (i) Pitt 2001[161]: Background medications: 25% of patients on BB, 41% on CCB, and 73% on nitrates

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**Table 13.4: ACE inhibitors+BB vs. BB**

Quality assessment							Summary of findings				
							No of patients		Effect		Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	ACE+BB	BB	Relative (95% CI)	Absolute	
<b>Exercise time (min) (follow-up 12 weeks; better indicated by higher values)</b>											
Klein 1990[162] (c)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	None	23	23	-	MD 0.2 higher (1.16 lower to 1.56 higher)	⊕⊕○○ LOW
<b>Time to 1mm ST segment depression (min) (follow-up 12 weeks; better indicated by higher values)</b>											
Klein 1990[162]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	None	23	23	-	MD 0.2 higher (1.3 lower to 1.7 higher)	⊕⊕○○ LOW

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- (a) Randomised, cross over, double blind, baseline comparisons made. 6% (2/31) lost to follow-up. Allocation concealment not reported, Intention to treat analysis not used.
- (b) 95% CI includes no effect and the upper and lower CI crosses the MID.
- (c) Drugs used: Benazepril 10 mg twice daily plus metoprolol OROS, 14/190 mg once daily or metoprolol OROS, 14/190 mg (release rate/total dose) once daily.

1 **Table 13.5: ACE inhibitors + background medication vs. nifedipine + background medication**

Quality assessment							Summary of findings				Quality
							No of patients		Effect		
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	ACE +background medication	Nifedipine + background medication	Relative (95% CI)	Absolute	
<b>Combined Cardiac events (follow-up 3 years) (f)</b>											
Yui 2004[163] (e,i)	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	106/822 (12.9%)	116/828 (14%)	RR 0.92 (0.72 to 1.18)	11 fewer per 1000 (from 39 fewer to 25 more)	⊕⊕○○ LOW
<b>sudden death or cardiac death (follow-up 3 years)</b>											
Yui 2004[163]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	6/822 (0.7%)	6/828 (0.7%)	RR 1.01 (0.33 to 3.11)	0 more per 1000 (from 5 fewer to 15 more)	⊕⊕○○ LOW
<b>MI (follow-up 3 years)</b>											
Yui 2004[163]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	13/822 (1.6%)	16/828 (1.9%)	RR 0.82 (0.4 to 1.69)	3 fewer per 1000 (from 12 fewer to 13 more)	⊕⊕○○ LOW
<b>Hospitalisation for angina pectoris (follow-up 3 years)</b>											
Yui 2004[163]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	56/822 (6.8%)	50/828 (6%)	RR 1.13 (0.78 to 1.63)	8 more per 1000 (from 13 fewer to 38 more)	⊕⊕○○ LOW
<b>Hospitalisation for HF (follow-up 3 years)</b>											
Yui 2004[163]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	9/822 (1.1%)	12/828 (1.4%)	RR 0.76 (0.32 to 1.78)	3 fewer per 1000 (from 10 fewer to 11 more)	⊕⊕○○ LOW
<b>Non cardiac death (follow-up 3 years)</b>											
Yui 2004[163]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	9/822 (1.1%)	6/828 (0.7%)	RR 1.51 (0.54 to 4.23)	4 more per 1000 (from 3 fewer to 23 more)	⊕⊕○○ LOW
<b>Total mortality (follow-up 3 years)</b>											
Yui 2004[163]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	15/822 (1.8%)	12/828 (1.4%)	RR 1.26 (0.59 to 2.67)	4 more per 1000 (from 6 fewer to 24 more)	⊕⊕○○ LOW
<b>Adverse events (follow-up 3 years) (g)</b>											
Yui 2004[163]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (c)	none	121/822 (14.7%)	76/828 (9.2%)	RR 1.6 (1.22 to 2.1)	55 more per 1000 (from 20 more to 101 more)	⊕⊕○○ LOW
<b>Withdrawal due to adverse effects (follow-up 3 years) (h)</b>											

Yui 2004[163]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (c)	none	72/822 (8.8%)	41/828 (5%)	RR 1.77 (1.22 to 2.56)	38 more per 1000 (from 11 more to 77 more)	⊕⊕○○ LOW
<b>Diabetes sub group (combined cardiac events) (follow-up 3 years)</b>											
Yui 2004 (Subgroup Diabetes)[164]	randomised trials	serious (d)	no serious inconsistency	no serious indirectness	serious (b)	none	26/173 (15%)	30/199 (15.1%)	RR 1 (0.61 to 1.62)	0 fewer per 1000 (from 59 fewer to 93 more)	⊕⊕○○ LOW
<b>Diabetes sub group (cardiac death or sudden death) (follow-up 3 years)</b>											
Yui 2004 (Subgroup Diabetes)[164]	randomised trials	serious (d)	no serious inconsistency	no serious indirectness	serious (b)	none	3/173 (1.7%)	1/199 (0.5%)	RR 3.45 (0.36 to 32.87)	12 more per 1000 (from 3 fewer to 160 more)	⊕⊕○○ LOW
<b>Diabetes sub group (MI) (follow-up 3 years)</b>											
Yui 2004 (Subgroup Diabetes)[164]	randomised trials	serious (d)	no serious inconsistency	no serious indirectness	serious (b)	none	4/173 (2.3%)	4/199 (2%)	RR 1.15 (0.29 to 4.53)	3 more per 1000 (from 14 fewer to 71 more)	⊕⊕○○ LOW
<b>Diabetes sub group (hospitalisation for angina pectoris) (follow-up 3 years)</b>											
Yui 2004 (Subgroup Diabetes)[164]	randomised trials	serious (d)	no serious inconsistency	no serious indirectness	serious (b)	none	12/173 (6.9%)	16/199 (8%)	RR 0.86 (0.42 to 1.77)	11 fewer per 1000 (from 47 fewer to 62 more)	⊕⊕○○ LOW
<b>Diabetes sub group (Hospitalisation for HF) (follow-up 3 years)</b>											
Yui 2004 (Subgroup Diabetes)[164]	randomised trials	serious (d)	no serious inconsistency	no serious indirectness	serious (b)	none	5/173 (2.9%)	8/199 (4%)	RR 0.72 (0.24 to 2.16)	11 fewer per 1000 (from 31 fewer to 47 more)	⊕⊕○○ LOW
<b>Diabetes sub group (Total mortality)</b>											
Yui 2004 (Subgroup Diabetes)[164]	randomised trials	serious (d)	no serious inconsistency	no serious indirectness	serious (b)	none	5/173 (2.9%)	2/199 (1%)	RR 2.88 (0.57 to 14.64)	19 more per 1000 (from 4 fewer to 137 more)	⊕⊕○○ LOW

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- (a) Randomised, open, blinded endpoint design, sample size calculation reported, Intention to treat analysis used. concealment of allocation not reported
- (b) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.
- (c) 95% CI around the pooled estimate of effect includes appreciable benefit or appreciable harm.
- (d) Randomised, open, blinded endpoint design, sample size calculation reported, Intention to treat analysis used. Allocation concealment not used.
- (e) Drugs used: nifedipine retard (long acting nifedipine 20-40 mg/day) OR an ACE inhibitor (Enalapril 5-10 mg/day, Imidapril 5-10 mg/day, or Lisinopril 10-20 mg/day)
- (f) Combined cardiac events (cardiac death or sudden death, MI, angina pectoris requiring hospitalisation, HF requiring hospitalisation, serious arrhythmia, performance of coronary interventions)
- (g) The major adverse events occurring in the nifedipine group were those related to vasodilatory effect, including hypotension, facial erythema, and hot flushes. On the other hand dry cough accounted for most of the adverse events occurring in the ACE inhibitor group.
- (h) The main reasons for withdrawal were vasodilatory effect in the nifedipine group and predominantly cough in the ACE inhibitor group.

## DRAFT FOR CONSULTATION

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- (i) Medications used before the observation period- nifedipine group (67%) and ACE group (65%) on nitrates; nifedipine group (21%) and ACE group (18%) on BB; nifedipine group (52%) and ACE group (49%) on CCB . If the anti anginal effect of the treatment was inadequate, long acting or short acting nitrates and/or BB were used concomitantly.

### 13.3.3 Economic evidence

No economic studies were identified on this question. We calculated the daily and annual cost of a standard treatment with the most used ACE inhibitor based on the unit cost reported in the BNF59[18].

**Table 13.6: Drug cost - ACE inhibitors**

	Additional cost per day (£)	Additional cost per year (£)
Ramipril tablets, 5mg, 1/day	0.07	25.6

The costs of adverse effects were not estimated.

### 13.3.4 Evidence statements

#### Clinical ACE inhibitors +background medication vs. placebo + background medication

**Braunwald 2004[160]:** Evidence from one RCT shows that there was no significant difference between ACE inhibitors and placebo for combined cardiac events (death from CV causes or non fatal MI) [RR 0.97 (0.84 to 1.12)], hospitalisation due to CHF. [RR 0.78 (0.61 to 1)] and death from CHF [RR 0.6 (0.31 to 1.13)]. (Follow-up mean 4.8 years)

**Braunwald 2004[160]; Pitt 2001[161]:** Evidence from 2 RCTs shows that there was no significant difference between ACE inhibitors and placebo for death from CV causes [RR 0.95 (0.77 to 1.18)], non fatal MI [RR 0.99 (0.83 to 1.17)], and death from non cardiovascular or unknown causes [RR 0.85 (0.69 to 1.04)]. [Follow-up 3- 4.8 years]

**Pitt 2001[161]:** Evidence from one RCT shows that there was no significant difference between ACE inhibitors and placebo for hospitalisation with unstable angina [RR 1.15 (0.78 to 1.69)] and all causes death [RR 0.99 (0.59 to 1.68)]. (Follow-up 36 months).

#### ACE inhibitors+BB vs. BB

**Klein 1990[162]:** Evidence from one underpowered RCT shows that there was no significant difference between ACE + BB compared to BB for exercise time (min) [MD 0.2 (-1.16 to 1.56)] and time to 1 mm ST segment depression (min) [MD 0.2 (-1.3 to

1.7)]. [Follow-up 12 weeks]

**ACE inhibitors+ background medication vs. nifedipine + background medication**

**Yui 2004[163]:** Evidence from one RCT shows that there was no significant difference between ACE inhibitor and nifedipine for combined cardiac events (cardiac death or sudden death, MI, angina pectoris requiring hospitalisation, HF requiring hospitalisation, serious arrhythmia, performance of coronary interventions) [RR 0.92 (0.72 to 1.18)], sudden death or cardiac death [RR 1.01 (0.33 to 3.11)], MI [RR 0.82 (0.4 to 1.69)], hospitalisation for angina pectoris [RR 1.13 (0.78 to 1.63)], hospitalisation for HF [RR 0.76 (0.32 to 1.78)], non cardiac death [RR 1.51 (0.54 to 4.23)] and total mortality [RR 1.26 (0.59 to 2.67)]. [Follow-up 3 years]

**Yui 2004 (Diabetes Subgroup)[164]:** Evidence from one RCT shows that there was no significant difference between ACE inhibitor and nifedipine in diabetes sub group of patients for combined cardiac events (cardiac death or sudden death, MI, angina pectoris requiring hospitalisation, HF requiring hospitalisation, serious arrhythmia, performance of coronary interventions) [RR 1 (0.61 to 1.62)], cardiac death or sudden death [RR 3.45 (0.36 to 32.87)], MI [RR 1.15 (0.29 to 4.53)], hospitalisation for angina pectoris [RR 0.86 (0.42 to 1.77)], hospitalisation for HF [RR 0.72 (0.24 to 2.16)] and total mortality [RR 2.88 (0.57 to 14.64)]. [Follow-up 3 years]

**Yui 2004[163]:** Evidence from one RCT shows that there were significantly more adverse events [RR 1.6 (1.22 to 2.1)] and more withdrawals due adverse events [RR 1.77 (1.22 to 2.56)] in the ACE inhibitor group compared to nifedipine group [Follow-up 3 years]

**Economic**

No economic evidence was found on this question. A simple cost analysis showed a low additional cost of adding ACE-inhibitors to standard treatment.

### 13.3.5 Recommendations and link to evidence

<b>Recommendation</b>	<b>Do not offer angiotensin-converting enzyme (ACE) inhibitors to manage stable angina. Offer ACE inhibitors to treat other conditions, as appropriate.</b>
<b>Relative values of different outcomes</b>	The GDG were interested in intermediate and longterm morbidity and mortality outcomes when evaluating the value of ACE inhibitors for people with stable angina.
<b>Trade off between clinical benefits and harms</b>	There is no symptomatic or prognostic benefit from use of ACE inhibitors in the management of stable angina. There was no evidence available for ARB's in the management of stable angina.
<b>Economic considerations</b>	Since the clinical review showed no benefit from treatment with ACE inhibitors, using these drugs would increase costs with no additional benefits.
<b>Quality of evidence</b>	Large RCTs were available to answer this question.  No economic evidence was available on this question.
<b>Other considerations</b>	The GDG recognised that many people with stable angina may be on ACE inhibitors for management of other cardiac related conditions and these patients should remain on treatment as appropriate.

## 13.4 Further secondary prevention approaches covered by other NICE Clinical Guidelines

The use of statins and the treatment of high blood pressure are the subjects of other NICE Clinical Guidelines. Listed below are the details of the interventions and the recommendations made in the NICE TA.

### 13.4.1 Statins—NICE Clinical Guideline 67 (March 2010)

*“Cardiovascular risk assessment and the modification of blood lipids for the primary and secondary prevention of cardiovascular disease”*

<b>Recommendation</b>	<b>Offer statin treatment in line with 'Lipid modification' (NICE clinical guideline 67).</b>
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**13.4.2 Hypertension- NICE Clinical Guideline 34 (June 2006)**

*“Management of hypertension in adults in primary care”*

<b>Recommendation</b>	<b>Offer treatment for high blood pressure in line with 'Hypertension' (NICE clinical guideline 34, currently being updated).</b>
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## 14 Risk scores

### 14.1 Introduction

The GDG were interested in whether there were scoring systems available that would predict adverse outcomes. Ideally clinicians would like to be able to predict which patients were likely to have an adverse outcome and to intervene in those patients. In the absence of clear evidence for benefit of pharmacological or revascularisation strategies this might mean providing more intensive education and rehabilitation and support programmes to help patients to engage in secondary prevention strategies.

In this chapter we address the following key clinical question:

In adults with stable angina which tables, equations, engines, models or scoring systems are most reliable/effective for prognostic-risk stratification in prediction of adverse cardiac outcomes?

Two risk scoring systems were found that have been developed to predict adverse outcomes in patients with stable angina. The two risk scoring systems are: ACTION score- derived from a clinical trial population (ACTION trial)[165] and Euro heart Angina score - derived from a large cohort population (Euro Heart survey[166]).

### 14.2 Clinical Evidence

The “Review Protocol” for this topic can be found in Appendix C, the “Search Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix E1, and the “Clinical Evidence Tables” in Appendix E2.

#### **Derivation of risk scores**

For each risk score, multivariate analysis of baseline characteristics was performed to ascertain those characteristics which were most strongly associated with adverse outcomes- death or MI in Euro heart Angina score; and death all causes, MI or disabling stroke in the ACTION score. Risk scores were generated from the coefficients with an appropriate number of points given for the presence of each risk factor.

The components of each of the risk scores are shown below:

- A. ACTION risk score for death, MI or disabling stroke at 4.9 years follow-up:

Age, left ventricular ejection fraction, smoking, white blood cell count, diabetes, casual blood glucose concentration, creatinine concentration, previous stroke, at least one attack a week, coronary angiographic findings (if available), lipid lowering treatment, QT interval, systolic blood pressure  $\geq 155$  mm Hg, number of drugs used for angina, previous MI, sex.

B. Euro heart Angina score for death or MI at one year follow-up:

co-morbidity, diabetes, duration of symptoms, severity of symptoms, resting electrocardiogram abnormalities, abnormal ventricular function

#### 14.2.1 ACTION risk score

##### Clayton 2005[165]

This study used data from the ACTION trial (a coronary disease trial investigating outcome with nifedipine GITS), which followed 7665 patients with stable symptomatic angina for a mean of 4.9 years, to develop a score for predicting the combined risk of death from any cause, MI or stroke.

**Participants:** The Model was based on 7311 patients with values for all variables in model, of who 1063 had the combined event of death, MI, or disabling stroke.

**Inclusion criteria In the ACTION trial:** Eligible patients had stable symptomatic angina requiring treatment and either previous MI or proved angiographic coronary artery disease. Patients without a previous MI or coronary angiography could participate only if there was a positive result on an exercise or perfusion test. Key exclusions were ejection fraction below 40%, clinically significant heart failure, major cardiovascular event or intervention within the past 3 months, planned coronary angiography or intervention, and known intolerance to dihydropyridines. The patients were recruited from outpatient cardiology clinics in Western Europe, Israel, Canada, Australia, and New Zealand.

**Outcomes and follow-up:** The outcome measures were death from any cause or MI or disabling stroke with a follow-up of 4.9 years.

**Statistical analysis:** Multivariate Cox proportional hazard models used for the outcome time to death, MI, or disabling stroke as adjudicated by the critical events committee, using patients who had no missing values for the predictor variables. Each variables strength of predictive contribution was expressed by its z score (the model co-efficient divided by its standard error) and quantified each variables predictive power as hazard ratio with 95% CI.

For each patient, the risk score was calculated by multiplying each coefficient in the final model by 10, then by the patient's variable value, and then summed up the results.

**Results:** Table 14.1 shows the 16 variables, with the risk scores and Cox regression coefficients that were in the final model as derived for 7311 patients (95%) with complete information

**Table 14.1: Predictors of death, MI, or disabling stroke for 7311 participants in the ACTION trial (Cox proportional hazard analysis) – figures are numbers (%)**

Risk factors	Death, MI or stroke **(n=1063)	No death, MI, or stroke (n=6248)	Z score*	Co-efficient	Contribution to risk score
Mean age SD (year)	66.5 (9.5)	63 (9.2)	10.77	0.55	0 when age ≤ 60 years or add per 10 years > 60 years
Mean SD (ejection fraction)	46.7 (6.6)	48.6 (6.3)	6.47	0.17	0 when ≤ 60 years or add per 5% < 60%
Smoking					
Never	260 (24)	1784 (29)	-	-	
Ex smoker	560 (53)	3417 (55)	1.54	0.12	Add if applicable
current	243 (23)	1047 (17)	6.12	0.60	Add if applicable
Mean (SD) white blood cells (10 <sup>9</sup> /l)	7.4 (2.5)	7 (1.8)	6.07	0.068	0 when ≤ 5109/l > 5
Diabetes					
No diabetes	848 (80)	5393 (86)	-	-	
Non- ID diabetes	167 (16)	727 (12)	1.06	0.13	Add if applicable
ID diabetes	48 (5)	128 (2)	5.61	0.85	Add if applicable
Mean (SD) glucose, no diabetes (mg/dl)	103 (26)	99 (20)	4.68	0.072	0 when ≤ 100 mg/dl or add per 10mg/dl > 100 mg/dl.
Mean (SD) glucose, non-ID diabetes (mg/dl)	189 (79)	168 (65)	3.36	0.032	0 when ≤ 100 mg/dl or add per 10mg/dl > 100 mg/dl.
Mean (SD) creatinine (mg/dl)	1.14 (0.25)	1.08 (0.21)	4.27	0.078	0 when ≤ 1.15 mg/dl or add per 0.1 mg/dl > 1.15 mg/dl.
Previous stroke	50 (5)	116 (2)	3.59	0.53	Add if yes
Angina attack ≥ 1 /week	364 (34)	1750 (28)	3.42	0.22	Add if applicable
Previous angiography					
Never done	350 (33)	1842 (29)	1.50	0.11	Add if applicable
0-2 vessel disease	421 (40)	3069 (49)	-	-	Add 0 if applicable
≥ 3 vessel disease	292 (27)	1337 (21)	3.23	0.25	Add if applicable
No lipid lowering therapy	406 (38)	1950 (31)	3.20	0.21	Add if applicable
QT interval (12 lead ECG) ≥ 430msec	238 (22)	1096 (18)	3.05	0.23	Add if applicable
Systolic blood pressure ≥ 155 mmHg	275 (26)	1097 (18)	2.84	0.21	Add if applicable
No of drugs for					

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angina					
0	8 (1)	53 (1)			
1	268(25)	1953 (31)	2.76	0.13	Add once for each drug used
2	626 (59)	3487 (56)			
3	161 (15)	755 (12)			
Previous MI	597 (56)	3118 (50)	2.16	0.14	Add if yes
Male	863 (81)	4944 (79)	1.87	0.16	Add if male

\*Z score- co-efficient divided by its SE. Larger values indicate more highly significant risk factor: z scores of 1.96, 2.58, 3.29 and 3.89 correspond to p=0.05, p=0.01, p=0.001 and p=0.0001.

\*\*The definition of stroke excluded events without lasting disability. MI- did not include patients with chest pain and raised troponin concentrations.

**Note:** Age was the strongest predictor. Male sex was of borderline significance (p=0.06) but was retained for completeness. Diabetes and stroke were the strongest predictors from clinical history. Patients with known three or more vessel disease had raised risk. Other predictors included were left ventricular ejection fraction, a prolonged QT interval, use of lipid lowering drugs, and the number of drugs used for angina (including past use of CCB).

The table below presents hazard ratios for the individual events of death, MI, and disabling stroke with the same variables as for the combined endpoint.

**Table 14.2: Predictors of death, MI, and disabling stroke (Cox proportional hazard analysis) - figures are hazard ratios (95% CI)**

Risk factor	Death, MI, or stroke (n=1063)	Death (n=569)	MI (n=495)	Stroke (n=170)
Age per 10 years >60	1.73 (1.57 to 1.92)	2.30 (2.01 to 2.64)	1.45 (1.25 to 1.69)	1.75 (1.37 to 2.24)
Ejection fraction per 5%<60	1.19 (1.13 to 1.25)	1.26 (1.17 to 1.35)	1.14 (1.06 to 1.23)	1.24 (1.09 to 1.41)
Smoking				
Never	1.00	1.00	1.00	1.00
Ex smoker	1.13 (0.97 to 1.32)	1.19 (0.96 to 1.48)	0.99 (0.79 to 1.24)	1.42 (0.95 to 2.13)
current	1.82 (1.50 to 2.20)	2.20 (1.69 to 2.85)	1.39 (1.05 to 1.84)	2.44 (1.49 to 3.99)
White blood cells per 10 <sup>9</sup> /l>5	1.07 (1.05 to 1.09)	1.09 (1.07 to 1.12)	1.05 (1.01 to 1.10)	1.00 (0.92 to 1.09)
Diabetes				
No diabetes	1.00	1.00	1.00	1.00
Non ID diabetes	1.14 (0.90 to 1.44)	0.93 (0.66 to 1.32)	1.14 (0.81 to 1.60)	1.75 (1.06 to 2.90)
ID diabetes	2.33 (1.74 to 3.14)	3.44 (2.40 to 4.94)	2.62 (1.75 to 3.93)	0.56 (0.14 to 2.29)
Glucose per 10 mg/dl >100† (no diabetes)	1.08 (1.04 to 1.11)	1.10 (1.06 to 1.14)	1.05 (1.00 to 1.10)	1.07 (0.98 to 1.15)
Glucose per 10 mg/dl >100† (non-ID diabetes)	1.03 (1.01 to 1.05)	1.04 (1.01 to 1.07)	1.03 (1.00 to 1.06)	1.03 (0.99 to 1.07)
Creatinine per 0.1 mg/dl >1.5	1.08 (1.04 to 1.12)	1.09 (1.04 to 1.14)	1.08 (1.02 to 1.14)	1.06 (0.97 to 1.16)
Previous stroke	1.70 (1.27 to 2.28)	1.74 (1.19 to 2.54)	1.50 (0.95 to 2.36)	4.28 (2.60 to 7.06)
Angina attack ≥1 /week	1.25 (1.10 to 1.42)	1.27 (1.07 to 1.51)	1.21 (1.00 to 1.46)	1.16 (0.84 to 1.61)
Previous angiography				
Never done	1.12 (0.97 to 1.30)	1.16 (0.95 to 1.41)	1.20 (0.96 to 1.49)	1.10 (0.77 to 1.58)
0-2 vessel disease	1.00	1.00	1.00	1.00

≥3 vessel disease	1.28(1.10 to 1.50)	1.14 (0.92 to 1.41)	1.50 (1.21 to 1.87)	1.06 (0.72 to 1.57)
No lipid lowering therapy	1.23 (1.08 to 1.40)	1.33 (1.12 to 1.58)	1.10 (0.91 to 1.33)	1.09 (0.79 to 1.51)
QT interval (12 lead ECG) ≥ 430msec	1.26 (1.08 to 1.45)	1.52 (1.26 to 1.84)	1.08 (0.87 to 1.35)	1.69 (1.22 to 2.36)
Systolic blood pressure ≥ 155 mmHg	1.23 (1.07 to 1.42)	1.18 (0.98 to 1.43)	1.09 (0.88 to 1.35)	1.69 (1.22 to 2.36)
For each additional drug for angina	1.14 (1.04 to 1.25)	1.09 (0.96 to 1.24)	1.20 (1.05 to 1.38)	1.21 (0.96 to 1.54)
Previous MI	1.15 (1.01 to 1.30)	1.10 (0.92 to 1.30)	1.16 (0.96 to 1.39)	1.01 (0.74 to 1.38)
Male	1.17 (0.99 to 1.39)	1.21 (0.96 to 1.52)	1.24 (0.97 to 1.59)	0.88 (0.59 to 1.30)

**Note:** Patterns of risk factors were broadly similar, though risk of stroke was more strongly linked to raised blood pressure but unrelated to white cell count, angiographic data, previous MI and sex.

**Limitations of the score:** The risk score did not seem to predict the nature of the event (death in 39%, myocardial infarction in 46%, and disabling stroke in 15%) or the incidence of angiography or revascularisation, which occurred in 29% of patients.

**Summary:** The risk score combined 16 routinely available variables: age, left ventricular ejection fraction, smoking, white blood cell count, diabetes, casual blood glucose concentration, creatinine concentration, previous stroke, at least one attack a week, coronary angiographic findings (if available), lipid lowering treatment, QT interval, systolic blood pressure ≥ 155 mm Hg, number of drugs used for angina, previous MI, and sex. The patients risk is calculated by using ACTION score which is a number in the range of 0 to 60.

#### 14.2.2 Euro heart angina score

##### Daly 2006[166]

The Euro heart survey of stable angina was designed as a prospective observational cohort study of patients presenting to cardiology services with stable angina. Participating centres were a mix of academic and non academic institutions, and hospitals with and without interventional and cardiac surgical facilities.

**Participants:** N=3031 patients enrolled from 156 centres in 34 countries.

**Inclusion criteria:** Patients attending cardiology services with a new presentation of stable angina were considered for enrolment, and consecutive patients in whom the cardiologist made a clinical diagnosis of stable angina caused by myocardial ischemia due coronary disease were included in the survey. Exclusion criteria included unstable angina, admission to hospital within 24 hours of assessment, myocardial infarction within one year, previous revascularisation, or a cause of angina other than coronary disease.

**Baseline characteristics:** The population was relatively young 61 years and 58% male. Most patients had mild to moderate symptoms of angina for 6 months or less before presentation to a cardiologist, although only 48 (1.7%) patients had symptoms for less than one month before cardiology assessment. 10496 (40%) of patients were in class 1.

At baseline 1 602 (47%) of patients were on aspirin, 1 429 (21%) patients on statins and 1 142 (38%) on BBs.

**Confirmation of coronary disease:** Coronary angiography was done at least once during follow-up in 1 253 (41%) patients. At the end of the follow-up period, approximately one third (n = 994) of patients had had coronary disease confirmed angiographically and a further third (n = 1 023) had negative investigations. One sixth of patients had no definitive diagnostic test to confirm the presence or absence of coronary disease

**Outcome:** The primary outcome of interest was death or non fatal MI.

**Follow-up:** The median duration of follow-up was -13 months (interquartile range 12-15 months).

**Statistical analysis:** Cox's proportional hazards models were used to determine the effects of clinical and investigative variables on the occurrence of death or non fatal MI in both univariate and multivariate analysis. Starting with clinical variables, stepwise regression was done (using entry/removal P value = 0.15) to determine the factors predictive of death or infarction during follow-up. Models were developed separately for clinical and investigative parameters and then for a combination of clinical and investigative parameters. Final model was refitted for all patients without missing values for the variables selected.

**Results:** The Euro heart Angina score involves six characteristics: co-morbidity, diabetes, severity of symptoms, duration of symptoms, resting electrocardiogram abnormalities, and abnormal ventricular function.

The major clinical events occurring during follow-up in the overall population with stable angina (N=3031) are shown in the table below.

**Table 14.3: Major clinical events occurring during follow-up in the overall population with stable angina**

Endpoint	No of events	Event rate (95% CI) per 100 patient
Death*	50	1.5 (1.1 to 1.9)
Non cardiovascular death	14(28%)	
Non fatal MI	48	1.4 (1.1 to 1.9)
Death and non fatal MI	93	2.3 (1.9 to 2.8)
Cerebro vascular event	34	1.1 (0.8 to 1.5)
Heart failure	49	1.5 (1.1 to 2.0)
Unstable angina	164	5.2 (4.4 to 6.0)
All cardiovascular events	328	10.3 (9.3 to 11.5)

\*of 50 deaths, the cause of death was classified as unknown or missing in 6 and cardiac or cardiovascular in 29.  
 Note: Comparisons with clinical trial populations with stable angina: The annual incidence of death in the survey was 1.5% and the incidence of non fatal MI was 1.4%. In the subgroup with proved coronary disease these rates were 1.8% and 3.2%. Estimates of annual mortality from modern clinical trials of secondary prevention, anti anginal treatment, or revascularisation range from 0.9% to 1.7%, with a higher mortality in populations with more severe symptoms. Reported annual incidences of non fatal MI range from 1.1% to 1.5%.

The table below shows the risk of death or myocardial infarction associated with baseline clinical characteristics and results of investigations. Previous myocardial infarction, signs of heart failure, or a past history of diabetes, hypertension, or any co-morbidity were significant predictors of adverse outcome, as were increasing severity of symptoms and shorter duration of symptoms. Resting electrocardiographic abnormalities (Q wave or ST/T wave changes) were associated with approximately double the risk of death or myocardial infarction, but positive non-invasive stress test results were not significantly associated with adverse outcome.

**Table 14.4: Unadjusted hazard ratio of death or MI associated with clinical and investigative parameters in general population with stable angina (n=3031)**

Clinical variables	Hazard ratio	P value
Age (per 1 year increment)	1.03 (1.01 to 1.05)	0.001
Sex (female v male)	1.19 (0.79 to 1.79)	0.40
Diabetes	2.40 (1.55 to 3.70)	<0.001
Hypertension	2.12 (1.29 to 3.48)	0.002
Hyperlipidaemia	1.00 (0.63 to 1.58)	0.99
Ever smoked	1.53 (1.00 to 2.36)	0.05

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Previous myocardial infarction	3.24 (1.72 to 6.13)	0.002
Comorbidity	2.98 (1.98 to 4.52)	<0.001
Symptom severity:		
Class II versus class I	2.34 (1.37 to 4.00)	0.0002
Class III versus class I	3.44 (1.80 to 6.55)	
Symptom duration >6 months	0.60 (0.39 to 0.94)	0.03
Signs of heart failure	2.67 (1.56 to 4.57)	0.001
Body mass index >30	0.82 (0.49 to 1.37)	0.43
Tertiary education	0.78 (0.40 to 1.52)	0.46
Investigative variables		
Left bundle branch block	1.50 (0.66 to 3.43)	0.34
Q wave	2.37 (1.38 to 4.06)	0.002
ST or T wave changes	2.26 (1.50 to 3.41)	<0.001
Ischaemic ECG changes	2.27 (1.50 to 3.43)	<0.001
Result of individual stress tests:		
Positive exercise ECG (n=2299)	1.44 (0.80 to 2.61)	0.22
Positive stress echocardiogram (n=119)	1.24 (0.24 to 6.40)	0.80
Positive perfusion scan (n=420)	3.55 (0.77 to 16.47)	0.07
Result of any stress test		
Positive test	1.50 (0.82 to 2.73)	<0.0001

No test done	4.42 (2.50 to 7.82)	
Echocardiography (before events):		
Abnormal left ventricular function	5.21 (3.19 to 8.49)	<0.001

The table below shows stepwise regression selected co-morbidity, diabetes, recent onset of symptoms, more severe symptoms, ST or T wave abnormalities on the resting electrocardiogram, not having any stress test done, and abnormal ventricular function as the variables most predictive of outcome

**Table 14.5: Clinical and investigative parameters independently predictive of death or MI, determined by using stepwise selection procedures in general population with stable angina\*\***

Clinical variables (n=2183)	Hazard ratio (95% CI)	P- value
Comorbidity	2.41 (1.49 to 3.91)	<0.001
Signs of heart failure	1.62 (0.85 to 3.07)	0.14
Previous myocardial infarction	2.19 (1.08 to 4.42)	0.03
Diabetes	2.03 (1.25 to 3.31)	0.004
Symptom duration >6 months	0.54 (0.33 to 0.87)	0.01
Symptom severity:		
Class II versus class I	1.95 (1.07 to 3.54)	0.005
Class III versus class I	2.65 (1.29 to 5.50)	
Investigative variables (n=2963)		
Stress testing:		
Positive test	1.43 (0.76 to 2.70)	0.0001
No stress test done	3.78 (2.04 to 7.00)	
Echocardiography:		
Abnormal left ventricular function	2.57 (1.62 to 4.08)	<0.0001
Electrocardiography:		
ST or T wave changes	1.63 (1.06 to 2.50)	0.03
Combined clinical and investigative variables		

(n=2528)		
Comorbidity	2.25 (1.43 to 3.56)	0.0008
Diabetes	1.95 (1.22 to 3.11)	0.007
Previous myocardial infarction	—	
Symptoms >6 months	0.48 (0.30 to 0.77)	0.002
Symptom severity:		
Class II versus class I	1.76 (1.00 to 3.09)	0.05
Class III versus class I	2.18 (1.10 to 4.33)	
ST or T wave changes	1.56 (0.99 to 2.45)	0.05
Stress test:		
Positive stress test result	1.29 (0.63 to 2.67)	<0.0001
No stress test done	3.48 (1.71 to 7.07)	
Abnormal left ventricular function	2.11 (1.29 to 3.46)	0.004

*\*\* As non performance of a test is not an objective measure of a patient but can be influenced by many physician related and non clinical factors. A further stepwise selection process was used to consider only the non invasive investigations that had been done. A positive versus negative or inconclusive non-invasive stress test result was not selected as a significant predictor of outcome when combined with information from echocardiography and resting echocardiography.*

In the model developed to derive the clinical risk score the final predictors of death or MI were co-morbidity, diabetes, severity of symptoms, duration of symptoms, resting electrocardiogram abnormalities, and abnormal ventricular function.

**Validity:** Applying the model developed on 75% of the population to the remaining 25% of the population gave a C-statistic for the angina score to predict outcome of 0.74.

Cox's proportional hazards models were used to determine the effects of clinical and investigative variables on the occurrence of death or non fatal MI in both univariate and multivariate analysis.

To develop a scoring system for predicting probability of death or infarction during the first year after presentation that was based only on objective information generally available to clinicians and not on whether a test was done a further multivariate model was developed without the stress test done/not done variable. The performance of the model was assessed by calculating the Harrells C-statistics (comparable to the area under the receiver operating characteristics curve).

**Table 14.6: Score for each factor to calculate risk score for patients presenting with Stable angina**

Risk factor	Score contribution
Comorbidity*	
No	0
Yes	86
Diabetes	
No	0
Yes	57
Angina score	
Class 1	0
Class 2	54
Class 3	91
Duration of symptoms	
>6 months	0
<6 months	80
Abnormal ventricular function	
No	0
Yes	114
ST depression or T wave inversion on resting electrocardiogram	
No	0
Yes	34

\*One or more of previous cerebrovascular event; hepatic disease defined as chronic hepatitis or cirrhosis, or other hepatic disease causing elevation of transaminases more than three times upper limit of normal; peripheral vascular disease defined as claudication either at rest or on exertion, amputation for arterial vascular insufficiency, vascular surgery (reconstruction or bypass) or angioplasty to the extremities, documented aortic aneurysm, or non-invasive evidence of impaired arterial flow; chronic renal failure defined as chronic dialysis or renal transplantation or serum creatinine greater than 200 mol/l; chronic respiratory disease defined as a diagnosis previously made by physician or patient receiving bronchodilators or FEV1<75%, arterial pO2<60%, or arterial pCO2>50% predicted in previous studies; chronic inflammatory conditions defined as a diagnosis of rheumatoid arthritis, systemic lupus erythematosus or other connective tissue diseases, polymyalgia rheumatica, and so on; malignancy defined as a diagnosis of malignancy within a year or active malignancy.

Limitations: Small sample. The Euro heart survey of stable angina population differs from a general selection of people with angina in the community, many of whom may not have a diagnosis, and differs from the overall primary care angina population in that they have been selected for specialist assessment. However, the population is comparatively less highly selected than those in randomised controlled trials. The score has not been validated so far in a stable angina population.

Summary: In the model developed to derive the clinical risk score the final predictors of death or MI were co-morbidity, diabetes, shorter duration of symptoms, increasing severity of symptoms, abnormal ventricular function, resting electrocardiographic changes, or not having any stress test done. Results of the non invasive stress tests did not significantly predict outcome in the population who had tests done. A score was constructed using the parameters predictive of outcome to estimate the probability of the death or myocardial infarction within one year of presentation of stable angina. Applying the model developed on 75% of the population to the remaining 25% of the population gave a C-statistic for the angina score to predict outcome of 0.74.

### 14.3 Economic evidence

No economic studies were found on this question.

### 14.4 Evidence statements

**Clinical** There was evidence from 2 studies[165,166] that reported the derivation of ACTION risk score and Euro heart Angina score. However, there was no evidence available that validated the ACTION risk score and Euro heart Angina score in a stable angina population.

**Economic** No economic evidence was found on this question.

### 14.5 Recommendations and link to evidence

<b>Recommendation</b>	<i>No recommendation was made</i>
<b>Relative values of different outcomes</b>	
<b>Trade off between clinical benefits and harms</b>	
<b>Economic considerations</b>	No economic evidence was identified. If routine clinical indicators are used additional consultation costs are unlikely.
<b>Quality of evidence</b>	<p>Both risk scores were derived from selected patient populations that may not be representative of the wider population of patients with stable angina. The Euroheart score was developed from 75% of the total Euroheart survey population (derivation cohort) and tested in the remaining 25% of the population.</p> <p>The population used to develop the ACTION score was derived from the randomized ACTION trial, which enrolled patients with previous MI, or angiographic or other evidence of coronary heart disease.</p> <p>The available risk scores have not been validated in populations other than the cohorts in which they were developed.</p>
<b>Other considerations</b>	The GDG recognised that given the low event rate in stable angina a large cohort is required when developing a predictive model in a general angina population. The GDG did not consider that the evidence was sufficient to recommend using clinical risk scores but acknowledged that the clinical factors identified in the Euroheart study can result in a poorer outcome.

## 15 Functional and anatomical investigations

### 15.1 Introduction

NICE Clinical Guideline 'Chest pain of recent onset' emphasizes the importance of clinical assessment in establishing a diagnosis in people with chest pain. When the diagnosis is uncertain functional tests for the demonstration of inducible myocardial ischaemia and anatomical tests to confirm the presence of obstructive coronary artery disease are also recommended.

In people with an established diagnosis of stable angina non-invasive functional testing has also been recommended before invasive coronary angiography or revascularisation procedures for detection of myocardial ischaemia, risk stratification, and selection of appropriate treatment.[167].

In this chapter we review whether functional or anatomical tests in people with an established diagnosis of stable angina provide incremental value for the prediction of adverse cardiovascular outcomes and/or influence management to improve outcome. To add incremental value a test must provide additional prognostic information over and above that provided by standard clinical variables alone. Studies that did not assess incremental prognostic value were excluded.

The following tests were assessed in this review:

- Exercise electrocardiography - 2 papers
- Stress echocardiography - 2 papers
- Myocardial Perfusion Imaging - 9 papers
- Ambulatory electrocardiography - 2 papers

Tests in patients with normal coronary arteries and chest pain

- Stress echocardiography - 1 paper

Exercise electrocardiography is carried out with an exercise treadmill or bicycle ergometer, with step-wise increases in workload and continuous electrocardiographic monitoring. The test is continued to maximal exercise tolerance or development of clinical and/or electrocardiographic evidence of myocardial ischaemia (ST segment depression).

Stress echocardiography is carried out during exercise stress on a treadmill or bicycle ergometer, or during pharmacological stress induced by intravenous administration of dobutamine or dipyridamole. Detection of new wall motion abnormalities on the echocardiogram during stress is interpreted as evidence of inducible myocardial ischaemia.

Myocardial perfusion scintigraphy requires intravenous administration of a radioactive tracer [labelled with thallium-201 or technetium-99m (as tetrofosmin or sestamibi)] that is taken up by myocardial cells. A gamma camera is used to image the distribution of the tracer within the myocardium and to detect abnormalities of myocardial perfusion before and after exercise or pharmacological stress. Myocardial perfusion scintigraphy was originally developed as a planar imaging technique, but more recently single photon emission computed tomography (SPECT) has facilitated acquisition of tomographic images of the myocardium. In addition, ECG gating synchronises image acquisition with cardiac contraction, which reduces cardiac motion artefacts and facilitates measurement of left ventricular ejection fraction.

Ambulatory electrocardiography involves continuous electrocardiography, usually over 24-48 hours, and allows detection of spontaneous symptomatic or asymptomatic episodes of ST segment depression (myocardial ischaemia) during normal daily activities.

We found no evidence assessing the incremental prognostic value of cardiac computed tomography, cardiac magnetic resonance stress imaging, or invasive coronary angiography in patients with stable angina.

In this evidence review studies were not combined in a meta-analysis, because all of the included studies were observational studies. Additionally there was poor reporting of results in studies and heterogeneity across studies. The review is presented narratively with details of the test, population, end points, follow-up, results, and evidence statements for each study.

The following criteria were taken into consideration to give an overall quality rating of the primary studies: representativeness of the cohort; loss to follow-up being unrelated to key characteristics sufficient to limit potential bias ; adequate measurement of outcome of interest in study participants; prospectiveness of the study; adjustment for confounding factors in the analysis and at least 10 events per factor in the analysis (the study had to have at least 8 to 10 events per factor and analysis was adjusted for at least 3 of 4 relevant factors in the analysis). However, if there were insufficient relevant factors taken into account, the quality of the study was downgraded. All these factors were taken into consideration to give an overall quality rating.

**Table 15.1: List of studies with test, test variables, clinical variables, and outcomes - for patients with stable angina**

Study	Tests	Test variables considered in the analysis	Clinical variables considered in the analysis	Outcomes	Follow-up
<b>Exercise Electrocardiography</b>					
Forslund 2000[168]	Exercise electrocardiography (bicycle ergometry)	<p>Univariate analysis:                      Exercise duration (s)                      Maximal heart rate during exercise (beats.min)                      Time to onset of chest pain (s)                      Time to 1 mm ST segment depression (s)                      Maximal ST segment depression (mm)                      Maximal ST segment depression (mm) after 2 min rest (mm)</p> <p>Multivariate analysis:                      Maximal ST segment depression                      ST segment depression after exercise                      Exercise duration.</p>	<p>Univariate analysis:                      Age, sex, smoking habits, hypertension, previous MI, congestive heart failure, diabetes, mellitus.</p> <p>Multivariate analysis:                      Sex, MI, history of hypertension and diabetes mellitus.</p>	<p>1) CV death                      2) CV death and MI</p>	Median 40 months (6 months to 75 months)
Sekhri 2008[169]	Exercise electrocardiography (treadmill ergometry)	<p>Univariate analysis:  <u>Resting ECG:</u>                      Abnormal axis                      Q waves                      Change in ST segment or T wave                      Left ventricular hypertrophy                      Bundle branch block  <u>Exercise ECG:</u>                      Exercise time (mins)                      Maximum workload                      % predicted heart rate                      Maximum blood pressure</p> <p>Multivariate analysis:  <u>Resting ECG:</u>                      Q waves                      Bundle branch block</p>	<p>Univariate analysis:                      Age (10 year increase)                      Sex (female vs. male)                      Typicality of chest pain                      Heart rate per second increase                      Systolic blood pressure                      Hypertension                      Diabetes                      Current smoker</p> <p>Multivariate analysis:                      Age (10 year increase)                      Sex (female vs. male)                      Typicality of chest pain                      Diabetes</p>	Composite of death due to coronary heart disease or non fatal acute coronary syndrome.	Median 2.46 years

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		Change in ST segment or T wave  <u>Exercise ECG:</u> Exercise time (min)			
<b>Exercise echocardiography</b>					
D'Andrea 2005[170]	Exercise stress echocardiography (bicycle ergometry)	Univariate and multivariate analysis: <u>Rest echo:</u> Rest WMSI <u>Exercise echo:</u> Positive ESE Peak WMSI Low workload Angina during ESE	Multivariable analysis included significant variables in univariate analysis: Age, hypercholesterolemia, cigarette smoking.	1) Cardiac death 2) Cardiac death and non fatal MI	Mean 46.9 months (range 12-60 months).
Elhendy 2004[171]	Exercise echocardiography (treadmill ergometry)	Univariate analysis: <u>Echocardiographic variables:</u> Wall motion abnormality during exercise New wall motion abnormality (ischaemia) Percent ischaemic segments Wall motion score index during exercise Mean motion score index  <u>Exercise test variables:</u> 85% age predicted heart rate Systolic blood pressure during exercise Rate pressure product during exercise Workload (METs) Exercise induced angina Ischaemic electrocardiographic changes  Multivariate analysis: - <u>Echocardiographic variables:</u> Wall motion abnormalities	Univariate analysis: Age, gender, diabetes mellitus, smoking.  Multivariate analysis included significant variables in univariate analysis: Age, gender, diabetes	1) Any cardiac event defined as coronary artery revascularisation, non fatal MI, and cardiac death) 2) Cardiac death and non fatal MI	Median 2.7 years (1 to 7.8 years)

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		<p><u>Exercise test variables:</u> Workload Ischaemic electrocardiographic changes</p>			
<b>Myocardial Perfusion Imaging</b>					
Groutars 2002[172]	Myocardial perfusion scintigraphy using technetium-99m tetrofosmin with bicycle ergometry	<p>Multivariate analysis: Abnormal SPECT (Summed stress score SSS &gt;3) Summed stress score (SSS) Summed difference score (SDS) Severe ischaemia (SDS &gt;12) *</p>	<p>Univariate analysis: History of MI, history of PTCA, history of CABG, type of chest pain (indeterminate, atypical angina, typical angina, shortness of breath), age and gender, hypercholesterolemia, smoking, diabetes mellitus, hypertension.</p> <p>Exercise variable: Post exercise test likelihood of coronary artery disease.</p> <p>Multivariate analysis included significant variables in univariate analysis: History of MI, history of PTCA, history of CABG, typical angina symptoms, age and gender.</p> <p>Exercise variable: Post exercise test likelihood of coronary artery disease.</p>	Death, caused by any cardiac disorder with underlying coronary artery disease, including sudden death (confirmed by review of death certificate or hospital chart), or non fatal MI	Mean 23±9 months
Elhendy 2005[173]	Myocardial perfusion scintigraphy (SPECT) using technetium-99m tetrofosmin with bicycle ergometry	<p>Univariate and multivariate analysis:  Reversible perfusion defects Fixed perfusion defects</p>	<p>Univariate and multivariate analysis: Age Male sex History of heart failure Diabetes mellitus Smoking</p>	<p>1) Death from any cause 2) Cardiac death and non fatal MI</p>	Mean 6±1.7 years
Stratmann 1992[174]	Dipyridamole thallium-201 scintigraphy	<p>Univariate analysis: Normal scan Abnormal scan Reversible defect ≥1 segment</p>	<p>Univariate analysis: Age Sex History of old MI History of congestive cardiac failure</p>	Cardiac event (development of unstable angina, occurrence of a nonfatal MI, or death	Mean 18±9 months

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		<p>≥3 segments Fixed defect ≥1 segment ≥3 segments Reversible and fixed defects</p> <p>Multivariate analysis: Fixed defect Abnormal scan</p>	<p>History of diabetes mellitus History of systemic hypertension History of peripheral vascular disease History of cigarette smoking Pre study coronary angiography Pre study CABG Pre study coronary angioplasty</p> <p>Multivariate analysis: History of MI History of peripheral vascular disease History of congestive heart failure Pre test CABG</p>	<p>resulting from a primary cardiac cause) and cardiac death.</p>	
Wiersma 2009[175]	Myocardial perfusion scintigraphy (SPECT) using several isotopes and bicycle or treadmill ergometry	<p>Univariate analysis: Abnormal rest ECG MPS: severe ischaemia</p> <p>Multivariate analysis MPS: severe ischaemia</p>	<p>Univariate analysis: Male gender CCS II/IV BMI≥29.9 kg/m<sup>2</sup> Age 65 years or older Previous MI Previous revascularisation Aspirin Statin Insulin</p> <p>Multivariable analysis: Insulin</p>	Cardiac death or non fatal MI	Mean 2.2±0.7 years
Stratmann 1994[176]	Myocardial perfusion scintigraphy (SPECT) using technitium-99m sestamibi and pharmacological (dipyridamole) stress	<p>Univariate analysis: Occurrence of dipyridamole-induced chest pain, or MIBI perfusion defects.</p> <p>Multivariate analysis: Abnormal scan Reversible defect Fixed defect Chest pain during test</p>	<p>Univariate analysis: Age, gender, history of previous MI, congestive heart failure, diabetes mellitus treated with medication, systemic hypertension, peripheral vascular disease, cigarette smoking, or pre-test coronary revascularisation. CAD documented by coronary angiography before or ≤2 months after dipyridamole testing. Q waves on the pre test</p>	Cardiac death or non fatal acute MI	Mean 13±5 months

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			<p>Electrocardiogram consistent with prior MI Electrocardiographic changes consistent with ischaemia</p> <p>Multivariate analysis: History of congestive heart failure History of diabetes mellitus CAD by coronary angiography Q waves on pre-test ECG</p>		
Stratmann 1994[177]	Myocardial perfusion scintigraphy (SPECT) using technetium-99m sestamibi and treadmill ergometry	<p>Univariate analysis:</p> <p><u>Exercise treadmill test:</u> Chest pain during exercise Ischaemic ST depression <math>\geq 2</math> mm Peak HR, beats per minute peak HR <math>\geq 85\%</math> of age predicted maximal Peak SBP, mm Hg Peak DP, beats-mm Hg/min <math>\times 103</math> Exercise duration (Sec) Exercise duration <math>\geq 360</math> sec</p> <p><u>MPS:</u> Abnormal scan Reversible defect Fixed defect Reversible and fixed defects</p> <p>Multivariate analysis: Abnormal scan Reversible defect Fixed defect Ischaemic ST depression</p>	<p>Univariate analysis: Age, sex, history of congestive heart failure, history of old MI, history of diabetes mellitus, history of systemic hypertension, history of peripheral vascular disease, history of cigarette smoking, CAD by coronary angiography, pre study revascularisation, Q wave on pre test ECG, medications</p> <p>Multivariate analysis: History of congestive heart failure History of MI History of diabetes mellitus.</p>	Cardiac death or non fatal MI	Mean $13 \pm 5$ months (range 1 to 24 months)
Poornima 2004[178]	Myocardial perfusion scintigraphy (SPECT)	<p>Univariate and bi-variate analysis: A global stress score (GSS) was obtained by</p>	<p>Univariate and bi-variate analysis: <u>Clinical score:</u></p>	Cardiac death, MI, late revascularisation.	Mean $7 \pm 1$ year

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	using thallium-201 and treadmill ergometry	adding the scores on all the stress short axis images.	A simple five-point scoring system was developed after consideration of 16 clinical and ECG variables. The variables included in the five point scoring were male gender, history of MI (clinical event and Q waves on ECG), diabetes, insulin use, and typical angina.		
Vanzetto 1999[179]	Myocardial perfusion scintigraphy (SPECT) using thallium-201 and treadmill ergometry	<p>Univariate analysis:  <u>ETT variables:</u>                      Maximal heart rate, bpm                      Percentage of MPRR                      Maximum workload, W                      Negative ETT                      Positive ETT                      Strongly positive ETT                      Non diagnostic ETT                      Maximum ST segment depression, mm</p> <p><u>SPECT variables:</u>                      Abnormal T1201 SPECT                      Mean number of abnormal segments                      Mean number of fixed segments                      Mean number of reversible segments</p> <p>Multivariate analysis-  <u>ETT variables:</u>                      Negative ETT                      Positive ETT                      Strongly positive ETT                      Non diagnostic ETT                      Maximum ST segment depression, mm</p> <p><u>SPECT variables:</u>                      Normal T1201 SPECT                      1 or 2 abnormal segments on T1201-SPECT                      ≥3 abnormal segments on T1201-SPECT</p>	<p>Univariate analysis:</p> <p>Age &gt;60 years, sex, patients with &gt;1 risk factor, previous history of MI, typical angina.                      Multivariate analysis included significant variables in univariate analysis:                      Age &gt;60 years, patients with &gt;1 risk factor, previous history of MI</p>	Overall mortality; cardiac mortality (sudden death or death of demonstrated cardiac origin); occurrence of MI	Mean 72± SD 18 months
Lima 2004[180]	Myocardial perfusion scintigraphy (SPECT) with technitium-99 m	<p>Univariate analysis:  <u>ETT variables:</u>                      Peak rate pressure product                      V02 (METS)</p>	<p>Univariate variables:                      Not specifically reported</p> <p>Multivariate analysis included significant</p>	Cardiac events (cardiac death, MI, or myocardial revascularisation)	Mean 34±15 months

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	and pharmacological (dipyridamole) or exercise stress	Peak heart rate (beats/min) Peak % MAPHR Peak systolic arterial pressure (mmHg) ETT duration (min)  MPS: abnormal scan (Perfusion defects) Left ventricular enlargement	variables in univariate analysis: Gender, pre scan likelihood of CAD.		
<b>Ambulatory Electrocardiography</b>					
Forslund 1999[181]	Ambulatory electrocardiography	Univariate analysis: No. of VPCs No. of segment depressions/24 hr Duration of ST segment depression/24 hr (min)  Multivariate analysis: ST segment depression over 24 hours.	Univariate analysis: Age, sex, smoking, hypertension, previous MI, congestive heart failure, diabetes mellitus  Multivariate analysis: Sex, previous MI, hypertension and diabetes.	CV death, non fatal MI, and revascularisation	Median 40 months (6 to 75 months)
Conti 1997[182]	Exercise test and Ambulatory electrocardiography	Univariate analysis: Number of ambulatory ECG episodes Mean heart rate and maximum change in heart rate on baseline ambulatory ECG monitoring Abnormal 12 lead electrocardiogram at rest.  Multivariate analysis: Exercise time Ambulatory ECG episodes	Univariate analysis: Mean heart rate and maximum change in heart rate on baseline ambulatory ECG monitoring, history of revascularisation, history of MI, history of congestive cardiac failure, family history of coronary artery disease before age 55, diabetes mellitus, demographic variables (age, gender, race), certain variables related to history and disease (stenosis 50% in 1,2 or 3 vessels), ejection fraction <50%, history of hypertension, abnormal 12 lead electrocardiogram at rest and history of smoking.  Multivariate analysis included variables in univariate analysis p<0.05 (specific variables not reported).	Death, MI or hospitalisation for ischaemic event.	1 year

<sup>1</sup>SSS was obtained by calculating the sum of the scores of the 20 segments of the stress technetium-tetrofosmin images. The SRS was calculated on a similar basis. The SDS was calculated as the sum of the differences between SSS and the SRS for each segment.

**Table 15.2: For patients with chest pain and normal coronary arteries (Cardiac syndrome X)**

Study	Tests	Test variables	Clinical variables considered in the analysis	Outcomes	Follow-up
<b>Stress echocardiography</b>					
Bigi 2002[183]	Pharmacological stress echocardiography (dobutamine or dipyridamole)	Univariate analysis: Positive SE Rest WMSI Peak WMSI  Multivariate analysis: Positive SE Rest WMSI Peak WMSI	Univariate analysis: Clinical Age sex Previous infarction hypertension Diabetes Hypercholesterolemia  Multivariate analysis included significant variables in univariate analysis: Hypertension	Target events were cardiac death, non fatal infarction, and unstable angina. Only the worst event was taken in to account for statistical analysis.	Mean 36 months

1

2 **15.2 Exercise Electrocardiography**3 **15.2.1 Clinical question**

4 In adults with stable angina what is the incremental value/effectiveness of exercise  
5 electrocardiography for prognostic risk stratification in prediction of adverse cardiac  
6 outcomes?

7 **15.2.2 Clinical evidence**

8 The “Review Protocol” for this topic can be found in Appendix C, the “Search  
9 Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
10 E1, and the “Clinical Evidence Tables” in Appendix E2.

11 Two papers Forslund 2000[168,169] assessed the incremental value of exercise  
12 electrocardiography for prediction of adverse cardiac outcomes.

13 **Forslund 2000[168]** (n=809) evaluated the prognostic value of exercise tolerance  
14 testing (ETT) among patients with chronic stable angina.

15 The end-points were cardiovascular death, and cardiovascular death+MI.  
16 Cardiovascular death was defined as death from acute MI, sudden death (within 2  
17 hours of onset of symptoms) or death from other vascular causes (e.g. fatal  
18 cerebrovascular disease, pulmonary emboli). At follow-up ranging from 6 to 75  
19 months (median 40 months) there were 32 cardiovascular deaths and 29 MIs.

20 Prognostic implications of results from exercise tests were assessed in a multivariate  
21 Cox model which included sex, previous MI, hypertension and diabetes mellitus. After  
22 adjustment for these variables, maximal ST depression during exercise, ST segment  
23 depression 2 min after exercise, and exercise duration all carried independent  
24 relationships to both cardiovascular death and the combined endpoint of  
25 cardiovascular death + MI.

26

27 **Table 15.3: Prognostic evaluation of exercise variables –multivariate analysis for CV death**

Prognostic factors	Odds ratio (95% CI)	p value
Maximal ST depression	1.450 (1.15 to 1.83)	0.0018
Maximal ST depression 1-2 mm	0.827 (0.30 to 2.30)	0.71
Maximal ST depression $\geq$ 2 mm	1.619 (0.73 to 3.59)	0.23
ST segment depression after exercise:	1.850 (1.43 to 2.39)	0.00

ST segment depression 1-2 mm	1.502 (0.63 to 3.59)	0.36
ST segment depression $\geq 2$ mm	5.180 (2.12 to 12.67)	0.0003
Exercise duration (male patients)	0.786 (0.69 to 0.90)	0.0006
Exercise duration 9-13 min	0.358 (0.16 to 0.82)	0.015
Exercise duration $\geq 13$ min	0.250 (0.08 to 0.77)	0.016

1

2 **Table 15.4: Prognostic evaluation of exercise variables – multivariate analysis for CV death**  
3 **+MI:**

Prognostic factors	Odds ratio (95% CI)	p value
Maximal ST depression	1.33 (1.12 to 1.58)	0.001
Maximal ST depression 1-2 mm	1.36 (0.66 to 2.80)	0.402
Maximal ST depression $\geq 2$ mm	2.06 (1.11 to 3.83)	0.02
ST segment depression after exercise:	1.54 (1.26 to 1.91)	0.00
ST segment depression 1-2 mm	1.59 (0.89 to 2.85)	0.11
ST segment depression $\geq 2$ mm	3.03 (1.46 to 6.31)	0.002
Exercise duration (male patients)	0.834 (0.76 to 0.92)	0.0002
Exercise duration 9-13 min	0.506 (0.28 to 0.92)	0.02
Exercise duration $\geq 13$ min	0.314 (0.14 to 0.71)	0.005

4

5 **Sekhri 2008[169]** (n=1422) evaluated the prognostic value of exercise  
6 electrocardiograms (ECG) among patients with suspected angina and no previous  
7 diagnosis of coronary artery disease.

8 The primary end point was a composite of death due to coronary heart disease or  
9 non-fatal acute coronary syndrome. There were a total of 353 events at 1 year and  
10 the median follow-up was 2.46 years.

Adjusted hazard ratios for three models were reported: basic clinical assessment, basic clinical assessment plus resting electrocardiogram (ECG), and basic clinical assessment plus resting ECG plus exercise ECG (table X). In the final models (clinical assessment plus resting ECG plus exercise ECG) the major contributors to the risk of the primary end point were typical symptoms and abnormalities on the exercise ECG, with age, sex, and diabetes making variable additional contributions depending on whether the summary ECG subset or detailed ECG subset were analysed.

In the summary ECG subset only the clinicians' assessment of ischaemia was recorded (positive, negative, or equivocal). In the detailed ECG subset, data recorded included exercise time, maximum workload, maximum heart rate, maximum blood pressure, diagnostic change in ST segment, arrhythmias, and reason for stopping (limiting symptoms, ST segment displacement of more than 1 mm 0.08 seconds after the J point, or target heart rate achieved).

**Table 15.5: Sekhri 2008[169], Multivariate analysis for coronary heart disease death + MI**

Covariate	Coefficient	Adjusted hazard ratio (95% CI)	P value
Clinical assessment with significant variables (whole cohort)			
Age (10 year increase)	0.26	1.30 (1.21 to 1.39)	<0.001
Sex (female v male)	-0.28	0.75 (0.64 to 0.89)	0.0008
Typical v atypical	1.13	3.09 (2.58 to 3.71)	<0.001
Non-cardiac v atypical chest pain	-0.38	0.68 (0.50 to 0.93)	
Diabetes	0.45	1.58 (1.28 to 1.94)	<0.001
Clinical assessment plus resting ECG (whole cohort)			
Age (10 year increase)	0.23	1.26 (1.17 to 1.35)	<0.001
Sex (female v male)	-0.27	0.76 (0.65 to 0.90)	0.0013
Typical v atypical chest pain	1.04	2.82 (2.34 to 3.40)	<0.001
Non-cardiac v atypical chest pain	-0.37	0.69 (0.50 to 0.95)	
Diabetes	0.41	1.50 (1.22 to 1.83)	0.0002

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		1.86)	
Q waves	0.57	1.77 (1.24 to 2.53)	0.0037
Bundle branch block	0.30	1.36 (0.95 to 1.94)	0.1089
Change in ST segment or T wave	0.45	1.57 (1.28 to 1.94)	<0.001
Clinical assessment plus resting ECG plus summary exercise ECG			
Age (10 year increase)	0.10	1.11 (1.00 to 1.22)	0.048
Sex (female v male)	-0.05	0.95 (0.76 to 1.18)	0.64
Typical v atypical chest pain	0.75	2.12 (1.66 to 2.71)	<0.001
Non-cardiac v atypical chest pain	-0.54	0.58 (0.29 to 1.19)	
Diabetes	0.36	1.44 (1.09 to 1.89)	0.0134
Q waves	0.75	2.12 (1.28 to 3.49)	0.051
Bundle branch block	-0.11	0.90 (0.40 to 2.02)	0.79
Change in ST segment or T wave	0.29	1.34 (1.01 to 1.79)	0.0078
Positive v negative exercise ECG	0.92	2.53 (1.95 to 3.30)	<0.001
Equivocal v negative exercise ECG	0.44	1.55 (1.06 to 2.28)	
Clinical assessment plus resting ECG plus detailed exercise ECG			
Age (10 years increase)	0.03	1.03 (0.85 to 1.25)	0.76
Sex (female v male)	-0.59	0.55 (0.37 to 0.83)	0.0036
Typical v atypical	0.90	2.45 (1.62 to	<0.001

chest pain		3.70)	
Non-cardiac v atypical chest pain	-0.52	0.59 (0.14 to 2.45)	
Diabetes	0.03	1.03 (0.63 to 1.70)	0.9023
Q waves	0.49	1.64 (0.64 to 4.18)	0.3338
Bundle branch block	0.42	1.53 (0.48 to 4.89)	0.5022
Change in ST segment or T wave	0.32	1.37 (0.83 to 2.27)	0.2264
Exercise time (minutes)	-0.15	0.86 (0.79 to 0.93)	0.0005
Diagnostic change in ST segment	0.81	2.26 (1.44 to 3.53)	0.0005

1  
2 **Summary:** Two moderate quality prognostic studies showed that exercise  
3 electrocardiography (ECG) offered incremental prognostic value in prediction of CV  
4 death, CV death + MI, and death due to CHD + non fatal ACS. However it should be  
5 noted that in one of the studies the study sample did not entirely represent the  
6 population of interest. Also both studies reported composite outcomes and may  
7 overemphasize the incremental prognostic value of the tests.

### 8 15.2.3 Economic evidence

9 No relevant studies were found. Studies reporting the cost per case detected were not  
10 included as this question was addressed in the Chest Pain Guideline (CG95).

11 We looked for the costs of the individual tests from UK sources. We found that the  
12 unit cost of exercise test is £69 (NHS Reference Costs 2008-09 – Diagnostic Services -  
13 Exercise Test (including Treadmill, etc.) / Stress Test)[23].

### 15 15.2.4 Evidence statements

#### Clinical

#### **Exercise electrocardiography**

**Forslund 2000[168]:** Evidence from one study shows that exercise electrocardiography offers incremental prognostic information in prediction of CV death and CV death +MI [follow-up median (median 40 months)].

**Sekhri 2008[169] :** Evidence from one study shows that exercise ECG variables are independent predictors of death due to

coronary heart disease or non-fatal acute coronary syndrome. [median follow-up 2.46 years].

**Economic**

No economic evidence was found on this question. A simple cost analysis showed that exercise electrocardiography has a cost of £69 per test.

1

2 **15.3 Exercise echocardiography**

3 **15.3.1 Clinical question**

4 In adults with stable angina what is the incremental value/effectiveness of exercise  
5 echocardiography for prognostic risk stratification in prediction of adverse cardiac  
6 outcomes?

7 **15.3.2 Clinical evidence**

8 The “Review Protocol” for this topic can be found in Appendix C, the “Search  
9 Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
10 E1, the “Clinical Evidence Tables” in Appendix E2, and the “Forest Plots” in Appendix  
11 F.

12 Two papers[170,171] assessed the incremental prognostic value of exercise  
13 echocardiography for prediction of adverse cardiac outcomes.

14 **D’Andrea[170] 2005 (n=607)** assessed the long term predictive values of supine  
15 bicycle stress echocardiography (ESE) in patients with low, intermediate and high  
16 pretest risk of cardiac events.

17 The primary outcomes were cardiac death, and cardiac death and non fatal MI at a  
18 mean follow-up of 46.9 months (range 12-60 months). During the follow-up there 48  
19 deaths (21.6%) and 34 acute non fatal MIs (15.3%).

20 **Table 15.6: Multivariate predictive value of clinical risk factors and exercise stress**  
21 **echocardiography (ESE) results for cardiac death**

Variables	Chi square (X <sup>2</sup> )	p value	variables selected (partial X <sup>2</sup> ; 95% CI; p)
Clinical	9.3	0.01	cigarette smoking (2.8; 1.8 to 4.1; <0.01)
Clinical +rest echo	11.8	0.001	rest WMSI* (3.0; 2.1 to 4.1 ;< 0.01)
Clinical +rest echo+ ESE:	37.9	0.00001	positive ESE (4.1; 3.6 to 4.4; <0.0001)

			Peak WMSI* (3.5; 2.8 to 4.1); <0.0001  Low workload (3.1; 2.7 to 3.7; <0.01)
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1 \*wall motion score index

2 **Table 15.7: Multivariate predictive value of clinical risk factors and exercise stress**  
 3 **echocardiography (ESE) results for cardiac death+MI**

Variables:	Chi-square (X <sup>2</sup> )	p value	variables selected (partial X <sup>2</sup> ; 95% CI; p)
Clinical	9.6	0.01	hypercholesterolemia (2.5; 1.6 to 3.3; <0.01)
Clinical +rest echo	12.5	0.001	rest WMSI (3.1; 2.4 to 3.8 ;< 0.01)
Clinical +rest echo+ ESE	39.6	0.00001	Positive ESE (4.5; 3.6 to 5.3 ;< 0.0001)  Peak WMSI (3.7 ; 2.6 to 4.4; <0.0001)  Angina during ESE (2.9; 2.3 to 3.8; <0.01)

4

5 Multivariate analysis identified ESE positive for ischaemia, peak WMSI, low  
 6 workload, as the strongest independent predictors of cardiac death. Multivariate  
 7 analysis identified positive ESE, peak WMSI, angina during the test and  
 8 hypercholesterolemia as independent determinants of cardiac death or MI. The results  
 9 demonstrate that predictive information provided by ESE is additional and  
 10 independent to that provided by clinical and rest echocardiographic data.

11

12 **Elhendy 2004[171]** (n=437) assessed the incremental value of exercise  
 13 echocardiography in risk stratification of patients with a high pre-test probability of  
 14 coronary artery disease. Exercise echocardiography was done during symptom  
 15 limited treadmill exercise testing (Bruce protocol 89%, Naughton protocol 6%,  
 16 modified Bruce protocol 5%) with 12 channel electrocardiographic monitoring.

17 The end points were 1) non fatal MI and cardiac death and 2) coronary artery  
 18 revascularization, non fatal MI, and cardiac death assessed at a median follow-up of  
 19 2.7 years (1 to 7.8 years). Cardiac events occurred in 68 patients (16%) and  
 20 included four cardiac deaths, 15 non fatal MIs, and 53 revascularisation procedures  
 21 (4 subsequently had non fatal MI).

1

2 **Table 15.8: Independent predictors of cardiac events using a three step multivariate analysis**  
 3 **model**

Parameters	Chi-square ( $X^2$ )	p-value*; model chi-square **
Clinical model		
Age	0.01	0.9; 36
Gender	14	0.0002
Diabetes mellitus	1.9	0.2
Clinical and exercise test model		
Ischaemic electrocardiographic changes	3.2	0.07; 62 ***
Workload	4.8	0.03
Clinical, exercise stress and echocardiography model		
Wall motion abnormalities		78 *****
In multi vessel regions****	13.4	0.0003
In single vessel region****	2.8	0.1

4

\*Chi square and p value based on final model.

5

\*\* Overall model chi-square at each phase of the modelling process

6

\*\*\*  $p=0.0001$  versus the clinical model.

7

\*\*\*\* The reference group consisted of subjects with no wall motion abnormalities

8

\*\*\*\*\*  $p=0.001$  versus the clinical plus exercise stress model.

9

10 In a multivariate analysis of clinical, exercise, and echocardiographic parameters,  
 11 independent predictors of cardiac death and non-fatal MI were Q waves on the  
 12 stress electrocardiogram and the presence of wall motion of abnormalities during  
 13 exercise in multi-vessel distribution. In a separate analysis of clinical, exercise and  
 14 echocardiographic variables for the prediction of all cardiac events, the addition of  
 15 echocardiographic data increased the chi-square for the model from 62 to 78  
 16 ( $p=0.0003$ ).

17 **Summary:** Two moderate quality studies showed that **exercise echocardiography**  
 18 offered incremental value in prediction of cardiac death, MI and coronary  
 19 revascularization. The outcomes of interest were adequately assessed in both studies,  
 20 but one of the studies used composite outcomes and did not report individual cardiac  
 21 outcomes separately. This may cause bias as it offers an exaggerated perception of  
 22 the incremental prognostic value of the tests.

1 **15.3.3 Economic evidence**

2 No relevant studies were found. Studies reporting the cost per case detected were not  
3 included as this question was addressed in the Chest Pain Guideline (CG95).

4 We looked for the costs of the individual tests from UK sources. We found that the  
5 unit cost of stress echocardiography is £435[184].

6

7 **15.3.4 Evidence statements**

**Clinical** **Exercise echocardiography**

**D'Andrea 2005[170]**: Evidence from one cohort study shows that exercise stress echocardiography offered incremental prognostic information in prediction of cardiac death. [follow-up mean 46.9 months ].

**Elhendy 2004[171]** : Evidence from one cohort study shows that exercise echocardiography offered incremental prognostic information in prediction of cardiac events (cardiac death, non fatal MI, coronary revascularization) in addition to clinical and exercise variables. [follow-up median 2.7 years (1 to 7.8 years)].

**Economic**

No economic evidence was found on this question. A simple cost analysis showed that stress echocardiography has a cost of £435 per test.

8 **15.4 Myocardial perfusion imaging**

9 **15.4.1 Clinical question**

10 In adults with stable angina what is the incremental value/effectiveness of Myocardial  
11 Perfusion Imaging for prognostic risk stratification in prediction of adverse cardiac  
12 outcomes?

13 **15.4.2 Clinical evidence**

14 The "Review Protocol" for this topic can be found in Appendix C, the "Search  
15 Strategies" in Appendix D, the "List of Included and Excluded Studies" in Appendix E1  
16 and the "Clinical Evidence Tables" in Appendix E2.

17 Nine studies assessing the incremental prognostic value of myocardial perfusion  
18 imaging were included in this review [172-180]

19 **Groutars 2002[172]** evaluated the incremental prognostic value of myocardial  
20 perfusion scintigraphy using technetium-99m tetrofosmin with bicycle ergometry in  
21 597 patients with known or suspected coronary artery disease. Three nuclear

1 variables were evaluated, including the summed stress score (SSS), the summed rest  
 2 score (SRS), and the summed difference score (SDS). The SSS was obtained by  
 3 calculating the sum of the scores of the 20 segments of the stress technetium-  
 4 tetrofosmin images. The SRS was calculated on a similar basis. The SDS was  
 5 calculated as the sum of the differences between SSS and the SRS for each segment.  
 6 An SDS score between 2 and 12 was defined as moderate myocardial ischaemia and  
 7 an SDS score of >12 as severe ischaemia. The endpoints were death, caused by any  
 8 cardiac disorder with underlying coronary artery disease, including sudden death  
 9 (confirmed by review of death certificate or hospital chart), or non fatal MI  
 10 (documented by appropriate electrocardiographic and cardiac enzyme changes)  
 11 assessed at a mean follow-up of 23±9 months. A total of 46 events occurred: 16  
 12 cardiac deaths and 30 non fatal MI.

13 Multivariate analysis included four different nuclear variables, the SSS, SDS,  
 14 abnormal SPECT, and severe ischaemia. Abnormal SPECT was defined as an SSS  
 15 greater than 3 and severe ischaemia as SDS greater than 12. Abnormal SPECT (HR  
 16 5.438, CI 1.882 to 15.72, p=0.002) and SSS (HR 1.019, CI 1.001 to 1.038,  
 17 p=0.035) were significant independent predictors of cardiac death and MI.

18

19 **Table 15.9: Groutars 2002[172], Multivariate analysis of nuclear variables**

	Events (n=46)	No event (n=551)	HR	95% CI	P
Abnormal SPECT (SSS summed stress score >3)	41 (89%)	278 (50%)	5.438	1.882 to 15.72	0.002
Summed stress score (SSS)	28±20	13±17	1.019	1.001 to 1.038	0.035
Summed difference score (SDS)	12±14	7±11	1.036	1.036	0.110
Severe ischaemia (SDS >12)	15 (33%)	96 (17%)	0.342	0.342	0.072

20

21

22 **Elhendy 2005[173] (N=455)** assessed the independent value of SPECT imaging using  
 23 technetium-99m tetrofosmin with bicycle ergometry in predicting death from any  
 24 cause, cardiac death, and cardiac death and non-fatal MI (defined by cardiac  
 25 enzyme levels and electrocardiographic changes) in patients with stable angina  
 26 pectoris. During a mean follow-up of 6±1.7 years 93 (20%) patients died. Death was

1 considered cardiac in 46 patients (10%) and non fatal MI occurred in 40 patients  
2 (9%).

3  
4 **Table 15.10: Predictors of outcome events by Cox models**

Parameter	Univariate [RR (95% CI)]	Multivariate [RR (95% CI)]
<b>All cause mortality</b>		
Age	1.05 (1.02 to 1.09)	1.05 (1.03 to 1.08)
Male sex	2.5 (1.5 to 3.1)	2.1 (1.3 to 3.4)
History of heart failure	5.1 (2.7 to 10)	2.7 (1.6 to 4.5)
Diabetes mellitus	2 (1.2 to 3.4)	2.2 (1.4 to 3.5)
Smoking	1.9 (1.2 to 3.1)	1.7 (1.1 to 2.6)
Reversible perfusion defects	2 (1.2 to 3.1)	1.9 (1.1 to 2.8)
Fixed perfusion defects	2.3 (1.3 to 4.1)	2 (1.2 to 3.1)
<b>Cardiac mortality</b>		
Age	1.04 (1.01 to 1.09)	1.04 (1.02 to 1.07)
Male sex	2.5 (1.2 to 3.4)	1.8 (1.2 to 3.8)
History of heart failure	7.3 (3.5 to 15)	4.2 (2.1 to 7)
Diabetes mellitus	2.3 (1.2 to 4.4)	1.7 (1.2 to 3.9)
Abnormal perfusion	2.9 (1.8 to 5.1)	2.5 (1.5 to 3.5)
<b>Cardiac death or non-fatal MI</b>		
Age	1.03 (1.01 to 1.06)	1.03 (1.01 to 1.06)
Male sex	2.2 (1.3 to 3.6)	2.3 (1.3 to 4)
History of heart failure	2.9 (1.7 to 4.9)	2.8 (1.6 to 4.9)
Diabetes mellitus	1.6 (1.1 to 2.8)	1.8 (1.1 to 3.1)
Hypertension	1.7 (1.1 to 2.6)	1.9 (1.2 to 3)
Reversible perfusion defects	2 (1.2 to 3.1)	1.7 (1.1 to 2.4)

5  
6 In a multivariate model, independent predictors of death were age, male sex;  
7 diabetes, history of heart failure; smoking and MPS variables- reversible perfusion  
8 defects and fixed perfusion defects.

1 **Stratmann 1992[174]** (N=373) evaluated the usefulness of thallium -201  
 2 scintigraphy with dipyridamole stress for predicting the occurrence of cardiac events in  
 3 patients with stable chest pain. The outcomes assessed were cardiac event  
 4 (development of unstable angina, nonfatal MI, or death resulting from a primary  
 5 cardiac cause) and cardiac death at a mean follow-up of  $18\pm 9$  months. Cardiac  
 6 events occurred in 59 patients during the follow-up period, including unstable angina  
 7 in 27, non fatal MI in 11, and cardiac death in 21.

8 Regression analysis showed that a history of previous CABG and the presence of a  
 9 fixed perfusion defect were the only independent predictors of a subsequent cardiac  
 10 event. The presence of a fixed perfusion defect and a history of peripheral vascular  
 11 disease were found to be independent predictors of cardiac death.

12 **Table 15.11: Stratmann 1992[174], Predictors of cardiac events**

All cardiac events	Chi square	P value
Fixed defect	4.09	0.04
Abnormal scan (presence of a reversible defect, a fixed defect, or both reversible and fixed defects)	2.20	0.13
History of old MI	2.88	0.09
History of congestive heart failure	2.46	0.11
Pretest CABG	3.87	0.04
<b>Cardiac death</b>		
Fixed defect	7.04	0.008
Abnormal scan	0.36	0.54
History of old MI	5.46	0.02
History of peripheral vascular disease	8.54	0.004

13  
 14 **Wiersma 2009[175]** (N=319) determined the prognostic value of myocardial  
 15 perfusion scintigraphy in a population with type 2 diabetes mellitus and mild stable  
 16 angina (CCS class I-II). The outcome assessed was cardiac death or spontaneous, non  
 17 procedure-related, non fatal MI. During a mean follow-up of  $2.2\pm 0.6$  years 14  
 18 patients had a non fatal MI or died from a cardiac cause. Multivariate analysis  
 19 identified the presence of severe myocardial ischaemia (SDS  $\geq 8$ ) (HR 5.45, 95% CI  
 20 1.89 to 15.71) and insulin use (HR 4.00 95% CI 1.25 to 12.75) as independent  
 21 predictors of cardiac events.

1 **Table 15.12: Wiersma 2009[175], Multivariable Analysis**

Characteristic	Present	Absent	HR (95% CI)
Insulin use	11/158	3/161	4.00 (1.25 to 12.75)
MPS: severe ischaemia	8/63	6/256	5.446 (1.89 to 15.71)

2

3 **Stratmann 1994[176] (N=534)** evaluated technitium-99 m sestamibi SPECT using  
4 dipyridamole stress for assessing risk of subsequent cardiac events in patients with  
5 stable chest pain who were unable to perform diagnostically useful levels of exercise  
6 stress. Cardiac events included non fatal MI and cardiac death, and occurred in 58  
7 patients at a mean follow-up of  $13 \pm 5$  months

8 Stepwise logistic regression was used to evaluate the independent predictive value of  
9 clinical and test variables. In Model 1, the only scintigraphic variable included was the  
10 presence of an abnormal scan. In the second model the scintigraphic variables  
11 entered were specific types of myocardial perfusion defects, either reversible or  
12 fixed. In the first model, the presence of an abnormal scan, a history of congestive  
13 heart failure or diabetes mellitus, Q waves on the pre test electrocardiogram, and an  
14 abnormal MIBI study were identified as independent predictors of cardiac events. In  
15 the second model, reversible and fixed myocardial perfusion defects retained  
16 independent predictive value for cardiac events, as did congestive heart failure, Q  
17 waves on the pre-test electrocardiogram and dipyridamole induced chest pain.

18

1 **Table 15.13: Stratmann 1994[176], multivariate analysis**

Multivariate analysis	RR (95% CI)
<b>Model I</b>	
Abnormal scan	5.8 (1.8 to 19) *
Chest pain during test	1.8 (0.9 to 3.6)
History of congestive heart failure	1.8 (1.1 to 3.1) *
History of diabetes mellitus	1.8 (1.0 to 3.1)
CAD by coronary angiography	1.3 (0.8 to 2.3)
Q waves on pre-test ECG	1.8 (1.0 to 3.1) *
<b>Model II</b>	
Reversible defect	2.1 (1.2 to 3.5) *
Fixed defect	1.8 (1.0 to 3.4) *
Chest pain during test	1.7 (0.8 to 3.5)
History of congestive heart failure	2.0 (1.1 to 3.5) *
History of diabetes mellitus	1.9 (1.1 to 3.2) *
CAD by coronary angiography	1.4 (0.8 to 2.3)
Q waves on pre-test ECG	(1.0 to 3.2) *

2

3 **Stratmann 1994[177] (n=548)** assessed the relative prognostic value of exercise stress  
4 with myocardial perfusion imaging in a large population of patients presenting for the  
5 evaluation of stable chest pain consistent with angina pectoris. The outcomes assessed were  
6 cardiac events (cardiac death or non fatal MI) at a mean follow-up  $13 \pm 5$  months (range 1  
7 to 24 months). During follow-up 24 patients had a cardiac event including non fatal MI in  
8 11 and death from a cardiac cause in 13.

9 In the first regression model, the only scintigraphic variable included in the analysis was the  
10 presence of an abnormal perfusion scan. In the second model, patients with an abnormal  
11 perfusion scan result were classified into those with either reversible or fixed perfusion  
12 defects.

13

1

2 **Table 15.14: Stratmann 1994[177] (Exercise MIBI imaging), Univariate & multivariate analysis**

Multivariate analysis	RR (95% CI)
<b>Multivariate analysis- Model I</b>	
Abnormal scan	11.9 (1.6 to 89.4)
Ischaemic ST depression	2.2 (0.9 to 5)
History of congestive heart failure	1.6 (0.6 to 4.2)
History of old MI	1.2 (0.5 to 2.8)
history of diabetes mellitus	1.5 (0.6 to 4.1)
<b>Multivariate analysis- Model II</b>	
Reversible defect	2.9 (1.2 to 7)
Fixed defect	1.4 (0.6 to 3.3)
Ischaemic ST depression	2.0 (0.8 to 4.6)
History of congestive heart failure	1.9 (0.7 to 5.2)
History of old MI	1.3 (0.6 to 3.2)
history of diabetes mellitus	1.6 (0.6 to 4.2)

3 \*In Model I, scintigraphic variable included 'abnormal scan'; In Model II, scintigraphic variables included were  
4 'reversible defect' and 'fixed defect'.  
5

6 **Poornima 2004[178]** (N=1,461) assessed the incremental value of SPECT using  
7 thallium-201 and treadmill ergometry in symptomatic patients with low-risk Duke  
8 treadmill scores ( $\geq 5$ ). Most of the patients had atypical angina (71%). The outcomes  
9 assessed were: 1) cardiac death, non-fatal MI, late revascularization and 2) cardiac  
10 death or non fatal MI at a mean follow-up of  $7 \pm 1$  years. The total number of events  
11 was 211 and included 30 deaths, 55 non fatal MIs and 124 revascularization  
12 procedures.  
13

14 **Table 15.15: Poornima 2004[178] Univariate analysis**

Univariate results	Chi square ( $X^2$ )	p-value
Clinical score (CS) <sup>1</sup>		
Cardiac death	41.9	0.0001
Cardiac death/MI	102.7	0.0001
Cardiac death/MI/ late	102.7	0.0001

revascularisation		
global stress score (GSS) <sup>2</sup>		
Cardiac death	24.9	0.0001
Cardiac death/MI	14.2	0.0002
Cardiac death/MI/ late revascularisation	65.6	0.0001

1

2 **Table 15.16: Poornima 2004[178] Bivariate analysis**

<b>Bivariate results (including both CS and GSS)</b>	<b>Chi square (X<sup>2</sup>) (Adjusted)</b>	<b>p-value</b>
Clinical score (CS) <sup>1</sup>		
Cardiac death	31	0.0001
Cardiac death/MI	40.5	0.0001
Cardiac death/MI/ late revascularisation	73.5	0.0001
global stress score (GSS) <sup>2</sup>		
Cardiac death	7.74	0.005
Cardiac death/MI	2.71	0.10
Cardiac death/MI/ late revascularisation	23.6	0.0001

3

4

1 Clinical score (CS): A simple five-point scoring system was developed after consideration of 16 clinical and ECG variables. The variables included in the five point scoring were male gender, history of MI (clinical event and Q waves on ECG), diabetes, insulin use, and typical angina.

5

6

7

2A global stress score (GSS) was obtained by adding the scores of perfusion on all the stress short axis images. A global rest score (GRS) was obtained by adding the scores of all the redistribution short axis images. A global difference score (GDS) was obtained by subtracting GSS from GRS.

8

9

10

The CS (clinical score) and GSS (Global stress score) were significant independent predictors of cardiac death.

11

12

**Vanzetto 1999[179]** (N=1137) evaluated the prognostic value of Thallium 201 SPECT and exercise treadmill test in patients with low to intermediate-likelihood of future cardiac events.

13

14

15

The outcomes assessed were mortality, cardiac mortality (sudden death or death of demonstrated cardiac origin) and occurrence of MI (on the basis of characteristic chest pain, ECG changes, and serum creatine kinase level >twice the upper limit of normal).

16

17

1 During follow-up ( $72 \pm 18$  months [11 days to 8 years]) 88 patients died, 46 from a  
 2 cardiac cause. MI occurred in 57 patients, 7 of whom died from a cardiac cause  $8 \pm 4$   
 3 months later.

4  
 5 Age ( $p=0.04$ ), exercise treadmill test ( $p=0.03$ ), and thallium-201 SPECT ( $p=0.003$ )  
 6 were independent predictors of overall mortality. Thallium-201 SPECT and exercise  
 7 treadmill test were independent predictors of cardiac death. Thallium-201 SPECT was  
 8 also predictive of future MI, whereas exercise treadmill test was not.

9 In multivariate analysis, SPECT was of incremental prognostic value over clinical and  
 10 exercise treadmill test data for predicting overall mortality and major cardiac events.

11  
 12 **Table 15.17: Multivariate predictors of cardiac death**

	Odds ratio	95% CI	P value
Age >60 years	1.78	1.02 to 3.11	0.05
Previous MI	3.50	2.06 to 5.96	0.006
Positive ETT	0.83	0.25 to 2.80	Ns
Strongly positive ETT	2.66	1.23 to 5.76	0.02
Non diagnostic ETT	2.48	1.31 to 4.69	0.006
1 or 2 abnormal segments on T1201 SPECT	2.20	0.97 to 4.98	0.08
$\geq 3$ abnormal segments on T1201 SPECT	4.83	2.22 to 9.54	0.001

13  
 14 **Table 15.18: Multivariate predictors of non fatal MI**

	Odds ratio	95% CI	P value
Presence of $\geq 1$ risk factor	2.50	1.50 to 4.17	0.03
Previous MI	2.89	1.78 to 4.69	0.01
Positive Exercise Treadmill Test (ETT)	1.14	0.60 to 2.18	Ns
Strongly positive Exercise Treadmill Test (ETT)	0.89	0.43 to 1.85	Ns
Non diagnostic Exercise Treadmill	0.93	1.54 to 1.60	Ns

Test (ETT)			
Maximum ST segment depression $\geq 2$	1.34	0.76 to 2.37	Ns
1 or 2 abnormal segments on T1201 SPECT	4.20	1.93 to 9.14	0.002
$\geq 3$ abnormal segments on T1201 SPECT	4.97	2.15 to 11.49	0.004

1

2 **Lima 2003[180] (N=328)** evaluated the value of pharmacological (dipyridamole) or  
 3 exercise stress SPECT with technitium-99m for risk stratification of patients aged  $\geq 75$   
 4 years with suspected CAD.

5 The outcomes assessed were cardiac death or MI, and cardiac death, MI or  
 6 myocardial revascularization. During follow-up, 56 patients had cardiac events  
 7 including 24 cardiac deaths, 11 non fatal MIs and 21 revascularization procedures.

8 Logistic regression analysis of clinical, exercise treadmill test and MPS data was used  
 9 to identify significant predictors of cardiac events, with separate models for cardiac  
 10 death, cardiac death and MI, and any cardiac event. For cardiac death, the MPS  
 11 result was the most significant predictor variable ( $\chi^2=17.7$ , 95% CI: 5.9 to 30.6,  
 12  $p=0.0001$ ), followed by LV enlargement ( $\chi^2=10.3$ , 95% CI: 2.26 to 46.7,  
 13  $p=0.0004$ ).

14 For cardiac death and MI MPS result was also the most predictive variable ( $\chi^2=12.9$ ,  
 15 95% CI 5.3 to 3.19,  $p=0.0001$ ), followed by male gender ( $\chi^2=3.7$ , 95% CI 1.5 to  
 16 8.9,  $p=0.0001$ ) and pharmacologic stress ( $\chi^2=2.8$ , 95% CI 1.15 to 6.4,  $p=0.03$ ).

17 The independent predictors of any cardiac event were an abnormal scan ( $\chi^2=18.7$ ,  
 18 95% CI 8.9 to 39.6,  $p=0.0001$ ) and male gender ( $\chi^2=2.6$ , 95% CI 1.3 to 5.2,  
 19  $p=0.009$ ).

20 **Summary:** Nine studies (2 high quality, 2 moderate quality, and 5 low quality)  
 21 showed that **Myocardial Perfusion Imaging** offered incremental prognostic value in  
 22 prediction of cardiac death, MI, and/or revascularization. Most of the studies were  
 23 not of high quality as they did not have sufficient number of events (for validity the  
 24 study should have at least 10 patients (continuous) or 10 events (dichotomous) per  
 25 variable). Some studies did not include relevant risk factors (e.g. CCS class, LV  
 26 function) and had short follow-up periods. Also many studies reported composite  
 27 outcomes as their primary endpoint, and the components of these outcomes have  
 28 been inconsistently defined, and inadequately reported. This may cause an  
 29 exaggerated perception of the incremental prognostic value of the test being  
 30 evaluated.

1 **15.4.3 Economic evidence**

2 No relevant studies were found. Studies reporting the cost per case detected were not  
3 included as this question was addressed in the Chest Pain Guideline (CG95).

4 We looked for the costs of the individual tests from UK sources. We found that the  
5 unit cost of MPS with SPECT is £293[185].

6 **15.4.4 Evidence statements**

**Clinical**

**Myocardial perfusion imaging:**

**Groutars 2002[172] (Myocardial SPECT using technetium-99m tetrofosmin):** Evidence from one study shows that myocardial perfusion scanning using technetium 99m tetrofosmin offered incremental prognostic information in prediction of cardiac death or non fatal MI [follow-up 2 years].

**Elhendy 2005[173] (Myocardial SPECT using technetium-99m tetrofosmin):** Evidence from one study shows that stress technetium-tetrofosmin myocardial perfusion imaging is an independent predictor of all cause mortality in patients with stable angina. [mean follow-up 6±1.7 years].

**Stratmann 1992[174] (Dipyridamole thallium-201 scintigraphy):** Evidence from one study shows that presence of a fixed perfusion defect during dipyridamole stress and a history of CABG are independent predictors of cardiac events. The presence of a fixed perfusion defect and a history of peripheral vascular disease were independent predictors of cardiac death [mean follow-up 18 months].

**Wiersma 2009[175] (Myocardial SPECT):** Evidence from one study shows that the presence of severe myocardial ischaemia and insulin use were independent predictors of cardiac death or non fatal MI [follow-up 2.2±0.6 years].

**Stratmann 1994[176] (Myocardial SPECT using technetium-99m sestamibi and dipyridamole stress):** Evidence from one study shows that reversible and fixed perfusion defects on SPECT, history of congestive heart failure, history of diabetes mellitus, and Q waves on pre-test ECG were independent predictors of cardiac death or MI [mean follow-up 13±5 months].

**Stratmann 1994[177] (Myocardial SPECT using technetium-99m sestamibi and exercise stress):** Evidence from one study shows that exercise perfusion abnormalities and reversible perfusion defects were independent predictors of cardiac death or non-fatal MI [mean follow-up 13±5 months].

**Poornima 2004[178] (Myocardial SPECT and treadmill ergometry):** Evidence from one prognostic study shows that the

clinical score (CS) and global stress score (GSS) were significant independent predictors of cardiac death, cardiac death or MI; and cardiac death, MI or revascularisation. The independent predictive power of CS appeared to be greater than that of GSS [follow-up 7±1 year].

**Vanzetto 1999[179] (Myocardial SPECT using thallium 201 and exercise treadmill test):** Evidence from one prognostic cohort study shows that exercise tolerance test and SPECT were independent predictors of overall mortality. Exercise tolerance test and SPECT were independent predictors of cardiac death and SPECT was an independent predictor of MI [Follow-up 72±18 months]

**Lima 2004[180] (Myocardial SPECT using technetium-99m and exercise treadmill test):** Evidence from one study shows that SPECT was an independent predictor of cardiac death or MI, and of total cardiac events (cardiac death, MI or myocardial revascularisation). [mean follow-up 34±15 months]

**Economic** No economic evidence was found on this question. A simple cost analysis showed that MPS with SPECT has a cost of £293 per test.

1

## 2 15.5 Ambulatory ECG

### 3 15.5.1 Clinical question

4 In adults with stable angina what is the incremental value/effectiveness of “exercise  
5 tests and ambulatory ECG” for prognostic risk stratification in prediction of adverse  
6 cardiac outcomes?

### 7 15.5.2 Clinical evidence

8 The “Review Protocol” for this topic can be found in Appendix C, the “Search  
9 Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
10 E1, and the “Clinical Evidence Tables” in Appendix E2.

11 Two papers assessed the incremental prognostic value of ambulatory ECG for  
12 prediction of adverse cardiac outcomes [181,182].

13 **Forslund 1999[181] (N=686)** investigated whether ambulatory ECG and exercise  
14 testing provide complementary prognostic information in patients with stable angina  
15 pectoris.

16 The outcomes assessed were CV death, non fatal MI, and revascularisation. CV death  
17 was defined as death from acute MI, sudden death, or death from other vascular  
18 diseases. The criteria for MI were typical clinical presentation, significant increase in  
19 cardiac enzymes, and/or development of a new Q wave on the electrocardiogram.  
20 During follow-up (median 40 months, range 6 to 75 months) 29 patients had CV  
21 death, 27 had a nonfatal MI, and 89 underwent revascularisation.

1 The duration of ST segment depression over 24 hours (log transformed) was  
 2 independently related to CV death (OR 1.23, 95% CI 1.04 to 1.46, p=0.018) and to  
 3 CV death+ MI (OR 1.13, 95% CI 1.00 to 1.27, p=0.050). The odds ratio for  
 4 revascularisation was 1.11 (CI 1.01 to 1.22, p=0.035), and for the composite  
 5 endpoint was 1.11 (CI 1.04 to 1.20, p=0.004).

6 **Conti 1997[182] (n=558)** assessed the prognostic value of exercise test and  
 7 ambulatory ECG among patients enrolled in the ACIP trial. The outcome event (a  
 8 composite of death, MI, or hospitalisation for ischaemic event at 1 year) occurred in  
 9 73 cases.

10  
 11 **Table 15.19A: Model 1: (=angina history, ischaemia guided therapy, revascularisation strategy  
 12 –all baseline variables with p<0.05) (n=548)**

Variable:	p value	RR; 99% CI
History of angina(within 6 weeks of randomization)	0.01	1.95
Exercise time	0.01	0.89
Ambulatory ECG episodes	0.39	1.03
Duration of ischaemia	0.33	1.00
Ischaemia guided strategy	0.32	0.76
Revascularisation strategy	0.04	0.55

13  
 14 **Table 15.19B: Model 2: (=angina history, ischaemia guided therapy, revascularisation strategy-  
 15 all baseline variables stepwise)**

Variable	p value	RR (99% CI interval)
History of angina (within 6 weeks of randomization)	0.008	2.00; 1.02 to 3.94
Exercise time	0.006	0.88; 0.78 to 0.99
Ambulatory ECG episodes	NA	
Duration of ischaemia	NA	
Ischaemia guided strategy	0.32;	0.76
Revascularisation strategy	0.04; 0.55	

16  
 17 The model indicates that a history of angina in the 6 weeks before randomization and  
 18 a short total time on exercise treadmill at baseline were highly significant predictors  
 19 of adverse events (death, MI, or hospitalization for iscahemic event) within 1 year.

1           **Summary:** Of the two studies (one moderate and one low quality) , one of the studies  
2 showed that **ambulatory ECG** offered incremental prognostic value in prediction of  
3 cardiac death and MI; and the other study showed that ambulatory ECG was not an  
4 independent predictor of death, MI or hospitalisation for ischaemic events) . The  
5 studies were not of high quality as they had very few events (for validity the study  
6 should have at least 10 patients (continuous) or 10 events (dichotomous) per variable).  
7 Also both studies reported composite outcomes as their primary endpoint instead of  
8 reporting individual cardiac outcomes.

9   **15.5.3       Economic evidence**

10           No relevant studies were found. Studies reporting the cost per case detected were not  
11 included as this question was addressed in the Chest Pain Guideline (CG95).

12           We looked for the costs of the individual tests from UK sources. We found that the  
13 unit cost of ambulatory ECG is £56 (NHS Reference Costs 2008-09 – Diagnostic  
14 Services – 24 Hour ECG/BP monitoring)[23].

15   **15.5.4       Evidence statements**

**Clinical                Ambulatory ECG**

**Forslund 1999[181]:** Evidence from one study shows that duration  
of ST segment depression during the ambulatory  
electrocardiograph was an independent predictor of CV death  
and CV death+MI.

**Conti 1997[182] [exercise test and ambulatory ECG]:** Evidence  
from one study shows that history of angina in the 6 weeks before  
randomisation and a short total time on exercise treadmill at  
baseline were statistically significant predictors of adverse events  
(death, MI or hospitalisation for ischaemic events) within 1 year.  
Angina during ambulatory ECG or stress test was not predictive of  
an adverse event [follow-up 1 year].

**Economic**                No economic evidence was found on this question. A simple cost  
analysis showed that ambulatory ECG has a cost of £56 per test.

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17

1 **15.6 Recommendations and link to evidence**

<b>Recommendation</b>	<b>Do not routinely perform functional tests for myocardial ischaemia or anatomical tests for obstructive coronary artery disease to stratify risk. [This recommendation partially updates recommendation 1.2 of 'Myocardial perfusion scintigraphy for the diagnosis and management of angina and myocardial infarction' (NICE technology appraisal guidance 73)].</b>
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**Relative values of different outcomes**

The review indicates that functional tests provide modest incremental prognostic information. The magnitude of the increment in prognostic value is however unclear. The GDG were interested in identifying prognostic information and whether acting on this prognostic information would be beneficial to patients. Since all patients are given anti-anginal drugs and secondary prevention measures, the ability to identify patients who would benefit from revascularisation is critical. However neither this review nor the evidence reviews examining medical and revascularisation strategies (see chapter 10 and 11) provided evidence to identify patients who receive prognostic benefit from revascularisation

**Economic considerations**

All of the tests considered in this review are associated with some cost but there is no evidence that routine functional or anatomical testing provides additional clinical benefit. Routine functional testing was therefore not considered cost-effective.

**Quality of evidence**

The study inclusion criteria varied widely and the study participants may not be representative of the wider population of people with stable angina.

The studies were generally small with relatively short follow-up times. Consequently most of the studies had relatively few outcome events and limited statistical power to reliably identify predictor variables. For validity of the results the analysis should have at least 10 patients (continuous) or 10 events (dichotomous) per variable.

The studies generally did not include all potentially important clinical predictors of risk in the univariate and multivariate analyses and it is therefore not possible to

accurately quantify the incremental predictive value of any of the functional tests.

Several of the studies used composite outcomes. The use of revascularization as a component of a composite outcome is problematic when assessing the prognostic value of a functional test because the test result may directly influence the likelihood that individual patients will undergo revascularization

**Other considerations**

The GDG discussed whether it was appropriate to routinely perform tests that would provide prognostic information but that would not influence treatment. The GDG agreed the need to inform patients of the purpose and potential therapeutic implications of all investigations, particularly those associated with risk. The GDG agreed that functional and anatomical testing for prognostic information alone was unlikely to be justified or appropriate for the majority of patients.

The GDG were aware of a historical understanding in cardiology that functional testing in people with a confirmed diagnosis of stable angina is important in clinical assessment, including risk stratification and decisions about treatment. This strategy is recommended by other groups. [167]

The GDG discussed evidence that did not fulfill the inclusion criteria for the review but is influential in the discussion within cardiology about the benefit of functional testing. One study reports evidence from a registry of 10627 patients (of whom 39.7% had angina) who underwent exercise or adenosine myocardial perfusion SPECT[186]. Myocardial revascularisation was carried out within six weeks of the scan in 671 patients, and 9956 patients were initially managed medically. All patients were followed for a mean of 1.9 years and multivariate modelling was used to assess the effect of the extent of inducible myocardial ischaemia on the relationship between treatment strategy (revascularisation or medical therapy) and cardiac mortality. Above a threshold of 10%-12.5% ischaemic myocardium revascularisation was associated with lower cardiac death rate than medical therapy.

In the nuclear substudy of COURAGE (n=314) percutaneous coronary intervention produced more effective resolution of ischaemia than optimal medical treatment, and in multivariate analyses reduction of ischaemia was associated with greater event-free

survival[187].

The GDG considered evidence from these studies to be hypothesis-generating rather than definitive evidence on which recommendations could be based. The GDG considered this area a high priority for further research.

The GDG were aware that people with a confirmed diagnosis of stable angina may have had a functional or anatomical test during diagnostic assessment and that functional testing can be part of the assessment when deciding on revascularization strategy.

<b>Recommendation</b>	<p><b>Review the results of any functional and/or anatomical tests performed at diagnosis when revascularisation is being considered (see 'Chest pain of recent onset', NICE clinical guideline 95).</b></p> <p><b>Offer coronary angiography to guide the revascularisation strategy if not recently completed during diagnosis. Additional non-invasive or invasive functional testing may be required. [This recommendation partially updates recommendation 1.2 of 'Myocardial perfusion scintigraphy for the diagnosis and management of angina and myocardial infarction' (NICE technology appraisal guidance 73)].</b></p> <p><b>Consider further investigation to confirm the diagnosis of stable angina if the lack of response to drug treatment raises uncertainty about the diagnosis (see 'Chest pain of recent onset', NICE clinical guideline 95).</b></p>
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**Other considerations**

Diagnosis of angina is not in the scope of this guideline and was included in NICE Guideline 'Chest pain of recent onset'. That guideline includes recommendations on use of functional and anatomical tests in diagnosis of angina. The GDG were aware that the results of these investigations would already be available for some people with stable angina.

Patients who had not had coronary angiography and who had not responded to optimal medical treatment would require angiography to evaluate the coronary artery anatomy before a decision on revascularisation could be made. The GDG discussed whether all patients would also require functional testing. The evidence (reviewed for Chest pain guideline and discussed by the Stable Angina GDG) indicated functional testing in patients at low and moderate likelihood of coronary

artery disease was valuable but that patients at high likelihood of coronary artery disease a strategy of functional testing before angiography was not cost effective.

The patients in this guideline have already been diagnosed as having stable angina and were judged by the GDG to be at high likelihood of having coronary artery disease. Invasive functional testing done at the time of angiography or non-invasive functional testing might be required to guide revascularisation strategy.

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## 2    **16    Rehabilitation**

### 3    **16.1 Introduction**

4           Cardiac rehabilitation programmes have been shown to be of benefit to people with  
5           cardiovascular disease in certain circumstances e.g. those who have had a myocardial  
6           infarction (Myocardial infarction: secondary prevention. NICE clinical guideline 48  
7           (2007). The GDG were interested in whether there was evidence that patients with  
8           stable angina would similarly benefit from cardiac rehabilitation programmes.

9           There is no universal definition of cardiac rehabilitation. Cardiac rehabilitation can be  
10          defined 'as the process by which patients with cardiac disease, in partnership with a  
11          multidisciplinary team of health professionals, are encouraged and supported to  
12          achieve and maintain optimal physical and psychosocial health'[188].

13          Cardiac rehabilitation is usually discussed in the context of patients who have had an  
14          acute event such as myocardial infarction or cardiac surgery. The process of  
15          rehabilitation can be generally divided in to 4 phases: inpatient care, the early post  
16          discharge period, exercise training, and finally long term follow up. Early phases of  
17          rehabilitation concentrate on helping patients resume previous activity levels and this  
18          may not be appropriate for people with chronic stable angina who have not had an  
19          acute event or procedure. Cardiac rehabilitation programmes following discharge  
20          post myocardial infarction generally include structured exercise programmes with  
21          educational and psychological support and advice on risk factors offered by health  
22          professionals. The final phase of the rehabilitation incorporates the long term  
23          maintenance of physical activity and lifestyle changes. There is therefore considerable  
24          overlap between cardiac rehabilitation, secondary prevention and the longer term  
25          routine medical care that cardiac patients require. This is particularly so for people  
26          with stable angina who may not have or require any inpatient treatment.

27          This evidence review has used broad criteria when considering what evidence should  
28          be included. The main criteria were that the patients had stable angina and had an  
29          active intervention that could be considered important for rehabilitation and/or  
30          secondary prevention. Outcomes sought were those which represented improvement in  
31          angina, cardiovascular outcomes and quality of life. Programmes where patients were  
32          given advice only e.g. to exercise, to change diet are included in the review on effect  
33          of lifestyle factors in chapter 17.

1 A total of 20 papers were included in this review, with the trials evaluating a range  
2 of rehabilitation programmes. The studies have been analysed and presented  
3 separately according to the following themes:

- 4 • Exercise only – 4 papers
- 5 • Health Education – 2 papers
- 6 • Stress management programmes– 4 papers
- 7 • Intensive lifestyle programme – 1 paper
- 8 • Yoga life style programme– 1 paper
- 9 • Nurse led cardiac rehabilitation programme – 1 paper
- 10 • Angina management programme – 1 paper
- 11 • Angina Plan- 2 paper

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13 Only 2 papers [189,190] included Phases 1 and 2. The majority of the papers  
14 examined phases 3 and 4 and included exercise, education and advice on risk  
15 factors.

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17 The main results of the review are presented according to the content of rehabilitation  
18 programmes and their relevant comparisons as follows:

19 **Exercise programmes**

- 20 • Intensive exercise programme vs. control for stable angina
- 21 • Exercise and placebo vs. placebo for stable angina
- 22 • Exercise and BB vs. BB for stable angina
- 23 • Exercise plus low fat diet vs. control for stable angina
- 24 • Exercise vs. PCI

25 **Health education**

- 26 • Health education vs. control for stable angina

27 **Stress management**

- 28 • Stress management vs. routine care control for stable angina

- 1           • Stress management + exercise vs. routine care control for stable angina

2           **Yoga life style**

- 3           • Yoga life style intervention programme vs. control for stable angina

4           **Intensive lifestyle programme**

- 5           • Intensive life style intervention programme vs. control for stable angina

6           **Nurse led cardiac rehabilitation**

- 7           • Nurse led cardiac rehabilitation vs. routine care for stable angina

8           **Angina management programme**

- 9           • Angina management programme (AMP) vs. control for stable angina

10          **Angina Plan**

- 11          • Angina Plan vs. education session for stable angina

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13          **16.2 Clinical Evidence**

14          **16.2.1 Exercise Programmes**

15           The “Review Protocol” for this topic can be found in Appendix C, the “Search  
16           Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
17           E1, the “Clinical Evidence Tables” in Appendix E2, and the “Forest Plots” in Appendix  
18           F.

19           There were 4 papers with 5 relevant comparisons (Intensive exercise programme vs.  
20           control; Exercise and placebo vs. placebo; Exercise and BBs vs. BB; Exercise plus low  
21           fat diet vs. control; PCI vs. exercise +medical therapy) evaluating effectiveness of  
22           exercise training programmes for stable angina. Of these 4 papers 2 papers  
23           compared the effectiveness of exercise training with medical therapy[191] and  
24           PCI[192]. These studies could be considered as treatment options rather than  
25           rehabilitation but are included here for simplicity.

26

1 **Table 16.1: Intensive exercise programme vs. control**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Exercise (1 year intensive)	Control	Relative (95% CI) <sup>9</sup>	Absolute	
<b>Max ST depression (mm) (follow-up 1 years; better indicated by lower values)</b>											
Todd 1990[193] (a,b,c,d,e)	randomised trials	serious (f)	no serious inconsistency	no serious indirectness	no serious imprecision	none	20	20	-	MD 0.2 higher (-0.43 to 0.83)	⊕⊕⊕O MODERATE
<b>Time to 1mm ST depression (s) (follow-up 1 years; better indicated by higher values)</b>											
Todd 1990[193] (a,b,c,d,e)	randomised trials	serious (f,g)	no serious inconsistency	no serious indirectness	serious imprecision (i)	none	20	20	-	MD 166 higher (-221.71 to 553.7)	⊕⊕⊕O LOW
<b>Treadmill time (s) (follow-up 1 years; better indicated by higher values)</b>											
Todd 1990[193] (a,b,c,d,e)	randomised trials	serious (f,h)	no serious inconsistency	no serious indirectness	serious imprecision (i)	none	20	20	-	MD 262 higher (66.64 to 590.64)	⊕⊕⊕O LOW

- 2 (a) The intervention is a one-year intensive exercise training programme. The training group undertook the Canadian Air force Programme for Physical Fitness. It is a brief (11
- 3 minutes) daily exercise programme of five callisthenic type exercises. Exercise levels increase in intensity each week to achieve a progressive increase in physical fitness.
- 4 (b) All study patients were given atenolol for 2 weeks and then atenolol was stopped 4 wks before they were randomised to the exercise or control group. The main comparison
- 5 is between the exercise training programme (n=20) and B-blockers (same patients at baseline) with regard to exercise tolerance. In addition, a further comparison is made
- 6 between the exercise training programme patients and those who did not receive the exercise programme. A modified Naughton protocol exercise program was used to
- 7 assess tolerance. Randomisation produced groups whose baseline measurements differed statistically in only one respect. The mean (SD) maximum ST depression for the
- 8 control group (1.5 (0.8) mm) was significantly less than that for the exercise group (1.9 (0.9) mm). Quite large variations in other variables were, however, not statistically
- 9 significant. Most notable among these differences was the time to 1 mm ST depression, which was twice as long in the controls as in the exercise group. The overall trend
- 10 was for the exercise group to be less fit, as judged by resting and submaximal heart rate, and to have more severe disease, judged by maximum heart rate and double
- 11 product, maximum ST depression, and double product ST threshold.
- 12 (c) All patients had an exercise test at baseline then received 100 mg atenolol daily for one week and had another exercise test thereafter. Atenolol was then withdrawn and
- 13 patients were randomised. With regards to exercise compared to β blockers the authors conclude that regular exercise training was as good as atenolol in antianginal
- 14 efficacy since both improved Max ST depression, time to 1 mm ST depression and treadmill time equally well.
- 15 (d) Within the exercise group maximum ST depression was (1.9±0.9 to 1.6±1.2, p<.05), time to 1 mm ST depression (37.4±36.9 to 88.1±66.8, p<.001) and total treadmill
- 16 time (741±356 to 1272±514, p<.001) improved significantly.
- 17 (e) All patients in the exercise group reported an improvement in their anginal symptoms.
- 18 (f) No information was reported for methods of randomisation, or concealment of allocation to investigators - small sample size
- 19 (g) Time to 1 mm ST depression increased significantly from baseline for the exercise, but not the control group. Change score statistics not provided
- 20 (h) Treadmill time increased significantly from baseline for the exercise, but not the control group (p<0.001). Change score statistics not provided
- 21 (i) 95% CI includes no effect and the upper and lower CI crosses the MID.
- 22
- 23

1 **Table 16.2: Exercise and placebo vs. placebo for stable angina**

Quality assessment							Summary of findings				
							No of patients		Effect		Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Exercise (and placebo)	Placebo	Relative (95% CI)	Absolute	
<b>Maximal working capacity kpm/min (follow-up 4 months; better indicated by higher values)</b>											
Malmborg 1974[191]	randomised trials	serious (a,b)	no serious inconsistency	no serious indirectness	no serious imprecision	none	8	8	-	MD 4 lower (43.5 lower to 35.5 higher)	⊕⊕⊕○ LOW
<b>Anginal attacks / week (follow-up 4 months; better indicated by lower values)</b>											
Malmborg 1974[191]	randomised trials	serious (a,b)	no serious inconsistency	no serious indirectness	serious imprecision (c)	none	8	8	-	MD 25 lower (82.38 lower to 32.38 higher)	⊕⊕⊕○ LOW
<b>Nitroglycerin tablets/ week (follow-up 4 months; better indicated by lower values)</b>											
Malmborg 1974[191]	randomised trials	serious (a,b)	no serious inconsistency	no serious indirectness	serious imprecision (c)	none	8	8	-	MD 4 higher (96.75 lower to 104.75 higher)	⊕⊕⊕○ LOW

(a) This is a small pilot study (n=29 with n=8 maximum in the 4 groups).  
 (b) It did not specify a primary outcome and did not perform a power calculation.  
 (c) 95% CI includes no effect and the upper and lower CI crosses the MID.

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1 **Table 16.3: Exercise and BBs vs. BB for stable angina**

Quality assessment							Summary of findings				
							No of patients		Effect		Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Exercise and BBs	BB	Relative (95% CI)	Absolute	
<b>Maximal working capacity kpm/min (follow-up 4 months; better indicated by higher values)</b>											
Malmborg 1974[191]	randomised trials	serious (a,b)	no serious inconsistency	no serious indirectness	no serious imprecision	none	6	7	-	MD 6 lower (55.60 lower to 43.60 higher)	⊕⊕⊕○ LOW
<b>Anginal attacks / week (follow-up 4 months; better indicated by lower values)</b>											
Malmborg 1974[191]	randomised trials	serious (a,b)	no serious inconsistency	no serious indirectness	serious imprecision (c)	none	6	7	-	MD 41 higher (1.93 to 83.93 higher)	⊕⊕⊕○ LOW
<b>Nitroglycerin tablets/ week (follow-up 4 months; better indicated by lower values)</b>											
Malmborg 1974[191]	randomised trials	serious (a,b)	no serious inconsistency	no serious indirectness	serious imprecision (c)	none	6	7	-	MD 58 higher (37.02 lower to 153.02 higher)	⊕⊕⊕○ LOW

(a) This is a small pilot study (n=29 with n=8 maximum in the 4 groups).  
 (b) It did not specify a primary outcome and did not perform a power calculation.  
 (c) 95% CI includes no effect and the upper and lower CI crosses the MID.

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**Table 16.4: Exercise plus low fat diet vs. control**

Quality assessment							Summary of findings				
							No of patients		Effect		Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Exercise + low fat diet	Control	Relative (95% CI)	Absolute	
<b>Cardiac mortality (follow-up 12 months)</b>											
Schuler 1992[194]	randomised trials	serious (a,b)	no serious inconsistency	no serious indirectness	serious imprecision (c)	none	2/56 (3.6%)	0/57 0%	RR 5.09 (0.25 to 103.66)	40 more per 1000 (from 20 fewer to 90 more)	⊕⊕⊕○ LOW
<b>Mortality (all) (follow-up 12 months)</b>											
Schuler 1992[194]	randomised trials	serious (a,b)	no serious inconsistency	no serious indirectness	serious imprecision (c)	none	2/56 (3.6%)	1/57 1.8%	RR 2.04 (0.19 to 21.82)	19 more per 1000 (from 15 fewer to 375 more)	⊕⊕⊕○ LOW
<b>Non-fatal MI (follow-up 12 months)</b>											
Schuler 1992[194]	randomised trials	serious (a,b)	no serious inconsistency	no serious indirectness	serious imprecision (c)	none	0/56 (0%)	2/57 3.5%	RR 0.2 (0.01 to 4.15)	28 fewer per 1000 (from 35 fewer to 110 more)	⊕⊕⊕○ LOW

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- (a) Only compliant and responsive subjects were selected for this study, so results are likely to be better than those which would be found in a general population of patients with angina.
- (b) More patients dropped out of the study before treatment was complete in the exercise group (29% vs. 9% in the control group). No allowance was made for this in analysis of final dataset. Therefore, the health benefits gained in the exercise group will be an overestimate.
- (c) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.

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**Table 16.5: Exercise vs. PCI**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Exercise	PCI	Relative (95% CI)	Absolute	
<b>Death of cardiac causes (follow-up 12 months)</b>											
Hambrecht 2004[192]	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	0/51 (0%)	0/50 (0%)	not pooled	not pooled	⊕⊕⊕O MODERATE
<b>Cerebrovascular accident (follow-up 12 months)</b>											
Hambrecht 2004[192]	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	serious imprecision (b)	none	2/51 (3.9%)	3/50 (6%)	RR 0.65 (0.11 to 3.75)	21 fewer per 1000 (from 53 fewer to 165 more)	⊕⊕⊕O LOW
<b>Revascularisation (follow-up 12 months)</b>											
Hambrecht 2004[192]	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	serious imprecision (b)	none	3/51 (5.9%)	10/50 (20%)	RR 0.29 (0.09 to 1.01)	142 fewer per 1000 (from 182 fewer to 2 more)	⊕⊕⊕O LOW
<b>Hospitalisation and coronary angiography owing to worsening angina (follow-up 12)</b>											
Hambrecht 2004[192]	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	serious imprecision (b)	none	1/51 (2%)	7/50 (14%)	RR 0.14 (0.02 to 1.1)	120 fewer per 1000 (from 137 fewer to 14 more)	⊕⊕⊕O LOW

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(a) Even though allocation concealment is reported the method of randomisation is not clearly described  
 (b) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm

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**Additional data:**

**(Todd 1990)[193]**

No. of participants: n= 40 (Exercise (n=20); Control (n=20)). All patients in the exercise group noted an improvement in their symptoms within 6-8 weeks of starting training. At one year six patients were symptom free during normal activities these six and two others had stopped taking all antianginal agents except sublingual glyceryl trinitrate.

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2     **16.2.2   Health Education**

3             The “Review Protocol” for this topic can be found in Appendix C, the “Search  
4             Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
5             E1, the “Clinical Evidence Tables” in Appendix E2, and the “Forest Plots” in Appendix  
6             F.

7             There were 2 papers comparing Health Education programmes with control for stable  
8             angina [195,196].

1 **Table 16.6: Health education vs. control**

Quality assessment							Summary of findings				
							No of patients		Effect		Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Health Education	Control	Relative (95% CI)	Absolute	
<b>Mortality (follow-up 2 years)</b>											
Cupples 1994[195]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious imprecision (d)	none	13/342 (3.8%) (a)	29/346 (8.4%)	RR 0.45 (0.24 to 0.86)	46 fewer per 1000 (from 12 fewer to 64 fewer)	⊕⊕⊕○ LOW
<b>Increase in frequency of exercise (follow-up 2 years)</b>											
Cupples 1994[195]	randomised trials	serious	no serious inconsistency	no serious indirectness	serious imprecision (d)	none	108/342 (31.6%)	63/346 (18.2%)	RR 1.73 (1.32 to 2.28)	133 more per 1000 (from 58 more to 233 more)	⊕⊕⊕○ LOW
<b>Nottingham Health Profile (follow-up 2 years; measured with: Nottingham Health Profile (NHP); better indicated by higher values)</b>											
O'Neill 1996[196]	randomised trials	serious (b,c)	no serious inconsistency	no serious indirectness	no serious imprecision	none	221 MD -7.64	212 MD -20.43	-	confidence interval cannot be calculated – missing standard deviations	□□□□ MODERATE

- 2 (a) 10/13 deaths in the intervention group and 28/29 deaths in the control group were due to cardiovascular causes  
 3 (b) The conclusions reached in the abstract do not match the statistics in the result section.  
 4 (c) The mean differences in overall NHP scores did not reach statistical significance (p=0.0659), but were described in the abstract as significant. Mean differences of two  
 5 subscales reached statistical significance. Physical Mobility (MD intervention -1.49 and MD control -6.19, p=0.0015) and Social Isolation (MD intervention +1.42 and MD  
 6 control -3.01, p=0.0408) in favour of the intervention group  
 7 (d) 95% CI around the pooled estimate of effect includes appreciable benefit or appreciable harm.

1        **Additional data:**

2        Cupples 1994[195]: Angina episodes per week

3        For this outcome SD not reported along with the mean values hence data was not  
4        analysed. Data reported as in the paper.

5        The mean number of episodes of angina per week in the intervention group  
6        decreased from 3.2 (95% CI 2.7 to 3.7) at baseline to 2.6 (1.7 to 3.5) at review  
7        ( $p=0.04$ ), but no significant change was seen in the control group 2.5 (2.1 to 2.9) at  
8        baseline and 2.14 (1.7 to 2.5) at review.

9        **16.2.3    Stress management**

10       The “Review Protocol” for this topic can be found in Appendix C, the “Search  
11       Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
12       E1, the “Clinical Evidence Tables” in Appendix E2, and the “Forest Plots” in Appendix  
13       F.

14       There were 4 papers with 2 relevant comparisons (stress management vs. routine care  
15       control and; stress management + exercise vs. routine care control) evaluating the  
16       effectiveness of stress management programmes for stable angina[197-200].

1 **Table 16.7: Stress management vs. routine care control**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Stress management	routine care control	Relative (95% CI)	Absolute	
<b>Frequency of angina (average no. of. daily attacks) (8 weeks) (follow-up 8 weeks; better indicated by lower values)</b>											
Bundy 1998[200]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	42	16	-	MD 0.00 higher (2.92 lower to 2.92 higher)	⊕⊕⊕○ MODERATE
<b>Average duration of angina per attack (mins) (8 weeks) (follow-up 8 weeks; better indicated by lower values)</b>											
Bundy 1998[200]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	42	16	-	MD 0.40 lower (4.70 lower to 3.90 higher)	⊕⊕⊕○ MODERATE
<b>Frequency of chest pain at rest (days per fortnight) (6 months) (follow-up 6 months; better indicated by lower values)</b>											
Gallacher 1997[198]	randomised trials	serious limitations (b)	no serious inconsistency	no serious indirectness	no serious imprecision	none	158	179	-	MD 0.59 lower (1.24 lower to 0.06 higher)	⊕⊕⊕⊕ MODERATE
<b>Frequency of chest pain on exertion (days per fortnight) (6 months) (follow-up 6 months; better indicated by lower values)</b>											
Gallacher 1997[198]	randomised trials	serious limitations (b)	no serious inconsistency	no serious indirectness	no serious imprecision	none	158	179	-	MD 0.54 lower (1.35 lower to 0.27 higher)	⊕⊕⊕⊕ MODERATE

2 (a) Allocation concealment not clear. N=120 patients were randomised but only data for 99 patients was included in the analysis. It is not clear how the excluded patients  
3 were distributed among the groups or if there were systematic differences in excluded patients between groups. This is a relatively small, short term study aimed at assessing  
4 stress mgt, exercise training, stress mgt + exercise training combined with a waiting list control group. Patients were male and all had angina. No primary outcome  
5 measures were specified. Rather the study measured exercise workload, anginal symptoms and glyceryltrinitrate usage. 17% of patients were excluded from the analysis  
6 because they had only partial outcome data. No description of these patients was given or the distribution among treatment groups.  
7 (b) This is a large (n=452), well conducted study. Analysis however, was performed on data for only 70% of patients in the SMP group and 80% of those in the control  
8 group. Randomisation method was well described. Blinding was not described but relevant study results are based on patient diaries and not investigator assessment

1 **Table 16.8: Stress management + exercise vs. routine care control**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Stress management + exercise	routine care control (8 weeks)	Relative (95% CI)	Absolute	
<b>Frequency of angina (average no. of daily attacks (follow-up 8 weeks; better indicated by lower values) (final scores)</b>											
Bundy 1998[200]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	20	16	-	MD 0.6 higher (2.97 lower to 4.17 higher)	⊕⊕⊕O MODERATE
<b>Duration of angina (min) (follow-up 8 weeks; better indicated by lower values) (final scores)</b>											
Bundy 1998[200]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	20	16	-	MD 4.4 lower (9.08 lower to 0.28 higher)	⊕⊕⊕O MODERATE
<b>Frequency of angina (follow-up 8 weeks; better indicated by lower values) (change scores)</b>											
Bundy 1994[199]	randomised trials	serious (b)	no serious inconsistency	no serious indirectness	serious imprecision (c)	none	14	15	-	MD 2.70 lower (5.98 lower to 0.58 higher)	⊕⊕⊕O LOW
<b>Duration of angina (follow-up 8 weeks; better indicated by lower values) (change scores)</b>											
Bundy 1994[199]	randomised trials	serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	none	14	15	-	MD 0.70 lower (1.06 to 0.34 lower)	⊕⊕⊕O MODERATE

- 2 (a) Allocation concealment not clear. N=120 patients were randomised but only data for 99 patients was included in the analysis. It is not clear how the excluded patients were  
3 distributed among the groups or if there were systematic differences in excluded patients between groups. This is a relatively small, short term study aimed at assessing  
4 stress mgt, exercise training, stress mgt + exercise training combined with a waiting list control group. Patients were male and all had angina. No primary outcome  
5 measures were specified. Rather the study measured exercise workload, anginal symptoms and glyceryltrinitrate usage. 17% of patients were excluded from the analysis  
6 because they had only partial outcome data. No description of these patients was given or the distribution among treatment groups
- 7 (b) No description of method of randomisation or of "blinding" reported. All patients randomised completed the study. This is a small study (n=29) which aims to evaluate the  
8 effects of Stress Management Training (SMT) compared to routine care (RC) on exercise tolerance, angina symptoms, medication use and anxiety. All patients completed  
9 the study and the intervention is well described. Follow-up was relatively short (8 weeks after study end) and the study did not specify a primary outcome. It simply reports  
10 results for all study outcomes measured. Only exercise tolerance was reported at 8 weeks follow up. The remaining outcomes (medication use, angina symptoms and  
11 anxiety) were only reported at baseline and at study end (8 weeks from start of treatment).
- 12 (c) 95% CI includes no effect and the upper and lower CI crosses the MID.

1        **Additional data**

2        **Amarosa-Tupler 1989[197]**

3        Number of angina incidents: Data not given but plotted on a line graph. No change in  
4        the weekly number of incidents of angina for the group which listened to the tape  
5        which contained information. Groups which listened to the tape containing relaxation  
6        and/or imagery instructions showed a marked decrease in the weekly number of  
7        angina incidents. When the subjects stopped listening to the tapes the incidents of  
8        chest pain remained low for 1 or 2 weeks, then began to increase. Pain intensity and  
9        number of medications: for the three groups with relaxation and/or imagery tapes,  
10       the results followed the same pattern as the number of weekly incidents of angina  
11       described previously, i.e. a decrease during the tape exposure followed by an  
12       increase.

13       **16.2.4    Yoga life style intervention programme**

14       The “Review Protocol” for this topic can be found in Appendix C, the “Search  
15       Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
16       E1, the “Clinical Evidence Tables” in Appendix E2, and the “Forest Plots” in Appendix  
17       F.

18       There was one paper[201] comparing Yoga Lifestyle programmes with control  
19       (conventional medical therapy) for Stable angina.

1 **Table 16.9: Yoga life style intervention programme vs. control**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Yoga life style intervention programme	Control (1 year)	Relative (95% CI)	Absolute	
<b>Mortality (follow-up 1 years)</b>											
Manchanda 2000[201]	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	0/21 (0%)	0/21 (0%)	not pooled	not pooled	⊕⊕⊕O MODERATE
<b>Angina episodes per week (follow-up 1 years; range of scores: -, better indicated by less)</b>											
Manchanda 2000[201] (b)	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	21	21	-	MD 3.3 lower (4.82 to 1.78 lower)	⊕⊕⊕O MODERATE
<b>Exercise duration (sec) (follow-up 1 years; range of scores: -, better indicated by less)</b>											
Manchanda 2000[201]	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	serious imprecision (d)	none	21	21	-	MD 39 higher (46.78 lower to 124.78 higher)	⊕⊕⊕O LOW
<b>ST segment depression (mm) (follow-up 1 years; range of scores: -, better indicated by less)</b>											
Manchanda 2000[201]	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	21	21	-	MD 2.52 lower (2.95 to 2.09 lower)	⊕⊕⊕O MODERATE
<b>Revascularisation (follow-up 1 years)</b>											
Manchanda 2000[201] (c)	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	serious imprecision (e)	none	1/21 (4.8%)	8/21 (38.1%)	RR 0.12 (0.02 to 0.91)	335 fewer per 1000 (from 34 fewer to 373 fewer)	⊕⊕⊕O LOW

- 2 (a) Strengths: prospective randomised ; no attrition ; independent observers blinded to treatment allocation ; good compliance Weaknesses: small sample size ; randomisation  
3 and allocation concealment methods unclear ; blinding not possible due to nature of intervention ; groups significantly different at baseline in number of anginal episodes  
4 and exercise duration  
5 (b) At baseline patients in yoga group had significantly more anginal episodes per week (6.7±3 vs. 4.1±2.1).  
6 (c) Only 1 in the yoga group needed revascularisation (PTCA) against 8 in the control group (2 PTCA and 6 CABG) (RR 5.45 p=0.001)  
7 (d) 95% CI includes no effect and the upper and lower CI crosses the MID.  
8 (e) 95% CI around the pooled estimate of effect includes appreciable benefit or appreciable harm.

1

2     **16.2.5   Intensive life style programme**

3           The “Review Protocol” for this topic can be found in Appendix C, the “Search  
4           Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
5           E1, the “Clinical Evidence Tables” in Appendix E2, and the “Forest Plots” in Appendix  
6           F.

7           There was one paper comparing Intensive lifestyle programme with control for Stable  
8           angina[202].

1 **Table 16.10: Intensive lifestyle programme vs. control for stable angina**

Quality assessment							Summary of findings				
							No of patients		Effect		Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Intensive lifestyle programme	control (5 years)	Relative (95% CI)	Absolute	
<b>Angina frequency (times per week) (follow-up 5 years; range of scores: -; better indicated by less)</b>											
Ornish 1998[202]	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	18	14	-	MD 0.7 higher (0.9 lower to 2.3 higher)	⊕⊕⊕O MODERATE
<b>Chest pain duration (min) (follow-up 5 years; range of scores: -; better indicated by less)</b>											
Ornish 1998[202]	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	18	14	-	MD 0.1 lower (1.64 lower to 1.44 higher)	⊕⊕⊕O MODERATE
<b>MI (follow-up 5 years)</b>											
Ornish 1998[202]	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	serious imprecision (b)	none	2/28 (7.1%)	4/20 (20%)	RR 0.36 (0.07 to 1.76)	128 fewer per 1000 (from 186 fewer to 152 more)	⊕⊕⊕O LOW
<b>PTCA (follow-up 5 years)</b>											
Ornish 1998[202]	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	serious imprecision (b)	none	8/28 (28.6%)	14/20 (70%)	RR 0.41 (0.21 to 0.78)	413 fewer per 1000 (from 154 fewer to 553 fewer)	⊕⊕⊕O LOW
<b>CABG (follow-up 5 years)</b>											
Ornish 1998[202]	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	serious imprecision (b)	none	2/28 (7.1%)	5/20 (25%)	RR 0.29 (0.06 to 1.33)	178 fewer per 1000 (from 235 fewer to 83 more)	⊕⊕⊕O LOW
<b>Death (follow-up 5 years)</b>											
Ornish 1998[202]	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	serious imprecision (b)	none	2/28 (7.1%)	1/20 (5%)	RR 1.43 (0.14 to 14.7)	21 more per 1000 (from 43 fewer to 685 more)	⊕⊕⊕O LOW

2 (a) Strengths -RCT conducted from 1986 to 1992 using a randomised invitational design. Quantitative coronary arteriograms were blindly analysed without knowledge of  
 3 group assignment. Baseline comparisons made. No loss to follow-up . Limitations- small sample size, Allocation concealment not reported.  
 4 (b) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.

1        **Additional data:**

2        **Ornish 1998[202]:**

3        There was significantly more cardiac hospitalisation in the control group (44/20)  
4        compared to intervention group (23/28) at 5 years ( $p < 0.001$ ). Cardiac  
5        hospitalisation included hospitalisation for MI, PTCA and CABG.

6

7        **16.2.6 Nurse led cardiac rehabilitation**

8

9        The “Review Protocol” for this topic can be found in Appendix C, the “Search  
10        Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
11        E1, the “Clinical Evidence Tables” in Appendix E2, and the “Forest Plots” in Appendix  
12        F.

13        There was one paper comparing nurse led cardiac rehabilitation with routine care for  
14        stable angina[190].

15

1 **Table 16.11: Nurse led cardiac rehabilitation vs. routine care for stable angina**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Nurse led cardiac rehab	routine care (6 months)	Relative (95% CI)	Absolute	
<b>Walking performance (Jenkins activity checklist for walking) (follow-up 6 months; range of scores: -; better indicated by more) (b)</b>											
Jiang 2007[190]	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	83	84	-	MD 2.01 higher (1.23 to 2.79 higher)	⊕⊕⊕○ MODERATE

- 2 (a) This is a relatively short term study of patients (n=167). Very little information is given about whether investigators were "blinded" to patients' allocation to intervention or  
 3 control group. Most of the outcomes measured in the study were not relevant to the review question for which this study was included. No description of routine care was  
 4 given or even if it included advice on diet, exercise and smoking cessation.
- 5 (b) Jenkins Activity check list used: There were 16 activities on the scale, ranging from walking from bed to bathroom to walking 6.5 km. Subjects were required to indicate  
 6 whether they had performed each activity in the previous 24 hour period. For scoring, the number of 'yes' responses was summed to provide an activity total score, ranging  
 7 from 0 to 16.

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**16.2.7 Angina management programme (AMP)**

The “Review Protocol” for this topic can be found in Appendix C, the “Search Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix E1, the “Clinical Evidence Tables” in Appendix E2, and the “Forest Plots” in Appendix F.

There was one paper comparing the Angina management programme (AMP) with waiting list control for stable angina[203].

**Intervention:** For the Angina Management Programme (AMP) patients attended the hospital for two mornings per week for eight weeks. The AMP included the following elements: Exercise - consisted of 10 movements designed to improve general fitness and flexibility. Number of repetitions increased as patients felt fitter up until "somewhat hard"; Psychological elements of the programme included : Stress management - using relaxation, breathing re-training, bio-feedback, yoga exercises and behaviour modification; Psychological status - a self help rehab programme designed to reverse beliefs known to predict poor psychological recovery from MI; Behavioural change - help to return to appropriate but abandoned activities using goal setting and pacing; and education - Patients received extensive information about CAD, secondary prevention and angina.

1 **Table 16.12: AMP vs. control for stable angina**

Quality assessment							Summary of findings				Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		
							Angina management programme (AMP)	control (at the end of 8 week treatment period)	Relative (95% CI)	Absolute	
<b>Mean no. of Episodes of angina per week (follow-up 8 weeks; range of scores: -; better indicated by less)</b>											
Lewin 1995[203]	randomised trial	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	34	31	-	MD 12.1 lower (18.65 to 5.55 lower)	⊕⊕⊕⊕ HIGH
<b>Severity of angina (self rated out of 100 with scores being worse) (follow-up 8 weeks; range of scores: -; better indicated by less)</b>											
Lewin 1995[203]	randomised trial	no serious limitations	no serious inconsistency	no serious indirectness	no serious imprecision	none	34	31	-	MD 11.7 lower (23.04 to 0.36 lower)	⊕⊕⊕⊕ HIGH
<b>Duration of angina (mins) (follow-up 8 weeks; range of scores: -; better indicated by less)</b>											
Lewin 1995[203]	randomised trial	no serious limitations	no serious inconsistency	no serious indirectness	no serious imprecision	none	34	31	-	MD 9.7 lower (25.8 lower to 6.4 higher)	⊕⊕⊕⊕ HIGH
<b>Disability (Sickness Impact Profile) (100 being completely medically dependent and 0 indicating no measurable impairment) (follow-up 8 weeks; range of scores: -; better indicated by less)</b>											
Lewin 1995[203]	randomised trial	no serious limitations	no serious inconsistency	no serious indirectness	no serious imprecision	none	34	31	-	MD 12.7 lower (17.71 to 7.69 lower)	⊕⊕⊕⊕ HIGH

2 (a) *Randomised. Allocation concealment reported. For investigator measured outcome such as the exercise tolerance test, results were analysed by a doctor not otherwise*  
 3 *involved in the trial and blinded to occasion and group. 5/39(13%) in treatment group and 7/38(18%) in the control group lost to follow-up. This paper reports*  
 4 *summary results of 5 small (n=16) trials which took place over 2 years. Each trial was exactly the same design. In total n=77 patients were randomised to the Angina*  
 5 *Management Programme (AMP) or to Waiting List Controls (WLC) for 8 weeks. After 8 weeks of being in the WLC group patients went on to the AMP for 8 weeks.*  
 6 *Further assessments were carried out for all patients at 4 months and 1 year. However, at the latter two time points all patients had had treatment with AMP. Therefore, the*  
 7 *only relevant results are for the initial 8 week controlled phase of the study. That is, there was no long term control group.*

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2 **16.2.8 Angina Plan**

3 The “Review Protocol” for this topic can be found in Appendix C, the “Search  
4 Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
5 E1, the “Clinical Evidence Tables” in Appendix E2, and the “Forest Plots” in Appendix  
6 F.

7 There were 2 papers comparing the Angina Plan with education only for stable  
8 angina[189,204].

9 ‘The Angina Plan’ consisted of a 70-page, patient-held ‘work-book’ and an audio-  
10 taped relaxation programme which was introduced to the patient during a 30 to 40-  
11 minute structured interview. Before commencing, the nurse asked the patient to  
12 complete a questionnaire designed to establish if he or she had any of the common  
13 misconceptions about angina. Any misconceptions were discussed with the patient to  
14 correct their understanding of the illness and to explain how such beliefs can lead to  
15 undue invalidism. The nurse then worked with the patient to identify all of his or her  
16 personal risk factors for coronary heart disease in the normal manner.

17 A method of gradually and systematically reducing these and increasing activity  
18 levels, ‘goal setting and pacing’ that we have developed in previous research with  
19 angina patients, was used to negotiate gradual return to abandoned activities or to  
20 increase the patients’ capacity for that activity. The same method was used to  
21 introduce lifestyle change; improved diet and walking.

22 Patients were asked to practice relaxation, using the audio cassette, for 20 minutes  
23 each day. The nurse contacted the patient with a brief phone call at the end of weeks  
24 1, 4, 8, and 12. Any success with the goals the patients had set was rewarded with  
25 praise and encouragement and they were asked if they wished to extend the goal.

26 The Plan also contained written information about the role of frightening thoughts and  
27 misconceptions in triggering adrenaline release and anxiety and how this can result in  
28 poor coping strategies (such as the ‘over activity-rest cycle’), as well as an  
29 explanation of the symptoms of hyperventilation and panic. Standard advice on risk  
30 factors, medication, and what to do in the event of a suspected heart attack were  
31 also included.

32 Educational sessions: The nurse identified the patients’ risk factors for coronary heart  
33 disease from the research clinic measurements and a personal history and discussed  
34 ways in which each of them could be reduced. Patients were invited to ask questions  
35 about each risk factor and about angina or heart disease in general. They were also  
36 encouraged to discuss how it had affected their lives. Any questions they had were  
37 answered in an honest and factual manner by the nurse. If she did not know the  
38 answer at the time then she found it later and telephoned or wrote to them.

39 Every patient was given a package of written information, designed for people with  
40 coronary heart disease and angina and produced by authoritative sources, including  
41 the British Heart Foundation, the Chest Heart and Stroke Association, and the Family  
42 Heart Association.

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2 **Zetta 2009[189]:**

3 N=218 (n=109- standard care) (n=109 Angina Plan)

4 Angina Plan – During a 45 minute in-hospital consultation the AP nurse completed an  
5 assessment and initiated the AP intervention, which was then facilitated over the next  
6 12 weeks. The patients' cardiac misconceptions were identified using the brief  
7 questionnaire within the AP pack at the start of the consultation to allow the nurse to  
8 proactively target and correct these misconceptions. Individual cardiovascular risk was  
9 assessed and advice on risk factor modification given. Participants received the AP,  
10 which included a patient-held 'work-book' and an audio taped relaxation and  
11 information programme. The work-book provided information on angina and its  
12 management, cardiovascular risk, relaxation, exercise and goal setting and pacing  
13 techniques. Over the following 12 weeks a method of 'goal setting and pacing' based  
14 on the principles of CBT was used by the AP facilitator introduce lifestyle changes and  
15 support recovery during telephone follow-up at weeks 1,4, 8 and 12 for all  
16 participants in the AP group.

17 Standard care – A minimal intervention by nurses during their admission which  
18 identified patients risk factors , provided advice on their condition and risk factor  
19 reduction where possible depending on staff workload and skill mix.

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**Table 16.13: Angina Plan vs. education session for stable angina**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Angina Plan	Education session (6 months) (change scores)	Relative (95% CI)	Absolute	
<b>Anxiety (HAD scale) (follow-up 6 months; better indicated by lower values) (f)</b>											
Lewin 2002[204], Zetta 2009[189]	randomised trials	no serious limitations (a)	serious inconsistency (b)	no serious indirectness	no serious imprecision	None	177	183	-	MD 0.16 lower (0.39 lower to 0.06 higher)	⊕⊕⊕○ MODERATE
<b>Depression (HAD scale) (follow-up 6 months; better indicated by lower values)</b>											
Lewin 2002[204], Zetta 2009[189]	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	177	183	-	MD 0.86 lower (1.07 to 0.66 lower)	⊕⊕⊕⊕ HIGH
<b>Angina attacks per week (angina diary- self reported) (follow-up 6 months; better indicated by lower values)</b>											
Lewin 2002[204]	randomised trials	no serious limitations (c)	no serious inconsistency	no serious indirectness	no serious imprecision	None	68	74	-	MD 2.57 lower (4.46 to 0.68 lower)	⊕⊕⊕⊕ HIGH
<b>Mean pain score (follow-up 6 months; better indicated by lower values)</b>											
Lewin 2002[204]	randomised trials	no serious limitations (c)	no serious inconsistency	no serious indirectness	no serious imprecision	None	68	74	-	MD 1.79 higher (3.5 lower to 7.08 higher)	⊕⊕⊕⊕ HIGH
<b>Mean duration of pain (follow-up 6 months; better indicated by lower values)</b>											
Lewin 2002[204]	randomised trials	no serious limitations (c)	no serious inconsistency	no serious indirectness	no serious imprecision	None	68	74	-	MD 2.43 lower (12.23 lower to 7.37 higher)	⊕⊕⊕⊕ HIGH
<b>Physical limitation (Seattle Angina questionnaire)(follow-up 6 months; better indicated by higher values) (g)</b>											
Lewin 2002[204]	randomised trials	no serious limitations (c)	no serious inconsistency	no serious indirectness	no serious imprecision	None	68	74	-	MD 9.85 higher (4.84 to 14.86 higher)	⊕⊕⊕⊕ HIGH
<b>Angina stability (Seattle Angina questionnaire) (follow-up 6 months; better indicated by higher values)</b>											
Lewin 2002[204]	randomised trials	no serious limitations (c)	no serious inconsistency	no serious indirectness	no serious imprecision	None	68	74	-	MD 4.56 higher (5.56 lower to 14.68 higher)	⊕⊕⊕⊕ HIGH
<b>Angina frequency (Seattle Angina questionnaire) (follow-up 6 months; better indicated by higher values)</b>											
Lewin 2002[204], Zetta 2009[189]	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	177	183	-	MD 3.78 higher (1.82 lower to 9.39 higher)	⊕⊕⊕⊕ HIGH
<b>Treatment satisfaction (Seattle Angina questionnaire) (follow-up 6 months; better indicated by higher values)</b>											
Lewin 2002[204]	randomised trials	no serious limitations (c)	no serious inconsistency	no serious indirectness	no serious imprecision	None	68	74	-	MD 1.94 lower (6.99 lower to 3.11 higher)	⊕⊕⊕⊕ HIGH

<b>Disease perception (Seattle Angina questionnaire) (follow-up 6 months; better indicated by higher values)</b>											
Lewin 2002[204], Zetta 2009[189]	randomised trials	no serious limitations (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	177	183	-	MD 2.86 higher (1.24 lower to 6.96 higher)	⊕⊕⊕⊕ HIGH
<b>Misconceptions/knowledge (follow-up 6 months; better indicated by lower values) (h)</b>											
Zetta 2009[189]	randomised trials	no serious limitations (d)	no serious inconsistency	no serious indirectness	no serious imprecision	None	109	109	-	MD 5.50 lower (7.39 to 3.61 lower)	⊕⊕⊕⊕ HIGH
<b>CLASP angina (follow-up 6 months; better indicated by lower values) (i)</b>											
Zetta 2009[189]	randomised trials	no serious limitations (d)	no serious inconsistency	no serious indirectness	no serious imprecision	None	109	109	-	MD 0.80 higher (0.01 lower to 1.61 higher)	⊕⊕⊕⊕ HIGH
<b>Physical limitation (SF-36) (follow-up 6 months; better indicated by higher values) (k)</b>											
Zetta 2009[189]	randomised trials	no serious limitations (d)	no serious inconsistency	no serious indirectness	no serious imprecision	None	109	109	-	MD 3.67 higher (2.31 lower to 9.65 higher)	⊕⊕⊕⊕ HIGH
<b>Energy and vitality (SF- 36) (follow-up 6 months; better indicated by higher values)</b>											
Zetta 2009[189]	randomised trials	no serious limitations (d)	no serious inconsistency	no serious indirectness	no serious imprecision	None	109	109	-	MD 4.52 higher (1.02 lower to 10.06 higher)	⊕⊕⊕⊕ HIGH
<b>Pain (SF-36) (follow-up 6 months; better indicated by higher values)</b>											
Zetta 2009[189]	randomised trials	no serious limitations (d)	no serious inconsistency	no serious indirectness	no serious imprecision	None	109	109	-	MD 11.87 higher (4.04 to 19.7 higher)	⊕⊕⊕⊕ HIGH
<b>GH perception (SF-36) (follow-up 6 months; better indicated by higher values)</b>											
Zetta 2009[189]	randomised trials	no serious limitations (d)	no serious inconsistency	no serious indirectness	no serious imprecision	None	109	109	-	MD 5.03 higher (0.12 to 9.94 higher)	⊕⊕⊕⊕ HIGH
<b>Change in health (SF-36) (follow-up 6 months; better indicated by higher values)</b>											
Zetta 2009[189]	randomised trials	no serious limitations (d)	no serious inconsistency	no serious indirectness	no serious imprecision	None	109	109	-	MD 5.25 higher (2.52 lower to 13.02 higher)	⊕⊕⊕⊕ HIGH
<b>SE1 QOL- DW QOL score (follow-up 6 months; better indicated by higher values) (e)</b>											
Zetta 2009[189]	randomised trials	no serious limitations (d)	no serious inconsistency	no serious indirectness	no serious imprecision	None	109	109	-	MD 1.70 higher (2.5 lower to 5.9 higher)	⊕⊕⊕⊕ HIGH

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(a) Randomised (Lewin 2002)[204]. Allocation concealment reported. Baseline and follow-up measures were collected, scored, and entered into the computer by research staff blinded to group allocation. 5/68 (7%) in the Angina Plan group and 7/74 (9%) in the Education Programme group lost to follow-up. The data were analysed by a medical statistician not otherwise involved in the research. The study had 80% power to detect a difference of 0.5 units on the Hospital Anxiety and Depression Scale. . However, the study acknowledges that the mean reduction in anxiety and depression is slight, even though for some patients it was profound. Follow up was 6 months so the study was not capable of determining if the observed benefits continue beyond this time. In Zetta 2009[189] random allocations were computer generated, allocated to permuted fixed blocks of 20 and stratified for site. The researcher was blinded to group allocation throughout the trial. ITT reported.

- 1 (b) 12 = 71%
- 2 (c) Randomised. Allocation concealment reported. Baseline and follow-up measures were collected, scored, and entered into the computer by research staff blinded to group
- 3 allocation. 5/68 (7%) in the Angina Plan group and 7/74 (9%) in the Education Programme group lost to follow-up. The data were analysed by a medical statistician not
- 4 otherwise involved in the research.. The study had 80% power to detect a difference of 0.5 units on the Hospital Anxiety and Depression Scale. However, the study
- 5 acknowledges that the mean reduction in anxiety and depression is slight, even though for some patients it was profound. Follow up was 6 months so the study was not
- 6 capable of determining if the observed benefits continue beyond this time.
- 7 (d) Random allocations were computer generated, allocated to permuted fixed blocks of 20 and stratified for site. The researcher was blinded to group allocation
- 8 throughout the trial. ITT reported.
- 9 (e) SEIQoL-DW (Schedule for the Evaluation of Individual Quality of Life-Direct weighting) is an interview based tool specifically designed for the assessment of individual
- 10 quality of life. Using the SEIQoL-DW participants define five areas that comprise individual 'quality of life'. These items are rated in terms of level of importance. An
- 11 overall score ranging from 0-100 is then calculated with higher scores reflecting better quality of life. The SEIQoL-DW is totally subjective and patient centred and
- 12 provides a relatively unique measure of quality of life.
- 13 (f) Hospital Anxiety and Depression scale (HADS): 14 item tool with 2, seven item subscales to measure anxiety and depression within a non psychiatric population. A score
- 14 from 0 to 3 for each item generated a total score (range 0 to 21 for each sub scale. Scores between 8 and 10 indicate borderline presence of anxiety or depression and
- 15 those above suggest that these states may be present.
- 16 (g) The Seattle Angina Questionnaire is a disease specific health related quality of life measure comprised of a 19 item questionnaire measuring five dimensions of coronary
- 17 artery disease: physical limitation, angina stability, anginal frequency, treatment satisfaction and disease perception. Each dimension is scored separately on a 0-100 scale
- 18 with higher scores indicating better functioning.
- 19 (h) Knowledge and misconceptions were assessed using the 14 item York Angina Beliefs Questionnaire. This uses a Likert scale response format ranging 'strongly agree' to
- 20 'strongly disagree'. Items targeted the cause, physiology and coping with angina. Summation and transformation of the item scores generated a scale total ranging from 0-
- 21 56 with higher numbers indicating more misconceptions.
- 22 (i) The Cardiovascular Limitations and Symptoms Profile (CLASP) measures nine physical and functional dimensions, including four symptom subscales (angina, shortness of
- 23 breath, tiredness, ankle swelling) and five subscales focusing on functional limitations (mobility, social life and leisure activities, activities within the home, concerns and
- 24 worries, sexual activity). Each of the nine subscales is scored separately to calculate a specific measure of impairment.
- 25 (j) The Short Form 36 Health Survey (SF-36) is a 36 item questionnaire assessing general health and QoL. The 8 dimensions of SF-36 (physical functioning, role limitations
- 26 caused by emotional problems, bodily pain, social functioning, mental health, role limitations caused by emotional problems, vitality-energy/fatigue and general health
- 27 perception) generates scores on each dimension between 0 and 100, with higher scores representing better health status.

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**16.3 Economic evidence**

Two studies were included, one comparing stent angioplasty with exercise training[192] and one comparing health education with control[205]. These are summarised in the economic evidence profile below. See also Economic Evidence Tables in Appendix G.

**Table 16.14: PCI vs. exercise - Economic study characteristics**

Study	Limitations	Applicability	Other Comments
Hambrech 2004[192]	Potentially serious limitations (a)	Partial applicability (b)	Based on a RCT included in our review (see 16.2.1)

- g) Limited follow-up (1 year). A breakdown of the cost items was not reported. A sensitivity analysis was not conducted. The study received an unconditional grant from Aventis.
- h) Study conducted in Germany. Effectiveness was not reported in terms of QALYs.

**Table 16.15: PCI vs. exercise - Economic summary of findings**

Study	Incremental cost (£)	Incremental effects	ICER	Uncertainty
Hambrech 2004[192]	1,502 (a)	- (b)	- (c)	No sensitivity analyses were performed.

- (d) 2003 GBP; cost of interventions including hospital charges, expenses for supervised training sessions, bicycle ergometer, coronary angiographies, and rehospitalisation. P value <0.001
- (e) Outcomes considered were deaths of cardiac causes, cerebrovascular accidents, and revascularisation. None of them was statistically significant.
- (f) An overall summary of cost-effectiveness was provided only in the text but the details of the effectiveness measure were not reported anywhere. To gain one CCS class the cost was £4,396 in the PCI group and £2,167 in the exercise group.

**Table 16.16: Health education vs. control - Economic study characteristics**

Study	Limitations	Applicability	Other Comments
O'Neill 1996[205]	Potentially serious limitations (a)	Partial applicability (b)	Based on a RCT included in our review (see 16.2.1). Funded by the Medical Research Council.

- (a) Not all the important outcomes were evaluated (e.g. angina symptoms, MI).
- (b) Relatively old study; medical treatment might have not been optimal at that time. Unclear what the control group received. Effectiveness was not reported in terms of QALYs.

**Table 16.17: Health education vs. control - Economic summary of findings**

Study	Incremental cost (£)	Incremental effects (deaths saved)	ICER	Uncertainty
O'Neill 1996[205]	39 (a)	4.6% (b)	NR	No sensitivity analyses were performed.

- (a) 1996 GBP; cost of intervention (staff time and travel-related costs), drugs, GP visits, hospital visits, tests and other treatments. Community care costs were excluded. Difference in costs was not statistically significant.
- (b) Not statistically significant.

## 1 16.4 Evidence statements

### Clinical A. Exercise programmes:

#### Intensive exercise programme vs. control

**Todd 1990[193]:** Evidence from one RCT shows that the differences between exercise group and control group at time of follow up were not significant for max ST depression (mm) [MD 0.2 higher (-0.43 to 0.83)], time to 1 mm ST depression (sec) [MD 166 higher (-221.71 to 553.71)] and treadmill time (sec) [MD 262 higher (-66.64 to 590.64)]. [1 year follow-up]

#### Exercise plus placebo vs. placebo

**Malmberg 1974[191]:** Evidence from one small scale pilot RCT shows that the differences in proportion of change in maximal working capacity kpm/min [MD 4 lower (-43.5 to 35.5)], angina attacks per week [MD 25 lower (-82.38 to 32.38)], and nitro-glycerine tablet intake per week [MD 4 higher (-96.75 to 104.75)] were not significantly different in an exercise group compared to a placebo group. [follow-up 4 months]

#### Exercise plus BB and vs. BB

**Malmberg 1974[191]:** Evidence from one small scale pilot RCT shows that the differences in proportion of change in maximal working capacity kpm/min [MD 6 lower (55.60 lower to 43.60 higher)], angina attacks per week [MD 41 higher (1.93 to 83.93 higher)] and nitro-glycerine tablet intake per week [MD 58 higher (37.02 lower to 153.02 higher)] were not significantly different in an exercise group compared to a placebo group. [follow-up 4 months] [moderate quality]

#### Exercise plus low fat diet vs. control

**Schuler 1992[194]:** Evidence from one RCT shows that cardiac mortality [RR 5.09 (0.25 to 103.66)], total mortality [RR 2.04 (0.19 to 21.82)] and non-fatal MI [RR 0.2 (0.01 to 4.15)] did not significantly differ between the an exercise + low fat diet compared to a control group [ follow-up 12 months]

#### Exercise programme vs. PCI

**Hambrecht 2004[192]:** Evidence from one RCT shows that there was no significant differences between Exercise group and PCI group for cerebrovascular accidents [RR 0.65 (0.11 to 3.75)], hospitalisation and also no significant difference in coronary angiography owing to worsening angina [RR 0.14 (0.02 to 1.1)] there were no deaths of

cardiac causes in either group [0/51 in Exercise group and 0/50 in PCI group]. There was a significantly higher proportion of patients needing revascularisation in the PCI group compared to the exercise group [RR 0.29 (0.09 to 1.01)]. [1 year follow-up].

## **B. Health Education**

### **Health education vs. control – mortality and frequency of exercise**

**Cupples 1994[195]:** Evidence from one RCT shows that total mortality was significantly lower in the health education group compared to control group [RR 0.45 (0.24 to 0.86)]. Significantly more patients in the health education group increased their frequency of exercise compared to the control group [RR 1.73 (1.32 to 2.28)]. [Follow-up 2 years]

### **Health education vs. control for stable angina – Quality of life**

**O'Neill 1996[196]:** Evidence from one RCT shows that the mean differences in overall Nottingham Health Profile (NHP) scores did not reach statistical significance ( $p=0.0659$ ) Mean differences of two NHP subscales reached statistical significance. Physical Mobility [MD intervention -1.49 and MD control -6.19,  $p=0.0015$ ] and Social Isolation [MD intervention +1.42 and MD control -3.01,  $p=0.0408$ ] with better self ratings associated with the intervention group. [follow-up 2 years ]

## **C. Stress management**

### **Stress management vs. routine care control**

**Bundy 1998[200]:** Evidence from one RCT shows that there was no significant difference between stress management and routine care control for frequency of angina [(MD 0.00 (-2.92 to 2.92)] and average duration of angina attack (mins) [MD -0.40 (-4.70 to 3.90)] [follow-up 8 weeks]

**Gallacher 1997[198]:** Evidence from one RCT shows that there was no significant difference between stress management and control, group for frequency of chest pain at rest (days per fortnight) [MD -0.59 (-1.24 to 0.06)] and frequency of chest pain on exertion (days per fortnight) [MD -0.54 (-1.35 to 0.27)] [Follow-up 6 months]

### **Stress management + exercise vs. routine care control**

**Bundy 1998 [200]:** Evidence from one RCT shows that there was no

significant difference between stress management along with exercise compared to routine care control for frequency of angina (average no. of daily attacks) [MD 0.6 (-2.97 to 4.17)] and duration of angina (min) [MD -4.4 (-9.08 to 0.28)] [finalscores]. [8 weeks follow-up]

**Bundy 1994 [199]:** Evidence from one RCT shows that there was no significant difference between stress management along with exercise compared to routine care control for frequency of angina (average no. of daily attacks) [MD -2.70 (-5.98 to 0.58)] . Duration of angina (min) was significantly lower in the stress management group compared to routine care control [MD -0.70 (-1.06 to -0.34)] [change scores]. [8 weeks follow-up]

#### **D. Yoga lifestyle**

##### **Yoga lifestyle vs. control**

**Manchanda 2000[201]:** Evidence from one RCT shows that there was no mortality in either the Yoga life style intervention programme and control group [0/21 in intervention and 0/21 in control group]. There was significantly fewer angina episodes per week in the Yoga intervention group compared to control group MD -3.3 (-4.82 to -1.782). There was no significant difference between yoga life style and control group for exercise duration (sec) [MD 39 (-46.78 to 124.78)]. ST-Segment depression was significantly lower in the Yoga Lifestyle group compared to control [MD -2.52 (-2.95 to -2.09)]. Revascularisation was significantly lower in the Yoga lifestyle compared to control group [RR 0.12 (0.02 to 0.91)] [1 year follow-up]

#### **E. Intensive lifestyle**

##### **Intensive style vs. control**

**Ornish 1998[202]:** Evidence from one RCT shows that there no significant difference between intensive lifestyle programme and control for angina frequency (times per week) [MD 0.7 (-0.9 to 2.3)], chest pain duration (min) [MD -0.1 (-1.64 to 1.44)] , MI [RR 0.36 (0.07 to 1.76)], CABG [RR 0.29 (0.06 to 1.33)] and death [RR 1.43 (0.14 to 14.7)] . There was significantly lower PTCA in the lifestyle programme compared to control [RR 0.41 (0.21 to 0.78)] [5 years follow-up]

## **F. Nurse led cardiac rehabilitation**

### **Nurse led cardiac rehab vs. routine care**

**Jiang 2007[190]:** Evidence from one RCT shows that 'walking performance' [measured using Jenkins Activity check list ] was significantly higher in the Nurse cardiac rehab group compared to control [MD 2.01 (1.23 to 2.79)] [6 months follow-up]

## **G. Angina management Programme (AMP)**

### **AMP vs. control**

**Lewin 1995[203]:** Evidence from one RCT shows that significantly fewer mean no. of episodes of angina per week in the AMP group compared to control [MD -12.1 (-18.65 to -5.55)], severity of angina was significantly lower in the AMP group compared to control [MD -11.7 (-23.04 to -0.36)], there was no significant difference between AMP and control group for duration of angina (mins) [MD -9.7 (-25.8 to 6.4)] and disability [measured by Sickness Impact Profile] was significantly lower in the AMP group compared to control [MD -12.7 (-17.71 to -7.69)] [follow-up – at the end of 8 weeks treatment period]

## **H. Angina Plan**

### **Angina Plan vs. education session**

**Lewin 2002[204]:** Evidence from one RCT shows that there was significantly greater reduction angina attacks per week (from angina diary of patients) [MD -2.57 (-4.46 to -0.68)], physical limitation (Seattle Angina Questionnaire) [MD 9.85 (4.84 to 14.86)], in the Angina Plan group compared to standard care/education session control group. There was no significant difference between angina plan and standard care/education session for mean duration of pain [MD -2.43 (-12.23 to 7.37)], mean pain score [MD 1.79 (-3.5 to 7.08)] , Angina stability (Seattle Angina Questionnaire) [MD 4.56 (-5.56 to 14.68)] treatment satisfaction (Seattle Angina Questionnaire) [MD -1.94 (-6.99 to 3.11)] [6 months follow-up]

**Lewin 2002[204]; Zetta 2009[189]:** Evidence from 2 RCTs shows that depression (HAD scale) was found to be significantly reduced in the

Angina Plan group compared to standard care group/education session [MD -0.86 (-1.07 to -0.66 )]. There was no significant between the Angina Plan and standard care/education session group for Anxiety (HAD scale) [MD 0.16 lower (0.39 lower to 0.06 higher)], angina frequency (Seattle Angina Questionnaire) [MD 3.78 higher (-1.82 to 9.39 )] , disease perception (Seattle Angina Questionnaire) [MD 3.51 (-1.64 to 8.66)] [6 months follow-up]

**Zetta 2009[189]:** Evidence from one RCT shows that significantly more patients in the Angina Plan group reported increased knowledge and less misconceptions compared to standard care/education session group [MD -5.50 (-7.39 to -3.61 lower)] and significant improved General Health perception (SF-36) [MD 5.03 (0.12 to 9.94)] in angina plan group compared to standard care/education session group . There was no significant difference between angina plan and standard care/education session group for CLASP angina [MD 0.80 (-0.01 to 1.61)], Physical function (SF-36) [MD 3.67 (-2.31 to 9.65)], energy and vitality (SF-36) [MD 4.52 (-1.02 to 10.06)], Pain (SF-36) [MD 11.87 (4.04 to 19.70)], change in health (SF-36) [MD 5.25 (-2.52 to 13.02)], SEI QoL-DW QoL Score [MD 1.70 (-2.50 to 5.90)]. [6 months follow-up]

**Economic** Exercise training reduces costs compared to PCI while health education does not generate additional costs compared to control. This evidence has potentially serious limitations and partial applicability.

## 1 16.5 Recommendations and link to evidence

<b>Recommendation</b>	<p><b>Assess the person's need for lifestyle advice (for example about exercise, stopping smoking, diet and weight control) and psychological support, and offer interventions as necessary.</b></p> <p><b>Address personal issues including:</b></p> <ul style="list-style-type: none"> <li>• <b>self-management skills such as pacing activities and goal setting</b></li> <li>• <b>dealing with stress or depression</b></li> <li>• <b>advice about physical exertion including sexual activity.</b></li> </ul>
<b>Relative values of different outcomes</b>	<p>The GDG were interested in whether cardiac rehabilitation programmes would influence mortality and morbidity outcomes as well as quality of life. The GDG recognised that intermediate outcomes such as change in diet and exercise may indicate potential benefit but considered that harder outcomes were required if they</p>

were to recommend rehabilitation outcomes to the NHS as standard treatment for people with stable angina.

**Trade off between clinical benefits and harms**

**Economic considerations**

Exercise training and health education could improve outcomes without creating additional costs.

**Quality of evidence**

The quality and quantity of evidence for comprehensive programmes of cardiac rehabilitation was not adequate to suggest these could be recommended for people with stable angina. In particular the number of patients included in the studies was small and the length of follow up was extremely short. Yoga/lifestyle programme, angina management programme and angina plan did result in a reduction in angina frequency.

In the yoga lifestyle programme[201] at one year, the yoga groups showed significant reduction in number of anginal episodes per week, improved exercise capacity and decrease in body weight, revascularisation procedures (coronary angioplasty or bypass surgery) were also less frequently required in the yoga group (one vs. eight patients RR 5.45 p=0.01). The study had a very small size (n=42) and also the follow-up of 1 year was not sufficient to determine if the observed benefits would continue at a longer follow-up. Also by nature of the interventions the study could not be blinded, and hence a placebo effect of yoga interventions cannot be excluded. Further, the study did not look at differential effects of yogic exercises, dietary control and aerobic exercises, and the study considered yoga lifestyle modification as a composite incorporating all the above mentioned components.

The study by Lewin 1995[203] on the Angina management programme showed that significantly fewer mean number of episodes of angina per week compared to control , severity of angina compared to control . However the study sample was very small (n=65 patients who completed the study) and the follow-up period of 8 weeks was very short to determine if the programme would sustain its effect at a longer follow-up.

The study by Cupples 1994[195] showed that the education programme reduced angina episodes per week and increased frequency of exercise in people with angina. The study also reports that percentage of patients who took drugs prophylactically increased

significantly by the end of the study, which could have caused a reduction in symptoms. Further, the study did not validate the patients reporting of their frequency of exercise and some reporting bias may have occurred. Nevertheless, the study was large (n=688) and well conducted.

The study by Lewin 2002[204] on the self-management programme- Angina Plan was a small (n=142), well conducted study. Most of the patients who received Angina Plan reported a reduction of three episodes of angina per week; this is clinically worthwhile reduction of nearly 50% from the baseline mean of seven episodes per week. The authors propose that increased activity levels and daily walking may have raised the Angina Plan patients' threshold for exercise induced pain. There was significant reduction in anxiety and depression; this reduction was slight, even though for some patients it was profound. Follow up was 6 months, so the study was not capable of determining if the observed benefits continue beyond this time.

The study by Zetta (2009)[189] recruited patients who were admitted to medical admission or coronary care units and were considered by the GDG not to be representative of people with chronic stable angina.

The economic evidence has potentially serious limitations and partial applicability.

### **Other considerations**

The GDG considered that the term cardiac rehabilitation and the traditional 4 phases of cardiac rehabilitation were not necessarily helpful in the context of people with stable angina. The GDG did not consider that the evidence indicated benefit for patients from comprehensive cardiac rehabilitation programmes. The evidence did not support any particular model of care for delivering individual interventions that patients might benefit from.

The self management programme (Angina plan) includes a brief, cognitive-behavioural programme comprising a 76-page patient-held workbook (contains information about risk factor reduction, stress management, angina management and how to use goal setting and pacing to increase activity safely), a tape or CD based relaxation programme, an advice tape to introduce the concepts in the Angina Plan to the patient before they see the facilitator, and a misconceptions questionnaire. The Angina Plan is introduced to the patient (and their partner) in an interview lasting thirty or forty minutes,

and followed up by four, ten to fifteen minute appointments or phone calls over three months.

The GDG considered that the components of the Angina plan were beneficial to people with stable angina but the evidence was not adequate to recommend the programme based on a small study sample with a short follow-up.

People with angina are likely to need a variety of interventions geared to understanding and coping with their diagnosis and helping them to engage in activities for secondary prevention. The GDG preferred the idea of a menu of health needs that may need to be addressed and patients should be directed to services they individually require. It is the GDG opinion that a tailored approach is cost-effective (i.e. offer only the rehabilitation components that are required rather than a comprehensive programme).

The GDG considered that many of the aspects of care that would be of benefit to people with stable angina are available via primary care and via services such as National Exercise Referral Scheme in Wales[206].

## 1 16.6 Research recommendation

2 The GDG recommended the following research question:

3 ➤ **Research question:** Is an 8-week, comprehensive, multidisciplinary, cardiac  
4 rehabilitation service more clinically and cost effective for managing stable  
5 angina than current clinical practice?

6 ➤ **Why this is important:** Cardiac rehabilitation programmes are an established  
7 treatment strategy for certain heart conditions, such as for people who have  
8 had a heart attack. However, there is no evidence to suggest that cardiac  
9 rehabilitation is clinically or cost effective for managing stable angina.  
10 Research to date has looked at short-term outcomes, such as a change in diet  
11 or exercise levels, but the effect on morbidity and mortality has not been  
12 studied. A randomised controlled trial is required to compare comprehensive  
13 cardiac rehabilitation with standard care in people with stable angina, with  
14 measures of angina severity (exercise capacity, angina frequency, use of a  
15 short-acting nitrate), and long-term morbidity and mortality as endpoints.

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## 2 **17 Lifestyle Adjustments**

### 3 **17.1 Introduction**

4 Lifestyle interventions such as exercise are known to have a positive effect on  
5 cardiovascular health. The GDG were interested in whether there were specific lifestyle  
6 interventions that would reduce mortality and morbidity in people with stable angina.

7 The aim of our evidence review was to look at programmes which modify lifestyle/CVD risk  
8 factors specifically for angina patients. The following lifestyle factors were considered for  
9 this review:

- 10 • Diet (including folic acid, vitamin E, C, beta carotene supplements, Omega 3-acid  
11 ethyl esters, Mediterranean diet, low saturated diet, low glycaemic diet, fruit and  
12 vegetables, fish diet)
- 13 • Physical activity

14 A total of 5 papers (3 RCTs and 2 cross over trials) have been included in this review. Three  
15 papers (2 RCTs and one cross over trial) evaluated the effectiveness of fish oil  
16 diet/capsules and two papers (one RCT and one cross over trial) evaluated the  
17 effectiveness of Vitamin E in people with stable angina. However we did not identify any  
18 papers looking at the following interventions in people with stable angina: Folic acid,  
19 Vitamin C, beta carotene supplements, Mediterranean diet, low saturated fat diet, and low  
20 glycaemic diet.

21 There was significant overlap between review of lifestyle interventions and review of  
22 rehabilitation programmes. The evidence relating to the effect of exercise primarily came  
23 from supervised programmes and these are therefore reported in the chapter on  
24 rehabilitation (chapter 16).

### 25 **17.2 Fish oils**

#### 26 **17.2.1 Clinical question**

27 What is the clinical /cost effectiveness of fish oils for reducing symptoms, morbidity,  
28 mortality and improving quality of life in stable angina patients?

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2 **17.2.2 Clinical evidence**

3 The “Review Protocol” for this topic can be found in Appendix C, the “Search  
4 Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
5 E1, the “Clinical Evidence Tables” in Appendix E2, and the “Forest Plots” in Appendix  
6 F.

7 There were 3 studies evaluating the effectiveness of fish oil. One RCT[207] and one  
8 cross over trial[208] (analysed as a parallel RCT) evaluated the effectiveness of fish  
9 oil capsules compared to placebo and one RCT[209] evaluated the effectiveness of  
10 both dietary fish advice and fish oil capsules compared to fruit advice and sensible  
11 eating.

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1 **Table 17.1: Fish oil capsules vs. placebo for stable angina (Follow-up at end of treatment period)**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Fish oil capsules	Placebo (Follow-up at end of treatment period)	Relative (95% CI)	Absolute	
<b>Anginal episodes per week (better indicated by lower values) (Follow-up at the end of 12 weeks treatment period)</b>											
Salachas 1994[207]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious imprecision (c)	none	20	19	-	MD 3 lower (54.01 lower to 48.01 higher)	⊕⊕OO LOW
<b>GTN consumption per week (better indicated by lower values) (Follow-up at the end of 12 weeks treatment period)</b>											
Salachas 1994[207]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious imprecision (c)	none	20	19	-	MD 1.99 lower (10.69 lower to 6.71 higher)	⊕⊕OO LOW
<b>Exercise test duration (min) (better indicated by higher values) (Follow-up at the end of 12 weeks treatment period)</b>											
Salachas 1994[207]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious imprecision (c)	none	20	19	-	MD 0.99 higher (2.01 lower to 3.99 higher)	⊕⊕OO LOW
<b>Number of anginal attacks per 30 days (better indicated by lower values) (Follow-up at the end of 12 weeks treatment period)</b>											
Aucamp 1993[208]	randomised trials	very serious (b)	no serious inconsistency	no serious indirectness	serious imprecision (c)	none	12	11	-	MD 9.2 lower (29.15 lower to 10.75 higher)	⊕⊕OO VERY LOW
<b>Duration of angina attacks per minute (better indicated by lower values) (Follow-up at the end of 12 weeks treatment period)</b>											
Aucamp 1993[208]	randomised trials	very serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	none	12	11	-	MD 0.4 lower (0.95 lower to 0.15 higher)	⊕⊕OO LOW
<b>Intensity of pain per attack per patient (on a 10 cm visual analogue scale) (better indicated by lower values) (Follow-up at the end of 12 weeks treatment period)</b>											
Aucamp 1993[208]	randomised trials	very serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	none	12	11	-	MD 1 lower (2.12 lower to 0.12 higher)	⊕⊕OO LOW
<b>No. of sublingual isosorbide dinitrate tablets taken per 30 days (better indicated by lower values) (Follow-up at the end of 12 weeks treatment period)</b>											
Aucamp 1993[208]	randomised trials	very serious (b)	no serious inconsistency	no serious indirectness	serious imprecision (c)	none	12	11	-	MD 0 higher (16.14 lower to 16.14 higher)	⊕⊕OO VERY LOW

- 2 (a) Randomised. Double blind. Allocation concealment not reported. Numbers lost to follow-up not reported. No ITT reported. Baseline comparison between groups not made.
- 3 (b) Placebo controlled cross-over trial. Single blind. 23 patients completed the trial: 11 patients taking placebo in phase 1 (group A) and 12 patients taking the active fish oil in
- 4 phase 1 (group B). Very little baseline characteristics reported. No ITT reported. No method of randomisation and allocation concealment reported. Very poorly reported
- 5 trial.
- 6 (c) 95% CI includes no effect and the upper and lower CI crosses the MID
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**Table 17.2: Fish advice (dietary fish advice + fish oil capsule) vs. fruit advice for stable angina (Follow-up after 3 to 9 yrs)**

Quality assessment							Summary of findings				Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		
							Fish advice (dietary fish advice + fish oil capsule)	Fruit advice (Follow-up 6 months after entering the trial)	Relative (95% CI)	Absolute	
<b>All death (Follow-up after 3 to 9 years)</b>											
Burr 2003[209]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious imprecision (b)	None	141/764 (18.5%)	133/779 (17.1%)	RR 1.08 (0.87 to 1.34)	14 more per 1000 (from 22 fewer to 58 more)	⊕⊕○○ LOW
<b>Cardiac death (Follow-up after 3 to 9 years)</b>											
Burr 2003[209]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious imprecision (b)	None	94/764 (12.3%)	72/779 (9.2%)	RR 1.33 (1 to 1.78)	31 more per 1000 (from 0 more to 72 more)	⊕⊕○○ LOW
<b>Sudden death (Follow-up after 3 to 9 years)</b>											
Burr 2003[209]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious imprecision (b)	None	42/764 (5.5%)	30/779 (3.9%)	RR 1.43 (0.9 to 2.26)	17 more per 1000 (from 4 fewer to 49 more)	⊕⊕○○ LOW

3 (a) Randomised. Baseline characteristics reported, Loss to follow-up not reported. ITT not reported. Allocation concealment not reported.  
4 (b) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.  
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1 **Table 17.3: Fish advice (dietary fish advice + fish oil capsule) vs. fish +fruit advice for stable angina (Follow-up after 3 to 9 yrs)**

Quality assessment							Summary of findings				Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		
							Fish advice (dietary fish advice + fish oil capsule) (Follow-up 6 months after entering the trial)	Fish +Fruit advice	Relative (95% CI)	Absolute	
<b>All death (Follow-up after 3 to 9 years)</b>											
Burr 2003[209]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious imprecision (b)	None	141/764 (18.5%)	142/807 (17.6%)	RR 1.05 (0.85 to 1.3)	9 more per 1000 (from 26 fewer to 53 more)	⊕⊕○○ LOW
<b>Cardiac death (Follow-up after 3 to 9 years)</b>											
Burr 2003[209]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious imprecision (b)	None	94/764 (12.3%)	86/807 (10.7%)	RR 1.15 (0.88 to 1.52)	16 more per 1000 (from 13 fewer to 55 more)	⊕⊕○○ LOW
<b>Sudden death (Follow-up after 3 to 9 years)</b>											
Burr 2003[209]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious imprecision (b)	None	42/764 (5.5%)	31/807 (3.8%)	RR 1.43 (0.91 to 2.25)	17 more per 1000 (from 3 fewer to 48 more)	⊕⊕○○ LOW

(a) Randomised. Baseline characteristics reported, Loss to follow-up not reported. ITT not reported. Allocation concealment not reported.

(b) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.

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1 **Table 17.4: Fish advice (dietary fish advice + fish oil capsule) vs. sensible eating (non -specific advice) for stable angina (Follow-up after 3 to 9 yrs)**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Fish advice (dietary fish advice + fish oil capsule)	Sensible eating (non -specific advice) (Follow-up 6 months after entering the trial)	Relative (95% CI)	Absolute	
<b>All deaths (Follow-up after 3 to 9 years)</b>											
Burr 2003[209]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious imprecision (b)	None	141/764 (18.5%)	109/764 (14.3%)	RR 1.29 (1.03 to 1.63)	41 more per 1000 (from 4 more to 90 more)	⊕⊕○○ LOW
<b>Cardiac death (Follow-up after 3 to 9 years)</b>											
Burr 2003[209]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious imprecision (b)	None	94/764 (12.3%)	67/764 (8.8%)	RR 1.4 (1.04 to 1.89)	35 more per 1000 (from 4 more to 78 more)	⊕⊕○○ LOW
<b>Sudden death (Follow-up after 3 to 9 years)</b>											
Burr 2003[209]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	None	42/764 (5.5%)	17/764 (2.2%)	RR 2.47 (1.42 to 4.3)	33 more per 1000 (from 9 more to 73 more)	⊕⊕⊕○ MODERATE

2 (a) Randomised. Baseline characteristics reported, Loss to follow-up not reported. ITT not reported. Allocation concealment not reported.  
 3 (b) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.

In order to attempt to explain the unexpected excess mortality associated with fish advice, ad hoc subgroup analyses were carried out by the study authors. The apparently adverse effect of fish advice was confined to the second phase of the trial (data not shown), when a much higher proportion of participants were given fish capsules than in the first phase. During this phase some of the participants in the fish advice group were sub randomised to receive fish oil capsules, so survival analysis was carried out to examine the effect on those sub randomised to capsules rather than to dietary fish advice.

**Table 17.5: Survival analysis of subjects advised on dietary fish or fish oil**

Outcome	Dietary fish (n=1109)	Fish oil capsules (n=462)
All death	n=198 (HR 1.13 (0.94 to 1.37) p=0.20	n=85 (HR 1.19 (0.92 to 1.54) p=0.19
Cardiac death	n=121 (HR 1.20 (0.93 to 1.53) p=0.16	n=59 (HR 1.45 (1.05 to 1.99) p=0.02
Sudden death	n=49 (HR 1.43 (0.95 to 2.15) p=0.08	n=24 (HR 1.84 (1.11 to 3.05); p=0.01

\*hazard ratios adjusted for age, smoking, previous MI, history of high blood pressure, diabetes, BMI, serum cholesterol, medication and fruit advice.

The hazard ratios for each mortality category were higher in the fish oil capsules than in the dietary fish group. The possibility was considered that dietary fish or fish oil could adversely interact with drugs commonly given for heart disease. Hazard ratios of cardiac deaths were calculated in relation to fish advice, with subjects classified in to those receiving and those not receiving various types of drugs at recruitment in to the trial. No evidence was found of any adverse interactions; treatment with BB showed a significant favourable interaction with fish advice.

### 17.2.3 Economic evidence

No economic studies were retrieved on this question.

### 17.2.4 Evidence statement

#### Clinical Fish oil capsule vs. placebo

**Salachas 1994[207]:** Evidence from one RCT shows that there was no significant difference between Fish oil capsules and placebo for number of anginal attacks per week [MD -3.00 [-54.01 to 48.01], GTN

consumption per week [MD -1.99 [-10.69 to 6.71] and exercise duration (min) [MD 0.99 [-2.01 to 3.99]. [Follow-up end of 12 weeks treatment period]

**Aucamp 1992[208]:** Evidence from one RCT shows that there was no significant difference between fish oil capsules and placebo for number of anginal attacks per 30 days [MD -9.20 [-29.15 to 10.75], duration of angina attacks per minute [MD -0.40 [-0.95 to 0.15], intensity of pain per attack per patient (on a 10 cm visual analogue scale)[MD -1.00 [-2.12 to 0.12], no. of sublingual isosorbide dinitrate tablets taken per 30 days [MD 0.00 [-16.14 to 16.14] [Follow-up at end of 12 weeks treatment period]

#### **Fish advice (dietary fish+ fish oil capsule) vs. fruit advice**

**Burr 2003[209]:** Evidence from one RCT shows that there was significantly higher cardiac death in the fish advice group [RR 1.33 [1.00 to 1.78] compared to fruit advice group; and there was no significant difference between fish advice and fruit advice group for all death [RR 1.08 [0.87 to 1.34] and sudden death [RR 1.43 [0.90 to 2.26] [Follow-up after 3 to 9 yrs]

#### **Fish advice (dietary fish+ fish oil capsule) vs. fish +fruit advice**

**Burr 2003[209]:** Evidence from one RCT shows that there was no significant difference between fish advice and fish+fruit advice for all death [RR 1.05 [0.85 to 1.30], cardiac death [RR 1.15 [0.88 to 1.52] and sudden death [RR 1.43 [0.91 to 2.25] [Follow-up after 3 to 9 yrs]

#### **Fish advice (dietary fish+ fish oil capsule) vs. sensible eating (non - specific advice)**

**Burr 2003[209]:** Evidence from one RCT shows that there was significantly lower all death [RR 1.29 [1.03 to 1.63], cardiac death [RR 1.40 [1.04 to 1.89] and sudden death [RR 2.47 [1.42 to 4.30] in the sensible eating group compared to fish advice group [Follow-up after 3 to 9 yrs]

**Economic** No economic studies were retrieved on this question.

1 **17.2.5 Recommendations and link to evidence**

<b>Recommendation</b>	<b>Do not offer fish oils to treat stable angina. Inform people that there is no evidence that they help stable angina.</b>
<b>Relative values of different outcomes</b>	<p>The outcomes considered as important during the development of the review protocol for lifestyle adjustments included exercise tolerance, mortality, angina frequency/severity, major cardiac events, hospitalisation, revascularisation, QoL.</p> <p>Evidence showed that there was no significant improvement in angina and exercise duration with the use of fish oil capsules.</p> <p>There was no improvement in outcomes when fish oil capsules (short term use) were compared with placebo.</p> <p>However fish oil capsules (long term use) when compared to fruit advice showed statistically significantly increased cardiac death and when compared to sensible eating showed statistically significantly higher all death, cardiac death and sudden death.</p>
<b>Trade off between clinical benefits and harms</b>	<p>There is no evidence of clinical benefits arising from the use of fish oils in stable angina patients and some evidence of harm when compared to advice on sensible eating</p>
<b>Economic considerations</b>	<p>The use of fish oils would generate costs without improving outcomes.</p>
<b>Quality of evidence</b>	<p>The evidence for outcomes was of moderate quality except for the cross over trial where evidence was low quality.</p>
<b>Other considerations</b>	

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3 **17.3 Vitamin E**

4 **17.3.1 Clinical question**

5 What is the clinical /cost effectiveness of Vitamin E for reducing symptoms, morbidity,  
6 mortality and improving quality of life in stable angina patients?

7

1    **17.3.2    Clinical evidence**

2           The “Review Protocol” for this topic can be found in Appendix C, the “Search  
3           Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
4           E1, the “Clinical Evidence Tables” in Appendix E2, and the “Forest Plots” in Appendix  
5           F.

6           There were 2 studies (one RCT and one cross over trial) evaluating the effectiveness  
7           of Vitamin E compared to placebo[210,211].

8

1 **Table 17.6: Vitamin E vs. placebo for stable angina (Follow-up at the end of treatment period)**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Vitamin E	Placebo (Follow-up at the end of treatment period)	Relative (95% CI)	Absolute	
<b>Improved anginal symptoms (Follow-up at the end of 9 week treatment period)</b>											
Anderson 1974[210]	randomised trials	very serious (a)	no serious inconsistency	no serious indirectness	serious imprecision (c)	none	5/18 (27.8%)	5/18 (27.8%)	RR 1 (0.35 to 2.87)	0 fewer per 1000 (from 181 fewer to 519 more)	⊕⊕○○ VERY LOW
<b>No change in anginal symptoms (Follow-up at the end of 9 week treatment period)</b>											
Anderson 1974[210]	randomised trials	very serious (a)	no serious inconsistency	no serious indirectness	serious imprecision (c)	none	13/18 (72.2%)	12/18 (66.7%)	RR 1.08 (0.7 to 1.67)	53 more per 1000 (from 200 fewer to 447 more)	⊕⊕○○ VERY LOW
<b>Slightly worse anginal symptoms (Follow-up at the end of 9 week treatment period)</b>											
Anderson 1974[210]	randomised trials	very serious (a)	no serious inconsistency	no serious indirectness	serious imprecision (c)	none	0/18 (0%)	1/18 (5.6%)	RR 0.33 (0.01 to 7.68)	37 fewer per 1000 (from 55 fewer to 371 more)	⊕⊕○○ VERY LOW
<b>Duration treadmill (min) (better indicated by higher values) (Follow-up end of 6 months treatment period)</b>											
Gillilan 1977[211]	randomised trials	serious (b)	no serious inconsistency	no serious indirectness	serious imprecision (d)	none	48	40	-	MD 0.18 higher (0.51 lower to 0.87 higher)	⊕⊕⊕○ LOW
<b>Angina attacks per week (better indicated by lower values) (Follow-up end of 6 months treatment period)</b>											
Gillilan 1977[211]	randomised trials	serious (b)	no serious inconsistency	no serious indirectness	serious imprecision (d)	none	48	48	-	MD 0.6 higher (4.04 lower to 5.24 higher)	⊕⊕⊕○ LOW
<b>Nitroglycerin consumption per week (better indicated by lower values) (Follow-up end of 6 months treatment period)</b>											
Gillilan 1977[211]	randomised trials	serious (b)	no serious inconsistency	no serious indirectness	serious imprecision (d)	none	48	48	-	MD 0.1 lower (5.38 lower to 5.18 higher)	⊕⊕⊕○ LOW

- (a) Randomised. Double blind. 33/40 completed 9 full weeks of records. In 5 cases (3 vitamin and 2 placebo) only 8 weeks of records could be used because one record card was incomplete or missing, in one (vitamin group) only 7 weeks of records were available, and one other patient (vitamin) withdrew from the study after 7 weeks because of persistent diarrhoea. allocation concealment not reported. Randomisation was not carried out properly, patients randomised after giving the intervention. Baseline characteristics not well reported. Only subjective data available. Blinding process unclear. ITT not reported.
- (b) Double blind cross over study. Blinding of outcome assessors. Baseline comparison between groups not reported. Method of randomisation and allocation concealment not reported. No ITT reported.
- (c) 95% CI around the pooled estimate of effect includes both: 1) no effect and 2) appreciable benefit or appreciable harm.
- (d) 95% CI includes no effect and the upper and lower CI crosses the MID.

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1           **Additional data from two studies :**

2           **Anderson 1974[210] (Vitamin E capsules vs. placebo)**

- 3           • **Nitroglycerin consumption:** Mean nitroglycerin consumption was higher in the  
4           vitamin E group from the start, and increased from 18.7 to 23.5 between the  
5           first and last weeks. In the placebo group the mean intake was 10.9 tablets in  
6           the first week and this declined to 6.4 in the last week (Standard deviation not  
7           reported). The authors report that these differences were largely due to one  
8           or two patients in each group who had a large initial intake and showed  
9           great variation. Thus the increase in the Vitamin E group was attributed  
10          entirely due to one patient, whose consumption of NTG averaged 180 tablets  
11          per week-more than that of the entire placebo group. Most of the patients in  
12          each group showed little change in NTG consumption during the trial.
- 13          • **Pain score:** The net pain score for the placebo group was lower than that for  
14          the vitamin group in 7 out of the 9 weeks. Comparing the last and first weeks,  
15          the overall mean change in score was -0.81 for the vitamin group and +0.17  
16          for the placebo group (Standard deviation not reported).
- 17          • **Side effects:** There were no side effects with Vitamin E reported by the  
18          patients. Headache and constipation were reportedly two patients who  
19          proved to have been on placebo.

20

21          **Gillilan 1977[211] (Vitamin E capsules vs. placebo)**

- 22          • There were 4 deaths during the study, two of which occurred suddenly at  
23          home (apparently cardiac death) and two of which occurred during  
24          hospitalisation for recurrent MI (established at autopsy).
- 25          • No deleterious side effects were observed resulting from the use of Vitamin E  
26          during the study. There were slightly more complaints of mild gastrointestinal  
27          disturbances during placebo phase (6%) than during vitamin E phase (4%). No  
28          exacerbation of hypertension, congestive heart failure, or skeletal-muscular  
29          complaints could be attributed to vitamin E therapy.

30

31   **17.3.3       Economic evidence**

32           No economic studies were retrieved on this question.

33

34   **17.3.4       Evidence statements**

**Clinical           Vitamin E vs. placebo**

**Anderson 1974[210]:** Evidence from one RCT shows that there

was no significant difference between Vitamin E and placebo for Improved anginal symptoms [RR 1.00 [0.35 to 2.87], no change in anginal symptoms [RR 1.08 [0.70 to 1.67], slightly worse anginal symptoms [RR 0.33 [0.01 to 7.68] [Follow-up at the end of 9 weeks treatment period]

**Gillilan 1977[211]:** Evidence from one RCT shows that there was no significant difference Vitamin E and placebo for duration treadmill (min) [MD 0.18 [-0.51 to 0.87], angina attacks per week [MD 0.60 [-4.04 to 5.24], and nitroglycerin consumption per week [MD -0.10 [-5.38 to 5.18] [Follow-up at the end of 6 months treatment period]

**Economic** No economic studies were retrieved on this question.

1 17.3.5 Recommendations and link to evidence

<b>Recommendation</b>	<b>Do not offer vitamin supplements to treat stable angina. Inform people that there is no evidence that they help stable angina.</b>
<b>Relative values of different outcomes</b>	The outcomes considered as important during the development of the review protocol for lifestyle adjustments included exercise tolerance, mortality, angina frequency/severity, major cardiac events, hospitalisation, revascularisation, QoL.  Evidence showed that there was no significant difference between Vitamin E and placebo for any of the anginal or exercise test outcomes.
<b>Trade off between clinical benefits and harms</b>	There is no evidence of clinical benefits arising from the use of Vitamin E in stable angina patients.
<b>Economic considerations</b>	The use of Vitamin E would generate costs without improving outcomes.
<b>Quality of evidence</b>	The available studies had short follow-up and evidence for outcomes was of low to moderate quality (based on GRADE). No evidence for other vitamin supplements in the treatment of stable angina has been identified that met our inclusion criteria for reviewing.
<b>Other considerations</b>	The GDG considered the evidence on Vitamin E which did not show any benefit and patients should be informed of this. No evidence was found for other supplements. The GDG considered that although no evidence did not mean there might not be a potential

benefit, the lack of any evidence of benefit was something that both patients and healthcare practitioners should be aware of as supplements are a cost either to patients themselves or to the health service and there is no evidence of benefit.

1

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## 2 **18 Pain Interventions and Refractory angina**

### 3 **18.1 Introduction**

4 Stable angina presents as chest pain. Interventions are primarily related to  
5 addressing cardiac work by for example improving blood flow by medical treatment  
6 or revascularisation and by addressing the progression of underlying coronary artery  
7 disease. Chronic refractory angina has been defined as angina that cannot be  
8 controlled with optimal medical therapy and where revascularisation is  
9 unfeasible[20]. The decision as to when revascularisation is unfeasible is a decision  
10 made by interventional radiologists and cardiac surgeons. Revascularisation will also  
11 carry risks and an informed patient may decide that these risks outweigh possible  
12 benefits. The current UK national chronic refractory angina group's definition of  
13 chronic refractory angina is, "Chronic stable angina that persists despite optimal  
14 medication and when revascularisation is unfeasible or where the risks are unjustified.  
15 Interventions directed towards pain rather than towards coronary artery disease have  
16 been used for people with 'refractory' angina.

17 The GDG choose not to make a decision on a definition of refractory angina. They  
18 considered that different definitions and inclusion criteria might have been used in  
19 different studies and considered it more appropriate to examine evidence for use of  
20 pain interventions in as wide a population of people with angina as possible. The  
21 evidence review therefore describes the populations included in each study. The GDG  
22 were addressed by Professor Michael Chester and Dr. Austin Leach from the National  
23 Angina Refractory Centre who also advised on the interventions to include in the  
24 evidence review.

25 The following pain interventions have been included in the review:

- 26 • TENS (Transcutaneous electric nerve stimulation),
- 27 • EECPP (Enhanced external counter pulsation)
- 28 • Acupuncture
- 29 • Self-pain management programmes

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- 1 The evidence review includes 1 paper on TENS (Transcutaneous Electric Nerve
- 2 Stimulation), 3 papers on EECp (Enhanced external counter pulsation) and 3 papers on
- 3 Acupuncture, 2 papers on self-management of pain. No studies were identified
- 4 evaluating the effectiveness of opioids in the management of people with angina.
- 5

1 **Table 18.1: Pain interventions – Summary of evidence**

Study	Intervention	Comparison	Duration of intervention	Study design	No of participants	Follow-up	Outcomes
Manheimer 1985[212]	TENS	Control group did not receive TENS	10 weeks. Three TENS treatment sessions of at least 1 hr each per day (morning, noon and evening)	RCT	N=23 (n=12 TENS and n=11). Severe angina pectoris (duration 1 to 20 years, functional class III or IV, NYHA). The antianginal pharmacological treatment taken at the beginning of the study was regarded as optimal. All patients had been considered for aortocoronary bypass surgery: one patient had undergone such a operation, five were waiting for surgery, and the remaining were being considered for surgical treatment.	After 2 weeks	Maximal total work during exercise was determined as a product of workload in watts and time in mins (W.min); ST segment depression during and after exercise; pain and dyspnea reported by the patient during and after exercise.; frequency of anginal attacks and consumption of short acting nitroglycerin per week.
Arora 1999[213]	EECP	Inactive counterpulsation (CP)	35 hours of (once or twice/day) of active counterpulsation over a 4 to 7-week period.	RCT	N = 139 (n=EECP 72, n=67 inactive counterpulsation. Chronic stable angina- CCS I, II or III.  More than 70% of patients in each group had CCS class II or III and over 70%	3 days after follow-up for angina pain counts, one week after treatment for exercise duration.	Exercise test, Anginal pain counts, Nitroglycerin use.

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					of each group had undergone prior CABG or angioplasty.		
Arora 2002[213]	EECP	Inactive counterpulsation (CP).	35 hours of (once or twice/day) of active counterpulsation over a 4 to 7-week period.	RCT	N = 139 (n=EECP 72, n=inactive counterpulsation 67); n=71 (36 in EECP and n=35 inactive CP). Chronic stable angina - CCS I, II or III	At end of treatment and 1 year after treatment	Health related quality of life (HQOL)
Loh 2008[214]	EECP	No comparison	A standard course of 35 one hour treatment sessions. The patients received a mean of 33.3±9.6 hours of treatment over a mean period of 48 days.	Before-After study	N=1427, CCS I, I, III angina. Anginal status: [CCS class I: 2.2% CCS class II: 8.6% CCS class III: 62.8% CCS class IV: 26.4%].  88% had prior PCI or CABG and 88% were unsuitable for further coronary intervention.	3 years (median 37 months)	Anginal status (CCS class), weekly angina episode, nitroglycerin use, QOL (using a simple 5 point scale where 1 represents the worst and 5 represents the best QOL), clinical events (PCI, CABG, MI, death, MACE (composite of death/MI/CABG/PCI) and hospitalisation.
Ballegaard 1990[215]	Acupuncture	Sham acupuncture.	Ten (10) treatments in the supine position within 3 weeks	RCT	N=49 (n=24 in genuine acupuncture and n=25 sham acupuncture).  Clinically stable exercise induced angina pectoris for more than 6 months (2 or more anginal attacks per week). All patients on medical treatment.	Just after the treatment period	Exercise test; no. of anginal attacks; activity at the time of the pain; nitroglycerin consumption (diaries); daily well being on an ordinal scale; global evaluation of the effect of the treatment on an ordinal scale:

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Ballegaard 1986[216]	Acupuncture	Sham acupuncture.	Seven (7) treatments in the supine position for 3 weeks	RCT	N=26 (n=13 in active acupuncture and n=13 in sham acupuncture).  Stable, medically resistant, exercise provoked angina pectoris (functional class II-IV NYHA). The patients were selected among 56 consecutive patients with a positive evaluation with regard to aortocoronary bypass surgery	Immediately after the 9 week treatment period.	Exercise tests variables (Exercise tolerance, difference in pressure rate product between rest and maximum exercise, maximal PRP during exercise, maximum ST depression and length of time maximum ST depression); anginal attacks, activity at the time of the pain attack and nitroglycerin consumption (from diaries); subjective global evaluation by the patient at the end of the trial : improvement of general well-being after treatment /no improvement of general well-being after treatment.
Richter 1991[217]	Acupuncture	Tablet placebo.	The treatment was given 3 times per week during the 4 week period.	RCT (cross over trial)	N=21 (cross over). Patients with stable effort angina and at least five anginal attacks per week during the last 6 months, inspite of intensive antianginal treatment.  Bypass surgery had been performed in 8 patients, in two of them repeatedly, while 5 patients were still waiting for operation.	Immediately after the 4 treatment period	Exercise test, self rating quality of life questionnaire, no. of anginal attacks.
McGillion	Chronic Angina	Waiting list	The psycho education	RCT	n=130 were	3 months from	Health Related Quality of Life

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2008[218]	Self-Management Program (CASMP)	control (offered entry into the next available CASMP once post-test measures were completed).	programme given in two-hour sessions weekly, over a six-week period by a registered nurse using a group format.		randomised, n=66 to the CASMP and n=64 to the waiting list control group.  Chronic stable angina patients.	start of treatment	(HRQL) which included the SF-36 and the SAQ (Seattle Angina Questionnaire)
Payne 1994[219]	Pain management programme	standard medical care	The pain management programme administered over three consecutive weekly sessions (length of sessions not reported).	RCT	N =52 (N=26 pain management treatment and N=26 controls).  Episodes of chest pain or discomfort in the previous 4 weeks in patients with diagnosed coronary artery disease.	6 months.	Pain frequency and intensity; frequency of NTG usage; mood and psychological distress.

1

2 **18.2 Transcutaneous electric nerve stimulation (TENS)**

3 **18.2.1 Clinical question**

4 What is the clinical/cost effectiveness of TENS in people with stable angina?

5

6 **18.2.2 Clinical evidence**

7 The “Review Protocol” for this topic can be found in Appendix C, the “Search  
8 Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
9 E1, the “Clinical Evidence Tables” in Appendix E2, and the “Forest Plots” in Appendix  
10 F.

11 There was one RCT[212] evaluating the effectiveness of TENS in patients with severe  
12 angina pectoris.

13 The “Review Protocol” for this topic can be found in Appendix C, the “Search  
14 Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
15 E1, the “Clinical Evidence Tables” in Appendix E2, and the “Forest Plots” in Appendix  
16 F.

17

1 **Table 18.2: TENS vs. control (no TENS) for stable angina – Quality assessment & Summary of findings**

Quality assessment							Summary of findings				Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		
							TENS vs. control (no TENS) (Follow-up 2 weeks after treatment)	control	Relative (95% CI)	Absolute	
Exercise tolerance (W.min) (follow-up 2 weeks; better indicated by higher values)											
Mannheimer 1985[212]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	11	10	-	MD 9 lower (170.42 lower to 152.42 higher)	⊕⊕○○ LOW
ST segment depression (mm) during exercise (follow-up 2 weeks; better indicated by lower values)											
Mannheimer 1985[212]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	11	10	-	MD 0.2 lower (1.36 lower to 0.96 higher)	⊕⊕○○ LOW
ST segment depression (mm) after exercise (follow-up 2 weeks; better indicated by lower values)											
Mannheimer 1985[212]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	11	10	-	MD 0.2 higher (0.97 lower to 1.37 higher)	⊕⊕○○ LOW
Frequency of angina attacks per week (follow-up 2 weeks; better indicated by lower values)											
Mannheimer 1985[212]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	11	10	-	MD 4 lower (21.98 lower to 13.98 higher)	⊕⊕○○ LOW
Nitroglycerin consumption per week (follow-up 2 weeks; better indicated by lower values)											
Mannheimer 1985[212]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	11	10	-	MD 17 higher (9.31 lower to 43.31 higher)	⊕⊕○○ LOW

2 (a) Randomised. Blind outcome assessment (ST segment changes were measured blindly by two independent observers). Method of randomisation not reported. Allocation  
3 concealment not reported. Small sample size. Loss to follow-up not reported. ITT not reported. No blinding of participants (not possible due to the kind of intervention)  
4 (b) Upper and lower confidence limit crosses the minimal important difference.

1

2 **18.2.3 Economic evidence**

3 No relevant economic evaluations comparing TENS with any other intervention were  
4 identified.

5

6 **18.2.4 Evidence statement**

**Clinical TENS vs. control**

**Mannheimer 1985[212]:** Evidence from one RCT shows that there was no significant difference between TENS treatment and control group for exercise tolerance (W.min) (MD -9.00 [-170.42, 152.42]); ST segment depression (mm) during exercise (MD -0.20 [-1.36, 0.96]); ST segment depression after exercise (MD 0.20 [-0.97, 1.37]); frequency of angina attack per week (MD -4.00 [-21.98, 13.98]); and nitroglycerin consumption per week (MD 17.00 [-9.31, 43.31]) [follow-up 2 weeks after treatment]

**Economic** No economic evidence was found.

7 **18.2.5 Recommendations and link to evidence**

<b>Recommendation</b>	<b>Do not offer transcutaneous electric nerve stimulation (TENS) to manage stable angina.</b>
Relative values of different outcomes	The outcomes considered as important during the development of the review protocol for pain interventions included: improvement in anginal symptoms (angina frequency and nitroglycerin consumption), mortality, exercise tolerance, major cardiac events, hospitalisation, revascularisation, QoL and adverse events. Three of these outcomes are included in the evidence identified on TENS. These include frequency of anginal attacks, exercise tolerance and nitroglycerin consumption. This evidence demonstrates that TENS is not clinically effective with respect to any of these three outcomes.
<b>Trade off between clinical benefits and harms</b>	There is no evidence of clinical benefits arising from the use of TENS in stable angina patients.
<b>Economic considerations</b>	No published health-economic evaluation of TENS was identified. The intervention is not cost-effective as it is associated with costs to the NHS without being effective

at improving the outcomes considered.

**Quality of evidence**

The available evidence was of low quality as assessed by GRADE with a very small sample size (n=23) and the follow-up period was too short (2 weeks) to detect any sustainable improvement in outcomes.

**Other considerations**

The GDG considered that current evidence base is weak and shows no effectiveness of TENS. TENS should not be used unless new evidence emerges that demonstrates TENS's clinical and cost-effectiveness in people with stable angina.

1

2 **18.3 Enhanced external counterpulsation (EECP)**

3 **18.3.1 Clinical question**

4 What is the clinical/cost effectiveness of EECP in people with stable angina?

5

6 **18.3.2 Clinical evidence**

7 The "Review Protocol" for this topic can be found in Appendix C, the "Search  
8 Strategies" in Appendix D, the "List of Included and Excluded Studies" in Appendix  
9 E1, the "Clinical Evidence Tables" in Appendix E2, and the "Forest Plots" in Appendix  
10 F.

11 There were 3 papers (1 RCT, one sub-study of the RCT and one Before-After study)  
12 evaluating the effectiveness of EECP in patients with chronic stable angina and  
13 refractory angina.

1 **Table 18.3: EECp vs. inactive CP for stable angina**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							EECP	inactive CP (Follow-up 3 days after treatment for angina pain counts, one week after treatment for exercise duration)	Relative (95% CI)	Absolute	
Exercise duration (sec) (change scores) (follow-up after 1 week) (follow-up 1 weeks; better indicated by higher values)											
Arora 1999[213]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious imprecision (d)	none	57	58	-	MD 16 higher (15.86 lower to 47.86 higher)	⊕⊕⊕ LOW
Time to >1mm ST segment depression (Sec) (change scores) (follow-up after 1 week) (follow-up 1 weeks; better indicated by higher values)											
Arora 1999[213]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	56	56	-	MD 41 higher (9.13 to 72.87 higher)	⊕⊕⊕ MODERATE
Angina episodes/day (change scores) (follow-up after 3 days) (follow-up 3 days; better indicated by lower values)											
Arora 1999[213]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious imprecision (d)	none	71	66	-	MD 0.24 lower (0.83 lower to 0.35 higher)	⊕⊕⊕ LOW
NTG use/day (change scores) (follow-up after 3 days) (follow-up 3 days; better indicated by lower values)											
Arora 1999[213]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	71	66	-	MD 0.22 lower (0.55 lower to 0.11 higher)	⊕⊕⊕ MODERATE
Adverse events (no. of patients) (up to the end of treatment) (c)											
Arora 1999[213]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	39/71 (54.9%)	17/66 (25.8%)	RR 2.13 (1.35 to 3.38)	291 more per 1000 (from 90 more to 613 more)	⊕⊕⊕ LOW

2 (a) Multicentre randomised study. Baseline characteristics reported. The EECp group and inactive CP group were not balanced at baseline, the patients in the EECp group had  
 3 significantly longer duration of angina and higher proportion of patients with previous MI. Allocation concealment reported. 2 /139 withdrew prior to first treatment. 1/66  
 4 in inactive CP and 12/71 in EECp lost to follow-up [more drop out from the EECp than the control group] . No data reported on long term outcomes especially cardiac

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- 1 mortality. Completed trial: N = 124: EECP, n= 59; Inactive CP ,n=65. ITT analysis used (but not for all outcomes). ITT was not reported for ST segment depression and  
2 exercise duration. This may overestimate the treatment effect. Data not well reported. Very short follow-up  
3 (b) Less than 300 events  
4 (c) The adverse experiences (device related) were: Paresthesia, edema, swelling, skin abrasion, bruise, blister, pain (legs, back). The adverse experiences (non device related)  
5 were: viral syndrome, anxiety, dizziness, tinnitus, GI disturbance, headache, blood pressure change, epistaxis, angina, other chest pain, A/V arrhythmia, heart rate change,  
6 respiratory.  
7 (d) 95% CI includes no effect and the upper and lower CI crosses the MID.

1 **ARORA 2002[220] (MUST EECP trial; EECP vs. control)**

2 This is a sub-study of the (MUST EECP trial assessing HQOL [Health related quality of  
3 life] at one year follow-up).

4 **Population:** N=139 (n=EECP 72, n=inactive counterpulsation n= 67) all male. Data  
5 was available only for n=71 (36 in EECP and n=35 inactive CP) [Hence there is a  
6 high risk that this sample is not representative of the study population]

7 **Outcome:** Health related quality of life (HQOL). Four primary outcomes were used in  
8 the analysis: the physical functioning, bodily pain and social functioning subscales of  
9 the SF-36, and QOL score. The 36 item Short-Form Health Survey (SF-36) and the  
10 cardiac version of the Quality of Life Index (QIL) used for measuring HQOL. The SF-  
11 36 comprises 36 items that yield 8 multi item scales that measure physical functioning,  
12 work role disability due to emotional problems, bodily pain, general health  
13 perceptions, vitality, social functioning, work role disability due to emotional  
14 problems, mental health, and a single item evaluation of change in health. The QIL is  
15 in 2 parts: Part 1 measures satisfaction with various aspects of life as they are  
16 impacted by the respondent's cardiac health. Part 2: Measures the importance of  
17 these same aspects of life to the respondent personally.

18 **Results:**

- 19 A. Baseline to end of treatment: Both EECP and inactive CP groups reported  
20 significant improvements in physical functioning, bodily pain, and cardiac specific  
21 health and functioning from baseline to end of treatment. The size of the  
22 improvement in HQOL parameters was always larger for the EECP than for  
23 inactive CP; however, this difference was only statistically significant for one of  
24 the four primary parameters: social functioning. Those in the EECP group reported  
25 a substantially greater increase in their abilities to participate in social activities  
26 with family and friends than did those in the inactive CP, who, on average,  
27 reported a decrease in social activity. [Values not reported]
- 28 B. Baseline to 1 year follow-up: At 1 year follow-up, the EECP group maintained  
29 statistically significant improvements in HQOL across all primary HQOL  
30 parameters, where as the inactive CP group only maintained a significant  
31 improvement in the physical functioning scale. At 1 year follow-up, improvements  
32 for the EECP group were significantly greater than those for the inactive CP group  
33 on 3 of 4 primary parameters: bodily pain, social functioning, and cardiac  
34 specific health and functioning [no values reported]

35  
36 **Loh 2008[214] (International EECP Patient Registry [IPER]):**

37 This is a Before-After study. This study is the 3 year follow-up of the patients in the  
38 International EECP Patient Registry (IEPR)

39 **Population:** N=1427. Five thousand patients from 99 American and 9 international  
40 centres were enrolled between Jan 1998 and July 2001. Consecutive patients from  
41 each centre who had at least 1 hour of EECP treatment were enrolled. The mean age

1 was 66.3±10.8 years and 72% were men.76% had multivessel coronary  
 2 disease.88% had prior PCI or CABG and 88% were unsuitable for further coronary  
 3 intervention. The Anginal status of patients was: CCS class I: 2.2%; CCS class II: 8.6%;  
 4 CCS class III: 62.8%; CCS class IV: 26.4%.

5 **Intervention:** EECp. A standard course of 35 one hour treatment sessions was  
 6 recommended. The patients received a mean of 33.3±9.6 hours of treatment over a  
 7 mean period of 48 days.

8 **Follow-up:** 3 years (median 37 months)

9 **Outcome:** The primary outcome measure was Anginal status (CCS class). The other  
 10 outcomes were weekly angina episode, nitroglycerin use , QOL (using a simple 5  
 11 point scale where 1 represents the worst and 5 represents the best QOL), clinical  
 12 events (PCI, CABG, MI, death, MACE (composite of death/MI/CABG/PCI) and  
 13 hospitalisation.

14 **Results:** Immediately post EECp, the proportion of patients who suffered from CCS  
 15 Class III/IV angina reduced from 89.2% to 24.9%, p<0.001. The CCS class improved  
 16 by at least 1 class in 77.9% of the patients and by 2 classes in 38%. 16.3% of  
 17 patients had no angina. These were sustained in 74% patients whose anginal status  
 18 was documented at 3 year follow-up. At 3 years, 36.4% of the patients had class II  
 19 or milder angina. The Cumulative 3 year repeat EECp and major cardiovascular event  
 20 rates: (Percentage (95% CI)) was: Repeat EECp: 22.5% (20.1% -24.9%); PCI: 16.4%  
 21 (14.3% -18.5%); CABG: 7.5% (6%-9%); MI: 11.8% (10%-13.7%); Death- 17%  
 22 (14.9%-19.1%); MACE: 40.8% (38.8%-43.5%). Of the patients who responded to  
 23 the QOL questionnaires there was sustained improvement in their QOL after 3 years,  
 24 p<0.001.(results reported graphically).

25

26 **18.3.3 Economic evidence**

27 One study[221] was included. This is summarised in the economic evidence profile below.  
 28 See also Economic Evidence Tables in Appendix G.

29

30 **Table 18.4: EECp vs. no treatment- Economic study characteristics**

Study	Limitations	Applicability	Other Comments
McKenna 2009[221]	Potentially serious limitations (a)	Direct applicability	Decision model based on the MUST-EECP trial, included in the review of clinical effectiveness.

31

32

33

34

35

a) The analysis was based on limited data (one small RCT). Utilities were obtained from an algorithm converting SF-36 to EQ-5D. Durability of benefits obtained from expert opinion. The model does not consider: the effect of the intervention on mortality or myocardial infarction, the cost of escalating medical treatment over time, costs associated with no intervention.

1 **Table 18.5: EECP vs. no treatment - Economic summary of findings**

Study	Incremental cost (£)	Incremental effects (QALYs)	ICER	Uncertainty
McKenna 2009[221]	4,750 (a)	0.255 (b)	£18,643/QALY	One-way sensitivity analysis: results were sensitive to the probability of sustaining QoL benefits over time and to the cost of EECP. Results were not sensitive to the rate of repeat EECP within two years or to the discount rates used. Worst/best case scenario: if QoL benefits from EECP are only sustained in the first year, the ICER was £63,000. If QoL benefits are sustained over a lifetime, the ICER becomes £5,830. Monte-Carlo simulation: EECP was cost-effective in 44.4% of the simulations.

2 (a) 2008 GBP. Costs included were capital cost of EECP machine, equipment replacement costs,  
 3 consumables, staffing costs, overheads, repeat operations. Cost of no treatment was assumed to be null.  
 4 Cost data were obtained from personal communication and price list of supplier.

5 (b) Quality of life improvements were calculated as EQ-5D scores using an algorithm to convert the SF-36  
 6 scores from the study into EQ-5D. Utilities after one year were estimated with expert elicitation  
 7 techniques (frequency chart).

8  
 9  
 10 **18.3.4 Evidence statements**

11 **Clinical EECP vs. inactive CP**

**Arora 1999[213] (MUST EECP trial):** Evidence from one RCT shows that time to >1 mm ST segment depression (sec) increased significantly in the EECP compared to inactive CP (MD 41.00 [9.13, 72.87]). Adverse events were significantly higher in the EECP group compared to inactive CP (RR 2.13 (1.35 to 3.38)). There was no significant difference between EECP and inactive CP for exercise duration (sec) (MD 16.00 [-15.86, 47.86]); angina episodes/day (MD -0.24 [-0.83, 0.35]) ; NTG use/day (MD -0.22 [-0.55, 0.11]) [follow-up 3 days after treatment for angina pain counts, one week after treatment for exercise duration].

**EECP vs. control**

**Arora 2002[220] (MUST EECP trial):** Evidence from one RCT shows that both EECP and inactive CP groups reported statistically significant improvements in physical functioning, bodily pain, and cardiac specific health and functioning from baseline to end of treatment. At 1 year

follow-up, the EECp group maintained statistically significant improvements in HQOL across all primary HQOL parameters, where as the inactive CP group only maintained a statistically significant improvement in the physical functioning scale. At 12 month follow-up, improvements in HQOL for the EECp were significantly greater than those for the inactive CP group on three of four primary parameters (SF-36 scale): bodily pain, social functioning and cardiac specific health and functioning but not physical functioning. [Follow-up 12 months]

**EECP Patient Registry (no comparison group)**

**Loh 2008[214] (International EECp Patient Registry [IPER])** : Evidence from one Before-After study shows that immediately post EECp, the proportion of patients who suffered from CCS Class III/IV angina reduced from 89.2% to 24.9%,  $p < 0.001$ . The CCS class improved by at least 1 class in 77.9% of the patients and by 2 classes in 38%. 16.3% of patients had no angina. These were sustained in 74% patients whose anginal status was documented at 3 year follow-up. Immediately post EECp, 76% of the patients experienced at least 50% reduction in frequency of angina. This was sustained at 3 year follow-up. Of the patients who responded to the QOL questionnaires there was sustained improvement in their QOL after 3 years,  $p < 0.001$  (no values reported) [follow-up 3 years]

**Economic** The cost-effectiveness of EECp is very uncertain depending on the sustained effectiveness of the intervention. This evidence is directly applicable but it has potentially serious limitations.

1 **18.3.5 Recommendations and link to evidence**

<b>Recommendation</b>	<b>Do not offer enhanced external counterpulsation (EECP) to manage stable angina.</b>
<b>Relative values of different outcomes</b>	<p>The outcomes considered as important during the development of the review protocol for pain interventions included: improvement in anginal symptoms (angina frequency and nitroglycerin consumption), mortality, exercise tolerance, major cardiac events, hospitalisation, revascularisation, QoL and adverse events.</p> <p>The RCT evidence from the MUST EECp trial showed that there was statistically significant improvement in one exercise test variable i.e. time to ST depression in the EECp group when compared to the control group (one week follow-up period). However the GDG did not consider this improvement as clinically significant. Furthermore there were more adverse events in the EECp</p>

group when compared to the control group over the 7 week treatment period.

The registry study (International EECF Patient Registry (IPER)) showed significant improvement in CCS angina class after 3 years. However after 3 years there was repeat EECF in 22.5% of patients, PCI in 16.4% patients, CABG in 7.5% of patients and death in 17% of patients.

**Trade off between clinical benefits and harms**

Adverse events were significantly higher in the EECF group when compared to the control group over the 7 week treatment period.

**Economic considerations**

There is high uncertainty over the cost-effectiveness of EECF in people with stable angina mainly due to the unknown long-term benefits of the intervention.

**Quality of evidence**

The available evidence on EECF is weak. It is based on one relatively small RCT (MUST EECF trial) and one poor quality registry study (International EECF Patient Registry (IPER)). Also there was no evidence available on the long-term safety of EECF.

In the MUST EECF trial was a small study with a high risk of bias. The randomization scheme was not explained. Inadequate randomization may result in unequal distribution of potential confounders, undermining the validity of study findings.

The short follow-up period (1 year) limits conclusions regarding the durability of treatment effects.

The IPER registry study had serious limitations, especially in the selection of patients i.e. only patients from centres with at least 80% compliance in follow-up data were included [n=5000 patients were enrolled. However, only patients from centres with at least 80% compliance in follow-up data submission were included (N=1427)].

The economic evidence was directly applicable but it had potentially serious limitations.

**Other considerations**

The GDG considered that people with angina which has not responded to drug or revascularisation options or for whom these options are inappropriate or undesirable represent a significant clinical problem. They considered it important, however, that interventions offered to these patients should have robust evidence base. Without such

an evidence base the GDG considered it misleading to offer such interventions to patients and it was more appropriate for healthcare professionals to acknowledge the limitations of interventions available and provide information, education and support for patients.

1 **18.4 Acupuncture**

2 **18.4.1 Clinical question**

3 What is the clinical/cost effectiveness of Acupuncture in people with stable angina?  
4

5 **18.4.2 Clinical evidence**

6 The “Review Protocol” for this topic can be found in Appendix C, the “Search  
7 Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
8 E1, the “Clinical Evidence Tables” in Appendix E2, and the “Forest Plots” in Appendix  
9 F.

10 There were 3 RCTs[215-217] evaluating the effectiveness of acupuncture in people  
11 with stable angina.

12 Data for the 3 RCTS could not be analysed as the standard deviations were not  
13 reported. Hence results have been reported narratively.

14 **Ballegaard 1990 [215]**

15 **Population:** N=49 (n=24 in genuine acupuncture and n=25 sham acupuncture). The  
16 Median age (years) of the patients was 67 yrs in genuine and 66 yrs in sham  
17 acupuncture.

18 **Intervention:** Genuine acupuncture. The genuine acupuncture was given according to  
19 traditional Chinese medicine, each patient receiving 10 treatments in the supine  
20 position within 3 weeks.

21 **Comparison:** Sham acupuncture.

22 **Outcome:** Exercise test; no. of anginal attacks; activity at the time of the pain;  
23 nitroglycerin consumption (diaries); daily well being on an ordinal scale, using the  
24 terms very good (given value 1), good (2), fair (3), not good (4), bad (5) ; global  
25 evaluation of the effect of the treatment on an ordinal scale: much improved,  
26 somewhat improved, slightly improved, unchanged, slightly worse, somewhat worse,  
27 much worse.

28 **Follow-up:** Immediately after treatment

1       **Results:**

2       A. Exercise test variables

3       There was no significant between group differences for any of the exercise variables.  
4       Exercise variables (genuine (n=24 ) vs. sham acupuncture (n=25): Exercise tolerance  
5       (%): median change +9 , range -25 to +184 vs. median change +4 (-16 to +135);  
6       Maximal PRP (%): -1 (-12 to +47) vs. +5 (-22 to +25); Delta PRP (%): + 3 (-38 to  
7       +145) vs. +4 (-28 to + 78); Time to ST segment depression (%): median change 0 (-  
8       42 to +100) vs. median change 0 (-40 to +40); Time to end of ST depression  
9       (%):median change +9 (-75 to +600) vs. median change 0 (-58 to +300); Maximum  
10       ST depression (mm): median change 0 (-1.0 to +0.5) vs. 0 median change (-1.0 to  
11       +1.5); Time with minimum 1 mm ST depression (%):median change +15 (-79 to +490)  
12       vs. median change +5 (-72 to +200); Time to onset of pain (%):median change +10  
13       (-32 to +107) vs. median change +10 (-39 to +55); Post exercise pain duration (%):  
14       median change 0 (-47 to +700) vs. median change 0 (-77 to +78).

15       B. Subjective variables

16       Within both groups there was a significant decrease in both anginal attack rate and  
17       nitroglycerin consumption. After treatment all patients receiving genuine acupuncture  
18       decreased nitroglycerin consumption (median change -54%, range -14 to -100%).  
19       Anginal attack rate was reduced in 13 of 14 patients (93%) (median change -41%,  
20       range +18 to -95%). Nitroglycerin consumption and anginal attack rate were  
21       reduced in 15 of 16 patients (94%) receiving sham acupuncture. The median change  
22       being -53% (range +20 to -100%) and -55% (range +23% to -100%) respectively.  
23       Daily well being was improved in 14 out of 23 (61%) in both groups (median  
24       improvement +1 arbitrary value in both groups). Concerning global evaluation, 75%  
25       of the patients treated by genuine acupuncture reported improvement in their general  
26       condition after the end of the treatment and 6m months later 67% still felt the  
27       improvement. Among those treated by sham acupuncture 84% reported improvement  
28       and 6 months later 72% still felt it.

29       **Ballegaard 1986[216]**

30       Population: N=26 (n=13 in active acupuncture and n=13 in sham acupuncture).

31       **Intervention:** Active acupuncture. During the 3 weeks treatment period all patients  
32       received seven treatments in the supine position.

33       **Control:** Sham acupuncture.

34       **Follow-up:** Immediately after the 9 week treatment period.

35       **Results:** Patients receiving genuine acupuncture had a significantly higher dPRP  
36       (Pressure rate product) than patients receiving sham acupuncture, respectively.  
37       [Maximal PRP (mmHgmin-1 ): 24.640 vs. 13.530 ; Delta PRP (mmHgmin-1 ): 12.580  
38       vs. 6.592]. There was no significant difference between genuine and sham  
39       acupuncture, respectively for : Exercise tolerance (Wmin): 550 (150 to 1300) vs. 256  
40       (100 to 1700); Time to maximal ST depression (min): 2 (0 to 7.5) vs. 2 (0 to 4.5); and  
41       Size of maximal ST depression (mm): 1 (0 to 3) vs. 1 (0 to 2); No. of anginal attacks  
42       per 3 weeks: 55 (8 to 168) vs. 66 (41 to 149); and nitroglycerin consumption (0.25

1 mg tablets per 3 weeks): 39 (1 to 193) vs. 30 (0 to 152). Six of the 12 patients in the  
2 active treatment group and one of 12 patients in the sham treatment group reported  
3 improvement in general well being after treatment ( $p=0.10$ ). No complications or  
4 adverse effects were observed. The study period consisted of: 3 weeks of pre  
5 treatment control; after randomisation 3 weeks of treatment, during which the patients  
6 received either active or sham acupuncture, and 3 weeks of post treatment control.

7 **Richter 1991[217]**

8 **Population:** N=21 (cross over study).

9 **Intervention:** Acupuncture. The treatment was given 3 times per week during the 4  
10 week period.

11 **Comparison:** Tablet placebo.

12 **Follow-up:** Immediately after the 4 week treatment period (2 weeks wash out period  
13 between the treatment periods)

14 **Results:** During acupuncture treatment, 14 patients showed a reduced number of  
15 anginal attacks compared with placebo. The no. of attacks was unchanged in the  
16 remaining 7 patients; no worsening was observed in any of the patients. In the whole  
17 group, the average number of anginal attacks/week was 12.1 during the run-in  
18 period, 6.1 during the acupuncture period and 10.6 during the placebo period. The  
19 differences between acupuncture and both run-in and placebo periods were  
20 statistically significant ( $p<0.01$ ). The results of the exercise tests did not show any  
21 significant difference in maximal physical performance at the end of the acupuncture  
22 period compared with placebo, the mean values being 104.2 W and 101.4 W  
23 respectively. However, maximal workload until onset of chest pain was significantly  
24 increased after acupuncture compared with placebo (94.3 W vs. 81.9 W,  $P<0.05$ ).  
25 Mean chest pain score at maximal workload improved significantly after acupuncture  
26 compared with placebo (mean 0.81 W and 1.38,  $p<0.01$ ). ST segment depression  
27 at maximal workload was significantly reduced after acupuncture compared with  
28 placebo (mean 0.71 mm vs. 1.03 mm,  $p<0.01$ ). Similar results were obtained for ST  
29 segment depression at maximal comparable workload (mean 0.63 mm vs. 0.87 mm,  
30  $p<0.01$ ). [Standard deviations not reported]. Concerning the self-rating life quality  
31 questionnaire, the score was significantly improved for chest pain, physical  
32 performance, peripheral coldness, pessimism, vertigo and relaxation ( $p<0.05$ ). The  
33 statistical significance could not be proved for anxiety, tiredness, sleep disturbances  
34 and gastro-intestinal symptoms. No adverse effect of acupuncture was observed.  
35 [mean values and standard deviations not reported]

36

37 **18.4.3 Economic evidence**

38 One study[222] focusing on the addition of acupuncture and self-education to medical  
39 treatment was found but it was excluded as it had serious limitations due to the study  
40 design (within-group comparison) and it was partially applicable (cost estimates from  
41 the USA).

42

1 18.4.4 Evidence statements

**Clinical**

**Acupuncture vs. sham acupuncture**

**Ballegaard 1990[215]:** Evidence from one RCT shows that there was no significant difference between genuine acupuncture and sham acupuncture for Exercise variables ; Anginal attack rate ; and nitroglycerin consumption [follow-up 3 days after treatment for angina pain counts, one week after treatment for exercise duration].

**Ballegaard 1986[216]:** Evidence from one RCT shows that compared to patients receiving sham acupuncture the patients receiving active acupuncture increased cardiac work capacity significantly,. There was no significant difference between the groups for exercise tolerance Time to maximal ST depression (min); Size of maximal ST depression (mm) : and Nitroglycerin consumption [Follow-up immediately after the treatment period]

**Acupuncture vs. placebo**

**Richter 1991[217]:** Evidence from one randomised cross over trial shows that compared to placebo treatment acupuncture significantly reduced anginal attacks per week); maximal workload until onset of chest pain was significantly increased after acupuncture compared with placebo chest pain at maximal workload improved significantly after acupuncture compared with placebo ST segment depression at maximal workload was significantly reduced after acupuncture compared with placebo and ST segment depression at maximal comparable workload was significantly reduced after acupuncture compared with placebo There was no significant difference in maximal physical performance at the end of the acupuncture period compared with placebo [follow-up immediately after the treatment period]

**Economic**

No economic evidence was included on this intervention.

2 18.4.5 Recommendations and link to evidence

<b>Recommendation</b>	<b>Do not offer acupuncture to manage stable angina.</b>
<b>Relative values of different outcomes</b>	The outcomes considered as important during the development of the review protocol for pain interventions included: improvement in anginal symptoms (angina frequency and nitroglycerin consumption), mortality, exercise tolerance, major cardiac events, hospitalisation, revascularisation, QoL and adverse events.

	<p>One RCT[217] showed some improvement in angina and exercise test variables when compared to tablet placebo. However there was no improvement in angina and exercise test variables in the two RCTs[215,216] where acupuncture was compared to sham acupuncture.</p>
<b>Trade off between clinical benefits and harms</b>	<p>There is no evidence of clinical benefits arising from the use of acupuncture in stable angina patients.</p>
<b>Economic considerations</b>	<p>No published health-economic evaluation of acupuncture was included. The intervention is not cost-effective as it generates costs without being effective at improving the outcomes considered.</p>
<b>Quality of evidence</b>	<p>Evidence was obtained from 3 low quality RCTs[215-217]. Each of these RCTs had small sample size (&lt;50 patients); outcomes were measured immediately after treatment with no longer term follow-up. The methodology of the trials was not well reported and the derived data was not analysable. Hence the GDG was not confident in the results of these trials.</p>
<b>Other considerations</b>	<p>The GDG considered that people with angina which has not responded to drug or revascularisation options or for whom these options are inappropriate or undesirable represent a significant clinical problem. They considered it important however, that interventions offered to these patients should have robust evidence base. The GDG did not consider that the evidence for acupuncture supported its use in people with angina. The GDG recognised that people with angina may have pain in the chest that arises from separately from ischaemic pain and that acupuncture may have some role in the treatment of other chest pains.</p>

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## 2 **18.5 Self management of pain**

### 3 **18.5.1 Clinical question**

4           What is the clinical/cost effectiveness of self management of pain in people with  
5           stable angina?

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1 **18.5.2 Clinical evidence**

2 The “Review Protocol” for this topic can be found in Appendix C, the “Search  
3 Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
4 E1, the “Clinical Evidence Tables” in Appendix E2, and the “Forest Plots” in Appendix  
5 F.

6 There were 2 RCTs[218,219] comparing Psycho educational programmes with  
7 control/standard medical care for self management of pain in stable angina.

8 **McGillion 2008[218] (CASMP vs. control)**

9 **Population:** n=130 were randomised, n=66 to the CASMP and n=64 to the waiting  
10 list control group.

11 **Intervention:** The Chronic Angina Self-Management Program (CASMP) is a  
12 standardized psycho education programme given in two-hour sessions weekly, over a  
13 six-week period.

14 The CASMP is an adaptation of Lorig et al.’s Chronic Disease Self-Management  
15 Program (CDSMP, 1999 Stanford University). The programme was delivered by a  
16 registered nurse using a group format (e.g., 8-15 patients) in a comfortable classroom  
17 setting. Key pain related content includes relaxation and stress management  
18 techniques, energy conservation, symptom monitoring and management techniques,  
19 medication review, seeking emergency assistance, diet, and managing emotional  
20 responses to cardiac pain. Programme sessions were offered both day and evening  
21 and participants were encouraged to bring a family member or friend if they wished.  
22 A facilitator manual specified the intervention protocol in detail to ensure consistent  
23 delivery of the CASMP across sessions.

24 **Comparison:** Waiting list control: The patients in this group were offered entry into  
25 the next available CASMP once post-test measures were completed.

26 **Outcomes:** The primary outcome was Health Related Quality of Life (HRQL) which  
27 included the SF-36 and the SAQ (Seattle Angina Questionnaire). The secondary  
28 outcome was enabling skill, reflected by CSA patients’ self-efficacy and  
29 resourcefulness to self-manage their pain.

30 **Follow-up:** 3 months

1 **Table 18.6: Chronic angina self management Program (CASMP) vs. control (Follow-up 3 months from start of treatment) for stable angina**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Chronic angina self management Program (CASMP)	control (Follow-up 3 months from start of treatment)	Relative (95% CI)	Absolute	
Physical functioning (SF-36) (range 0-100 -higher score better functioning) (change scores) (follow-up 3 months; range of scores: 0-100; better indicated by higher values)											
McGillion 2008[218]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	57	60	-	MD 5.98 higher (2.59 to 9.37 higher)	⊕⊕00 LOW
Role physical functioning (SF-36) (change scores) (range 0-100) (follow-up 3 months; range of scores: 0-100; better indicated by higher values)											
McGillion 2008[218]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	57	60	-	MD 1.6 higher (2.5 lower to 5.7 higher)	⊕⊕⊕0 MODERATE
Bodily pain (SF-36) (change scores) (range 0-100) (follow-up 3 months; range of scores: 0-100; better indicated by higher values)											
McGillion 2008[218]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	57	60	-	MD 2.3 higher (0.94 lower to 5.54 higher)	⊕⊕⊕0 MODERATE
General Health (SF-36) (change scores) (0-100) (follow-up 3 months; range of scores: 0-100; better indicated by higher values)											
McGillion 2008[218]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	57	60	-	MD 3.87 higher (1.3 to 6.44 higher)	⊕⊕00 LOW
Angina frequency (SAQ) (range 0-100- higher scores better functioning) (change scores) (follow-up 3 months; range of scores: 0-100; better indicated by higher values)											
McGillion 2008[218]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	57	60	-	MD 9.2 higher (1.48 to 16.92 higher)	⊕⊕⊕0 MODERATE
Angina stability (SAQ) (range 0-100) (change scores) (follow-up 3 months; range of scores: 0-100; better indicated by higher values)											
McGillion 2008[218]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	57	60	-	MD 15.1 higher (4.11 to 26.09 higher)	⊕⊕⊕0 MODERATE
Disease perception (SAQ) (range 0-100) (change scores) (follow-up 3 months; range of scores: 0-100; better indicated by higher values)											
McGillion 2008[218]	randomised trials	serious(a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	57	60	-	MD 6.6 higher (1.18 lower to	⊕⊕⊕0 MODERATE

										14.38 higher)	
Physical limitation (SAQ) (range 0-100) (change scores) (follow-up 3 months; range of scores: 0-100; better indicated by higher values)											
McGillion 2008[218]	randomised trials	serious(a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	57	60	-	MD 5.5 higher (0.24 lower to 11.24 higher)	⊕⊕⊕○ MODERATE
Treatment satisfaction (SAQ) (range 0-100) (change scores) (follow-up 3 months; range of scores: 0-100; better indicated by higher values)											
McGillion 2008[218]	randomised trials	serious(a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	57	60	-	MD 4.9 higher (3.05 lower to 12.85 higher)	⊕⊕⊕○ MODERATE
Self-Efficacy to manage disease (Self-efficacy Scale )range scores 10- 100 -higher scores better) (change scores) (follow-up 3 months; range of scores: 10-100; better indicated by higher values)											
McGillion 2008[218]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	serious (b)	none	57	60	-	MD 8.6 higher (2.76 to 14.44 higher)	⊕⊕○○ LOW

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(a) Randomised. Allocation concealment reported. 9/66 (14%) in the intervention group and 4/64 (6%) in the control group. There are more patients in the intervention group who were lost to follow-up but there are no systematic differences between the two groups with respect to loss of participants. The study follow-up period was limited to three months after baseline for both groups. ITT used. No blinding of participants and outcome assessors.

(b) 2 Lower CI crosses MID.

1        **Additional data:**

2        **Payne 1994[219] (Self Pain management programme vs. control)**

3        Data was not analysed for the following study as it was poorly reported:

4        **Population:** n=52. Participants were 52 male veterans (26 in the treatment and 26  
5        controls). To qualify for the study, patients were required to meet the following  
6        criteria: (a) diagnosis of CAD, or positive diagnostic evaluation, such as an exercise  
7        stress test, thallium 201 scan or coronary angiogram (b) self report of at least 4  
8        episodes of chest pain or discomfort in the previous 4 weeks (c) 18-65 yrs of age (d)  
9        no hospitalisation within past 30 days (e) no current physical disorder associated with  
10       severely disabling symptoms or a recent change in symptoms (f) no history of heart  
11       valve replacement (g) no history of cardiac transplant surgery.

12       **Intervention:** A pain management programme administered over three consecutive  
13       weekly sessions (length of sessions not reported). The goals were to 1) educate  
14       patients regarding the role of psychological factors in pain and pain control and 2)  
15       teach participants an integrated set of self management skills to modify cognitions,  
16       behaviours and affective responses considered likely to adversely impact on the  
17       experience of chest pain. Specific skills taught included pacing of physical activities  
18       (e.g. taking scheduled breaks), modification of dysfunctional, stress engendering  
19       thoughts using cognitive reframing and problem solving techniques, and relaxation  
20       training via diaphragmatic breathing.

21       **Control:** Received standard medical care

22       **Follow-up:** 6 months.

23       **Primary outcomes:** No primary or secondary outcomes specified. Outcomes included:  
24       pain frequency and intensity; frequency of NTG usage; mood and psychological  
25       distress.

26       **Results:** There were no significant differences between groups with regard to pain  
27       frequency, pain intensity, psychological and other factors at 6 months. Actual data for  
28       results not reported.

29       **18.5.3       Economic evidence**

30       No economic studies were found on this question.

31       **18.5.4       Evidence statements**

**Clinical                    Self management programme vs. control**

**McGillion 2008[218]:** Evidence from one RCT shows that Physical  
         functioning (SF-36) (MD 5.98 [2.59, 9.37]) , General Health (SF-  
         36) (MD 3.87 [1.30, 6.44]), Angina frequency (SAQ) (MD 9.20  
         [1.48, 16.92], Angina stability (SAQ) (MD 15.10 [4.11,26.09]);  
         and self-efficacy to manage disease (self-efficacy scale) (MD 8.60  
         [2.76, 14.44]) were significantly improved in the CASMP  
         compared to control . There was no significant difference between

CASMP and control for Role physical functioning (SF-36) (MD 1.60 [-2.50, 5.70]); bodily pain (SF-36) (MD 2.30 [-0.94, 5.54]);disease perception (SAQ) (MD 6.60 [-1.18, 14.38]) ; physical limitation (SAQ) (MD 5.50 [-0.24,11.24]) and treatment satisfaction (SAQ) (MD 4.90 [-3.05, 12.85]) [Follow-up 3 months from start of treatment]

**Payne 2004[219]:** Evidence from one RCT shows that there were no significant differences between Pain management programme compared to control (standard care) with regard to pain frequency, pain intensity, psychological and other factors. (actual values for results not reported). [Follow-up 6 months]

**Economic**

No economic evidence was available on this question.

1 18.5.5 Recommendations and link to evidence

<p><b>Recommendation</b></p>	<p><b>Offer people whose stable angina has not responded to drug treatment and/or revascularisation comprehensive re-evaluation and advice, which may include:</b></p> <ul style="list-style-type: none"> <li>• exploring the person's understanding of their condition</li> <li>• exploring the impact of symptoms on the person's quality of life</li> <li>• reviewing the diagnosis and considering non-ischaemic causes of pain</li> <li>• reviewing drug treatment and considering future drug treatment and revascularisation options</li> <li>• explaining how the person can manage the pain themselves</li> <li>• acknowledging the limitations of future treatment</li> <li>• specific attention to the role of psychological factors in pain</li> <li>• development of skills to modify cognitions and behaviours associated with pain.</li> </ul>
<p><b>Relative values of different outcomes</b></p>	<p>Quality of Life outcomes were considered to be most important in assessing the effectiveness of self-management including various outcomes measured by the SF-36 health survey (physical functioning, bodily pain and general health), as well as those of the Seattle Angina Questionnaire (angina frequency and stability, disease perception, physical limitation and treatment satisfaction) and self-efficacy to manage disease. One RCT[218] showed statistically significant improvements in some Quality of Life variables including physical functioning, general health, angina frequency and stability, and self-efficacy to manage disease.</p>
<p><b>Trade off between clinical benefits and harms</b></p>	<p>The studies reviewed do not provide a report on harms arising from self-management. The GDG considered it unlikely that significant harms would occur from involvement in a self-management programme.</p>

**Economic considerations**

The small increase in staff time cost is likely to be offset by the improvement in quality of life shown by the clinical review.

**Quality of evidence**

Clinical evidence on self-management of pain that met our inclusion criteria for reviewing was obtained from one moderate quality RCT (n=117) and one low quality RCT (n=52) study as assessed by GRADE.

McGillion 2008[218] conducted a small RCT of a psycho education programme Chronic Angina Self-Management Program (CASMP) in which those treated were compared to patients in a waiting list control group. The study found statistically reliable short-term improvements in some components of the HRQL for those who participated in the CASMP as compared to the control group. However, the follow-up period was limited to three months after baseline hence the long-term sustainability of the observed intervention effects is not known. Due to the nature of the treatment, the patients undergoing EECF could not be blinded, increasing the likelihood of the placebo effect. Further, all psycho education sessions were delivered by a single facilitator increasing the threat to external validity.

Payne 1994[219] conducted a very small RCT evaluating a pain management programme + standard medical care compared with standard medical care alone. It found that there were short-term reductions in self-report of number of chest pain episodes in treated subjects but these were not evident at 6 month follow-up. The study however had a high risk of bias which would make the results unreliable.

No economic evidence was included on this intervention.

**Other considerations**

The GDG made a recommendation on intervention for patients whose angina has not responded to treatment or for whom revascularization is undesirable or inappropriate using the information presented by Professor Michael Chester and the evidence from the reviews on self management strategies.

The evidence for self-management strategies comes from two studies[218,219]. These programmes included a range of self management skills to modify cognitions, behaviours and affective responses considered likely to adversely impact on the experience of chest pain. Specific skills taught included components such as pacing of physical activities (e.g. taking scheduled breaks), modification of dysfunctional, stress engendering thoughts

using cognitive reframing and problem solving techniques, and relaxation training via diaphragmatic breathing, energy conservation, symptom monitoring and management techniques, medication review, seeking emergency assistance, diet, and managing emotional responses to cardiac pain.

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## 1 **19 Cardiac syndrome X**

### 2 **19.1 Introduction**

3 Cardiac syndrome X can be defined as angina in the presence of normal coronary arteries.  
4 Diagnostic criteria may also include evidence of ischaemia. The term microvascular angina is  
5 also used as it is thought that the pathology may lie within the microvasculature.  
6 Abnormalities of endothelial function have also been described.  
7

8 The GDG were interested in the efficacy of standard anti-anginal drug treatment and  
9 drugs for secondary prevention for people with syndrome X and for the evidence on  
10 benefit of rehabilitation programmes. This chapter reports on the results of these questions:

- 11 A. What is the clinical /cost effectiveness of using standard anti-angina drug therapy  
12 (short acting nitrates, BB,CCB, long acting nitrates, ACE/ARBs, nicorandil, Ivabradine,  
13 Ranolazine,) and /or drugs for secondary prevention in people with syndrome X.
- 14 B. What is the clinical/cost effectiveness and safety of cardiac rehabilitation  
15 programmes for people with syndrome X?
- 16 C. What is the incremental value/effectiveness of anatomical/functional tests for  
17 prognostic risk stratification in prediction of adverse cardiac outcomes in people with  
18 cardiac syndrome X?

19 The studies included in the review are all of patient with exertional angina who had positive  
20 exercise tests and normal coronary arteries on angiography.

### 21 **19.2 Clinical/Cost effectiveness of standard anti-anginal drug therapy for** 22 **management of syndrome X**

23 This review explores use of standard anti-anginal drug therapies for treating angina  
24 patients who have normal coronary arteries (cardiac syndrome X). This evidence  
25 review included a total of 7 papers. No economic evidence was available to assess  
26 cost-effectiveness; therefore this review focuses only on clinical effectiveness.

27 The results of the review have been analysed based on the type of drug involved  
28 (BBs, CCBs, nitrates, nicorandil, aminophylline, ACE inhibitors) and whether they were  
29 compared to placebo or to each other.

1 The main outcomes analysed were number of ischemic episodes, duration of ischemic  
2 episodes, exercise duration, time to 1 mm-ST segment depression and consumption of  
3 nitroglycerin tablets.

4

#### 5 **19.2.1 Clinical Evidence**

6 The “Review Protocol” for this topic can be found in Appendix C, the “Search  
7 Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
8 E1, the “Clinical Evidence Tables” in Appendix E2, and the “Forest Plots” in Appendix  
9 F.

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11 The results of the review are presented as follows:

12

- BBs vs. placebo

13

- CCBs vs. placebo

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- BB vs. CCB

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- BB vs. CCB in people with pressure-rate product variation <1050

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- BB vs. CCB in people with pressure-rate product variation >1050

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- BB vs. nitrates

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- CCB vs. nitrates

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- Nicorandil vs. placebo

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- Aminophylline vs. nitroglycerin

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- Angiotensin-converting enzyme inhibitors + statins vs. placebo

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1 **Table 19.1: BBs vs. placebo for Cardiac Syndrome X**

Quality assessment							Summary of findings				
							No of patients		Effect		Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	BB	control	Relative (95% CI)	Absolute	
<b>ischemic episodes - propranolol vs. placebo (follow-up 7 days; range of scores: -, better indicated by less)</b>											
Bugiardini 1989[223] (c)	randomised trial (b)	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	16	16	-	MD3.2 lower (4.13 to 2.27 lower)	⊕⊕⊕○ LOW
<b>ischemic duration (min) - propranolol vs. placebo (follow-up 7 days; range of scores: -, better indicated by less)</b>											
Bugiardini 1989[223]	randomised trial (b)	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	16	16	-	MD 25 lower (34.15 to 15.85 lower)	⊕⊕⊕○ LOW

(a) Randomisation and allocation concealment unclear, small sample size

(b) Crossover design

(c) Propranolol 120-160mg daily (optimal dose for each patient determined 2-3 weeks before the double blind study ; beta blockade occurred at 120mg a day in 6 patients and at 160mg in 10)

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1 **Table 19.2: CCBs vs. placebo for Cardiac Syndrome X**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							CCBs	control	Relative (95% CI)	Absolute	
<b>ischemic episodes (verapamil vs. placebo ; verapamil or nifedipine vs. placebo ) (follow-up 7-28 days; range of scores: -; better indicated by less)</b>											
Bugiardini 1989[223] (c) Cannon 1985[224] (d)	randomised trial (a)	serious (b)	serious inconsistency (e)	no serious indirectness	no serious imprecision	none	38	38		MD 0.6 lower (1.81 lower to 0.61 higher) (e)	⊕⊕⊕○ LOW
<b>ischemia duration (min) (verapamil vs. placebo; - verapamil or nifedipine vs. placebo) (follow-up 7-28 days; range of scores: -; better indicated by less)</b>											
Bugiardini 1989[223] (c) Cannon 1985[224] (d)	randomised trial (a)	serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	none	38	38	-	MD 0.74 higher (0.55 lower to 2.04 higher) (f)	⊕⊕⊕○ MODERATE
<b>Nitroglycerin tablets consumption - verapamil or nifedipine vs. placebo (follow-up 28 days; range of scores: -; better indicated by less)</b>											
Cannon 1985[224] (d)	randomised trial (a)	serious (b)	no serious inconsistency	no serious indirectness	serious imprecision (f)	none	22	22	-	MD 18 lower (41.74 lower to 5.74 higher)	⊕⊕⊕○ LOW
<b>presence of chest pain during exercise - verapamil or nifedipine vs. placebo (follow-up 28 days)</b>											
Cannon 1985[224] (d)	randomised trial (a)	serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	none	9/25 (36%)	16/22 (72.7%)	RR 0.49 (0.28 to 0.89)	371 fewer per 1,000 (from 80 fewer to 524 fewer)	⊕⊕⊕○ MODERATE

- (a) Crossover design
- (b) Randomisation and allocation concealment unclear, small sample size
- (c) Propanolol 120-160mg daily (optimal dose for each patient determined 2-3 weeks before the double blind study ; beta blockade occurred at 120mg a day in 6 patients and at 160mg in 10)
- (d) The drug and dosage used were determined from the unblinded lead-in phase: 17 patients received verapamil, 40-160mg 4 times a day (mode 80) and 9 patients received nifedipine 10-30mg 4 times a day (mode 10)
- (e) There was substantial heterogeneity (I<sup>2</sup>=71%) indicating that these results must be carefully interpreted
- (f) 95% CI includes no effect and the upper and lower CI crosses the MID.

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1 **Table 19.3: BBs vs. CCBs for Cardiac Syndrome X**

Quality assessment							Summary of findings				Quality
							No of patients		Effect		
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	BBs	CCBs	Relative (95% CI)	Absolute	
<b>Number of anginal episodes (per 4 weeks per patient) (propranolol vs. verapamil; atenolol vs. amlodipine) (follow-up 1-4 weeks; range of scores: -; better indicated by less)</b>											
Bugiardini 1989[223] Lanza 1999 (c)	randomised trial (a)	serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	none	26	26	-	MD 2.71 lower (3.6 to 1.83 lower)	⊕⊕⊕O MODERATE
<b>Chest pain episodes duration (min) ( propranolol vs. verapamil ; atenolol vs. amlodipine ) (follow-up 1-4 weeks; range of scores: -; better indicated by less)</b>											
Bugiardini 1989[223] Lanza 1999 (c)	randomised trial (a)	serious (b)	serious inconsistency (d)	no serious indirectness	no serious imprecision	none	26	26	-	MD 17.66 lower (24.35 to 10.97 lower)	⊕⊕⊕O LOW
<b>severity of chest pain (scale 1-5) - atenolol vs. amlodipine (follow-up 4 weeks; range of scores: -; better indicated by less)</b>											
Lanza 1999[225] (c)	randomised trial (a)	serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	none	10	10	-	MD 2 lower (15.26 lower to 11.26 higher)	⊕⊕⊕O LOW
<b>quality of life (scale 0-100 mm) - atenolol vs. amlodipine (follow-up 4 weeks; range of scores: -; better indicated by less)</b>											
Lanza 1999[225] (c)	randomised trial (a)	serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	none	10	10	-	MD 8 higher (15.73 lower to 31.73 higher)	⊕⊕⊕O LOW

- (a) Crossover design
- (b) Unclear randomisation and allocation concealment, small sample size
- (c) Bugiardini 1989[223]: propranolol 120-160mg/day (optimal dose for each patient determined 2-3 weeks before the double blind study ; beta blockade occurred at 120mg a day in 6 patients and at 160mg in 10). Lanza 1999[225]: atenolol 100mg/day, amlodipine 10mg/day
- (d) There was substantial heterogeneity (I<sup>2</sup>=86%) indicating that these results must be carefully interpreted

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1 **Table 19.4: BBs vs. CCBs in patients with pressure-rate product variation <1050 for Cardiac Syndrome X**

Quality assessment							Summary of findings				Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		
							BBs	CCBs in patients with pressure-rate product variation <1050	Relative (95% CI)	Absolute	
<b>exercise duration (sec) - acebutolol vs. verapamil in patients with pressure-rate product variation &gt;1050 (follow-up 4 weeks; range of scores: -; better indicated by less)</b>											
Romeo 1988[226] (c)	randomised trial (a)	serious (b)	no serious inconsistency	no serious indirectness	serious imprecision (d)	none	15	15	-	MD 44 lower (113.48 lower to 25.48 higher)	⊕⊕⊕○ LOW

- (a) Crossover design
- (b) Randomisation, allocation concealment and blinding not reported, small sample size
- (c) Acebutolol 400mg a day, verapamil 80mg 4 times a day
- (d) 95% CI includes no effect and the upper and lower CI crosses the MID.

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8 **Table 19.5: BBs vs. CCBs in patients with pressure-rate product variation >1050 for Cardiac Syndrome X**

Quality assessment							Summary of findings				Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		
							BBs	CCBs in patients with pressure-rate product variation >1050	Relative (95% CI)	Absolute	
<b>exercise duration (sec) - acebutolol vs. verapamil in patients with pressure-rate product variation &lt;1050 (follow-up 4 weeks; range of scores: -; better indicated by less)</b>											
Romeo 1988[226] (c)	randomised trial (a)	serious (b)	no serious inconsistency	no serious indirectness	serious imprecision (d)	none	15	15	-	MD 0 higher (52.48 lower to 52.48 higher)	⊕⊕⊕○ LOW

- (a) Crossover design
- (b) Randomisation, allocation concealment and blinding not reported, small sample size
- (c) Acebutolol 400mg a day, verapamil 80mg 4 times a day
- (d) 95% CI includes no effect and the upper and lower CI crosses the MID.

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1 **Table 19.6: BBs vs. nitrates for Cardiac Syndrome X**

Quality assessment							Summary of findings				Quality
							No of patients		Effect		
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	BBs	Nitrates	Relative (95% CI)	Absolute	
<b>Number of anginal episodes (per 4 weeks per patient) - atenolol vs. ISMN (follow-up 4 weeks; range of scores: -; better indicated by less)</b>											
Lanza 1999[225] (c)	randomised trial (a)	serious (b)	no serious inconsistency	no serious indirectness	serious imprecision (d)	none	10	10	-	MD 9 lower (24.84 lower to 6.84 higher)	⊕⊕⊕○ LOW
<b>Chest pain episodes duration (min) - atenolol vs. ISMN (follow-up 4 weeks; range of scores: -; better indicated by less)</b>											
Lanza 1999[225] (c)	randomised trial (a)	serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	none	10	10	-	MD 3 higher (6.15 lower to 12.15 higher)	⊕⊕⊕○ LOW
<b>severity of chest pain (scale 1-5) - atenolol vs. ISMN (follow-up 4 weeks; range of scores: -; better indicated by less)</b>											
Lanza 1999[225] (c)	randomised trial (a)	serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	none	10	10	-	MD 0.2 higher (0.85 lower to 1.25 higher)	⊕⊕⊕○ LOW
<b>quality of life (scale 0-100 mm) - atenolol vs. ISMN (follow-up 4 weeks; range of scores: -; better indicated by less)</b>											
Lanza 1999[225] (c)	randomised trial (a)	serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	none	10	10	-	MD 29 higher (4.44 to 53.56 higher)	⊕⊕⊕○ LOW

- (a) Crossover design
- (b) Randomisation and allocation concealment unclear, small sample size
- (c) Atenolol 100mg/day, ISMN 50mg/day
- (d) 95% CI includes no effect and the upper and lower CI crosses the MID

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1 **Table 19.7: CCBs vs. nitrates for Cardiac Syndrome X**

Quality assessment							Summary of findings				Quality
							No of patients		Effect		
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	CCBs	Nitrates	Relative (95% CI)	Absolute	
<b>Number of anginal episodes (per 4 weeks per patient) - amlodipine vs. ISMN (follow-up 4 weeks; range of scores: -; better indicated by less)</b>											
Lanza 1999[225] (c)	randomised trial (a)	serious (b)	no serious inconsistency	no serious indirectness	serious imprecision (d)	none	10	10	-	MD 2 lower (21.28 lower to 17.28 higher)	⊕⊕⊕○ LOW
<b>Chest pain episodes duration (min) - amlodipine vs. ISMN (follow-up 4 weeks; range of scores: -; better indicated by less)</b>											
Lanza 1999[225] (c)	randomised trial (a)	serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	none	10	10	-	MD 5 higher (6.39 lower to 16.39 higher)	⊕⊕⊕○ LOW
<b>severity of chest pain (scale 1-5) - amlodipine vs. ISMN (follow-up 4 weeks; range of scores: -; better indicated by less)</b>											
Lanza 1999[225] (c)	randomised trial (a)	serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	none	10	10	-	MD 0.4 higher (0.57 lower to 1.37 higher)	⊕⊕⊕○ LOW
<b>quality of life (scale 0-100 mm) - amlodipine vs. ISMN (follow-up 4 weeks; range of scores: -; better indicated by less)</b>											
Lanza 1999[225] (c)	randomised trial (a)	serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	none	10	10	-	MD 21 higher (1.81 lower to 43.81 higher)	⊕⊕⊕○ LOW

- (a) Crossover design
- (b) Randomisation and allocation concealment unclear, small sample size
- (c) Amlodipine 10mg/day, ISMN 50mg/day
- (d) 95% CI includes no effect and the upper and lower CI crosses the MID.

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1 **Table 19.8: Nicorandil vs. placebo for Cardiac Syndrome X**

Quality assessment							Summary of findings				Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		
							Nicorandil	control	Relative (95% CI)	Absolute	
<b>Time to 1mm ST-segment depression (sec) (follow-up 2 weeks; range of scores: -; better indicated by less)</b>											
Chen 1997[227] (c)	randomised trial (a)	serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	none	13	13	-	MD 69 higher (0.24 to 137.76 higher)	⊕⊕⊕○ LOW
<b>maximum ST-segment depression (mm) (follow-up 2 weeks; range of scores: -; better indicated by less)</b>											
Chen 1997[227] (c)	randomised trial (a)	serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	none	13	13	-	MD 0.4 lower (0.99 lower to 0.19 higher)	⊕⊕⊕○ LOW
<b>Total exercise duration (sec) (follow-up 2 weeks; range of scores: -; better indicated by less)</b>											
Chen 1997[227] (c)	randomised trial (a)	serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	none	13	13	-	MD 38 higher (16.85 lower to 92.85 higher)	⊕⊕⊕○ LOW

- (a) Crossover design
- (b) Randomisation, allocation concealment and blinding unclear, small sample size
- (c) Nicorandil 5mg 3 times a day

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6 **Table 19.9: Aminophylline vs. nitroglycerine for Cardiac Syndrome X**

Quality assessment							Summary of findings				Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		
							Aminophylline	Nitroglycerine	Relative (95% CI)	Absolute	
<b>Time to 1mm ST depression (follow-up 5min post nitroglycerin or 90min post aminophylline; range of scores: -; better indicated by less)</b>											
Radice 1996[228] (c)	randomised trial (a)	serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	none	20	20	-	MD 1.9 higher (0.88 to 2.92 higher)	⊕⊕⊕○ LOW

- (a) Crossover design
- (b) Randomisation, allocation concealment and blinding unclear, small sample size
- (c) Aminophylline 400mg or nitroglycerin (sublingual) 0.3mg administered once

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2 **19.2.2 Economic evidence**

3 No economic studies were identified on this question.

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5 **19.2.3 Evidence statements**

**Clinical BBs vs. placebo for cardiac syndrome X**

**Bugiardini 1989[223]:** Evidence from one RCT shows that there was significantly lower number of ischemic episodes [MD -3.2 (-4.13 to -2.27)] and smaller ischemic duration (min) [MD -25 (-34.15 to -15.85)] in the BBs group than in the placebo group. [7-day follow-up].

**CCBs vs. placebo for cardiac syndrome X**

**Bugiardini 1989[223]; Cannon 1985[224]:** Evidence from two RCTs shows that there was no significant difference between CCBs and placebo for number of ischemic episodes [MD -0.6 (-1.81 to 0.61)] and ischemic duration [MD 0.74 (-0.55 to 2.04)]. [follow-up 7-28 days]

**Cannon 1985[224]:** Evidence from one RCT shows that there was no significant difference between CCBs and placebo for consumption of nitroglycerin tablets [MD -18 (-41.74 to 5.74)] but patients in the CCBs group had significantly less chest pain during exercise compared to those receiving placebo [RR 0.49 (0.028 to 0.89)]. [follow-up 28 days]

**BBs vs. CCBs for cardiac syndrome X**

**Bugiardini 1989[223]; Lanza 1999[225]:** Evidence from two RCTs shows that there was a significantly lower number of anginal episodes [MD -2.71 (-3.6 to -1.83)] and shorter chest pain episode duration (min) [MD -17.66 (-24.35 to -10.97)] in the BBs compared to CCBs group. [follow up 1-4 weeks]

**Lanza 1999[225]:** Evidence from one RCT shows that there was no significant difference in severity of chest pain [MD -0.2 (-1.17 to 0.77)] and quality of life [MD 8 (-15.73 to 31.73)] between BBs and CCBs [follow up 4 weeks]

**BBs vs. CCBs in patients with pressure-rate product variation <1050**

**Romeo 1988[226]:** Evidence from one RCT shows that there was no significant difference between BBs and CCBs for exercise duration (sec) [MD -44 (-113.48 to 25.48)] in patients with pressure-rate product variation <1050 [follow up 4 weeks]

**BBs vs. CCBs in patients with pressure-rate product variation >1050**

**Romeo 1988[226]:** Evidence from one RCT shows that there was no

significant difference between BBs and CCBs for exercise duration (sec) [MD 0 (-52.48 to 52.48)] in patients with pressure-rate product variation >1050 [follow up 4 weeks]

**BBs vs. nitrates for cardiac syndrome X**

**Lanza 1999[225]:** Evidence from one RCT shows that there was no significant difference between BBs and nitrates for number of anginal episodes [MD -9 (-24.84 to 6.84)], chest pain duration (min) [MD 3 (-6.15 to 12.15)], severity of chest pain [MD0.2 (-0.85 to 1.25)]. Quality of life was significantly improved in the BBs group compared to the nitrates group [MD 29 (4.44 to 53.56)] [follow up 4weeks]

**CCBs vs. nitrates for cardiac syndrome X**

**Lanza 1999[225]:** Evidence from one RCT shows that there was no significant difference between CCBs and nitrates for number of anginal episodes [MD - (-21.28 to 17.28)], chest pain duration (min) [MD 5 (-6.39 to 16.39)], severity of chest pain [MD 0.4 (-0.57 to 1.37)] or quality of life [MD 21 (-1.81 to 43.81)] [follow up 4weeks]

**Nicorandil vs. placebo for cardiac syndrome X**

**Chen 1997[227]:** Evidence from one RCT shows that time to 1mm ST segment depression (sec) was significantly longer in the Nicorandil group compared to placebo [MD 69 (0.24 to 137.76)], and there was no significant difference between Nicorandil and placebo for maximum ST segment depression (mm) [MD-0.4 (-0.99 to 0.19)] and total exercise duration (sec) [MD38 (-16.85 to 92.85)]. [follow up 2 weeks]

**Aminophylline vs. nitroglycerine for cardiac syndrome X**

**Radice 1996[228]:** Evidence from one RCT shows that there was a significant increase in time to 1mm ST depression [MD 1.9 (0.88 to 2.92)] in the aminophylline group compared to the nitroglycerin group [follow up 5-90min after administration of drug]

**Economic** No economic evidence was found on this question.

1 **19.2.4 Recommendations and link to evidence**

<b>Recommendation</b>	<p><b>In people with angiographically normal coronary arteries and continuing anginal symptoms, consider a diagnosis of cardiac syndrome X.</b></p> <p><b>Continue drug treatment for stable angina only if it improves the symptoms of the person with suspected cardiac syndrome X.</b></p>
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**Relative values of different outcomes**

The evidence available in this review only included evidence for limited outcomes over short periods of time. Longer term morbidity and mortality outcomes were not available.

Evidence from placebo controlled trials indicated improvement of ischaemic episodes and duration over a short time period.

**Trade**

**off between clinical benefits and harms**

**Economic considerations**

No economic evidence was found on this question.

**Quality of evidence**

The evidence for available outcomes was of moderate quality.

**Other considerations**

Syndrome X is a diagnosis made following investigation with coronary angiography.

Patients are therefore already likely to be taking or to have tried one or more standard anti-anginal drugs. The GDG made a consensus recommendation that patients who were receiving symptomatic benefit should stay on anti-anginal drugs if they were benefiting from them. The evidence does not support use of standard anti-anginal drugs for any longer term benefit.

1 **19.3 Drugs for secondary prevention for people with syndrome X**

2 The use of aspirin, statins and ACE inhibitors have resulted in significant benefits for  
3 many people with cardiac conditions. The GDG were interested in whether these  
4 drugs were beneficial to patients who do not have evidence of coronary artery  
5 disease but have angina type pain and evidence of ischaemia. Studies were found  
6 examining the benefit of statins and a combination of statins and ace inhibitors.

7 **19.3.1 Clinical evidence**

8 The "Review Protocol" for this topic can be found in Appendix C, the "Search  
9 Strategies" in Appendix D, the "List of Included and Excluded Studies" in Appendix  
10 E1, the "Clinical Evidence Tables" in Appendix E2, and the "Forest Plots" in Appendix  
11 F

1 **Table 19.10: Statins vs. placebo for Cardiac Syndrome X**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Statins	Placebo	Relative (95% CI)	Absolute	
<b>Total exercise time (Sec) (follow-up 3 months; range of scores: -, better indicated by more)</b>											
Kayikcioglu 2003[229] (d)	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	19	19	-	MD 78 higher (11.17 lower to 167.17 higher)	⊕⊕⊕○ MODERATE
<b>Time to 1mm ST depression (Sec) (follow-up 3 months; range of scores: -, better indicated by more)</b>											
Kayikcioglu 2003[229]; Fabian 2004[230] (c)	randomised trial	serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	none	39	39	-	MD 48.36 lower (60.71 to 36.02 lower)	⊕⊕⊕○ MODERATE
<b>Hospitalisation for worsening of angina (follow-up 3 months)</b>											
Kayikcioglu 2003[229]	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	1/19 (5.3%)	1/19 (5.3%)	RR 1 (0.07 to 14.85)	0 fewer per 1000 (from 49 fewer to 734 more)	⊕⊕⊕○ MODERATE

- (a) Single blind, randomised, baseline comparisons made. Allocation concealment not reported, 0.5% drop out, intention to treat analysis not reported.
- (b) Fabian 2004[230]: Randomised, baseline comparisons made. Allocation concealment not reported, blinding not reported, drop out rate not reported, intention to treat analysis not reported. Kayikcioglu 2003[229]: Single blind, randomised, baseline comparisons made. Allocation concealment not reported, 0.5% drop out, intention to treat analysis not reported.
- (c) Drug dosage: Fabian 2004[230] - Simvastatin 20 mg/day
- (d) Drug dosage: Pravastatin 40 mg/day.

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1 **Table 19.11: Angiotensin-Converting Enzyme Inhibitors + statins vs. placebo for Cardiac Syndrome X**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Angiotensin-Converting Enzyme Inhibitors and statins	placebo	Relative (95% CI)	Absolute	
<b>Seattle Angina Questionnaire angina frequency score (follow-up 6 months; range of scores: -; better indicated by less)</b>											
Pizzi 2004[231] (a)	randomised trial	no serious limitations	no serious inconsistency	no serious indirectness	no serious imprecision	none	22	23	-	MD 19.7 higher (12.51 to 26.89 higher)	⊕⊕⊕⊕ HIGH
<b>Seattle Angina Questionnaire Quality of life score (follow-up 6 months; range of scores: -; better indicated by less)</b>											
Pizzi 2004[231] (a)	randomised trial	no serious limitations	no serious inconsistency	no serious indirectness	no serious imprecision	none	22	23	-	MD 24.6 higher (18.38 to 30.82 higher)	⊕⊕⊕⊕ HIGH
<b>Seattle Angina Questionnaire summary score (follow-up 6 months; range of scores: -; better indicated by less)</b>											
Pizzi 2004[231] (a)	randomised trial	no serious limitations	no serious inconsistency	no serious indirectness	no serious imprecision	none	22	23	-	MD 20.9 higher (15.5 to 26.3 higher)	⊕⊕⊕⊕ HIGH
<b>Peak exercise time (s) (follow-up 6 months; range of scores: -; better indicated by less)</b>											
Pizzi 2004[231] (a)	randomised trial	no serious limitations	no serious inconsistency	no serious indirectness	no serious imprecision	none	22	23	-	MD 67.2 higher (19.27 to 115.13 higher)	⊕⊕⊕⊕ HIGH
<b>ST depression (mV) (follow-up 6 months; range of scores: -; better indicated by less)</b>											
Pizzi 2004[231] (a)	randomised trial	no serious limitations				none	22	23	-	MD 0.09 lower (0.44 lower to 0.26 higher)	⊕⊕⊕⊕ HIGH
<b>Flow-mediated Dilation of brachial artery (%) (follow-up 6 months; range of scores: -; better indicated by less)</b>											
Pizzi 2004[231] (a)	randomised trial	no serious limitations	no serious inconsistency	no serious indirectness	no serious imprecision	none	22	23	-	MD 1.9 higher (1.04 to 2.76 higher)	⊕⊕⊕⊕ HIGH

2 (a) Drug dosage: ramipril (10mg/d) and atorvastatin (40mg/d)

1 **19.3.2 Economic evidence**

2 No economic studies were identified on this question.

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4 **19.3.3 Evidence statements**

**Clinical**

**Statins for cardiac syndrome X**

**Kayikcioglu 2003[229]; Fabian 2004[230]:** Evidence from 2 underpowered RCTs shows that Time to 1mm ST depression (sec) was significantly longer in the statins group compared to placebo [MD -48.36 (-60.71 to -36.02). [Follow-up 3 months]

**Kayikcioglu 2003[229]:** Evidence from one underpowered RCT shows that there was no significant difference between Statins and placebo for Total exercise time (sec) [MD 78 (-11.17 to 167.17)] and hospitalisation for worsening of angina [RR 1 (0.07 to 14.85)] [Follow-up 3 months].

**ACE Inhibitors + Statins for cardiac syndrome X**

**Pizzi 2004[231]:** Evidence from one RCT shows that angina frequency [MD 19.70 [12.51, 26.89]], Quality of Life [MD 24.60 [18.38, 30.82]], peak exercise time [MD 67.20 [19.27, 115.13]] and flow mediated dilation of brachial artery [MD1.90 [1.04, 2.76]] were significantly improved in the ACE inhibitors + statins group compared to placebo. There was no significant difference between groups for ST segment depression [MD -0.09 [-0.44 to 0.26]]. [ follow up 6 months]

**Economic**

No economic evidence was found on this question.

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6 **19.3.4 Recommendations and link to evidence**

<b>Recommendation</b>	<b>Do not routinely offer drugs for the secondary prevention of cardiovascular disease to people with suspected cardiac syndrome X.</b>
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**Relative values of different outcomes**

The GDG were interested in morbidity and mortality outcomes for interventions for people with syndrome X. They were aware however that this evidence was unlikely to be available and accepted evidence on short term outcomes.

**Trade off between clinical benefits and harms**

**Economic considerations**      Secondary prevention was not shown to add any benefits in people with suspected cardiac syndrome X. Therefore it is unlikely that this therapy is cost-effective.

**Quality of evidence**

**Other considerations**      No evidence was found examining the benefit of aspirin or ace inhibitors in people with syndrome X. The GDG considered that given the lack of evidence and potential risks and cost of using these drugs they should not be offered to people with syndrome X. The outcomes for statins versus placebo were ECG changes only and the GDG did not consider this adequate evidence to recommend use of statins. Quality of Life and angina score outcomes were available for combination of statin and ace inhibitor versus placebo but the study was small.

1    **19.4 Clinical/cost-effectiveness and safety of non-pharmacological**  
2                                    **treatments for syndrome X**

3    **19.4.1      Clinical Evidence**

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5                    The “Review Protocol” for this topic can be found in Appendix C, the “Search  
6                    Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
7                    E1, the “Clinical Evidence Tables” in Appendix E2, and the “Forest Plots” in Appendix  
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1 **Table 19.12: Exercise programme + symptoms monitoring vs. symptoms monitoring for Cardiac Syndrome X**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							Cardiac rehabilitation exercise programme + symptoms monitoring	symptoms monitoring	Relative (95% CI)	Absolute	
<b>HADS total score (follow-up 8 weeks; range of scores: -; better indicated by less)</b>											
Asbury 2008[232] (b)	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	32	32	-	MD 1.4 higher (1.14 lower to 3.94 higher)	⊕⊕⊕○ MODERATE
<b>SF36 physical functioning (follow-up 8 weeks; range of scores: -; better indicated by more)</b>											
Asbury 2008[232]	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	32	32	-	MD 1.8 higher (8.48 lower to 12.08 higher)	⊕⊕⊕○ MODERATE
<b>SF-36 pain (follow-up 8 weeks; range of scores: -; better indicated by more)</b>											
Asbury 2008[232]	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	32	32	-	MD 1.3 higher (9.15 lower to 11.75 higher)	⊕⊕⊕○ MODERATE
<b>SF-36 general health (follow-up 8 weeks; range of scores: -; better indicated by more)</b>											
Asbury 2008[232]	randomised trial	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	32	32	-	MD 3.9 higher (5.86 lower to 13.66 higher)	⊕⊕⊕○ MODERATE
<b>Symptom frequency (better indicated by lower values)</b>											
Asbury 2008[232]	randomised trials	serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	32	32	-	MD 2.6 lower (4.1 to 1.1 lower)	⊕⊕⊕○ MODERATE

(a) Small pilot study

(b) Cardiac rehabilitation: 8-week group-based phase III CR exercise programme: outpatient cardiovascular exercise programme designed to improve aerobic conditioning, functional capacity, muscular strength, endurance and flexibility. Each class was approx 80minutes long

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1 **Table 19.13: Physical training vs. normal activity for Cardiac Syndrome X**

Quality assessment							Summary of findings				Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		
							Cardiac rehabilitation physical training	normal activity	Relative (95% CI)	Absolute	
<b>Distance walked (m) (follow-up 8 weeks; range of scores: -; better indicated by more)</b>											
Tyni-Lenne 2002[233] (c)	randomised trial	very serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	7	7	-	MD 42 higher (7.79 lower to 91.79 higher)	⊕⊕○○ LOW
<b>peak heart rate (beats/min) (follow-up 8 weeks; range of scores: -; better indicated by less)</b>											
Tyni-Lenne 2002[233]	randomised trial	very serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	7	7	-	MD 4 lower (18.61 lower to 10.61 higher)	⊕⊕○○ LOW
<b>exertion (Borg RPE) (follow-up 8 weeks; range of scores: -; better indicated by less)</b>											
Tyni-Lenne 2002[233]	randomised trial	very serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	7	7	-	MD 1 lower (3.67 lower to 1.67 higher)	⊕⊕○○ LOW
<b>pain onset (min) after exercise (follow-up 8 weeks; range of scores: -; better indicated by more)</b>											
Eriksson 2000[234]	randomised trial	very serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	none	7	10	-	MD 2 higher (0.31 lower to 4.31 higher)	⊕⊕○○ LOW
<b>max pain (Borg CR-10 after exercise) (follow-up 8 weeks; range of scores: -; better indicated by less)</b>											
Eriksson 2000[234]	randomised trial	very serious (b)	no serious inconsistency	no serious indirectness	no serious imprecision	none	7	10	-	MD 1 lower (1.97 to 0.03 lower)	⊕⊕○○ LOW

- (a) Very small sample size, unclear randomisation and allocation concealment methods
- (b) Very small sample size, no description of randomisation, allocation concealment or blinding
- (c) Physical programme: outpatient group-based under supervision by physical therapist. Endurance training on cycle ergometer 3 times a week for 8 weeks at the intensity of 50% of the peak work rate achieved in VO2 max test. The training was 30 minutes.

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1 **Table 19.14: Physical training vs. relaxation therapy for Cardiac Syndrome X**

Quality assessment							Summary of findings				Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		
							cardiac rehabilitation physical training	relaxation therapy	Relative (95% CI)	Absolute	
<b>Distance walked (m) (follow-up 8 weeks; range of scores: -; better indicated by less)</b>											
Tyni-Lenne 2002[233]	randomised trial (b)	very serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	7	7	-	MD 22 higher (28.3 lower to 72.3 higher)	⊕⊕○○ LOW
<b>peak heart rate (beats/min) (follow-up 8 weeks; range of scores: -; better indicated by less)</b>											
Tyni-Lenne 2002[233]	randomised trial	very serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	7	7	-	MD 11 lower (28.29 lower to 6.29 higher)	⊕⊕○○ LOW
<b>exertion (Borg RPE) (follow-up 8 weeks; range of scores: -; better indicated by less)</b>											
Tyni-Lenne 2002[233]	randomised trial	very serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	7	7	-	MD 1 lower (4.14 lower to 2.14 higher)	⊕⊕○○ LOW

(a) Very small sample size, unclear randomisation and allocation concealment methods

(b) Interventions in the study: physical programme: outpatient group-based under supervision by physical therapist. Endurance training on cycle ergometer 3 times a week for 8 weeks at the intensity of 50% of the peak work rate achieved in VO2 max test. The training was 30minutes. Relaxation training consisted of a modified Jacobson approach and autogenous training for one hour at a time.

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6 **Table 19.15: Relaxation therapy vs. normal activity for Cardiac Syndrome X**

Quality assessment							Summary of findings				Quality
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		
							cardiac rehabilitation relaxation therapy	normal activity	Relative (95% CI)	Absolute	
<b>Distance walked (m) (follow-up 8 weeks; range of scores: -; better indicated by less)</b>											
Tyni-Lenne 2002[233] (b)	randomised trial	very serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	7	7	-	MD 20 higher (28.72 lower to 68.72 higher)	⊕⊕○○ LOW
<b>peak heart rate (beats/min) (follow-up 8 weeks; range of scores: -; better indicated by less)</b>											
Tyni-Lenne 2002[233]	randomised trial	very serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	7	7	-	MD 7 higher (6.98 lower to 20.98 higher)	⊕⊕○○ LOW
<b>exertion (Borg RPE) (follow-up 8 weeks; range of scores: -; better indicated by less)</b>											
Tyni-Lenne 2002[233]	randomised trial	very serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	7	7	-	MD 0 higher (2.67 lower to 2.67 higher)	⊕⊕○○ LOW

(a) Very small sample size, unclear randomisation and allocation concealment methods

(b) Relaxation training consisted of a modified Jacobson approach and autogenous training for one hour at a time

1 **Table 19.16: Exercise + relaxation training vs. exercise training for Cardiac Syndrome X**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							cardiac rehabilitation exercise + relaxation training	exercise training	Relative (95% CI)	Absolute	
<b>pain onset after exercise (min) (follow-up 8 weeks; range of scores: -; better indicated by more)</b>											
Eriksson 2000[234] (b)	randomised trial	very serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	7	10	-	MD 0 higher (2.34 lower to 2.34 higher)	⊕⊕○○ LOW
<b>max pain (Borg CR-10) after exercise (follow-up 8 weeks; range of scores: -; better indicated by less)</b>											
Eriksson 2000[234]	randomised trial	very serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	7	10	-	MD 1 higher (0.05 lower to 2.05 higher)	⊕⊕○○ LOW

- (a) Very small sample size, no description of randomisation, allocation concealment or blinding
- (b) Outpatient activity in outpatient setting supervised by physical therapist. Body awareness training consisted of body and mind relaxation performed twice a week for 8 weeks. Exercise training was performed on cycle ergometer 3 times a week for 8 weeks. Training was 30minutes and intensity was 50% of peak work rate determined at onset of study

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7 **Table 19.17: Exercise + relaxation training vs. normal activity for Cardiac Syndrome X**

Quality assessment							Summary of findings				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	No of patients		Effect		Quality
							cardiac rehabilitation exercise + relaxation training	normal activity	Relative (95% CI)	Absolute	
<b>pain onset after exercise (min) (follow-up 8 weeks; range of scores: -; better indicated by more)</b>											
Eriksson 2000[234] (b)	randomised trial	very serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	7	7	-	MD 3 higher (0.69 to 5.31 higher)	⊕⊕○○ LOW
<b>max pain (Borg CR-10) after exercise (follow-up 8 weeks; range of scores: -; better indicated by less)</b>											
Eriksson 2000[234]	randomised trial	very serious (a)	no serious inconsistency	no serious indirectness	no serious imprecision	none	7	7	-	MD 0 higher (0.97 lower to 0.97 higher)	⊕⊕○○ LOW

- (a) Very small sample size, no description of randomisation, allocation concealment or blinding
- (b) Outpatient activity in outpatient setting supervised by physical therapist. Body awareness training consisted of body and mind relaxation performed twice a week for 8 weeks. Exercise training was performed on cycle ergometer 3 times a week for 8 weeks. Training was 30minutes and intensity was 50% of peak work rate determined at onset of study

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1 **Additional data:**

2 **Psychological treatment vs. control for Cardiac syndrome X:**

3 **Potts 1999[235]**

4 The data from the study could not be analysed as SD for the eman values were not  
5 reported.

6 N=60 (n=34 immediate treatment and n=26 waiting control)

7 Intervention: Psychological treatment package consisting of education, relaxation,  
8 breathing training, graded exposure to activity and exercise, and the use of thought  
9 diaries to record and challenging automatic thoughts about heart disease.

10 Groups met weekly for 4 weeks, then every 2 weeks for a further 4 weeks. Each  
11 session lasted 2 hours, with a short break. Subjects were asked to practice various  
12 exercises at home between sessions, and to report their progress at the beginning of  
13 subsequent sessions. Treatment was broadly behavioural in orientation, based on a  
14 manual developed via an initial pilot group, and was supplemented by written  
15 material given to subjects at each session.

16 Control group was assigned to a waiting period before being reassessed and then  
17 entering treatment.

18 **Results:**

19 Treatment was associated with a significantly greater reduction in chest pain episode  
20 frequency (-3 vs. 0; p=0.01), than waiting control group. There was no significant  
21 difference between the treatment and control groups in changes in chest pain severity  
22 (-5.9 vs. 0.8; NS) or duration (min) (-1.6 vs. -0.5; NS,) although there were non  
23 significant trends to improvement in the treatment group, and the range of variation  
24 was very wide. Treatment was also associated with significant reductions in both the  
25 anxiety (-1.5 vs.0; p=0.05) and depression (-2 vs. 0; p=0.05) subscales of the HAD\*,  
26 the total disability score of the SIP\*\* (6.5 vs. 1.4; p=0.05), and two of the 4 subscales  
27 of the NHP \*\*\* (pain: 5 vs.0; p=0.05 and energy: -24 vs.0; p=0.01). Exercise  
28 duration (min) improved significantly. (1.3 vs. 0.1; p=0.5).

29 Note: All above values are medians, negative values indicating reductions.

30 \*Hospital anxiety Depression scale (HAD)-A 14 item inventory covering non somatic  
31 symptoms of anxiety and depression, intended for use in medical populations. It yields  
32 separate scores for anxiety and depression, with cut offs indicating caseness above  
33 11.

34 \*\*Sickness Impact Profile (SIP) – A 136 item inventory yielding measures of the impact  
35 of illness on various domains of everyday life, as well as an overall disability score.

36 \*\*\*Nottingham Health Profile (NHP) – A 24 item inventory quantifying the  
37 impairments due to illness in six areas.

38

- 1 **19.4.2 Economic evidence**  
2 No economic studies were found on this question.

3

- 4 **19.4.3 Evidence statements**

**Clinical Exercise programme + symptoms monitoring vs. symptoms monitoring for cardiac syndrome X**

**Asbury 2008[232]:** Evidence from one RCT shows that there was significantly lower symptom frequency in the exercise programme +symptom monitoring group compared to control [MD -2.6 (-4.1 to -1.1)]. No significant difference was found for the other outcomes (HADS total score, SF-36 physical function, SF-36 pain, SF-36 general health). [follow-up 8 weeks]

**Physical training vs. normal activity for cardiac syndrome X**

**Tyni-Lenne 2002[233]; Eriksson 2000[234]:** Evidence from two RCT shows that max pain after exercise was significantly reduced in the physical training group compared to the normal activity group [MD-1.00 [-1.97, -0.03]]. There was no significant difference between physical training and normal activity for all other outcomes (distance walked [MD [42.00 [-7.79, 91.79]], peak heart rate [MD -4.00 [-18.61, 10.61]], exertion [MD -1.00 [-3.67, 1.67]], pain onset after exercise [MD 2.00 [-0.31, 4.31]]). [follow-up 8 weeks]

**Physical training vs. relaxation therapy for cardiac syndrome X**

**Tyni-Lenne 2002[233]:** Evidence from one RCT shows that there was no significant difference between physical training and relaxation therapy for distance walked [MD 22 (-28.3 to 72.3)], peak heart rate [MD -11 (-28.29 to 6.29)] and exertion [MD -1 (-4.14 to 2.14)] [follow up 8 weeks]

**Relaxation therapy vs. normal activity for cardiac syndrome X**

**Tyni-Lenne 2002[233]:** Evidence from one RCT shows that there was no significant difference between relaxation therapy and normal activity for distance walked [MD 20 (-28.72 to 68.72)] , peak heart rate [MD 7 (-6.98 to 20.98)] and exertion [MD 0 (-2.67 to 2.67)] [follow up 8 weeks]

**Exercise +relaxation training vs. exercise training for cardiac syndrome X**

**Eriksson 2000[234]:** Evidence from one RCT shows that there was no significant difference between exercise +relaxation training and exercise training for pain onset after exercise [MD 0.00 (-2.34 to 2.34)] and max pain after exercise [MD 1 (-0.05 to 2.05)] [follow up 8 weeks]

**Exercise +relaxation training vs. normal activity for cardiac syndrome X**

**Eriksson 2000[234]:** Evidence from one RCT shows that there was no

significant difference between exercise +relaxation training and normal activity for pain onset after exercise [MD 3 (0.69to 5.31)] and max pain after exercise [MD 0.00 (-0.97 to 0.97)] [follow up 8 weeks]

**Economic** No economic evidence was found on this question.

1 **19.4.4 Recommendations and link to evidence**

<b>Recommendation</b>	<b>No recommendation was made</b>
<b>Relative values of different outcomes</b>	When considering the value of rehabilitation for people with syndrome X, the GDG were interested in improvements in quality of life as well as longer term outcomes such as angina frequency and morbidity and mortality.
<b>Trade off between clinical benefits and harms</b>	
<b>Economic considerations</b>	Rehabilitation is associated with important costs. The clinical evidence review did not indicate effectiveness of programmes of rehabilitations and programmes are therefore not likely to be cost effective.
<b>Quality of evidence</b>	The GDG considered that the evidence available was of the benefit of exercise and different exercise programmes for people with syndrome X and did not address the benefit of comprehensive cardiac rehabilitation programmes. The quality of evidence available was low.
<b>Other considerations</b>	<p>The GDG did not make a recommendation about cardiac rehabilitation for people with syndrome X.</p> <p>The quality of evidence was low but one moderate quality evidence study did suggest that exercise was beneficial but the review did not support any particular exercise programme over normal activity. The GDG did not consider that exercise was harmful to people with syndrome X and that it was important people with syndrome X are given positive encouragement to take part in exercise and to be as active as possible. The study by Potts 1999[235]reported only mean data but suggested that people with syndrome X benefit might benefit programmes including attention to beliefs about angina. The GDG considered that people with syndrome X would be similar to those with stable angina in their needs for and response to appropriate information and support tailored to their individual needs.</p>

1

## 2 **19.5 Stress echocardiography in people with cardiac syndrome X**

### 3 **19.5.1 Clinical question**

4 In adults with cardiac Syndrome X (i.e. those with chest pain and normal coronary  
5 arteries) what is the incremental value/effectiveness of functional tests for prognostic  
6 risk stratification in prediction of adverse cardiac outcomes?

### 7 **19.5.2 Clinical evidence**

8 The “Review Protocol” for this topic can be found in Appendix C, the “Search  
9 Strategies” in Appendix D, the “List of Included and Excluded Studies” in Appendix  
10 E1, and the “Clinical Evidence Tables” in Appendix E2.

11

12 **Bigi 2002[183]** (N=125) assessed the incremental prognostic value of dobutamine  
13 and dipyridamole stress echocardiography in patients with known or suspected  
14 coronary artery disease.

15 The outcome events were cardiac death, non fatal infarction, and unstable angina  
16 assessed at a mean follow-up of 36 months (range 6 to 80).

17 Target events occurred in 9 patients: 2 cardiac deaths, 5 non fatal MI, and 2  
18 hospitalisations for unstable angina. Six of the 9 patients with cardiac events had  
19 positive stress echocardiography.

20 Hypertension, positive stress echocardiography, and peak wall motion score index  
21 were multivariate predictors of outcome, but stress echocardiography provided an  
22 87.5% increase in the global chi-square ( $p < 0.001$ ). The event free survival of  
23 patients with positive stress echocardiography was significantly lower compared with  
24 those with negative test (Hazard ratio 4.7 95% CI 1.3 to 47)

25

1 **Table 19.18: Bigi 2002[183], Multivariate predictors of outcome**

Variables	Chi-square	Odds ratio	95% CI	P-value
Clinical				
Hypertension	5.7	13	1.6 to 105	0.01
Echocardiographic				
Positive SE	3.8	3.6	1 to 14	0.05
Peak WMSI	8.1	5.0	1.6 to 15	0.004

2

3 **Summary:** One low quality study showed that **stress echocardiography** offered  
4 incremental prognostic value in prediction of adverse cardiac outcomes (cardiac  
5 death, non fatal infarction or unstable angina) in people with chest pain and normal  
6 or slightly narrowed coronary arteries. The results should be considered with caution  
7 as the study had very few events, small sample size, and a short follow-up period.

8

9 **19.5.3 Economic evidence**

10 No relevant studies were found. Studies reporting the cost per case detected were not  
11 included as this question was addressed in the Chest Pain Guideline (CG95).

12 We looked for the costs of the individual tests from UK sources. We found that the  
13 unit cost of stress echocardiography is £435[184].

14

15 **19.5.4 Evidence statements****Clinical** **Stress echocardiography**

**Bigi 2002[183]:** Evidence from one study shows that stress echocardiography offers incremental prognostic information in prediction of cardiac outcomes (cardiac death, non fatal infarction or unstable angina) in patients with chest pain and normal or slightly narrowed coronary arteries. [Mean follow-up 36 months (range 6 to 80)].

**Economic** No economic evidence was found on this question. A simple cost analysis showed that stress echocardiography has a cost of £435 per test.

16

1 **19.5.5 Recommendations and link to evidence**

<b>Recommendation</b>	<b>No recommendation was made</b>
<b>Relative values of different outcomes</b>	
<b>Trade off between clinical benefits and harms</b>	
<b>Economic considerations</b>	No health economic evidence was available but the cost of testing is significant
<b>Quality of evidence</b>	One low quality study was found
<b>Other considerations</b>	The GDG agreed not to make a recommendation. The care of people with cardiac syndrome X is difficult. The diagnosis is made after angiography. The evidence does not support routine use of stress ECHO but the GDG recognized that further investigation may have a role in individual patients.

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