2013 ESC guidelines on the management of stable coronary artery disease

The Task Force on the management of stable coronary artery disease of the European Society of Cardiology

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<td>201TI</td>
<td>thallium 201</td>
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<td>ABCB1</td>
<td>ATP-binding cassette sub-family B member 1</td>
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<td>ABI</td>
<td>ankle-brachial index</td>
</tr>
<tr>
<td>ACC</td>
<td>American College of Cardiology</td>
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<tr>
<td>ACCF</td>
<td>American College of Cardiology Foundation</td>
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<td>ACCF</td>
<td>American College of Cardiology Foundation</td>
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<tr>
<td>ACE</td>
<td>angiotensin converting enzyme</td>
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<td>ACIP</td>
<td>Asymptomatic Cardiac Ischaemia Pilot</td>
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<td>ACS</td>
<td>acute coronary syndrome</td>
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<tr>
<td>ADA</td>
<td>American Diabetes Association</td>
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<td>ADP</td>
<td>adenosine diphosphate</td>
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<td>AHA</td>
<td>American Heart Association</td>
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<tr>
<td>ARB</td>
<td>angiotensin II receptor antagonist</td>
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<tr>
<td>ART</td>
<td>Arterial Revascularization Trial</td>
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<td>ASCOT</td>
<td>Anglo-Scandinavian Cardiac Outcomes Trial</td>
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<td>ASSERT</td>
<td>Asymptomatic atrial fibrillation and Stroke Evaluation in pacemaker patients and the atrial fibrillation Reduction atrial pacing Trial</td>
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<td>atrioventricular</td>
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<td>BIMA</td>
<td>bilateral internal mammary artery</td>
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<tr>
<td>BMI</td>
<td>body mass index</td>
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<tr>
<td>BMS</td>
<td>bare metal stent</td>
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<tr>
<td>BNP</td>
<td>B-type natriuretic peptide</td>
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<tr>
<td>BP</td>
<td>blood pressure</td>
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<tr>
<td>b.p.m.</td>
<td>beats per minute</td>
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<td>CABG</td>
<td>coronary artery bypass graft</td>
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<td>Clopidogrel for High Atherothrombotic Risk and Ischaemic Stabilization, Management and Avoidance</td>
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<td>CI</td>
<td>confidence interval</td>
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<td>CKD</td>
<td>chronic kidney disease</td>
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<td>Chronic Kidney Disease Epidemiology Collaboration</td>
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<td>CMR</td>
<td>cardiac magnetic resonance</td>
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<td>CORONARY</td>
<td>The CABG Off or On Pump Revascularization Study</td>
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<td>CPG</td>
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<td>DANAMI</td>
<td>Danish trial in Acute Myocardial Infarction</td>
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<td>DAPT</td>
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<td>DBP</td>
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<td>EACTS</td>
<td>European Association for Cardiothoracic Surgery</td>
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<td>EECP</td>
<td>enhanced external counterpulsation</td>
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<td>EASD</td>
<td>European Association for the Study of Diabetes</td>
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<td>ECG</td>
<td>electrocardiogram</td>
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<td>Echo</td>
<td>echocardiogram</td>
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<td>ESC</td>
<td>European Society of Cardiology</td>
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<td>Evaluation of XIENCE PRIME or XIENCE V vs. Coronary Artery Bypass Surgery for Effectiveness of Left Main Revascularization</td>
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<td>Fractional Flow Reserve vs. Angiography for Multivessel Evaluation</td>
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<td>FDA</td>
<td>Food &amp; Drug Administration (USA)</td>
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<td>FFR</td>
<td>fractional flow reserve</td>
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<td>GFR</td>
<td>glomerular filtration rate</td>
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<td>HbA1c</td>
<td>glycated haemoglobin</td>
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<td>HDL</td>
<td>high density lipoprotein</td>
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<td>high density lipoprotein cholesterol</td>
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<td>HR</td>
<td>hazard ratio</td>
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<td>HRT</td>
<td>hormone replacement therapy</td>
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<td>hs-CRP</td>
<td>high-sensitivity C-reactive protein</td>
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<td>HU</td>
<td>Hounsfield units</td>
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<td>ICA</td>
<td>invasive coronary angiography</td>
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<td>IMA</td>
<td>internal mammary artery</td>
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<td>IONA</td>
<td>Impact Of Nicorandil in Angina</td>
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<td>ISCHEMIA</td>
<td>International Study of Comparative Health Effectiveness with Medical and Invasive Approaches</td>
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<td>IVUS</td>
<td>intravascular ultrasound</td>
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<td>JASP</td>
<td>Japanese Stable Angina Pectoris</td>
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<td>KATP</td>
<td>ATP-sensitive potassium channels</td>
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<td>LAD</td>
<td>left anterior descending</td>
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<td>LBBB</td>
<td>left bundle branch block</td>
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<td>LIMA</td>
<td>Left internal mammary artery</td>
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<td>LDL</td>
<td>low density lipoprotein</td>
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<td>low density lipoprotein cholesterol</td>
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<td>LM</td>
<td>left main</td>
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<td>left main stem</td>
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<td>left ventricular ejection fraction</td>
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<td>left ventricular hypertrophy</td>
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<td>MACE</td>
<td>major adverse cardiac events</td>
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<td>MASS</td>
<td>Medical, Angioplasty, or Surgery Study</td>
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<td>non-steroidal anti-inflammatory drugs</td>
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<td>Occluded Artery Trial</td>
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<td>optimal medical therapy</td>
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<td>polyunsaturated fatty acid</td>
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<td>quality of life</td>
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<td>systolic blood pressure</td>
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<td>SCAD</td>
<td>stable coronary artery disease</td>
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<td>single internal mammary artery</td>
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<td>TC</td>
<td>total cholesterol</td>
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1. Preamble

Guidelines summarize and evaluate all evidence available, at the time of the writing process, on a particular issue with the aim of assisting physicians in selecting the best management strategies for an individual patient with a given condition, taking into account the impact on outcome, as well as the risk–benefit ratio of particular diagnostic or therapeutic means. Guidelines are not substitutes but are complements for textbooks, and cover the ESC Core Curriculum topics. Guidelines and recommendations should help physicians to make decisions in their daily practice; however, the final decisions concerning an individual patient must be made by the responsible physician(s).

A great number of Guidelines have been issued in recent years by the European Society of Cardiology (ESC) as well as by other societies and organisations. Because of the impact on clinical practice, quality criteria for the development of guidelines have been established in order to make all decisions transparent to the user. The recommendations for formulating and issuing ESC Guidelines can be found on the ESC website (http://www.escardio.org/guidelines-surveys/esc-guidelines/about/Pages/rules-writing.aspx). ESC Guidelines represent the official position of the ESC on a given topic and are regularly updated.

Members of this Task Force were selected by the ESC to represent professionals involved with the medical care of patients with this pathology. Selected experts in the field undertook a comprehensive review of the published evidence for the diagnosis, management and/or prevention of a given condition according to the ESC Committee for Practice Guidelines (CPG) policy. A critical evaluation of diagnostic and therapeutic procedures was performed, including assessment of the risk—benefit ratio. Estimates of expected health outcomes for larger populations were included, where data exist. The level of evidence and the strength of recommendation of particular treatment options were weighed and graded according to predefined scales, as outlined in Tables 1 and 2.

The experts of the writing and reviewing panels completed Declaration of Interest forms where real or potential sources of conflicts of interest might be perceived. These forms were compiled into one file and can be found on the ESC website (http://www.escardio.org/guidelines). Any changes in declarations of interest that arise during the writing period must be notified to the ESC and updated. The Task Force received its entire financial support from the ESC, without any involvement from healthcare industry.

The ESC CPG supervises and co-ordinates the preparation of new Guidelines produced by Task Forces, expert groups or consensus panels. The Committee is also responsible for the endorsement process of these Guidelines. The ESC Guidelines undergo extensive review by the CPG and external experts. After appropriate revisions, they are approved by all the experts involved in the Task Force. The finalized document is approved by the CPG for publication in the European Heart Journal.

The task of developing ESC Guidelines covers not only the integration of the most recent research, but also the creation of educational tools and implementation programmes for the recommendations. To implement the guidelines, condensed pocket editions, summary slides, booklets with essential messages, electronic versions for digital applications (smartphones etc.) are produced. These versions are abridged and thus, if needed, one should always refer to the full
2. Introduction

These guidelines should be applied to patients with stable known or suspected coronary artery disease (SCAD). This condition encompasses several groups of patients: (i) those having stable angina pectoris or other symptoms felt to be related to coronary artery disease (CAD) such as dyspnea; (ii) those previously symptomatic with known obstructive or non-obstructive CAD, who have become asymptomatic with treatment and need regular follow-up; (iii) those who report symptoms for the first time and are judged to already be in a chronic stable condition (for instance because history-taking reveals that similar symptoms were already present for several months). Hence, SCAD defines the different evolutionary phases of CAD, excluding the situations in which coronary artery thrombosis dominates clinical presentation (acute coronary syndromes).

However, patients who have a first or recurrent manifestation of angina but can be categorized as having a low-risk acute coronary syndrome (ACS) according to the current ACS guidelines of the ESC (no recurrence of chest pain, no signs of heart failure, no abnormalities in the resting electrocardiogram (ECG), no rise in markers of myocardial necrosis (preferably troponin) and hence are not candidates for swift intervention) should also be managed according to the algorithms presented in these Guidelines. Although routine screening of asymptomatic patients is discouraged, these guidelines can also be applied to asymptomatic patients presenting for further evaluation due to an abnormal test. The scope of the present Guidelines, therefore, spans from asymptomatic individuals to patients after stabilisation of an ACS.

The traditional understanding of SCAD is that of a disease causing exercise- and stress-related chest symptoms due to narrowings of ≥50% in the left main coronary artery and ≥70% in one or several of the major coronary arteries. Compared with the previous version of the Guidelines, the present edition considers not only such atherosclerotic narrowings, but also microvascular dysfunction and coronary vasospasm in the diagnostic and prognostic algorithms; the present Guidelines also distinguish diagnostic testing from prognostic assessment; they give increased importance to the pre-test probability (PTP) of disease strongly influencing the diagnostic algorithms and they take into account recent advances in technology, the importance of physiological assessment of CAD in the catheterisation laboratory and the increasing evidence that the prognostic benefit of revascularization may be less than has been traditionally expected.

In order to limit the length of the printed text, additional information, tables, figures and references are available as web addenda at the ESC website (www.escardio.org).

3. Definitions and pathophysiology (see web addenda)

Stable coronary artery disease is generally characterized by episodes of reversible myocardial demand/supply mismatch, related to ischaemia or hypoxia, which are usually inducible by exercise, emotion or other stress and reproducible—but, which may also be occurring spontaneously. Such episodes of ischaemia/hypoxia are commonly associated with transient chest discomfort (angina pectoris). SCAD also includes the stabilized, often asymptomatic, phases that follow an ACS.

Because the transition from unstable to stable syndromes is a continuum, without a clear boundary, angina at rest caused by coronary vasospasm may be regarded within the scope of SCAD, as in the present document or, conversely, within the scope of ACS as in some, but not in other, ACS guidelines. Recent use of ultrasensitive troponin tests has shown that episodes of minute troponin release—below the threshold for acute myocardial infarction—often occur in patients with stable CAD and this has been shown to have prognostic implications, also demonstrating the continuum of CAD subgroups.

The various clinical presentations of SCAD (see also section 6.1) are associated with different underlying mechanisms that mainly include: (i) plaque-related obstruction of epicardial arteries; (ii) focal or diffuse spasm of normal or plaque-diseased arteries; (iii) microvascular dysfunction and (iv) left ventricular dysfunction caused by prior acute myocardial necrosis and/or hibernation (ischaemic cardiomyopathy) (Table 3). These mechanisms may act singly or in combination. However, stable coronary plaques with and without previous revascularization may also be completely clinically silent. Additional information on the relationship between symptoms and underlying disease mechanisms, the histology of epicardial lesions, the definitions and pathogenesis of vasospasm, the

Table 2  Levels of evidence

<table>
<thead>
<tr>
<th>Level of evidence A</th>
<th>Data derived from multiple randomized clinical trials or meta-analyses.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Level of evidence B</td>
<td>Data derived from a single randomized clinical trial or large non-randomized studies.</td>
</tr>
<tr>
<td>Level of evidence C</td>
<td>Consensus of opinion of the experts and/or small studies, retrospective studies, registries.</td>
</tr>
</tbody>
</table>

The traditional understanding of SCAD is that of a disease causing exercise- and stress-related chest symptoms due to narrowings of ≥50% in the left main coronary artery and ≥70% in one or several of the major coronary arteries. The present Guidelines also distinguish diagnostic testing from prognostic assessment; they give increased importance to the pre-test probability (PTP) of disease strongly influencing the diagnostic algorithms and they take into account recent advances in technology, the importance of physiological assessment of CAD in the catheterisation laboratory and the increasing evidence that the prognostic benefit of revascularization may be less than has been traditionally expected.

In order to limit the length of the printed text, additional information, tables, figures and references are available as web addenda at the ESC website (www.escardio.org).
definition of microvascular dysfunction and ischaemic cardiomyopathy is available in sections 3.1–3.5 of the web addenda.

Myocardial ischaemia and hypoxia in SCAD are caused by a transient imbalance between blood supply and metabolic demand. The consequences of ischaemia occur in a predictable temporal sequence that involves:

1. Increased H+ and K+ concentration in the venous blood that drains the ischaemic territory
2. Signs of ventricular diastolic and subsequently systolic dysfunction with regional wall motion abnormalities
3. Development of ST–T changes
4. Cardiac ischaemic pain (angina).10

This sequence explains why imaging techniques based on perfusion, metabolism or wall motion are more sensitive than an ECG or symptoms in detecting ischaemia. Angina is ultimately caused by the release of ischaemic metabolites—such as adenosine—that stimulate sensitive nerve endings, although angina may be absent even with severe ischaemia owing, for instance, to impaired transmission of painful stimuli to the cortex and other as-yet-undefined potential mechanisms.11

The functional severity of coronary lesions can be assessed by measuring coronary flow reserve (CFR) and intracoronary artery pressures (fractional flow reserve, FFR). More detailed descriptions can be found in the web addenda.

### 4. Epidemiology

As SCAD is so multifaceted, its prevalence and incidence have been difficult to assess and numbers vary between studies, depending on the definition that has been used. For epidemiologic purposes, stable angina is essentially a diagnosis based on history and therefore relies on clinical judgement. The Rose angina questionnaire has a specificity of ~80–95%, but its sensitivity varies substantially from 20–80% when compared with clinical diagnosis, ECG findings and coronary angiography.

The prevalence of angina in population-based studies increases with age in both sexes, from 5–7% in women aged 45–64 years to 10–12% in women aged 65–84 and from 4–7% in men aged 45–64 years to 12–14% in men aged 65–84.13 Interestingly, angina is more prevalent in middle-aged women than in men, probably due to the higher prevalence of functional CAD—such as microvascular angina—in women, whereas the opposite is true in the elderly.

Available data suggest an annual incidence of uncomplicated angina pectoris of 1.0% in male western populations aged 45–65 years, with a slightly higher incidence in women under the age of 65.13,16 There is a steep increase with age and the incidence in men and women 75–84 years of age reaches almost 4%.16 The incidence of angina varies in parallel with observed international differences in CAD mortality.16,17

Temporal trends suggest a decrease in the annual death rate due to CAD.18 However, the prevalence of a history of diagnosed CAD does not appear to have decreased, suggesting that the prognosis of those with established CAD is improving. Improved sensitivity of diagnostic tools may additionally contribute to the contemporary high prevalence of diagnosed CAD.

Epidemiological data on microvascular angina and vasospastic angina are missing. However, recent clinical data suggest that abnormal coronary vasomotion is present in two-thirds of patients who suffer from stable angina but have no coronary stenoses at angiography.19

### 5. Natural history and prognosis

In many patients, early manifestations of CAD are endothelial dysfunction and microvascular disease. Both are associated with an increased risk of complications from CAD.20–22

Contemporary data regarding prognosis can be derived from clinical trials of anti-anginal and preventive therapy and/or revascularization, although these data are biased by the selected nature of the populations studied. From these, estimates for annual mortality rates range from 1.2–2.4% per annum, with an annual incidence of cardiac death between 0.6 and 1.4% and of non-fatal myocardial infarction (MI) between 0.6% in the Second Randomized Intervention Treatment of Angina (RITA-2) and 2.7% in the Clinical Outcomes Utilizing Revascularization and Aggressive Drug Evaluation (COURAGE) trials.23 These estimates are consistent with observational registry data.13,29

### Table 3  Main features of stable coronary artery disease

<table>
<thead>
<tr>
<th>Pathogenesis</th>
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<tbody>
<tr>
<td>Stable anatomical atherosclerotic and/or functional alterations of epicardial vessels and/or microcirculation</td>
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</table>

<table>
<thead>
<tr>
<th>Natural history</th>
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<tbody>
<tr>
<td>Stable symptomatic or asymptomatic phases which may be interrupted by ACS</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Mechanisms of myocardial ischaemia</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fixed or dynamic stenoses of epicardial coronary arteries; Microvascular dysfunction; Focal or diffuse epicardial coronary spasm; The above mechanisms may overlap in the same patient and change over time</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Clinical presentations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Effort induced angina caused by: • epicardial stenoses; • microvascular dysfunction; • vasconstriction at the site of dynamic stenosis; • combination of the above.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Rest angina caused by:</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Vasospasm (focal or diffuse) • epicardial focal; • epicardial diffuse; • microvascular; • combination of the above.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Asymptomatic:</th>
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</thead>
<tbody>
<tr>
<td>because of lack of ischaemia and/or of LV dysfunction; despite ischaemia and/or LV dysfunction.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Ischaemic cardiomyopathy</th>
</tr>
</thead>
</table>

ACS = acute coronary syndrome; LV = left ventricular; SCAD = stable coronary artery disease.
However, within the population with stable CAD, an individual’s prognosis can vary considerably, depending on baseline clinical, functional and anatomical characteristics. This is exemplified in the Reduction of Atherothrombosis for Continued Health (REACH) registry,30 which included very high-risk patients, many with peripheral arterial disease or previous MI and almost 50% with diabetes. Consequently, annual mortality rate was as high as 3.8% in this population30, whereas patients with non-obstructive plaques within the coronary arteries have an annual mortality rate of only 0.63%.

Prognostic assessment is an important part of the management of patients with SCAD. On the one hand, it is important to reliably identify those patients with more severe forms of disease, who may have an improvement in outcome with more aggressive investigation and—potentially—intervention, including revascularization. On the other hand, it is also important to identify those patients with a less severe form of disease and a good prognosis, thereby avoiding unnecessary invasive and non-invasive tests and revascularization procedures.

Conventional risk factors for the development of CAD—hypertension,31 hypercholesterolaemia,35 diabetes,36 sedentary lifestyle,37 smoking,34,38 and a family history—have an adverse influence on prognosis in those with established disease, presumably through their effect on the progression of atherosclerotic disease processes. However, appropriate treatment can reduce these risks.40–42 An elevated resting heart rate is also indicative of coronary disease.43 In general, the outcome is worse in patients with reduced left ventricular ejection fraction (LVEF) and heart failure, a greater number of diseased vessels, more proximal locations of coronary stenoses, greater severity of lesions, more extensive ischaemia, more impaired functional capacity, older age, significant depression and more severe angina.44–47

Prognostic assessments are conducted simultaneously, rather than separately, and many of the investigations used for diagnosis also offer prognostic information. However, for the purpose of clarity, the processes of obtaining diagnostic and prognostic information are dealt with separately in this text.

6. Diagnosis and assessment (see web addenda)

The diagnosis and assessment of SCAD involves clinical evaluation, including identifying significant dyslipidaemia, hyperglycaemia or other biochemical risk factors and specific cardiac investigations such as stress testing or coronary imaging. These investigations may be used to confirm the diagnosis of ischaemia in patients with suspected SCAD, to identify or exclude associated conditions or precipitating factors, assist in stratifying risk associated with the disease and to evaluate the efficacy of treatment. In practice, diagnostic and prognostic assessments are conducted simultaneously, rather than separately, and many of the investigations used for diagnosis also offer prognostic information. However, for the purpose of clarity, the processes of obtaining diagnostic and prognostic information are dealt with separately in this text.

6.1 Symptoms and signs (see web addenda)

A careful history remains the cornerstone of the diagnosis of chest pain. In the majority of cases, it is possible to make a confident diagnosis on the basis of the history alone, although physical examination and objective tests are often necessary to confirm the diagnosis, exclude alternative diagnoses,48 and assess the severity of underlying disease.

The characteristics of discomfort-related to myocardial ischaemia (angina pectoris) may be divided into four categories: location, character, duration and relationship to exertion and other exacerbating or relieving factors. The discomfort caused by myocardial ischaemia is usually located in the chest, near the sternum, but may be felt anywhere from the epigastrium to the lower jaw or teeth, between the shoulder blades or in either arm to the wrist and fingers.

The discomfort is often described as pressure, tightness or heaviness; sometimes strangled, constricting or burning. It may be useful to directly ask the patient for the presence of ‘discomfort’ as many do not feel ‘pain’ or ‘pressure’ in their chest. Shortness of breath may accompany angina, and chest discomfort may also be accompanied by less-specific symptoms such as fatigue or faintness, nausea, burning, restlessness or a sense of impending doom. Shortness of breath may be the sole symptom of SCAD and it may be difficult to differentiate this from shortness of breath caused by bronchopulmonary disease.

The duration of the discomfort is brief—no more than 10 min in the majority of cases and more commonly even minutes or less—but chest pain lasting for seconds is unlikely to be due to angina. An important characteristic is the relationship to exercise, specific activities or emotional stress. Symptoms classically appear or become more severe with increased levels of exertion—such as walking up an incline or against a breeze or in cold weather—and rapidly disappear within a few minutes when these causal factors abate. Exacerbations of symptoms after a heavy meal or after waking up in the morning are classical features of angina. Angina may be reduced with further exercise (walk-through angina) or on second exertion (warm-up angina).99 Buccal or sublingual nitrates rapidly relieve angina. The angina threshold—and hence symptoms—may vary considerably from day to day and even during the same day.

Definitions of typical and atypical angina have been previously published and are summarized in Table 4.50 Atypical angina is most frequently chest pain resembling that of typical angina in location and character, that is responsive to nitrates but has no precipitating factors. Often, the pain is described as starting at rest from a low level of intensity, which slowly intensifies, remains at its maximum for up to 15 min and then slowly decreases in intensity. This characteristic description should alert the clinician to the possibility that coronary vasospasm is present.51 Another atypical presentation is pain of anginal location and quality, which is triggered by exertion...

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**Table 4 Traditional clinical classification of chest pain**

<table>
<thead>
<tr>
<th>Type of Angina</th>
<th>Characteristics</th>
</tr>
</thead>
<tbody>
<tr>
<td>Typical (definite)</td>
<td>Meets all three of the following characteristics:</td>
</tr>
<tr>
<td></td>
<td>• substernal chest discomfort of characteristic quality and duration;</td>
</tr>
<tr>
<td></td>
<td>• provoked by exertion or emotional stress;</td>
</tr>
<tr>
<td></td>
<td>• relieved by rest and/or nitrates within minutes.</td>
</tr>
<tr>
<td>Atypical (probable)</td>
<td>Meets two of these characteristics.</td>
</tr>
<tr>
<td>Non-anginal chest pain</td>
<td>Lacks or meets only one or none of the characteristics.</td>
</tr>
</tbody>
</table>
but occurs some time after exertion and may be poorly responsive to nitrates. This presentation is often seen in patients with microvascular angina.52

Non-anginal pain lacks the characteristic qualities described, may involve only a small portion of the left or right hemithorax, and last for several hours or even days. It is usually not relieved by nitroglycerin (although it may be in the case of oesophageal spasm) and may be provoked by palpation. Non-cardiac causes of pain should be evaluated in such cases.49

The Canadian Cardiovascular Society classification is widely used as a grading system for stable angina,53 to quantify the threshold at which symptoms occur in relation to physical activities (Table 5). It is, however, important to keep in mind that the grading system explicitly recognizes that rest pain may occur in all grades as a manifestation of associated and superimposed coronary vasospasm.5 It is also important to remember that the class assigned is indicative of the maximum limitation and that the patient may do better on other days.

Patients with chest pain are often seen in general practice. Applying a well-validated prediction rule containing the five determinants [viz. age/sex (male ≥ 55 years, female ≥ 65 years); known vascular disease; patient assumes pain is of cardiac origin; pain is worse during exercise and pain is not reproducible by palpation; one point for each determinant] leads to accurate ruling-out of CAD at the maximum limitation and that the patient may do better on other days.

Before any testing is considered one must assess the general health, comorbidities and quality of life (QoL) of the patient. If assessment suggests that revascularization is unlikely to be an acceptable option, further testing may be reduced to a clinically indicated minimum and appropriate therapy should be instituted, which may include a trial of anti-anginal medication even if a diagnosis of SCAD has not been fully demonstrated.

Basic (first-line) testing in patients with suspected SCAD includes standard laboratory biochemical testing (Table 6), a resting ECG (Table 8), possibly ambulatory ECG monitoring (if there is clinical suspicion that symptoms may be associated with a paroxysmal arrhythmia) (Table 10), resting echocardiography (Table 9) and, in selected patients, a chest X-ray (CXR) (Table 11). Such testing can be done on an outpatient basis.

### Table 5 Classification of angina severity according to the Canadian Cardiovascular Society

<table>
<thead>
<tr>
<th>Class</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>Ordinary activity does not cause angina such as walking and climbing stairs. Angina with strenuous or rapid or prolonged exertion at work or recreation.</td>
</tr>
<tr>
<td>II</td>
<td>Slight limitation of ordinary activity. Angina on walking or climbing stairs rapidly, walking or stair climbing after meals, or in cold, wind or under emotional stress, or only during the first few hours after awakening. Walking more than two blocks on the level and climbing more than one flight of ordinary stairs at a normal pace and in normal conditions.</td>
</tr>
<tr>
<td>III</td>
<td>Marked limitation of ordinary physical activity. Angina on walking one to two blocks* on the level or one flight of stairs in normal conditions and at a normal pace.</td>
</tr>
<tr>
<td>IV</td>
<td>Inability to carry on any physical activity without discomfort — angina syndrome may be present at rest.</td>
</tr>
</tbody>
</table>

*Equivalent to 100–200 m.
or post-glucose challenge glycaemia have been shown to predict adverse outcome in SCAD, independently of conventional risk factors.59 Finally, glycated haemoglobin (HbA1c) predicts outcome in diabetics, as well as in non-diabetic subjects.60,61 Patients with diabetes should be managed according to the ESC/European Association for the Study of Diabetes (EASD) Guidelines on diabetes.57

Fasting lipid profile, including total cholesterol (TC), high density lipoprotein (HDL) cholesterol, low density lipoprotein (LDL) cholesterol and triglycerides should also be evaluated in all patients with suspected or established ischaemic disease, including stable angina, to establish the patient’s risk profile and ascertain the need for treatment.62 The lipid profile and glycaemic status should be re-assessed periodically to determine efficacy of treatment and, in non-diabetic patients, to detect new development of diabetes (Table 7). There is no evidence to support recommendations for the frequency of re-assessment of these risk factors. Consensus suggests annual measurement.62

Renal dysfunction may occur in association with hypertension, diabetes or renovascular disease and has a negative impact on prognosis in patients with stable angina pectoris.63–65 Hence, baseline renal function should be evaluated with estimation of the glomerular filtration rate (GFR) using a creatinine (or cystatin C)-based method such as the Cockcroft–Gault,66 Modification of Diet in Renal Disease (MDRD),67 or Chronic Kidney Disease Epidemiology Collaboration (CKD-EPI) formulas.68

If there is a clinical suspicion of CAD instability, biochemical markers of myocardial injury—such as troponin T or troponin I—should be measured, preferably using high sensitivity or ultrasensitive assays. If troponin is elevated, further management should follow the non-ST-elevation acute coronary syndrome (NSTE-ACS) guidelines.1 As troponins have a central role in identifying unstable patients,5,7 it is recommended that troponin measurements be performed in every patient hospitalised for symptomatic SCAD.

Table 6  Blood tests in assessment of patients with known or suspected SCAD in order to optimize medical therapy

<table>
<thead>
<tr>
<th>Recommendations</th>
<th>Class a</th>
<th>Level b</th>
<th>Ref. c</th>
</tr>
</thead>
<tbody>
<tr>
<td>If evaluation suggests clinical instability or ACS, repeated measurements of troponin preferably using high sensitivity or ultrasensitive assays are recommended to rule out myocardial necrosis associated with ACS.</td>
<td>I</td>
<td>A</td>
<td>73, 74</td>
</tr>
<tr>
<td>Full blood count including haemoglobin and white cell count is recommended in all patients.</td>
<td>I</td>
<td>B</td>
<td>75</td>
</tr>
<tr>
<td>It is recommended that screening for potential T2DM in patients with suspected and established SCAD is initiated with HbA1c and fasting plasma glucose and that an OGTT is added if HbA1c, and fasting plasma glucose are inconclusive.</td>
<td>I</td>
<td>B</td>
<td>57, 58, 76</td>
</tr>
<tr>
<td>Creatinine measurement and estimation of renal function (creatinine clearance) are recommended in all patients.</td>
<td>I</td>
<td>B</td>
<td>77</td>
</tr>
<tr>
<td>A fasting lipid profile (including LDL) is recommended in all patients.</td>
<td>I</td>
<td>C</td>
<td>-</td>
</tr>
<tr>
<td>If indicated by clinical suspicion of thyroid disorder assessment of thyroid function is recommended</td>
<td>I</td>
<td>C</td>
<td>-</td>
</tr>
<tr>
<td>Liver function tests are recommended in patients early after beginning statin therapy</td>
<td>I</td>
<td>C</td>
<td>-</td>
</tr>
<tr>
<td>Creatine kinase measurement are recommended in patients taking statins and complaining of symptoms suggestive of myopathy</td>
<td>I</td>
<td>C</td>
<td>-</td>
</tr>
<tr>
<td>hs-CRP and prognosis among patients with SCAD sufficiently uncertain that no recommendation can be made to routinely measure this parameter.69</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

ACS = acute coronary syndrome; BNP = B-type natriuretic peptide; HbA1c = glycated haemoglobin; LDL = low density lipoprotein; NT-proBNP = N-terminal pro B-type natriuretic peptide; SCAD = stable coronary artery disease; T2DM = type 2 diabetes mellitus.

a Class of recommendation.

b Level of evidence.

c Reference(s) supporting class I (A + B) and Ila + IIb (A + B) recommendations.

d For details please refer to dyslipidaemia guidelines.62

Table 7  Blood tests for routine re-assessment in patients with chronic stable coronary artery disease

<table>
<thead>
<tr>
<th>Recommendations</th>
<th>Class a</th>
<th>Level b</th>
<th>Ref. c</th>
</tr>
</thead>
<tbody>
<tr>
<td>Annual control of lipids, glucose metabolism (see recommendation 3 in Table 6) and creatinine is recommended in all patients with known SCAD.</td>
<td>I</td>
<td>C</td>
<td>-</td>
</tr>
</tbody>
</table>

SCAD = stable coronary artery disease.

a Class of recommendation.

b Level of evidence.

c Reference(s) supporting class I (A + B) and Ila + IIb (A + B) recommendations.

Very low levels of troponin can be detected in many patients with SCAD when high-sensitive assays are employed. These levels are usually below the levels defined as being elevated. Although there is some prognostic value associated with the amount of troponin found in stable patients,8,9 troponin does not have enough independent prognostic value to recommend systematic measurement in out-of-hospital patients with SCAD. Elevated levels of high-sensitivity C-reactive protein (hs-CRP) have also been reported to be associated with an increased event risk in patients with SCAD. However, a recent analysis of 83 studies found multiple types of reporting and publication bias, making the magnitude of any independent association between hs-CRP and prognosis among patients with SCAD sufficiently uncertain that no recommendation can be made to routinely measure this parameter.69
Although there may be some additional prognostic value in other biomarkers, there is insufficient evidence to recommend the routine use of natriuretic peptides, haemostasis markers or genetic testing in the management of patients with SCAD (for additional information see web addenda).70–72

6.2.1.2 Resting electrocardiogram

All patients with suspected CAD should have a resting 12-lead ECG recorded. A normal resting ECG is not uncommon, even in patients with severe angina, and does not exclude the diagnosis of ischaemia. However, the resting ECG may show signs of CAD, such as previous MI or an abnormal repolarization pattern. An ECG will establish a baseline for comparison in future situations.

The ECG may assist in clarifying the differential diagnosis if taken in the presence of pain, allowing detection of dynamic ST-segment changes in the presence of ischaemia. An ECG during chest pain and immediately afterwards is always useful and can be diagnostic in patients with vasospasm, since ST segment shifts tend to be at least partially reversible once spasm is relieved. The ECG may also show other abnormalities such as left ventricular hypertrophy (LVH), left or right bundle branch block (LBBB or RBBB), pre-excitation, arrhythmias, or conduction defects. Such information may be helpful in defining the mechanisms responsible for chest pain (atrial fibrillation may be associated with chest discomfort without epicardial coronary disease)70 in selecting appropriate further investigations, or in tailoring individual patient treatment. The resting ECG also has a role in risk stratification, as outlined later.

6.2.1.3 Echocardiography at rest (see web addenda)

Resting two-dimensional and Doppler transthoracic echocardiography provide information on cardiac structure and function. Although left ventricular (LV) function is often normal in these patients, regional wall motion abnormalities may be detected, which increase the likelihood of CAD. Furthermore other disorders, such as valvular heart disease (aortic stenosis) or hypertrophic cardiomyopathy, can be ruled out as an alternative cause of symptoms. Finally, global ventricular function, an important prognostic parameter in patients with SCAD,29,79 can be measured. Echocardiography is particularly useful in patients with murmurs80, previous MI or symptoms/signs of heart failure.

Once resting echocardiography has been performed, ultrasound of the carotid arteries using an appropriate probe may be added by clinicians trained in the examination.81,82 The detection of increased intima-media thickness and/or plaques establishes the presence of atherosclerotic disease, with consequent implications for preventive therapy,37 and increases the pre-test probability of CAD in subsequent diagnostic tests.83

Tissue Doppler imaging and strain rate measurements may also be helpful in detecting heart failure with preserved EF as an explanation for physical activity-associated symptoms.84 Impaired diastolic filling is the first sign of active ischaemia and may point to the presence of microvascular dysfunction in patients who complain about shortness of breath, as a possible angina equivalent.85,86

Although the diagnostic yield of echocardiography in patients with angina is mainly concentrated in specific subgroups, estimation of ventricular function is important in all patients for risk stratification (see section 6.4). Hence, echocardiography (or alternative methods of assessment of ventricular function if echocardiography is of insufficient quality) should be performed in all patients with a first presentation with symptoms of SCAD.

There is no indication for repeated use of resting echocardiography on a regular basis in patients with uncomplicated SCAD in the absence of a change in clinical status.

6.2.1.4 Cardiac magnetic resonance at rest

Cardiac magnetic resonance (CMR) may also be used to define structural cardiac abnormalities and evaluate ventricular function.87 Use of
Ambulatory ECG monitoring adds important diagnostic information over and above that provided by the stress test. There is no good evidence to support the routine deployment of ambulatory ECG monitoring as a tool for refined prognostication.

Ambulatory monitoring, however, has a role in patients in whom arrhythmias or vasospastic angina are suspected (equipment for ST-segment evaluation required).

### 6.2.3 Principles of diagnostic testing

Interpretation of non-invasive cardiac tests requires a Bayesian approach to diagnosis. This approach uses clinicians’ pre-test estimates (termed pre-test probability (PTP)) of disease along with the results of diagnostic tests to generate individualized post-test disease probabilities for a given patient. The PTP is influenced by the prevalence of the disease in the population studied, as well as clinical features (including the presence of CV risk factors) of an individual. Major determinants of PTP are age, gender and the nature of symptoms.

Sensitivity and specificity are often used to describe the accuracy of a given diagnostic method, but they incompletely describe how a test performs in the clinical setting. First, some diagnostic methods may perform better in some patients than in others—such as coronary computed tomography angiography (CTA), which is sensitive to heart rate, body weight and the presence of calcification. Second, although sensitivity and specificity are mathematically independent from the PTP, in clinical practice many tests perform better in low-risk populations; in the example used above, coronary CTA will have higher accuracy values when low-likelihood populations—which are younger and have less coronary calcium—are subjected to the examination.

Because of the interdependence of PTP (the clinical likelihood that a given patient will have CAD) and the performance of the available diagnostic methods (the likelihood that this patient has disease if the test is positive, or does not have disease if the test is negative), recommendations for diagnostic testing need to take into account the PTP. Testing may do harm if the number of false test results is higher than the number of correct test results. Non-invasive, imaging-based diagnostic methods for CAD have typical sensitivities and specificities of approximately 85% (Table 12). Hence, 15% of all diagnostic results will be false and, as a consequence, performing no test at all will provide fewer incorrect diagnoses in patients with a PTP below

### Table 11 Chest X-ray for initial diagnostic assessment of SCAD

<table>
<thead>
<tr>
<th>Recommendations</th>
<th>Class</th>
<th>Level</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>CXR is recommended in patients with atypical presentation or suspicion of pulmonary disease.</td>
<td>I</td>
<td>C</td>
<td>-</td>
</tr>
<tr>
<td>CXR should be considered in patients with suspected heart failure.</td>
<td>IIA</td>
<td>C</td>
<td>-</td>
</tr>
</tbody>
</table>

CXR = chest X-ray.

Class of recommendation.

Level of evidence.

Reference(s) supporting class I (A + B) and IIA (A + B recommendations.)

### Table 10 Ambulatory electrocardiogram monitoring for initial diagnostic assessment of stable coronary artery disease

<table>
<thead>
<tr>
<th>Recommendations</th>
<th>Class</th>
<th>Level</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ambulatory ECG monitoring is recommended in patients with SCAD and suspected arrhythmia.</td>
<td>I</td>
<td>C</td>
<td>-</td>
</tr>
<tr>
<td>Ambulatory ECG monitoring should be considered in patients with suspected vasospastic angina.</td>
<td>IIA</td>
<td>C</td>
<td>-</td>
</tr>
</tbody>
</table>

ECG = electrocardiogram; SCAD = stable coronary artery disease.

Class of recommendation.

Level of evidence.

Reference(s) supporting class I (A + B) and IIA (A + B) recommendations.

CMR is recommended in patients in whom, despite the use of echo contrast agents, transthoracic echocardiography is unable to answer the clinical question (usually because of a restricted acoustic window) and who have no contra-indications for CMR.

### 6.2.2 Three major steps used for decision-making

These guidelines recommend a stepwise approach for decision making in patients with suspected SCAD. The process begins with a clinical assessment of the probability that SCAD is present in a particular patient (determination of PTP; Step 1) (see below). Step 1 is followed by non-invasive testing to establish the diagnosis of SCAD or non-obstructive atherosclerosis (typically by performing carotid ultrasound) in patients with an intermediate probability of disease (Step 2). Once the diagnosis of SCAD has been made, optimal medical therapy (OMT) is instituted and stratification for risk of subsequent events (referred to as ‘event risk’ in the following text) is carried out (Step 3)—usually on the basis of available non-invasive tests—in order to select patients who may benefit from invasive investigation and revascularization. Depending on the severity of symptoms, early invasive coronary angiography (ICA) may be performed with appropriate invasive confirmation of the significance of a stenosis (FFR) and subsequent revascularization, bypassing non-invasive testing in Steps 2 and 3.

The PTP is influenced by the prevalence of the disease in the population studied, as well as clinical features (including the presence of CV risk factors) of an individual. Major determinants of PTP are age, gender and the nature of symptoms.

Sensitivity and specificity are often used to describe the accuracy of a given diagnostic method, but they incompletely describe how a test performs in the clinical setting. First, some diagnostic methods may perform better in some patients than in others—such as coronary computed tomography angiography (CTA), which is sensitive to heart rate, body weight and the presence of calcification. Second, although sensitivity and specificity are mathematically independent from the PTP, in clinical practice many tests perform better in low-risk populations; in the example used above, coronary CTA will have higher accuracy values when low-likelihood populations—which are younger and have less coronary calcium—are subjected to the examination.

Because of the interdependence of PTP (the clinical likelihood that a given patient will have CAD) and the performance of the available diagnostic methods (the likelihood that this patient has disease if the test is positive, or does not have disease if the test is negative), recommendations for diagnostic testing need to take into account the PTP. Testing may do harm if the number of false test results is higher than the number of correct test results. Non-invasive, imaging-based diagnostic methods for CAD have typical sensitivities and specificities of approximately 85% (Table 12). Hence, 15% of all diagnostic results will be false and, as a consequence, performing no test at all will provide fewer incorrect diagnoses in patients with a PTP below
15% (assuming all patients to be healthy) or a PTP above 85% (assuming all patients to be diseased). In these situations, testing should only be done for compelling reasons. This is the reason why this Task Force recommends no testing in patients with (i) a low PTP <15% and (ii) a high PTP >85%. In such patients, it is safe to assume that they have (i) no obstructive CAD or (ii) obstructive CAD.

The low sensitivity of the exercise ECG—only 50% (despite an excellent specificity of 90%, values obtained from studies avoiding verification bias)—is the reason why the number of false test results will become higher than the number of correct test results in populations with a PTP >65%. Therefore, this Task Force recommends not employing the exercise stress test in such higher-risk populations for diagnostic purposes. However, the test may nevertheless provide valuable prognostic information in such populations.

In this new version of the Guidelines, more weight is given to testing based systematically on consideration of pre-test probabilities.

This Task Force selected the most recent estimates of CAD prevalences as the basis of these Guidelines’ clinical algorithm, as discussed in the web addenda and shown in Table 13. The web addenda also contains more information about changes from the previous Stable Angina guidelines of the ESC and the reasons why ECG exercise testing was kept in the algorithm.

If the pain is clearly non-anginal other diagnostic testing may be indicated to identify gastrointestinal, pulmonary or musculoskeletal causes of chest pain (Figure 1). Nevertheless, these patients should also receive risk factor modification based on commonly applied risk charts such as SCORE (http://www.heartscore.org/Pages/welcome.aspx) or the Framingham risk score (http://hp2010.nhlbihin.net/atpiii/calculator.asp). Patients with suspected SCAD, in whom comorbidities make revascularization inadvisable, should be treated medically but pharmacologic stress imaging may be an option if it appears necessary to verify the diagnosis. Patients with a reduced left ventricular ejection fraction (LVEF) of <50% and typical angina are at high risk for cardiovascular events (see later in the text) and they should be offered ICA without previous testing (see Figure 1).

Patients in whom anginal pain may be possible but who have a very low probability of significant CAD <15% should have other cardiac causes of chest pain excluded and their CV risk factors adjusted, based on risk score assessment. No specific non-invasive stress testing should be performed. In patients with repeated, unprovoked attacks of chest pain only at rest, vasospastic angina should be considered and diagnosed, and treated appropriately (see below). Patients with an intermediate PTP of 15–85% should undergo further non-invasive testing. In patients with a clinical PTP >85%, the diagnosis of CAD should be made clinically and further testing will not improve accuracy. Further testing may, however, be indicated for stratification of risk of events, especially if no satisfactory control of symptoms is possible with initial medical therapy (Figure 1). In patients with severe angina at a low level of exercise and those with a clinical constellation indicating a high event risk, proceeding directly to ICA is a reasonable option. Under such circumstances, the indication for revascularization should depend on the result of intra-procedural fractional flow reserve (FFR) testing when indicated.

The very high negative predictive value of a coronary CTA showing no stenoses can reassure patients and referring physicians that instituting medical therapy and not proceeding to further testing or invasive therapies is a good strategy. This makes the test potentially useful, especially for patients at low intermediate PTPs (Figure 2). One should remember that there may be overdiagnosis of stenoses in patients with Agatston scores >400, and it seems prudent to call a coronary CTA ‘unclear’ if severe focal or diffuse calcifications

### Table 12 Characteristics of tests commonly used to diagnose the presence of coronary artery disease

<table>
<thead>
<tr>
<th>Test</th>
<th>Sensitivity (%)</th>
<th>Specificity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Exercise ECG</td>
<td>45–50</td>
<td>85–90</td>
</tr>
<tr>
<td>Exercise stress echocardiography</td>
<td>80–85</td>
<td>80–88</td>
</tr>
<tr>
<td>Exercise stress SPECT</td>
<td>73–92</td>
<td>63–87</td>
</tr>
<tr>
<td>Dobutamine stress echocardiography</td>
<td>79–83</td>
<td>82–86</td>
</tr>
<tr>
<td>Dobutamine stress MRI</td>
<td>79–88</td>
<td>81–91</td>
</tr>
<tr>
<td>Vasodilator stress echocardiography</td>
<td>72–79</td>
<td>92–95</td>
</tr>
<tr>
<td>Vasodilator stress SPECT</td>
<td>90–91</td>
<td>75–84</td>
</tr>
<tr>
<td>Vasodilator stress MRI</td>
<td>67–94</td>
<td>61–85</td>
</tr>
<tr>
<td>Coronary CTA</td>
<td>95–99</td>
<td>64–83</td>
</tr>
<tr>
<td>Vasodilator stress PET</td>
<td>81–97</td>
<td>74–91</td>
</tr>
</tbody>
</table>

CAD = coronary artery disease; CTA = computed tomography angiography; ECG = electrocardiogram; MRI = magnetic resonance imaging; PET = positron emission tomography; SPECT = single photon emission computed tomography.

### Table 13 Clinical pre-test probabilities in patients with stable chest pain symptoms

<table>
<thead>
<tr>
<th>Age</th>
<th>Typical angina</th>
<th>Atypical angina</th>
<th>Non-anginal pain</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Men</td>
<td>Women</td>
<td>Men</td>
</tr>
<tr>
<td>30–39</td>
<td>59</td>
<td>28</td>
<td>29</td>
</tr>
<tr>
<td>40–49</td>
<td>69</td>
<td>37</td>
<td>38</td>
</tr>
<tr>
<td>50–59</td>
<td>77</td>
<td>47</td>
<td>49</td>
</tr>
<tr>
<td>60–69</td>
<td>84</td>
<td>58</td>
<td>59</td>
</tr>
<tr>
<td>70–79</td>
<td>89</td>
<td>68</td>
<td>69</td>
</tr>
<tr>
<td>&gt;80</td>
<td>93</td>
<td>76</td>
<td>78</td>
</tr>
</tbody>
</table>

ECC = electrocardiogram; PTP = pre-test probability; SCAD = stable coronary artery disease.

* Probabilities of obstructive coronary disease shown reflect the estimates for patients aged 35, 45, 55, 65, 75 and 85 years.

- Groups in white boxes have a PTP <15% and hence can be managed without further testing.
- Groups in blue boxes have a PTP of 15–65%.
- Groups in light red boxes have PTPs between 66–85%.
- In groups in dark red boxes the PTP is >85% and one can assume that SCAD is present. They need risk stratification only.

- Groups in blue boxes have a PTP of 15–65%. They could have an exercise ECG if feasible as the initial test. However, if local expertise and availability permit a non-invasive imaging based test for ischaemia this would be preferable given the superior diagnostic capabilities of such tests. In young patients radiation issues should be considered.
- Groups in light red boxes have PTPs between 66–85% and hence should have a non-invasive imaging functional test for making a diagnosis of SCAD.
- In groups in dark red boxes the PTP is >85% and one can assume that SCAD is present. They need risk stratification only.
prevent an unambiguous identification of the vessel lumen (see Figure 2). To obtain optimal results, published professional standards need to be meticulously adhered to.\textsuperscript{111} With these caveats in mind, coronary CTA may be considered to be an alternative to ischaemia testing, especially in patients with chest pain symptoms at intermediate PTPs lower than 50%.\textsuperscript{112}

### 6.2.4 Stress testing for diagnosing ischaemia

#### 6.2.4.1 Electrocardiogram exercise testing

Because of its simplicity and widespread availability, treadmill or bicycle exercise testing, using 12-lead ECG monitoring, remains a useful option (Table 14) in patients with suspected SCAD and a PTP (15–65%) at which the test performs well (see above). A detailed description of the exercise procedure, its interpretation, the influence of drugs and other factors on test performance, and test performance in special groups can be found in the previous version of these Guidelines on the ESC website.\textsuperscript{3}

The main diagnostic ECG abnormality during ECG exercise testing consists of a horizontal or down-sloping ST-segment depression $\geq 0.1$ mV, persisting for at least 0.06–0.08s after the J-point, in one or more ECG leads. It is worth noting that, in about 15% of patients, diagnostic ST-segment changes appear only during the recovery
phase. The test also provides additional information, such as heart rate response, blood pressure response, symptoms, and workload achieved, which has both diagnostic and prognostic relevance.

To obtain maximal diagnostic information from exercise ECG testing, the latter should be symptom/sign-limited and performed without the influence of anti-ischaemic drugs. There are numerous reviews and meta-analyses of the performance of exercise ECG for the diagnosis of coronary disease, showing variable diagnostic yield according to the threshold selected for the diagnosis. Using exercise ST-depression ≥0.1 mV or 1 mm to define a positive test, the reported sensitivities and specificities for the detection of significant CAD (usually diameter stenoses ≥50%) range between 23–100% (mean 68%) and 17–100% (mean 77%), respectively. Restricting the analysis to those studies designed to avoid work-up bias, sensitivities between 45–50% and specificities of 85–90% were reported (Table 12). Adding cardiopulmonary exercise testing may improve sensitivity significantly, but this combination of tests is not widely used.

It is important to remember that these numbers are valid only in patients without significant ECG abnormalities at baseline. Exercise ECG testing is not of diagnostic value in the presence of LBBB, paced rhythm and Wolff-Parkinson-White syndrome, in which cases the ECG changes are not interpretable. Additionally, false-positive results are more frequent in patients with abnormal resting ECG in the presence of LVH, electrolyte imbalance, intraventricular conduction abnormalities, atrial fibrillation, and use of digitalis.

Exercise ECG testing is also less sensitive and specific in women. However, a recent randomized trial, comparing an initial diagnostic strategy of exercise nuclear myocardial perfusion imaging (MPI) with standard exercise treadmill testing, in symptomatic women with suspected CAD and preserved functional capacity who were able to exercise, did not show an incremental benefit of the more expensive MPI strategy on clinical outcomes.

In some patients, the exercise ECG may be inconclusive; for example, when 85% of maximum heart rate is not achieved in the absence of symptoms or signs of ischaemia, when exercise is

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**Figure 2** Non-invasive testing in patients with suspected SCAD and an intermediate pre-test probability. CAD = coronary artery disease; CTA = computed tomography angiography; CMR = cardiac magnetic resonance; ECG = electrocardiogram; ICA = invasive coronary angiography; LVEF = left ventricular ejection fraction; PET = positron emission tomography; PTP = pre-test probability; SCAD = stable coronary artery disease; SPECT = single photon emission computed tomography.

- Consider age of patient versus radiation exposure.
- In patients unable to exercise use echo or SPECT/PET with pharmacologic stress instead.
- CMR is only performed using pharmacologic stress.
- Patient characteristics should make a fully diagnostic coronary CTA scan highly probable (see section 6.2.5.1.2) consider result to be unclear in patients with severe diffuse or focal calcification.
- Proceed as in lower left coronary CTA box.
- Proceed as in stress testing for ischaemia box.
limited by orthopaedic or other non-cardiac problems, or when ECG changes are equivocal. In these patients, an alternative non-invasive imaging test with pharmacologic stress should be selected (Figure 2). In patients who are appropriately selected (Figure 2), coronary CTA is another option. Furthermore, a ‘normal’ ECG stress test in patients taking anti-ischaemic drugs does not rule out significant coronary disease.

Exercise stress testing can also be useful to evaluate the efficacy of medical treatment or after revascularization, or to assist prescription of exercise after control of symptoms. For these indications, exercise stress testing should be performed on treatment to evaluate control of ischaemia or effort performance. The effect of routine periodic exercise testing on patient outcomes has not been formally evaluated.

6.2.4.2 Stress imaging (see web addenda)

6.2.4.2.1 Stress echocardiography. Stress echocardiography is performed with exercise (treadmill or bicycle ergometer) or with pharmacological agents. Exercise provides a more physiological data, such as exercise time and workload, as well as information about changes in heart rate, blood pressure and ECG. Thus, exercise is the test of choice when feasible (Table 15).

### Table 14 Performing an exercise electrocardiogram for initial diagnostic assessment of angina or evaluation of symptoms

<table>
<thead>
<tr>
<th>Recommendations</th>
<th>Class</th>
<th>Level</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Exercise ECG is recommended as the initial test for establishing a diagnosis of SCAD in patients with symptoms of angina and intermediate PTP of CAD (Table 13, 115–116), free of anti-ischaemic drugs, unless they cannot exercise or display ECG changes which make the ECG non-evaluable.</td>
<td>IIA</td>
<td>C</td>
<td></td>
</tr>
<tr>
<td>Stress imaging is recommended as the initial test option if local expertise and availability permit.</td>
<td>I</td>
<td>B</td>
<td>117–120</td>
</tr>
<tr>
<td>Exercise ECG should be considered in patients on treatment to evaluate control of symptoms and ischaemia.</td>
<td>IIa</td>
<td>C</td>
<td>-</td>
</tr>
<tr>
<td>Exercise ECG in patients with ≥0.1 mV ST-depression on resting ECG or taking digitalis is not recommended for diagnostic purposes.</td>
<td>III</td>
<td>C</td>
<td>-</td>
</tr>
</tbody>
</table>

**Table 15 Use of exercise or pharmacologic stress testing in combination with imaging**

<table>
<thead>
<tr>
<th>Recommendations</th>
<th>Class</th>
<th>Level</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>An imaging stress test is recommended as the initial test for diagnosing SCAD if the PTP is between 66–85% or if LVEF is &lt;50% in patients without typical angina.</td>
<td>I</td>
<td>B</td>
<td>143, 144</td>
</tr>
<tr>
<td>An imaging stress test is recommended in patients with resting ECG abnormalities which prevent accurate interpretation of ECG changes during stress.</td>
<td>I</td>
<td>B</td>
<td>117, 145</td>
</tr>
<tr>
<td>Exercise stress testing is recommended rather than pharmacologic testing whenever possible.</td>
<td>I</td>
<td>C</td>
<td>-</td>
</tr>
<tr>
<td>An imaging stress test should be considered in symptomatic patients with prior revascularization (PCI or CAGB).</td>
<td>IIa</td>
<td>B</td>
<td>146, 147</td>
</tr>
<tr>
<td>An imaging stress test should be considered to assess the functional severity of intermediate lesions on coronary arteriography.</td>
<td>IIa</td>
<td>B</td>
<td>148, 149</td>
</tr>
</tbody>
</table>

**Table 15 Use of exercise or pharmacologic stress testing in combination with imaging**

CAD = coronary artery disease; ECG = electrocardiogram; PTP = pre-test probability; SCAD = stable coronary artery disease.

- **Class of recommendation.**
- **Level of evidence.**
- **Reference(s) supporting levels of evidence.**

On the other hand, a pharmacological test is preferred when there is already a significant resting wall motion abnormality (dobutamine for viability assessment) and/or if the patient is unable to exercise adequately. Until recently, stress echocardiography relied on indubitable wall thickening abnormalities as a marker of ischaemia (supply–demand mismatch). As most data on diagnostic accuracy were obtained using this standard, there is a caveat, in that the values for sensitivity and specificity assumed in these guidelines (Table 12) rely heavily on old studies, carried out at a time when contrast media were not broadly utilized in clinical practice.

The pharmacological agent of choice to produce supply-demand mismatch is dobutamine. Myocardial contrast echocardiography, which utilizes microbubbles, allows assessment of myocardial perfusion, which provides information beyond wall thickening assessment during both vasodilator and isotropic stress echocardiography. This approach, however, is not widely employed clinically.

Contrast agents must be used in all patients undergoing all forms of stress echocardiography when two or more continuous segments (17 segment LV model) are not well visualised at rest. The use of contrast during stress echocardiography not only enhances image quality, but improves reader confidence and enhances
accuracy for the detection of CAD.\textsuperscript{122,124} Tissue Doppler imaging and strain rate imaging may also improve the diagnostic performance of stress echocardiography by improving the capability of echocardiography to detect ischaemia beyond wall motion assessment.\textsuperscript{125}

6.2.4.2.2 Myocardial perfusion scintigraphy (single photon emission computed tomography and positron emission tomography). Technetium-99m ($^{99m}$Tc) radiopharmaceuticals are the most commonly used tracers, employed with single photon emission computed tomography (SPECT) in association with a symptom-limited exercise test on either a bicycle ergometer or a treadmill (Table 15). Thallium 201 ($^{201}$Tl) is associated with a higher radiation and is less commonly used today. New SPECT cameras reduce radiation and/or acquisition time significantly.\textsuperscript{126}

Regardless of the radiopharmaceutical or camera used, SPECT perfusion scintigraphy is performed to produce images of regional tracer uptake, which reflect relative regional myocardial blood flow. With this technique, myocardial hypoperfusion is characterized by reduced tracer uptake during stress, in comparison with the uptake at rest. Increased uptake of the myocardial perfusion agent in the lung field identifies stress-induced ventricular dysfunction in patients with severe and extensive CAD.\textsuperscript{127} As with all stress imaging techniques, SPECT perfusion also provides a more sensitive prediction of the presence of CAD than the exercise ECG (Table 12). Transient ischaemic dilatation and reduced post-stress ejection fraction (EF) are important non-perfusion predictors of severe CAD.

Pharmacological stress testing with perfusion scintigraphy is indicated in patients who are unable to exercise adequately or may be used as an alternative to exercise stress. Adenosine may precipitate bronchospasm in asthmatic individuals by activating A\textsubscript{1}, A\textsubscript{2B} and A\textsubscript{3} receptors in addition to activation of the A\textsubscript{2A} adenosine receptor, which produces hyperaemia. This limitation exists irrespective of the imaging technique used but, in such cases, dobutamine or regadenoson,\textsuperscript{128} a selective A\textsubscript{2A} receptor agonist, may be used as an alternative stressor.

MPI using positron emission tomography (PET) is superior to SPECT imaging for the detection of SCAD in terms of image quality, interpretative certainty and diagnostic accuracy.\textsuperscript{129} However, SPECT scanners and imaging radiotracers are more widely available and less expensive than PET scanners and positron-emitting radiotracers (e.g. $^{82}$Rb, $^{13}$N-ammonia).\textsuperscript{130} Hence, as compared with the other stress imaging techniques, PET is less commonly used for diagnosing SCAD. PET has the unique ability to quantify blood flow in mL/min/g, which allows detecting microvascular disease.\textsuperscript{131}

6.2.4.2.3 Stress cardiac magnetic resonance. Cardiac magnetic resonance (CMR) stress testing, in conjunction with a dobutamine infusion, can be used to detect wall motion abnormalities induced by ischaemia.\textsuperscript{132} The technique has been shown to have a comparable safety profile to dobutamine stress echocardiography (DSE).\textsuperscript{133,134} Dobutamine stress CMR may be useful in patients with sub-optimal acoustic windows,\textsuperscript{132,135} especially those in whom pharmacologic perfusion imaging using adenosine is contra-indicated (Table 15).

Perfusion CMR is more widely used than dobutamine stress CMR. Recent studies have confirmed the good diagnostic accuracy of CMR perfusion imaging at 1.5 Tesla (T), as compared with nuclear perfusion imaging.\textsuperscript{101,136}

Details regarding stress and imaging protocols were recently reviewed.\textsuperscript{137} Analysis is either visual, to identify low signal areas of reduced perfusion, or with computer assistance to determine the up-slope of myocardial signal increase during the first pass. Quantitative CMR perfusion measurements demonstrate good correlations with FFR measurements.\textsuperscript{138} Although not widely available, the use of high-strength magnets at 3.0 T provides higher diagnostic accuracy, as compared with 1.5 T machines.\textsuperscript{139,140}

6.2.4.2.4 Hybrid techniques. Hybrid SPECT/CT, PET/CT and PET/CMR imaging are now available at a few selected centres. Hybrid imaging is a novel technique combining functional and anatomical aspects, which holds much promise for future clinical application. The limited evidence available today indicates a higher diagnostic accuracy, as compared with single techniques.\textsuperscript{141} Initial reports also point to the prognostic value of hybrid imaging.\textsuperscript{142}

6.2.5 Non-invasive techniques to assess coronary anatomy

6.2.5.1 Computed tomography

Spatial resolution and temporal resolution, as well as volume coverage of modern multidetector row CT systems, are sufficient to allow robust imaging of the coronary arteries in many patients.\textsuperscript{150} Radiation dose is a matter of concern and special measures need to be undertaken to avoid unnecessarily high radiation doses when CT is used for coronary artery imaging.\textsuperscript{151} CT imaging of the coronary arteries can be performed without contrast injection (coronary calcium scoring) or after intravenous injection of iodinated contrast (coronary CTA).

6.2.5.1.1 Calcium scoring. Multidetector row CT permits the detection of coronary calcification in non-contrast enhanced data sets. By consensus, pixels above a threshold of 130 Hounsfield units (HU) are defined as representing coronary calcium. Calcified lesions are usually quantified using the ‘Agatston score’.\textsuperscript{152}

With the exception of patients with renal failure—who may have medial calcification—coronary calcium is exclusively a consequence of coronary atherosclerosis. The amount of calcium correlates roughly to the total amount of atherosclerosis present in the coronary arteries,\textsuperscript{153} but correlation with the degree of luminal narrowing is poor. Even with severe calcification, luminal stenosis is not necessarily present and a ‘zero’ calcium score cannot be used to rule out coronary artery stenoses in symptomatic individuals (Table 16), especially when young and with acute symptoms.\textsuperscript{154}

6.2.5.1.2 Coronary computed tomography angiography. After intravenous injection of contrast agent, CT can visualize the coronary artery lumen. Adequate technology (at least 64-slice CT) and patient selection, as well as careful patient preparation, are mandated. According to expert consensus, only patients with adequate breath holding capabilities, without severe obesity, with a favourable calcium score (e.g. Agatston score \(< 400\)) and distribution, in sinus rhythm and with a heart rate of 65 beats per minute (b.p.m.) or less (preferably 60 b.p.m. or less), should be considered for coronary CTA.\textsuperscript{151} If necessary, the use of short-acting $\beta$-blockers or other heart rate-lowering medication is recommended.

Since the specificity of coronary CTA decreases with increasing amounts of coronary calcium,\textsuperscript{103,155,156} and the prevalence of coronary artery stenosis was found to be high in symptomatic individuals with an Agatston score \(< 400\),\textsuperscript{157} it is reasonable not to proceed with coronary CTA if the calcium score exceeds 400.\textsuperscript{154} However, on a patient level, per-segment calcification has a stronger influence on diagnostic accuracy than calcium,\textsuperscript{156} and the influence of calcium...
on the accuracy of coronary CTA is less pronounced in low heart rates and for modern CT systems. In the event that a calcium score is not obtained and calcifications are only seen on the completed coronary CTA scan, it may be prudent to refrain from stenosis quantification in areas of extensive calcifications and call the test ‘unclear’ (see Figure 2).

In patients with suspected CAD, multicentre studies using 64-slice CT have demonstrated sensitivities of 95–99% and specificities of 64–83% (Table 12) as well as negative predictive values of 97–99% for the identification of individuals with at least one coronary artery stenosis by ICA. Meta-analyses of smaller trials confirm a high sensitivity (98–99%) and negative predictive value (99–100%), paired with lower specificity (82–89%) and positive predictive value (91–93%). In a multicentre study, which included patients with previously known CAD, previous PCI and MI, diagnostic accuracy was lower (sensitivity 85% and specificity 90%). Severe coronary calcium negatively impacts the accuracy of coronary CTA. Also, coronary CTA remains less reliable in patients with coronary stents, due to artefacts caused by metal and the limited spatial resolution of CT. The assessment of coronary artery bypass grafts (CABG) is highly accurate while the evaluation of native coronary vessels in post-bypass patients is difficult and prone to false positive findings.

Whilst prospective trials—which have randomized patients to the use or non-use of coronary CTA looking at hard clinical endpoints in stable chest pain patients—are currently not available (just as for the other imaging techniques), registry data confirm an excellent diagnostic performance of coronary CTA, such as CT-FFR need further validation.

6.2.5.2 Magnetic resonance coronary angiography

Coronary MR angiography allows for non-invasive visualization of the coronary arteries without exposing the patient to ionizing radiation. A recent small, multicentre study showed sensitivity, specificity and positive and negative predictive values of 88, 72, 71 and 88%, respectively, in a patient-based analysis. However, long imaging times, lower spatial resolution and operator dependency remain major limitations. Advantages of the technique include evaluation of overall cardiac anatomy and function in the same examination. However, at present, MR coronary arteriography must still be regarded primarily as a research tool and is not recommended for routine clinical practice in the diagnostic evaluation of SCAD.

6.3 Invasive coronary angiography (see web addenda)

Non-invasive testing can establish the likelihood of the presence of obstructive coronary disease with an acceptable degree of certainty. Thus, ICA will only rarely be necessary in stable patients with suspected CAD, for the sole purpose of establishing or excluding the diagnosis. Such situations may arise in patients who cannot undergo stress imaging techniques in patients with reduced LVEF <50% and typical angina (see Figure 1) or in those patients with special professions, such as pilots, due to regulatory issues. ICA may, however, be indicated following non-invasive risk stratification for determination of options for revascularization. In patients who have a high PTP and severe symptoms, or a clinical constellation suggesting high event risk, early ICA without previous non-invasive judgement (especially a positive stress test result when clinical judgement speaks against the presence of severe stenoses) if ICA would otherwise be chosen to rule out CAD (Table 16).

Table 16 Use of coronary computed tomography angiography for the diagnosis of stable coronary artery disease

<table>
<thead>
<tr>
<th>Recommendations</th>
<th>Class a</th>
<th>Level b</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coronary CTA should be considered as an alternative to stress imaging techniques for ruling out SCAD in patients within the lower range of intermediate PTP for SCAD in whom good image quality can be expected.</td>
<td>IIa</td>
<td>C</td>
</tr>
<tr>
<td>Coronary CTA should be considered in patients within the lower range of intermediate PTP for SCAD after a non conclusive exercise ECG or stress imaging test or who have contraindications to stress testing in order to avoid otherwise necessary invasive coronary angiography if fully diagnostic image quality of coronary CTA can be expected.</td>
<td>IIa</td>
<td>C</td>
</tr>
<tr>
<td>Coronary calcium detection by CT is not recommended to identify individuals with coronary artery stenosis.</td>
<td>III</td>
<td>C</td>
</tr>
<tr>
<td>Coronary CTA is not recommended in patients with prior coronary revascularization.</td>
<td>III</td>
<td>C</td>
</tr>
<tr>
<td>Coronary CTA is not recommended as a ‘screening’ test in asymptomatic individuals without clinical suspicion of coronary artery disease.</td>
<td>III</td>
<td>C</td>
</tr>
</tbody>
</table>

CTA = computed tomography angiography; ECG = electrocardiogram; PTP = pre-test probability; SCAD = stable coronary artery disease.

Class of recommendation.

Level of evidence.
6.4 Stratification for risk of events

The long-term prognosis of SCAD depends upon a number of factors, such as clinical and demographic variables, LV function, the result of stress testing and coronary anatomy as determined by angiographic techniques.

When discussing risk stratification in patients with SCAD, event risk refers primarily to the risk of CV death and MI, although in some studies even wider combinations of CV endpoints are employed. As all-cause death is more precisely defined than other weaker endpoints—including MI—these guidelines stratify event risk according to this hard endpoint. The process of risk stratification serves to identify patients at high event risk who will benefit from revascularization beyond the amelioration of symptoms.

The definition of the high event risk group of patients who will benefit from revascularization has changed from the previous version of these Guidelines. Previously, identification of high event risk was solely based on the Duke treadmill score and a >2% annual risk of cardiac death was felt to be the threshold beyond which coronary angiography was recommended to identify the need for revascularization. This value was based on the CV mortality in the placebo arms of studies in ‘high-risk’ populations, such as in the diabetic Microalbuminuria, cardiovascular, and renal sub-study of the Heart Outcomes Prevention Evaluation study (MICRO-HOPE) and the Impact Of Nicorandil in Angina (IONA) studies, where the annualized CV mortality rates were >2%.

In these Guidelines, patients with an annual mortality >3% are defined as high event risk patients. As shown in the web addenda, both ischaemia- and anatomy-oriented indices come to similar conclusions in identifying which patients are at such high event risk with medical treatment alone that revascularization procedures become beneficial in terms of prognosis. Therefore, in these Guidelines, it is the goal of a event risk-driven diagnostic strategy to identify patients with an annual mortality >3% per year.

For the purpose of these Guidelines, low event risk patients are those with an annual mortality <1% per year, similar to the definition chosen in the previous edition. The intermediate event risk group has an annual mortality of ≥1% but ≤3% per year (Table 17).

The risk assessment sequence can be described as:

1. Risk stratification by clinical evaluation
2. Risk stratification by ventricular function
3. Risk stratification by response to stress testing
4. Risk stratification by coronary anatomy

Event risk stratification generally follows a pyramidal structure, with all patients having event risk stratification by clinical evaluation as the most basic requirement, proceeding to assessment of ventricular function by resting echocardiography and, in the majority, to non-invasive assessment of ischaemia/coronary anatomy (which is usually obtained in the process of making a diagnosis of SCAD, as discussed above). ICA for risk stratification will only be required in a selected subgroup of patients.

Table 17 Definitions of risk for various test modalities

<table>
<thead>
<tr>
<th>Test Modality</th>
<th>High Risk</th>
<th>Intermediate Risk</th>
<th>Low Risk</th>
</tr>
</thead>
<tbody>
<tr>
<td>Exercise stress ECG†</td>
<td>CV mortality &gt;3%/year. CV mortality between 1 and 3%/year. CV mortality &lt;1%/year.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ischaemia imaging</td>
<td>Area of ischaemia &gt;10% (&gt;10% for SPECT; limited quantitative data for CMR – probably ≥2/16 segments with new perfusion defects or &gt;3 dobutamine-induced dysfunctional segments; ≥3 segments of LV by stress echo). Area of ischaemia between 1 to 10% or any ischaemia less than high risk by CMR or stress echo. No ischaemia.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Coronary CTA‡</td>
<td>Significant lesions of high risk category (three-vessel disease with proximal stenoses, LM, and proximal anterior descending CAD). Significant lesion(s) in large and proximal coronary artery(ies) but not high risk category. Normal coronary artery or plaques only.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

CAD = coronary artery disease; CMR = cardiac magnetic resonance; CTA = computed tomography angiography; CV = cardiovascular; ECG = electrocardiogram; ICA = invasive coronary angiography; LM = left main; PTP = pre-test probability; SPECT = single photon emission computed tomography.

† For detailed explanation on rationale for risk stratification scheme see web addenda.
‡ From nomogram (see web addenda, Figure W1) or http://www.cardiology.org/tools/medcalc/duke/
6.4.1 Event risk stratification using clinical evaluation
Clinical history and physical examination can provide important prognostic information. The ECG can be conveniently incorporated into the event risk stratification at this level and the results of the laboratory tests discussed in the previous section may modify event risk estimation further. Diabetes, hypertension, current smoking and elevated TC (untreated or elevated despite treatment) have been shown to be predictive of adverse outcome in patients with SCAD or other populations with established coronary disease.178 Increasing age is an important factor to consider, as are the presence of chronic kidney- or peripheral vascular disease,65,179 prior MI,180 symptoms and signs of heart failure,180,181 and the pattern of occurrence (recent onset or progressive) and severity of angina, particularly if unresponsive to therapy.45,182 However, this information is too complex to be placed into a clinically useful event risk score for patients with SCAD and the recommendation is therefore to use the data—especially the severity of angina—to modulate decisions made on the basis of PTP and non-invasive ischaemia/anatomy evaluation of the prognosis (Figure 3).

6.4.2 Event risk stratification using ventricular function
The strongest predictor of long-term survival is LV function. In patients with SCAD as LVEF declines, mortality increases. In the Coronary Artery Surgery Study (CASS) registry, the 12-year survival rates of patients with EF $\geq 50\%$, 35–49% and $< 35\%$ were 73, 54 and 21%, respectively ($P < 0.0001$).183 Hence, a patient with an LVEF $<50\%$ is already at high risk for CV death (annual mortality $>3\%$), even without accounting for additional event risk factors, such as the extent of ischaemia. As a reduced LVEF $<50\%$ confers such an important increase in event risk, it may be important not to miss obstructed vessels causing ischaemia in such patients.184,185 Hence, stress imaging should be employed instead of the exercise ECG (Figure 2).

Although the likelihood of preserved ventricular systolic function is high in patients with a normal ECG, a normal CXR and no history of prior MI,186 asymptomatic ventricular dysfunction is not uncommon.187 Therefore, as already discussed above, a resting echocardiogram is recommended in all patients with suspected SCAD (Table 18).
6.4.3 Event risk stratification using stress testing

Symptomatic patients with suspected or known CAD should undergo stress testing to perform event risk stratification and use this as the basis for therapeutic decisions if they are candidates for coronary revascularization (Table 19). However, no randomized trials have been published demonstrating a better outcome for patients randomized to event risk stratification by stress testing, as compared with those without, and the evidence base therefore consists of observational studies only. As most patients will have undergone some form of diagnostic testing anyway, these results can also be used for event risk stratification. Patients with a high PTP >85%, who do not need diagnostic testing, should undergo stress imaging for event risk stratification purposes and the indication for revascularization should be discussed, considering the patient’s risk of events, as appropriate (Figure 3). If patients with a PTP >85% have early ICA for symptomatic reasons, additional FFR may be required for event risk stratification as appropriate (Figure 3). For guidance about stress imaging for identifying myocardial viability we refer to the ESC Guidelines on heart failure.

The prognosis for patients with a normal ECG and a low clinical risk for severe CAD is excellent. In one study in which 37% of outpatients referred for non-invasive testing met the criteria for low event risk, 182 <1% had left main stem (LMS) artery disease or died within 3 years. Lower-cost options, such as treadmill testing, should therefore be used, whenever possible, for initial event risk stratification, and those at high event risk should be referred to coronary arteriography.

The prognostic exercise testing markers include exercise capacity, BP response and exercise-induced ischaemia (clinical and ECG). Maximum exercise capacity is a consistent prognostic marker. This measure is at least partly influenced by the extent of rest ventricular dysfunction and the amount of further LV dysfunction induced by exercise.188 However, exercise capacity is also affected by age, general physical condition, comorbidities and psychological state. Exercise capacity may be measured by maximum exercise duration, maximum metabolic equivalent (MET) level achieved, maximum workload achieved, in Watts, maximum heart rate and double (rate–pressure) product. The specific variable used to measure exercise capacity is less important than the inclusion of this marker in the assessment.

The Duke treadmill score is well validated, combining exercise time, ST-deviation and angina during exercise to calculate the patient’s event risk (for more information and a web based tool for calculating the Duke treadmill score see web addenda). High event risk patients with an annual mortality >3% can also be identified using the Duke risk calculator (http://www.cardiology.org/tools/medcalc/duke/).

### Table 18 Risk stratification by resting echocardiography quantification of ventricular function in stable coronary artery disease

<table>
<thead>
<tr>
<th>Recommendations</th>
<th>Class</th>
<th>Level</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Resting echocardiography is recommended to quantify LV function in all patients with suspected SCAD.</td>
<td>I</td>
<td>C</td>
<td></td>
</tr>
</tbody>
</table>

LV = left ventricular; SCAD = stable coronary artery disease.

### Table 19 Risk stratification using ischaemia testing

<table>
<thead>
<tr>
<th>Recommendations</th>
<th>Class</th>
<th>Level</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Risk stratification is recommended based on clinical assessment and the result of the stress test initially employed for making a diagnosis of SCAD.</td>
<td>I</td>
<td>B</td>
<td>109, 206–209</td>
</tr>
<tr>
<td>Stress imaging for risk stratification is recommended in patients with a non-conclusive exercise ECG</td>
<td>I</td>
<td>B</td>
<td>210</td>
</tr>
<tr>
<td>Risk stratification using stress ECG (unless they cannot exercise or display ECG changes which make the ECG non-evaluable) or preferably stress imaging if local expertise and availability permit is recommended in patients with stable coronary disease after a significant change in symptom level.</td>
<td>I</td>
<td>B</td>
<td>210–212</td>
</tr>
<tr>
<td>Stress imaging is recommended for risk stratification in patients with known SCAD and a deterioration in symptoms if the site and extent of ischaemia would influence clinical decision making.</td>
<td>I</td>
<td>B</td>
<td>146, 213–215</td>
</tr>
<tr>
<td>Pharmacological stress with echocardiography or SPECT should be considered in patients with LBBB.</td>
<td>IIa</td>
<td>B</td>
<td>216–218</td>
</tr>
<tr>
<td>Stress echocardiography or SPECT should be considered in patients with paced rhythm.</td>
<td>IIa</td>
<td>B</td>
<td>219, 220</td>
</tr>
</tbody>
</table>

ECG = electrocardiogram; LBBB = left bundle branch block; SCAD = stable coronary artery disease; SPECT = single photon emission computed tomography.

a Class of recommendation.
b Level of evidence.
c Reference(s) supporting levels of evidence.
d Stress imaging has usually been performed for establishing a diagnosis of SCAD in most of these patients.
Early coronary arteriography should be considered in LV myocardium (with stress-induced reversible perfusion deficits by coronary CTA, is associated with increased mortality (risk ratio 1.77 when compared with individuals without any detectable plaque). However, the actual clinical utility of coronary CTA wall imaging for event risk stratification, beyond the detection of significant coronary stenosis, remains currently uncertain.

Large prospective trials have established the prognostic value of coronary CTA, both for the presence and extent of coronary luminal stenoses and for the presence of non- obstructive coronary atherosclerotic plaque. A strong predictive value has been demonstrated, independent of traditional risk factors, concerning mortality and the occurrence of major CV events. Importantly, event rates are very low in the absence of any plaque (0.22–0.28% per annualized death rate). In patients with coronary plaque but without stenosis, the death rate is higher but remains below 0.5%, confirming the excellent prognosis conferred by the absence of coronary stenosis on CT scans. In contrast, patients with left main stenosis or proximal triple vessel disease have a univariate hazard ratio for all-cause mortality of 10.52, suggesting that annual mortality for coronary CTA-defined stenoses is similar to that found in ICA studies.

Due to potential overestimation of obstructive coronary disease by coronary CTA, it may be prudent to perform additional ischaemia testing before sending for ICA a high event risk patient who is not very symptomatic on the basis of anatomy visualised by coronary CTA alone (Table 20).

Despite the recognized limitations of ICA to identify vulnerable plaques the extent, severity of luminal obstruction and location of coronary disease on coronary arteriography have been convincingly demonstrated to be important prognostic indicators in patients with angina (Table 20).

Several prognostic indices have been used to relate severity of disease to the risk of subsequent cardiac events; the simplest and most widely used is the classification of disease into one-vessel, two-vessel, three-vessel, or left main (LM) stem CAD. In the CASS registry of medically treated patients, the 12-year survival rate of patients with normal coronary arteries was 91%, compared with 74% for those with one-vessel disease, 59% for those with two-vessel disease and 50% for those with three-vessel disease (P < 0.001). Patients with severe stenosis of the LM coronary artery have a poor prognosis when treated medically. The presence of severe proximal left anterior descending (LAD) disease also significantly reduces the survival rate. The 5-year survival rate with three-vessel disease plus >95% proximal LAD stenosis was reported to be 59%, compared with a rate of 79% with three-vessel disease without LAD stenosis. However, it should be appreciated that, in these ‘older’ studies, preventive therapy was not at the level of current recommendations regarding both lifestyle and drug therapy. Accordingly, absolute estimates of event risk derived from these studies probably overestimate the risk of future events. Annual mortality rates corresponding to certain angiographic scenarios can be found in the web addenda figure W3.

More information on event risk stratification using intravascular ultrasound or optical coherence tomography and the invasive measurement of the functional severity of coronary lesions can be found in the web addenda of this document.
6.5 Diagnostic aspects in the asymptomatic individual without known coronary artery disease (see web addenda)

In an effort to lower the high burden of coronary deaths in asymptomatic adults, numerous measurements of risk factors and risk markers, as well as stress tests, are often performed as screening investigations. Details on the value of the various attempts to achieve this goal can be found in the new European Guidelines on prevention. The key messages of these Guidelines with respect to testing in asymptomatic individuals without known CAD are summarized in the web addenda of this document. The recent American College of Cardiology Foundation/American Heart Association (ACCF/AHA) guidelines for assessment of CV risk in asymptomatic adults give recommendations that are almost identical to those of the new European Guidelines. These recommendations were adapted for the purpose of these Guidelines (Table 21).

There are no data on how to manage asymptomatic patients who receive stress testing and have a pathologic test result, beyond the recommendations listed in these Guidelines. However, the principles of risk stratification, as described above for symptomatic patients, also apply to these individuals. Thus, patients at low and intermediate risk should receive preventive treatment as outlined in the European Guidelines on cardiovascular disease prevention in clinical practice. Only patients at high event risk, based on the result of a stress test performed without proper indication (for definitions of CAD = coronary artery disease; CMR = cardiac magnetic resonance; CT = computed tomography; CV = cardiovascular; MPI = myocardial perfusion imaging; SCORE = systematic coronary risk evaluation.

<table>
<thead>
<tr>
<th>Recommendations</th>
<th>Class a</th>
<th>Level b</th>
<th>Ref. c</th>
</tr>
</thead>
<tbody>
<tr>
<td>In asymptomatic adults with hypertension or diabetes a resting ECG should be considered for CV risk assessment.</td>
<td>IIa</td>
<td>C</td>
<td>-</td>
</tr>
<tr>
<td>In asymptomatic adults with diabetes 40 years of age and older, measurement of coronary calcium using CT may be considered for CV risk assessment.</td>
<td>IIb</td>
<td>B</td>
<td>226, 227</td>
</tr>
<tr>
<td>In asymptomatic adults with diabetes or asymptomatic adults with a strong family history of CAD or when previous risk assessment testing suggests high risk of CAD, such as a coronary artery calcium score of 400 or greater stress imaging tests (MPI, stress echocardiography, perfusion CMR) may be considered for advanced CV risk assessment.</td>
<td>IIb</td>
<td>C</td>
<td>-</td>
</tr>
<tr>
<td>In low- or intermediate-risk (based on SCORE) asymptomatic adults stress imaging tests are not indicated for further CV risk assessment.</td>
<td>III</td>
<td>C</td>
<td>-</td>
</tr>
</tbody>
</table>
see Table 17), should be considered for ICA. It is important to remember that data demonstrating improved prognosis following appropriate management are still lacking.

Persons whose occupations impact on public safety (e.g. airline pilots, lorry or bus drivers) or who are professional or high-profile athletes not uncommonly undergo periodic exercise testing for assessment of exercise capacity and evaluation of possible heart disease, including CAD. Although there are insufficient data to justify this approach, these evaluations are done for medico-legal reasons in some cases. The threshold for adding imaging to standard exercise electrocardiography in such persons may properly be lower than in the average patient. Otherwise, the same considerations as discussed above for other asymptomatic persons apply for these individuals.

### 6.6 Management aspects in the patient with known coronary artery disease

The clinical course of patients with known SCAD may continue to be stable or be complicated by phases of instability, MI and heart failure. Revascularization(s) may become necessary in the course of the disease. Recommendations for the management of patients in these clinical situations are given in the respective guidelines. There are no randomized trials evaluating the impact on outcome of different strategies for the follow-up of patients with SCAD. In particular, there are currently no data suggesting that any form of follow-up stress testing improves outcome in asymptomatic patients. However re-assessment of the prognosis, following an initial evaluation documenting a low event risk status (Figure 3), may be considered after the expiration of the period for which the test is valid and the patient’s prognosis becomes less well established and potentially less favourable (Table 22). A period of 3 years has been suggested in previous guidelines, although the mean validity period of a normal SPECT myocardial perfusion study is even longer in patients without known CAD (approximately 5.5 years). In contrast, the validity period in patients with known CAD is shorter and adversely modulated by clinical risk factors, such as age, female gender and the presence of diabetes. Thus, clinical judgement is required for determining the need for repeated stress testing, which should be performed using the same stress and imaging techniques.

By consensus, the following recommendations can be made:

### 6.7 Special diagnostic considerations: angina with ‘normal’ coronary arteries (see web addenda)

Since the beginning of ICA it has been known that many patients, especially women, who undergo this procedure because of symptoms of chest pain or shortness of breath with exertion felt to be inappropriate by patient and/or physician, do not have significant obstructive CAD. These patients often present with one of the following types of chest pain, each of which is associated with a different pathology:

1. **Angina with mostly typical features** (although duration may be prolonged and relationship to exercise somewhat inconsistent), which is often associated with abnormal results of stress tests and often represents angina due to microvascular disease (microvascular angina).

2. **Pain, which has typical features of angina in terms of location and duration but occurs predominantly at rest (atypical angina), which may be due to coronary spasm (vasospastic angina).**

3. **Pain that involves a small portion of the left hemithorax, lasts for several hours or even days, is not relieved by nitroglycerin and may be provoked by palpatation (non-anginal pain, often musculoskeletal in origin).**

For the clinicopathological correlation of symptoms with coronary anatomy, please consult the web addenda of this document. Patients with microvascular angina often have a typical constellation of classical atherosclerotic risk factors and represent a large group of patients who undergo a variety of non-invasive stress tests, and even repeated ICA, with the intention of revascularization. Microvascular disease may co-exist in patients with angiographically significant stenoses (≥70%). These patients probably belong to the group of approximately 20% of patients whose symptoms persist unchanged or have shown only minor amelioration after successful revascularization.

In contrast, patients with vasospastic angina predominantly experience angina at rest, which may also lead to emergency coronary angiograms. The rationale for the angiogram is not to miss a potentially treatable occlusive or near-occlusive lesion in these patients, who

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**Table 22 Re-assessment in patients with stable coronary artery disease**

<table>
<thead>
<tr>
<th>Recommendations</th>
<th>Class</th>
<th>Level</th>
</tr>
</thead>
<tbody>
<tr>
<td>Follow-up visits are recommended every 4–6 months in the first year following institution of therapy for SCAD which may be extended to 1 year afterwards. Visits</td>
<td>I</td>
<td>C</td>
</tr>
<tr>
<td>An annual resting ECG is recommended and an additional ECG if a change in anginal status occurred or symptoms suggesting an arrhythmia appeared or medication has been changed which might alter electrical conduction.</td>
<td>I</td>
<td>C</td>
</tr>
<tr>
<td>An exercise ECG or stress imaging if appropriate is recommended in the presence of recurrent or new symptoms once instability has been ruled out.</td>
<td>I</td>
<td>C</td>
</tr>
<tr>
<td>Reassessment of the prognosis using stress testing may be considered in asymptomatic patients after the expiration of the period for which the previous test was felt to be valid (“warranty period”).</td>
<td>IIb</td>
<td>C</td>
</tr>
<tr>
<td>Repetition of an exercise ECG may only be considered after at least 2 years following the last test (unless there is a change in clinical presentation).</td>
<td>IIb</td>
<td>C</td>
</tr>
</tbody>
</table>

ECG = electrocardiogram; SCAD = stable coronary artery disease.

*a Class of recommendation.

*b Level of evidence.
may present as ST-elevation acute coronary syndrome (ACS), non-ST-elevation MI or unstable angina.

Of course, thoracic pain may also be due to gastro-oesophageal reflux disease, musculo-skeletal problem, aortic disease or pericardial disease. A detailed discussion of the management of this group with non-anginal pain is beyond the scope of these Guidelines.

### 6.7.1 Microvascular angina

#### 6.7.1.1 Clinical picture (see web addenda)

Primary coronary microvascular disease should be suspected by exclusion in patients with sufficiently typical chest pain in whom, despite abnormalities of the ECG and/or stress test results indicative of myocardial ischaemia, coronary angiography fails to show fixed or dynamic obstructions in epicardial coronary arteries. Microvascular disease may also occur in the setting of specific diseases, such as hypertrophic cardiomyopathy or aortic stenosis, and this is defined as secondary coronary microvascular disease (which is not addressed in these Guidelines).

Arterial hypertension, either with or without associated ventricular hypertrophy, is frequently encountered in the population with chest pain and ‘normal coronary arteries’. The consequence of coronary microvascular disease—which is still often called ‘hypertensive heart disease’ but is similarly encountered in patients with diabetes or a strong family history of vascular disease—is a reduced coronary flow reserve (CFR) and later interstitial and perivascular fibrosis, resulting in impaired diastolic dysfunction. Even later in the course of the disease, epicardial plaques and stenoses may develop and eventually dominate the clinical picture.

#### 6.7.1.2 Pathogenesis and prognosis (see web addenda)

More details of the clinical presentation, the pathogenesis and prognosis of coronary microvascular disease are discussed in the web addenda of these Guidelines.

#### 6.7.1.3 Diagnosis and management of coronary microvascular disease (see web addenda)

Diagnosis and management of patients with microvascular angina represent a complex challenge. The diagnosis may be made if a patient with exercise-induced angina has normal or non-obstructed coronary arteries by arteriography (coronary CTA or ICA), but objective signs of exercise-induced ischaemia (ST-depression on exercise ECG, ischaemic changes) by MPI. Usually no wall motion abnormalities can be induced during DSE (Table 23). It is necessary to differentiate this pain from non-cardiac chest pain. Diffuse coronary artery spasm, pronounced in the distal epicardial coronary arteries and probably extending into the microvasculature, may be provoked by intracoronary injection of acetylcholine in a substantial proportion of patients with typical coronary microvascular disease. The clinical presentation of patients with microvascular disease differs from those with purely vasospastic angina, since the former usually have exercise-related symptoms in addition to symptoms at rest.

Invasive and non-invasive methods of supporting the diagnosis of microvascular disease (and supporting also some of the recommendations below) are discussed in the web addenda of this document.

### 6.7.2 Vasospastic angina

#### 6.7.2.1 Clinical picture

Patients with vasospastic angina present with typically located anginal pain, which occurs at rest but does not—or occurs only occasionally—with exertion. Such pain typically occurs at night and in the early morning hours. If the chest pain is severe, this may lead to hospital admission. Nitrates usually relieve the pain within minutes. Angina at rest caused by spasm is often observed in patients with otherwise stable obstructive atherosclerosis, while spasm-induced effort angina can occasionally occur in patients with non-obstructive atherosclerosis.

#### 6.7.2.2 Pathogenesis and prognosis (see web addenda)

These aspects of vasospastic angina are discussed in the web addenda of this document.

#### 6.7.2.3 Diagnosis of vasospastic angina

##### 6.7.2.3.1 Electrocardiography

The ECG during vasospasm is classically described as showing ST-elevation. Angiographically, these patients usually show focal occlusive spasm (Prinzmetal’s angina or variant angina). Most patients with coronary vasospasm, however, angiographically show distally pronounced diffuse subtotal vasospasm, which is usually associated with ST-depression. This form of spasm is usually associated with microvascular spasm and is found in patients presenting with microvascular and resting angina. In other patients, no ST-segment shift is seen during provoked vasospasm. As attacks of vasospasm tend to resolve themselves quickly, 12-lead ECG documentation is often difficult. Repeated 24-h ECG monitoring may be able to capture ST-segment shifts associated with anginal symptoms in these patients.
6.7.2.3.2 Coronary arteriography
Although the demonstration of ST-elevation at the time of angina and a normal coronary arteriogram make the diagnosis of variant angina highly likely, there is often uncertainty about the diagnosis in less well-documented or clinically less straightforward cases.
Spontaneous spasm during coronary arteriography is only occasionally observed in patients with symptoms suggestive of vasospastic angina. Hence, provocation tests are commonly used to demonstrate the presence and also the type of coronary vasospasm. Hyperventilation and the cold pressor test have only a rather limited sensitivity for the detection of coronary spasm. Thus, acetylcholine injections into the coronary artery are nowadays used in most centres for provocation of coronary spasm (Table 24). Acetylcholine is injected in incremental doses up to 200 μg, separated by intervals. Intra-coronary ergonovine provocation at incremental doses of up to 60 μg gives similar results.
Coronary spasm may be focal or diffuse. Lumen reductions between 75–99% when compared with the diameter following nitroglycerin injection are defined as spasm in the literature, but severe chest pain and ST-segment depression may also occur without epicardial spasm. The latter phenomenon, which has been termed microvascular spasm, is often seen in patients with a history of microvascular angina. Lumen reductions <30% are commonly seen in non-spastic coronary segments and may represent the ‘physiological’ constrictor response to high-dose acetylcholine or to ergonovine provocation.
Acetylcholine or ergonovine provocation of coronary spasm is a safe test, provided that the agent is infused selectively into the left coronary artery or the right coronary artery. Non-invasive intravenous ergonovine provocative testing has also been described, with echocardiographic or perfusion scintigraphy supplementing electrocardiographic monitoring, increasing the sensitivity and specificity of these tests. However, as fatal complications may occur with intravenous injection of ergonovine, due to prolonged spasm involving multiple vessels, the intracoronary route is preferred. Provocative testing with intravenous ergonovine is not recommended in patients without known coronary anatomy, nor in patients with high-grade obstructive lesions on coronary arteriography.

7. Lifestyle and pharmacological management
7.1 Risk factors and ischaemia management
7.1.1 General management of stable coronary artery disease patients
The aim of the management of SCAD is to reduce symptoms and improve prognosis. The management of CAD patients encompasses lifestyle modification, control of CAD risk factors, evidence-based pharmacological therapy and patient education. Lifestyle recommendations are described in recent ESC guidelines.

7.1.2 Lifestyle modifications and control of risk factors
7.1.2.1 Smoking
Smoking is a strong and independent risk factor for CVD and all smoking, including environmental smoking exposure, must be avoided in all patients with CVD. The benefits of smoking cessation have been extensively reported and quitting smoking is potentially the most effective of all preventive measures, being associated with a reduction in mortality of 36% after MI. Clinicians treating patients with CAD can take advantage of the unique situation and emphasize that the risk of future CAD events can be dramatically reduced by smoking cessation. Thus, smoking status should be assessed systematically (including passive smoking) and all smokers advised to quit and offered cessation assistance. Quitting smoking is complex because smoking is both pharmacologically and psychologically highly addictive. Advice, encouragement and pharmacological aid consistently improve success rates. Nicotine replacement therapy is safe in patients with CAD and should routinely be offered. Bupropion and varenicline have been found safe to use in patients with stable CAD in some studies, although the safety of varenicline has recently been questioned in a meta-analysis, being associated with a small but statistically significant increase in CVD.

7.1.2.2 Diet (Table 25)
A healthy diet reduces CVD risk. Cornerstones of a healthy diet are summarized below. Energy intake should be limited to the amount of energy needed to maintain (or obtain) a healthy weight—that is, a BMI <25 kg/m². In general, when following the rules for a healthy diet, no dietary supplements are needed. N-3 polyunsaturated fatty acid (PUFA) consumption, mainly from oily fish, is potentially associated with beneficial effects on cardiac risk factors, notably reduction in triglycerides, but not all randomized, controlled trials have shown reductions in CV events. Thus current recommendations are to increase PUFA intake through fish consumption, rather than from supplements. Recently, the largest study ever conducted with a so-called ‘Mediterranean’ diet, supplemented with extra-virgin olive oil...
7.1.2.3 Physical activity

Regular physical activity is associated with a decrease in CV morbidity and mortality in patients with established CAD and physical activity should be incorporated into daily activities. Aerobic exercise should be offered to patients with known CAD, usually as part of a structured cardiac rehabilitation program, with the need for an evaluation of both exercise capacity and exercise-associated risk. Patients with previous acute MI, CABG, percutaneous coronary intervention (PCI), stable angina pectoris or stable chronic heart failure should undergo moderate-to-vigorous intensity aerobic exercise training ≥3 times a week and for 30 min per session. Sedentary patients should be strongly encouraged to start light-intensity exercise programs after adequate exercise-related risk stratification. In patients with significant CAD who are not candidates for revascularization, exercise training may offer an alternative means of symptom alleviation and improved prognosis.

7.1.2.4 Sexual activity

Sexual activity is associated with an exercise workload of up to 6 METS (1 MET = approximately 3.5 mL oxygen consumption/kg/min) depending on the type of activity. Sympathetic activation is intrinsic to sexual arousal and heart rate and blood pressure (BP) response may be higher than expected from the level of exercise. Sexual activity may thus trigger ischaemia, and nitroglycerin prior to sexual intercourse may be helpful as in other physical activity.

Patients with mild angina, successful coronary revascularization and New York Heart Association (NYHA) functional Class I heart failure generally do not need specific evaluation before resuming sexual activity. Patients with more symptomatic heart disease, including moderate angina, may be guided by an exercise stress test as a means of assessing risk and reassuring the patient. Exercise training should be advocated to improve exercise capacity and reduce myocardial oxygen consumption during sexual activity.

Erectile dysfunction (ED) is associated with cardiac risk factors and is more prevalent in patients with CAD. The common denominator between erectile dysfunction and CAD is endothelial dysfunction and antihypertensive medication —in particular β-blockers and thiazides—increases the risk of erectile dysfunction.

Lifestyle and pharmacological intervention—including weight loss, exercise training, smoking cessation and statin treatment—ameliorates ED. Pharmacological therapy with phosphodiesterase type 5 (PDE5) inhibitors (sildenafil, tadalafil and vardenafil) are effective, safe and well tolerated in men with stable CAD. Low-risk patients, as defined above, can usually receive PDE5 inhibitors without cardiac work-up. However, use of nitric oxide donors, i.e. all of the preparations of nitroglycerin as well as isosorbide mononitrate and isosorbide dinitrate, are absolute contra-indications to the use of PDE5 inhibitors because of the risk of synergistic effects on vasodilation, causing hypotension and haemodynamic collapse. PDE5 inhibitors are not recommended in patients with low blood pressure, with severe heart failure (NYHA III–IV), refractory angina or recent CV events. Patients must be informed about the potentially harmful interactions between PDE5 inhibitors and nitrates. If a patient on a PDE5 inhibitor develops chest pain, nitrates should not be administered in the first 24 hours (sildenafil, vardenafil) to 48 hours (tadalafil).

7.1.2.5 Weight management

Both overweight and obesity are associated with an increased risk of death in CAD. Weight reduction in overweight and obese people is recommended in order to achieve favourable effects on BP, dyslipidaemia and glucose metabolism. The presence of sleep apnoea symptoms should be carefully assessed, especially in obese patients. Sleep apnoea has been associated with an increase in CV mortality and morbidity.

7.1.2.6 Lipid management

Dyslipidaemia should be managed according to lipid guidelines with pharmacological and lifestyle intervention. Patients with established CAD are regarded as being at very high risk for cardiovascular events and statin treatment should be considered, irrespective of low density lipoprotein (LDL) cholesterol (LDL-C) levels. The goals of treatment are LDL-C below 1.8 mmol/L (<70 mg/dL) or >50% LDL-C reduction when target level cannot be reached. In the majority of patients this is achievable through statin monotherapy. Other interventions (e.g. fibrates, resins, nicotinic acid, ezetimibe) may lower LDL cholesterol but no benefit on clinical outcomes has been reported for these alternatives. Although elevated levels of triacylglycerides and low HDL cholesterol (HDL-C) are associated with increased CVD risk, clinical trial evidence is insufficient to specify treatment targets, which should be regarded as not indicated.

For patients undergoing PCI for SCAD, high dose atorvastatin has been shown to reduce the frequency of peri-procedural MI in both statin-naïve patients and patients receiving chronic statin therapy. Thus reloading with high intensity statin before PCI may be considered.

7.1.2.7 Arterial Hypertension

Particular attention should be given to control of elevated BP but thresholds for the definition of hypertension by 24-h ambulatory and home BP monitoring differ from those measured at office or clinic (see Table 26). Elevated BP is a major risk factor for CAD as well as heart failure, cerebrovascular disease and renal failure. There is sufficient evidence to recommend that systolic BP (SBP)
be lowered to <140 mmHg and diastolic BP (DBP) to <90 mmHg in SCAD patients with hypertension. Based on current data, it may be prudent to recommend lowering SBP/DBP to values within the range 130–139/80–85 mmHg. BP targets in diabetes are recommended to be <140/85 mmHg (see below).37,273

7.1.2.8 Diabetes and other disorders (see also chapter 9 and web addenda)

Diabetes mellitus is a strong risk factor for CV complications, increases the risk of progression of coronary disease and should be managed carefully, with good control of glycated haemoglobin (HbA1c) to <7.0% (53 mmol/mol) generally and <6.5%–6.9% (48–52 mmol/mol) on an individual basis. Glucose control should be based on individual considerations, depending on the patient’s characteristics including age, presence of complications and diabetes duration.

As for other disorders, attention to management of risk factors is recommended, including weight management, exercise recommendations and statin treatment with an LDL-C target of 1.8 mmol/L (<70 mg/dL) in diabetic patients with angiographically proven CAD.62 The traditional treatment goal for BP in diabetes, i.e. below 130 mmHg, is not supported by outcome evidence in trials and has been difficult to achieve in the majority of patients. Thus, the BP target in patients with CAD and diabetes is to be <140/85 mmHg. An angiotensin converting enzyme (ACE) inhibitor or renin-angiotensin receptor blocker should always be included because of the renal protective effects.37,274,275

Patients with chronic kidney disease (CKD) are at high risk and particular care should be taken to address risk factors and achieve BP and lipid targets. Statins are generally well tolerated in CKD stages 1–2 (GFR >60–89 mL/min/1.73 m^2) whereas, in CKD stages 3–5, statins with minimal renal excretion should be chosen (atorvastatin, fluvastatin, pitavastatin, rosuvastatin).62

7.1.2.9 Psychosocial factors

Depression, anxiety and distress are common in patients with CAD. Patients should be assessed for psychosocial distress and appropriate care offered. Refer for psychotherapy, medication or collaborative care in the case of clinically significant symptoms of depression, anxiety and hostility. This approach can reduce symptoms and enhance quality of life, although evidence for a definite beneficial effect on cardiac endpoints is inconclusive.37

7.1.2.10 Cardiac rehabilitation

A comprehensive risk-reduction regimen, integrated into comprehensive cardiac rehabilitation, is recommended to patients with CAD.37,276 Cardiac rehabilitation is commonly offered after MI or recent coronary intervention, but should be considered in all patients with CAD, including those with chronic angina. Exercise-based cardiac rehabilitation is effective in reducing total- and CV mortality and hospital admissions,276 whereas effects on total MI or revascularization (CABG or PCI) are less clear, especially in the long term.277,278

Evidence also points towards beneficial effects on health-related quality of life (QoL). In selected sub-groups, centre-based cardiac rehabilitation may be substituted for home-based rehabilitation, which is non-inferior. Patient participation in cardiac rehabilitation remains far too low, particularly in women, the elderly and the socio-economically deprived, and could benefit from systematic referral.

7.1.2.11 Influenza vaccination

An annual influenza vaccination is recommended for patients with CAD, especially the elderly.279,280

7.1.2.12 Hormone replacement therapy

For decades, evidence from epidemiological and laboratory studies led us to believe that circulating oestrogens had a beneficial effect on the risk of CVD and that this could be transferred to the benefits of hormone replacement therapy (HRT). However, results from large randomized trials have not supported this; on the contrary, HRT increases the risk of CVD in women above the age of 60.281

The mechanisms are unclear and, if instituted at an earlier age (i.e. at the time of menopause) in women with intact vascular endothelium and few CV risk factors, the effect of HRT is still debated.282 However, HRT is at present not recommended for primary or secondary prevention of CVD.

7.1.3 Pharmacological management of stable coronary artery disease patients

7.1.3.1 Aims of treatment

The two aims of the pharmacological management of stable CAD patients are to obtain relief of symptoms and to prevent CV events.

Relief of anginal symptoms: rapidly acting formulations of nitroglycerin are able to provide immediate relief of the angina symptoms once the episode has started or when the symptom is likely to occur (immediate treatment or prevention of angina). Anti-isaemic drugs—but also lifestyle changes, regular exercise training, patient education and revascularization—all have a role to play in minimizing or eradicating symptoms over the long term (long-term prevention).

To prevent the occurrence of CV events: efforts to prevent MI and death in coronary disease focus primarily on reducing the incidence of acute thrombotic events and the development of ventricular dysfunction. These aims are achieved by pharmacological or lifestyle interventions which: (i) reduce plaque progression; (ii) stabilize plaque, by reducing inflammation and (iii) prevent thrombosis, should plaque rupture or erosion occur. In patients with severe lesions in coronary arteries supplying a large area of jeopardized myocardium, a combined pharmacological and revascularization strategy

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**Table 26** Blood pressure thresholds for definition of hypertension with different types of blood pressure measurement (adapted from Umpierrez et al. 2012 273).

<table>
<thead>
<tr>
<th>BP Type</th>
<th>SBP (mmHg)</th>
<th>DBP (mmHg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Office BP</td>
<td>140</td>
<td>90</td>
</tr>
<tr>
<td>Home BP</td>
<td>135</td>
<td>85</td>
</tr>
<tr>
<td>Ambulatory BP</td>
<td>130</td>
<td>80</td>
</tr>
<tr>
<td>24-h</td>
<td>135</td>
<td>85</td>
</tr>
<tr>
<td>Daytime (awake)</td>
<td>135</td>
<td>85</td>
</tr>
<tr>
<td>Nighttime (asleep)</td>
<td>120</td>
<td>70</td>
</tr>
</tbody>
</table>

BP = blood pressure; DBP = diastolic blood pressure; SBP = systolic blood pressure.
Nitroglycerin spray acts more rapidly. Nitroglycerin can be used in syncop, lying down enhances venous return and heart work) and angina, acting by their active component nitric oxide (NO) and by the reduction of preload.

**Short-acting nitrates for acute effort angina.** Sublingual nitroglycerin is the standard initial therapy for effort angina. When angina starts, the patient should rest sitting (standing promotes syncopa, lying down enhances venous return and heart work) and take sublingual nitroglycerin (0.3–0.6 mg) every 5 min until the pain goes or a maximum of 1.2 mg has been taken within 15 min. Nitroglycerin spray acts more rapidly. Nitroglycerin can be used prophylactically when angina can be expected, such as after a meal, emotional stress, sexual activity and in colder weather.

**Isosorbide dinitrate** (5 mg sublingually) helps to abort anginal attacks for about 1 h. Because the dinitrate requires hepatic conversion to the mono-nitrate, the onset of anti-anginal action (within 3–4 min) is slower than with nitroglycerin. After oral ingestion, haemodynamic and anti-anginal effects persist for several hours, conferring longer protection against angina than sublingual nitroglycerin.

**Long-acting nitrates for angina prophylaxis.** Long-acting nitrates are not continuously effective if regularly taken over a prolonged period without a nitrate-free or nitrate-low interval of about 8–10 hours (tolerance). Worsening of endothelial dysfunction is a potential complication of long-acting nitrates, hence the common practice of the routine use of long-acting nitrates as first line therapy for patients with effort angina needs re-evaluation.

**Isosorbide dinitrate** (oral preparation) is frequently given for the prophylaxis of angina. In a crucial placebo-controlled study, exercise duration improved significantly for 6–8 h after single oral doses of 15–120 mg isosorbide dinitrate, but for only 2 h when the same doses were given repetitively four times daily, despite much higher plasma isosorbide dinitrate concentrations during sustained than during acute therapy. With the extended-release formulation of isosorbide dinitrate, eccentric twice-daily dosing, with 40 mg in the morning, repeated 7 hours later, was not superior to placebo in a large multicentre study. Thus prolonged therapy with isosorbide dinitrate is not evidence-based.

**Mononitrates** have similar dosage and effects to those of isosorbide dinitrate. Nitrate tolerance—likewise a potential problem—can be prevented by changes in dosing and timing of administration, as well as by using slow-release preparations. Only twice-daily rapid-release preparations or very high doses of slow-release mononitrate—also twice daily—give sustained anti-anginal benefit.

**Transdermal nitroglycerin patches** fail to cover 24 h during prolonged use. A discontinuous administration at 12 h intervals allows on and off effects to start within minutes and last 3–5 h. There are no efficacy data for second or third doses during chronic administration.

**Nitrates should not be given with PDE5 inhibitors.** Combined with dihydropyridines (DHPs) to control angina. Combination therapy of β-blockers with verapamil and diltiazem should be avoided because of the risk of bradycardia or AV block (Table 27).

**Nitrate side-effects.** Hypotension is the most serious, and headache the most common side-effect of nitrates. Headaches (aspirin may relieve these) may facilitate loss of compliance, yet often pass over. Failure of therapy. Apart from non-compliance, treatment failure includes nitric oxide resistance and nitrate tolerance.

**Nitrate drug interactions.** Many are pharmacodynamic, including potentiation of vasodilator effects with calcium channel blockers (CCBs). Note that serious hypotension can occur with the selective PDE5 inhibitors (sildenafil and others) for erectile dysfunction or for the treatment of pulmonary hypertension. Sildenafil decreases the BP by about 8.4/5.5 mmHg and by much more with nitrates. In the case of inadvertent PDE5–nitrate combinations, emergency α-adrenergic agonists or even norepinephrine may be needed. Nitrates should not be given with α-adrenergic blockers. In men with prostatic problems, taking tamsulosin (α1A and α1D blocker), nitrates can be given.

**β-blockers** act directly on the heart to reduce heart rate, contractility, atrioventricular (AV) conduction and ectopic activity. Additionally, they may increase perfusion of ischaemic areas by prolonging the diastole and increasing vascular resistance in non-ischaemic areas. In post-MI patients, β-blockers achieved a 30% risk reduction for CV death and MI. Thus β-blockers may also be protective in patients with SCAD, but without supportive evidence from placebo-controlled clinical trials. However, a recent retrospective analysis of the REACH registry suggested that, in patients with either CAD risk factors only, known prior MI, or known CAD without MI, the use of β-blockers was not associated with a lower risk of cardiovascular events. Although propensity score matching was used for the analysis, the demonstration lacks the strength of a randomized evaluation. Among other limitations, most of the β-blocker trials in post-MI patients were performed before the implementation of other secondary prevention therapies, such as statins and ACE inhibitors, leaving uncertainty regarding their efficacy when added to modern therapeutic strategies. β-Blockers are clearly effective in controlling exercise-induced angina, improving exercise capacity and limiting both symptomatic as well as asymptomatic ischaemic episodes. Regarding angina control, β-blockers and CCBs are similar. β-Blockers can be combined with dihydropyridines (DHPs) to control angina. Combination therapy of β-blockers with verapamil and diltiazem should be avoided because of the risk of bradycardia or AV block (Table 27).

The most widely used β-blockers in Europe are those with predominant β1-blockade, such as metoprolol, bisoprolol, atenolol or nebivolol. Carvedilol, a non-selective β1+β2 blocker, is also often used. All of these reduce cardiac events in patients with heart failure. In summary, there is evidence for prognostic benefits from the use of β-blockers in post-MI patients, or in heart failure. Extrapolation from these data suggests that β-blockers may be the first-line anti-anginal strategy in stable CAD patients without contraindications. Nebivolol and bisoprolol are partly secreted by the kidney, whereas carvedilol and metoprolol are metabolized by the liver, hence being safer in patients with renal compromise.

**Calcium channel blockers.** Calcium antagonists (i.e. CCBs) act chiefly by vasodilation and reduction of the peripheral vascular resistance. CCBs are a heterogeneous group of drugs that can chemically be classified into the DHPs and the non-DHPs, their common pharmacological property being selective inhibition of L-channel opening in vascular smooth muscle and in the myocardium.
**Table 27** Major side-effects, contra-indications, drug–drug interactions (DDI) and precautions of anti-ischaemic drugs. (List is not exhaustive: refer to summary of products characteristics for details.)

<table>
<thead>
<tr>
<th>Drug class</th>
<th>Side effects</th>
<th>Contraindications</th>
<th>DDI</th>
<th>Precautions</th>
</tr>
</thead>
</table>
| Short-acting and long-acting nitrates[133] | • Headache  
• Flushing  
• Hypotension  
• Syncope and postural hypotension  
• Reflex tachycardia  
• Methaemoglobinemia | • Hypertrophic obstructive cardiomyopathy | • PDE5 inhibitors (sildenafil or similar agents)  
• α-adrenergic blockers  
• CCBs | - |
| B-blockers[290, 293, 302, b] | • Fatigue, depression[304]  
• Bradycardia  
• Heart block  
• Bronchospasm  
• Peripheral vasoconstriction  
• Postural hypotension  
• Impotence  
• Hypoglycaemia/mask hypoglycaemia | • Low heart rate or heart conduction disorder  
• Cardiogenic shock  
• Asthma  
• COPD caution; may use cardioselective B-blockers if fully treated by inhaled steroids and long-acting β-agonists[100]  
• Severe peripheral vascular disease  
• Decompensated heart failure  
• Vasospastic angina | • Heart-rate lowering CCB  
• Sinus-node or AV conduction depressors | • Diabetics  
• COPD[110] |
| CCBs: heart-rate lowering[293, 294] | • Bradycardia  
• Heart conduction defect  
• Low ejection fraction  
• Constipation  
• Gingival hyperplasia | • Low heart rate or heart rhythm disorder  
• Sick sinus syndrome  
• Congestive heart failure  
• Low BP | • Cardiodepressant (B-blockers, flecaïnide)  
• CYP3A4 substrates | - |
| CCBs: Dihydropyridines[27, 305, 331] | • Headache  
• Ankle swelling  
• Fatigue  
• Flushing  
• Reflex tachycardia | • Cardiogenic shock  
• Severe aortic stenosis  
• Obstructive cardiomyopathy | • CYP3A4 substrates | - |
| Ivabradine[307] | • Visual disturbances  
• Headache, dizziness  
• Bradycardia  
• Atrial fibrillation  
• Heart block | • Low heart rate or heart rhythm disorder  
• Allergy  
• Severe hepatic disease | • QTc prolonging drugs  
• Macrolide antibiotics  
• Anti-HIV  
• Anti-fungal | • Age >75 years  
• Severe renal failure |
| Nicorandil[317] | • Headache  
• Flushing  
• Dizziness, weakness  
• Nausea  
• Hypotension  
• Oral, anal, gastrointestinal ulceration | • Cardiogenic shock  
• Heart failure  
• Low blood pressure | • PDE5 inhibitors (Sildenafil or similar agents) | - |
| Trimetazidine[315, 316] | • Gastric discomfort  
• Nausea  
• Headache  
• Movement disorders | • Allergy  
• Parkinson disease  
• Tremors and movement disorders  
• Severe renal impairment | • None reported  
• Moderate renal impairment  
• Elderly | - |
| Ranolazine[317, 318] | • Dizziness  
• Constipation  
• Nausea  
• QT prolongation | • Liver cirrhosis  
• CYP450 substrates (digoxin, simvastatin, cyclosporine)  
• QTc prolonging drugs | - |
| Allopurinol[223] | • Rash  
• Gastric discomfort | • Hypersensitivity  
• Mercaptopurine / Azathioprine | • Severe renal failure | - |

AV = atrioventricular; CCBs = calcium channel blockers; CHF = congestive heart failure; COPD = chronic obstructive pulmonary disease; DDI = Drug-Drug Interactions; HIV = Human Immunodeficiency Virus; PDE5 = phosphodiesterase type 5.

*Very frequent or frequent; may vary according to specific drugs within the therapeutic class.

| a | Atenolol, metoprolol CR, bisoprolol, carvedilol. |
Distinctions between the DHPs and non-DHPs are reflected in different binding sites on the calcium channel pores and in the greater vascular selectivity of the DHP agents (amlodipine, nifedipine, felodipine).

The non-DHPs, by virtue of nodal inhibition, tend to reduce the heart rate (heart rate-lowering agents, verapamil and diltiazem) and explain the anti-anginal properties.

† Non-dihydropyridine (heart rate-lowering calcium channel blockers)

Verapamil. Among CCBs, verapamil has a large range of approved indications, including all varieties of angina (effort, vasospastic, unstable), supraventricular tachycardias and hypertension. Indirect evidence suggests good safety but with risks of heart block, bradycardia and heart failure. Compared with metoprolol, the anti-anginal activity was similar.298 Compared with atenol in hypertension with CAD, verapamil gave less new diabetes, fewer anginal attacks,303 and less psychological depression.304

β-Blockade combined with verapamil is not advised (due to risk of heart block): instead, use DHP-β-blockade.

Diltiazem. Diltiazem, with its low side-effect profile, has advantages, compared with verapamil, in the treatment of effort angina.295 Like verapamil, it acts by peripheral vasodilation, relief of exercise-induced coronary constriction, a modest negative inotropic effect and sinus node inhibition. There are no outcome studies comparing diltiazem and verapamil. As with verapamil, combination with β-blockade, as well as the use in patients with CAD and left ventricular dysfunction, is not advised.

† Dihydropyridines

Long-acting nifedipine. This agent is a powerful arterial vasodilator with few serious side-effects. Long-acting nifedipine is especially well-tested in hypertensive anginal patients when added to β-blockade.27 In ACTION, a large placebo-controlled trial long-acting nifedipine in SCAD proved to be safe and reduced the need for coronary angiography and cardiovascular interventions.27 Contra-indications to nifedipine are few (severe aortic stenosis, obstructive cardiomyopathy, or heart failure) and careful combination with β-blockade is usually feasible and desirable. Vasodilatory side-effects include headache and ankle oedema.

<table>
<thead>
<tr>
<th>Indication</th>
<th>Class</th>
<th>Level</th>
<th>Ref.</th>
<th>C</th>
</tr>
</thead>
<tbody>
<tr>
<td>General considerations</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Optimal medical treatment indicates at least one drug for angina/ischaemia relief plus drugs for event prevention.</td>
<td>I</td>
<td>C</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>It is recommended to educate patients about the disease, risk factors and treatment strategy.</td>
<td>I</td>
<td>C</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>It is indicated to review the patient’s response soon after starting therapy.</td>
<td>I</td>
<td>C</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>Angina/ischaemia† relief</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Short-acting nitrates are recommended.</td>
<td>I</td>
<td>B</td>
<td>3, 329</td>
<td></td>
</tr>
<tr>
<td>First-line treatment is indicated with β-blockers and/or calcium channel blockers to control heart rate and symptoms.</td>
<td>I</td>
<td>A</td>
<td>3, 331</td>
<td></td>
</tr>
<tr>
<td>For second-line treatment it is recommended to add long-acting nitrates or ivabradine or nicorandil or ranolazine, according to heart rate, blood pressure and tolerance.</td>
<td>IIa</td>
<td>B</td>
<td>177, 307, 3, 199, 284, 286, 308, 319-321, 328</td>
<td></td>
</tr>
<tr>
<td>For second-line treatment, trimetazidine may be considered.</td>
<td>IIb</td>
<td>B</td>
<td>313, 315</td>
<td></td>
</tr>
<tr>
<td>According to comorbidities/tolerance it is indicated to use second-line therapies as first-line treatment in selected patients.</td>
<td>I</td>
<td>C</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>In asymptomatic patients with large areas of ischaemia (&gt;10%) β-blockers should be considered.</td>
<td>IIa</td>
<td>C</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>In patients with vasospastic angina, calcium channel blockers and nitrates should be considered and beta-blockers avoided.</td>
<td>IIa</td>
<td>B</td>
<td>3, 365</td>
<td></td>
</tr>
<tr>
<td>Event prevention</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Low-dose aspirin daily is recommended in all SCAD patients.</td>
<td>I</td>
<td>A</td>
<td>333, 334, 366</td>
<td></td>
</tr>
<tr>
<td>Clopidogrel is indicated as an alternative in case of aspirin intolerance.</td>
<td>I</td>
<td>B</td>
<td>335</td>
<td></td>
</tr>
<tr>
<td>Statins are recommended in all SCAD patients.</td>
<td>I</td>
<td>A</td>
<td>62</td>
<td></td>
</tr>
<tr>
<td>It is recommended to use ACE inhibitors (or ARBs) if presence of other conditions (e.g. heart failure, hypertension or diabetes).</td>
<td>I</td>
<td>A</td>
<td>348, 349, 351, 352</td>
<td></td>
</tr>
</tbody>
</table>

ACE = angiotensin converting enzyme; SCAD = stable coronary artery disease.

a Class of recommendation.

b Level of evidence.
c Reference(s) supporting levels of evidence.
d No demonstration of benefit on prognosis.

Table 28 Pharmacological treatments in stable coronary artery disease patients

Dihydropyridines

Long-acting nifedipine. This agent is a powerful arterial vasodilator with few serious side-effects. Long-acting nifedipine is especially well-tested in hypertensive anginal patients when added to β-blockade.27 In ACTION, a large placebo-controlled trial long-acting nifedipine in SCAD proved to be safe and reduced the need for coronary angiography and cardiovascular interventions.27 Contra-indications to nifedipine are few (severe aortic stenosis, obstructive cardiomyopathy, or heart failure) and careful combination with β-blockade is usually feasible and desirable. Vasodilatory side-effects include headache and ankle oedema.

Diltiazem. Diltiazem, with its low side-effect profile, has advantages, compared with verapamil, in the treatment of effort angina.295 Like verapamil, it acts by peripheral vasodilation, relief of exercise-induced coronary constriction, a modest negative inotropic effect and sinus node inhibition. There are no outcome studies comparing diltiazem and verapamil. As with verapamil, combination with β-blockade, as well as the use in patients with CAD and left ventricular dysfunction, is not advised.
Amlodipine. The very long half-life of amiodipine and its good tolerability make it an effective once-a-day anti-anginal and antihypertensive agent, setting it apart from agents that are taken either twice or three times daily. Side-effects are few; mainly ankle oedema. In patients with CAD and normal blood pressure, amiodipine reduced CV events in a 24-month trial. Exercise-induced ischaemia is more effectively reduced by amiodipine than by the β-blocker atenolol and the combination is even better.

However, the CCB–β-blocker combination is often underused, even in some studies reporting ‘optimally treated’ stable effort angina.

Others. Felodipine, lacidipine and lercanidipine share the standard properties of other long-acting DHPs.

7.1.3.3.4 Ivabradine. Ivabradine is a heart rate-lowering agent selectively inhibiting the sinus node (If) pacemaking current, thereby decreasing the myocardial oxygen demand without effect on inotropism or BP. It was approved by the European Medicines Agency (EMA) for therapy of chronic stable angina in patients intolerant to—or inadequately controlled by—β-blockers and whose heart rate exceeded 60 b.p.m. (in sinus rhythm). Ivabradine was as effective as atenolol or amiodipine in patients with SCAD; adding ivabradine 7.5 mg twice daily to atenolol therapy gave better control of angina. Occasional side-effects include oral, intestinal and perioral analgesia. In patients with prior angina enrolled in the Morbidity-Mortality Evaluation of the Iα Inhibitor Ivabradine in Patients With Coronary Artery Disease and Left Ventricular Dysfunction (BEAUTIFUL) trial, ivabradine reduced the composite primary endpoint of CV death, hospitalization with MI and HF, and reduced hospitalization for MI. The effect was predominant in patients with a heart rate ≥70 bpm. Ivabradine is thus an effective anti-anginal agent, alone or in combination with β-blockers.

7.1.3.3.5 Nicorandil. Nicorandil is a nitraterelative of nicotinamide that can be used for the prevention and long-term treatment of angina, and may be added after β-blockers and CCBs. It is EMA-but not FDA-approved. Nicorandil dilates epicardial coronary arteries and stimulates ATP-sensitive potassium channels (KATP) in vascular smooth muscle. In the prospective Impact Of Nicorandil in Angina (IONA) study, over a mean of 1.6 years in 5126 patients with SCAD, CV events were reduced by 14% (relative risk 0.86; P = 0.027). However, symptom relief was not reported. Long-term use of oral nicorandil may stabilize coronary plaque in patients with stable angina. Occasional side-effects include oral, intestinal and peri-anal ulceration.

7.1.3.3.6 Trimetazidine. Trimetazidine is an anti-ischaemic metabolic modulator, similar to propranolol in doses of 20 mg thrice daily. The heart rate and rate pressure product at rest and at peak exercise remained unchanged in the trimetazidine group, thus showing a non-mechanical anti-ischaemic action.

Trimetazidine (35 mg twice daily) added to beta-blockade (atenolol) improved effort-induced myocardial ischaemia, as reviewed by the EMA in June 2012, and remains contra-indicated in Parkinson’s disease and motion disorders [such as tremor (shaking), muscle rigidity and walking disorders and restless leg syndrome]. In diabetic persons, trimetazidine improved HbA1c and glycaemia, while increasing forearm glucose uptake. Trimetazidine has not been evaluated in large outcome studies in SCAD patients.

7.1.3.3.7 Ranolazine. Ranolazine is a selective inhibitor of late sodium current with anti-ischaemic and metabolic properties. Doses of 500–2000 mg daily reduced angina and increased exercise capacity without changes in heart rate or BP. The EMA approved ranolazine in 2009 for add-on treatment in stable angina in patients inadequately controlled by—or intolerant to—first-line agents (beta-blockers and/or calcium antagonists). In the 6560 patients of the Metabolic Efficiency with Ranolazine for Less Ischemia in Non-ST-Elevation Acute Coronary Syndromes: Thrombolysis In Myocardial Infarction (MERLIN-TIMI 36) trial presenting with recent non-ST-elevation ACS (NSTE-ACS), ranolazine therapy showed no overall benefit. In patients with prior chronic angina enrolled in the MERLIN trial, ranolazine reduced recurrent ischaemia [hazard ratio (HR) 0.78; P = 0.002] in those studied after the coronary event, ranolazine reduced the incidence of newly increased HbA1c by 32%. In the recent TERISA study (Type 2 Diabetes Evaluation of Ranolazine in Subjects With Chronic Stable Angina), ranolazine reduced episodes of stable angina in 949 diabetes patients already receiving one or two anti-anginal drugs and led to less use of sublingual nitroglycerin, and the benefits appeared more prominent in patients with higher rather than lower HbA1c levels. These results suggest that this drug can be added to other well-established anti-anginal drugs, in particular in patients with higher HbA1c levels, who may also more often rely on medical management.

Ranolazine plasma levels increase with cytochrome P3A (CYP3A) inhibitors (diltiazem, verapamil, macrolide antibiotics, grapefruit juice). Ranolazine clearance is reduced by renal and hepatic impairment. Ranolazine increases QTc, and should therefore be used carefully in patients with QT prolongation or on QT-prolonging drugs.

7.1.3.3.8 Allopurinol. Allopurinol, an inhibitor of xanthine oxidase that reduces uric acid in persons with gout, is also anti-anginal. There is limited clinical evidence but, in a randomized crossover study of 65 patients with SCAD, allopurinol 600 mg/day increased times to ST-segment depression and to chest pain. In renal impairment, such high doses may have toxic side-effects. In optimally treated SCAD patients, allopurinol reduced vascular oxidative stress, while in heart failure patients it conserved ATP.

7.1.3.3.9 Molsidomine. This direct NO donor has anti-ischaemic effects similar to those of isosorbide dinitrate. The long-acting once-daily 16 mg formulation is as effective as 8 mg twice daily.

7.1.3.4 Patients with low blood pressure

Anti-anginal drugs should be started at very low doses, with preferential use of drugs with no-or limited impact on BP, such as ivabradine (in patients with sinus rhythm), ranolazine or trimetazidine.

7.1.3.5 Patients with low heart rate

Several studies have shown that increased resting heart rate is a strong independent risk factor for adverse outcome in patients with SCAD. There is a linear relationship between resting heart rate and major cardiovascular events, with a persistent decrease in CV risk with lower heart rate. A clinical benefit has been demonstrated of heart rate reduction using various drugs. Although lowering the heart rate < 60 b.p.m. is an important goal in the treatment of SCAD, patients presenting with low heart rate should be treated differently. Heart rate lowering drugs (β-blockers, ivabradine, heart rate lowering CCBs) should be avoided or used with caution and, if needed, started at very low doses. Anti-anginal drugs without heart lowering effects should preferably be given.
7.2 Event prevention

7.2.1 Antiplatelet agents

Antiplatelet agents decrease platelet aggregation and may prevent formation of coronary thrombus. Due to a favourable ratio between benefit and risk in patients with stable CAD and its low cost, low-dose aspirin is the drug of choice in most cases and clopidogrel may be considered for some patients. The use of antiplatelet agents is associated with a higher bleeding risk.

7.2.1.1 Low-dose aspirin

Aspirin remains the cornerstone of pharmacological prevention of arterial thrombosis. It acts via irreversible inhibition of platelet cyclooxygenase-1 (COX-1) and thus thromboxane production, which is normally complete with chronic dosing ≥75 mg/day. Contrary to the antiplatelet effects, the gastrointestinal side-effects of aspirin increase at higher doses. The optimal risk–benefit ratio appears to be achieved with an aspirin dosage of 75–150 mg/day.332–334

7.2.1.2 P2Y12 inhibitors

P2Y12 inhibitors, including thienopyridines, act as antagonists of the platelet adenosine diphosphate (ADP) receptor P2Y12, thereby inhibiting platelet aggregation. The major study supporting the use of thienopyridine in stable coronary patients is the Clopidogrel vs. Aspirin in Patients at Risk of Ischaemic Events (CAPRIE) trial, which showed an overall benefit of clopidogrel as compared with aspirin (with also a favourable safety profile) in preventing CV events in three categories of patients with previous MI, previous stroke or peripheral vascular disease (PVD).335 The clopidogrel benefit was driven by the peripheral vascular disease (PVD) sub-group and the dose of aspirin with which it was compared (325 mg/day) may not be the safest dose. Clopidogrel should thus be proposed as a second-line treatment, especially for aspirin-intolerant CVD patients. Prasugrel and ticagrelor are new P2Y12 antagonists that achieve greater platelet inhibition, compared with clopidogrel.336,337 Prasugrel and ticagrelor are both associated with a significant reduction of CV outcomes as compared with clopidogrel in ACS patients,338,339 but no clinical studies have evaluated the benefit of these drugs in SCAD patients. After unstable angina or myocardial infarction without ST-segment elevation when patients are stabilized and medically managed, there are no data supporting a beneficial effect of intensified platelet inhibition.340

7.2.1.3 Combination of antiplatelet agents

Dual antiplatelet therapy combining aspirin and a thienopyridine is the standard of care for patients with ACS, including after the acute phase, when the patients are stabilized, or in SCAD patients who have undergone elective PCI.133,339,342 However, in the Clopidogrel for High Atherothrombotic Risk and Ischemic Stabilization, Management, and Avoidance (CHARISMA) study,143 dual antiplatelet therapy did not confer benefit in patients with stable vascular disease or at risk of atherothrombotic events, although a significant benefit was observed in a post-hoc analysis of patients with documented atherothrombotic disease, and in particular in coronary patients with a prior history of MI.144 Combined antiplatelet therapy was also recently tested with an antagonist of protease activated receptor type 1 (PAR-1).145 The primary efficacy endpoint—a composite of CV death, MI or stroke—was significantly reduced with vorapaxar in addition to standard antiplatelet therapy in patients with stable atherosclerosis, and this benefit was particularly evident in the post-MI group of patients.345 However, it increased the risk of moderate or severe bleeding, including intracranial haemorrhage. Altogether, on the basis of these post-hoc analyses, combined antiplatelet therapy may be beneficial only in selected patients at high risk of ischaemic events, but cannot be recommended systematically in SCAD patients.

7.2.1.4 Poor response to antiplatelet agents

There is a wide variation in response to antiplatelet therapy and a great interest has recently emerged in the use of functional and/or genetic assays to guide such treatment. High platelet reactivity on aspirin and/or clopidogrel treatment results from multiple factors, including non-compliance, accelerated platelet turnover, drug interactions, patient characteristics (such as age, gender, diabetes) and single nucleotide polymorphisms [cytochrome P450 2C19 (CYP2C19*2), ATP-binding cassette sub-family B member 1 (ABCB1) for clopidogrel]. The influence of genetic variants on the response to antiplatelet agents, especially clopidogrel, has been well established in patients with ACS and planned PCI, but not in patients with stable CAD.346 However, there is currently no recommendation to perform genetic testing in patients with stable CAD. Platelet function testing in SCAD patients undergoing PCI is not recommended as a routine (see chapter 8).147

7.2.2 Lipid-lowering agents (see lipid management, above)

Patients with documented CAD are regarded as being at very high risk and should be treated with statins, in line with recommendations in the ESC/European Atherosclerosis Society Guidelines for the management of dyslipidaemia.52 The treatment target is LDL-C <1.8 mmol/L and/or >50% reduction if the target level cannot be reached.

7.2.3 Renin-angiotensin-aldosterone system blockers

Angiotensin converting enzyme inhibitors reduce total mortality, MI, stroke and heart failure among specific subgroups of patients, including those with heart failure,348–350 previous vascular disease alone,351–353 or high-risk diabetes.354 Hence, it is appropriate to consider ACE inhibitors for the treatment of patients with SCAD, especially with co-existing hypertension, LVEF ≤40%, diabetes or CKD, unless contra-indicated.

However, not all clinical trials have demonstrated that the ACE inhibitors reduce all-cause mortality, CV mortality, non-fatal MI, stroke and heart failure in patients with atherosclerosis and preserved LV function.351,352,353 In SCAD patients with hypertension, a combination therapy consisting of an ACE inhibitor and a DHP CCB, such as perindopril/amlopidine in the Anglo-Scandinavian Cardiac Outcomes Trial (ASCOT) trial and benazepril/amlopidine in the Avoiding Cardiovascular Events Through Combination Therapy in Patients Living With Systolic Hypertension (ACCOMPLISH) trial, is preferred.356,357 In contrast, adding an angiotensin II receptor antagonist (ARB) to an ACE inhibitor was associated with more adverse events, without an increase in benefit.358,359

Hence, ARB treatment may be an alternative therapy for patients with SCAD when ACE inhibition is indicated but not tolerated. There
are, however, no clinical outcome studies showing a beneficial effect of ARB in SCAD.

Aldosterone blockade with spironolactone or eplerenone is recommended for use in post-MI patients without significant renal dysfunction or hyperkalaemia, who are already receiving therapeutic doses of an ACE inhibitor and a β-blocker, have an LVEF ≤40% and have either diabetes or heart failure.360

7.3 Other drugs

7.3.1 Analgesics

The use of selective cyclooxygenase-2 (COX-2) inhibitors and traditional non-selective non-steroidal anti-inflammatory drugs (NSAIDs) has been associated with an increased risk for CV events in recent clinical trials in arthritis and cancer prevention and are not recommended.361–363 In patients at increased CV risk in need of pain relief, it is therefore recommended to commence with acetaminophen or aspirin at the lowest efficacious dose, especially for short-term needs.

If adequate pain relief requires the use of NSAIDs, these agents should be used in the lowest effective doses and for the shortest possible duration. In patients with atherosclerotic vascular disease—and in SCAD in particular—NSAID treatment should, when this is indicated for other reasons, be combined with low-dose aspirin to ensure effective platelet inhibition.

7.4 Strategy

Figure 4 summarizes the medical management of SCAD patients. This common strategy might be adjusted according to patient comorbidities, contra-indications, personal preference and drug costs. The medical management consists of a combination of at least a drug for angina relief plus drugs to improve prognosis, as well as use of sublingual nitroglycerin for chest pain management. It is recommended that either a β-blocker or a CCB to a short-acting nitrate be added as first-line treatment to control heart rate and symptoms. If the symptoms are not controlled, it is advised to switch to the other option (CCB or β-blocker) or to combine a β-blocker and a DHP CCB. The combination of a heart-lowering CCB with a β-blocker is not advised. Other anti-anginal drugs might be used as a second-line treatment when symptoms are not satisfactorily controlled. In selected patients with intolerance or contra-indications to both β-blockers and CCBs, second-line drugs can be used as a first-line treatment. The event prevention is optimally achieved by the prescription of antiplatelet agents and statins. In selected patients, the use of ACE inhibitors or ARBs can be considered.

7.5 Treatment of particular forms of SCAD

7.5.1 Microvascular angina

All patients with microvascular angina should achieve optimal coronary risk factor control. Symptomatic treatment is empirical because of the limited knowledge of its causes. Furthermore, the results of available therapeutic trials cannot be accepted as conclusive because of variable patient selection, small sample size, inadequate design and lack of demonstration of clinical improvement of microvascular disease.

Traditional anti-ischaemic drugs are the first step in medical treatment.52 Short-acting nitrates can be used to treat anginal attacks, but often they are only partially effective. β-Blockers seem a rational approach because the dominant symptom is effort-related angina; they were indeed found to improve symptoms in several studies and should constitute the first choice of therapy, particularly in patients

<table>
<thead>
<tr>
<th>Angina relief</th>
<th>Event prevention</th>
</tr>
</thead>
<tbody>
<tr>
<td>1st line</td>
<td></td>
</tr>
<tr>
<td>• Beta-blockers or CCB-heart rate ↓</td>
<td></td>
</tr>
<tr>
<td>• Consider CCB-DHP if low heart rate or intolerance/contraindications</td>
<td></td>
</tr>
<tr>
<td>• Consider Beta-blockers + CCB-DHP if CCS Angina &gt; 2</td>
<td></td>
</tr>
<tr>
<td>May add or switch (1st line for some cases)</td>
<td></td>
</tr>
<tr>
<td>2nd line</td>
<td></td>
</tr>
<tr>
<td>Ivabradine</td>
<td>• Lifestyle management</td>
</tr>
<tr>
<td>Long-acting nitrates</td>
<td>• Control of risk factors</td>
</tr>
<tr>
<td>Nicorandil</td>
<td>+ Educate the patient</td>
</tr>
</tbody>
</table>
| Ranolazine    | • Aspirine
| Trimetazidine | • Statins |
| a              | • Consider ACEI or ARBs |

Figure 4 Medical management of patients with stable coronary artery disease. ACEI = angiotensin converting enzyme inhibitor; CABG = coronary artery bypass graft; CCB = calcium channel blockers; CCS = Canadian Cardiovascular Society; DHP = dihydropyridine; PCI = percutaneous coronary intervention.

aData for diabetics.

bIf intolerance, consider clopidogrel.
with evidence of increased adrenergic activity (e.g., high heart rate at rest or during low-workload exercise).

Calcium antagonists and long-acting nitrates have shown variable results in clinical trials and are more helpful when used in addition to β-blockers in the case of insufficient control of symptoms. Calcium antagonist, however, can be first-line therapy in patients with a significant variable threshold of effort angina. In patients with persisting symptoms despite optimal anti-ischaemic drug therapy, several other treatments have been proposed. ACE inhibitors (and possibly ARBs) may improve microvascular function by counteracting the vasoconstrictor effects of angiotensin II; they have improved symptoms and exercise results in small trials and can be helpful, particularly in patients with hypertension or diabetes mellitus. α-Adrenergic antagonists may decrease sympathetic-mediated vasoconstriction and may be considered in individual patients, although clinical benefits have usually been disappointing. Improvement of exercise capacity has been observed in a small trial with nicorandil. Improvement of anginal symptoms, probably mediated primarily by improvement of endothelial function, has been reported with statins and with oestrogen replacement treatment.

In patients with angina refractory to various combinations of the previous medications, other forms of treatment can be proposed. Xanthine derivatives (aminophylline, babiphenyline) can be added to anti-ischaemic drug treatment to reduce angina by adenosine receptor blockade; adenosine is indeed a major mediator of cardiac ischaemic pain (see Table 29). New anti-ischaemic drugs such as ranolazine or ivabradine have shown good effects in some patients with microvascular angina. Finally, in case of refractory angina, additional interventions may be discussed (see section 9 on refractory angina).

In patients with microvascular angina, the susceptibility of symptoms to medical treatment is extremely variable and experimentation of different drug combinations, is needed before establishing satisfactory symptom control.

### 7.5.2 Treatment of vasospastic angina

All patients with vasospastic angina should achieve optimal coronary risk factor control, in particular through smoking cessation and aspirin. A drug-related cause (e.g., cocaine or amphetamines) should be systemically researched and managed if detected. Chronic preventive treatment of vasospastic angina is mainly based on the use of CCBs. Average doses of these drugs (240–360 mg/day of verapamil or diltiazem, 40–60 mg/day of nifedipine) usually prevent spasm in about 90% of patients. Long-acting nitrates can be added in some patients to improve the efficacy of treatment and should be scheduled to cover the period of the day in which ischaemic episodes most frequently occur, in order to prevent nitrate tolerance. β-Blockers should be avoided, as they might favour spasm by leaving α-mediated vasoconstriction unopposed by β-mediated vasodilation.

In about 10% of cases, coronary artery spasm is refractory to standard vasodilator therapy, although refractoriness is usually limited to brief periods in most patients. Very high doses of calcium antagonists and nitrates usually prevent transient ischaemic episodes in these critical periods. In the very rare patients in whom even this treatment is insufficient, the addition of anti-adrenergic drugs like guanethidine or clonidine might be helpful. PCI with stent implantation at the site of spasm (even in the absence of significant stenosis) as well as chemical or surgical sympathectomy have also been reported but are not recommended. Because of the high prevalence of silent ischaemic episodes and possible arrhythmias, 24-hour ambulatory ECG monitoring can be used to verify the treatment efficiency.

Implantation of an automatic cardioverter defibrillator or of a pacemaker is indicated in patients with ischaemia-related life-threatening tachyarrhythmias or bradyarrhythmias, respectively, when coronary spasm presents a poor or uncertain response to medical therapy.

## 8. Revascularization

### 8.1 Percutaneous coronary intervention

Advances in techniques, equipment, stents and adjuvant therapy have established PCI as a routine and safe procedure in patients with SCAD and suitable coronary anatomy. The mortality risk associated with the procedure in SCAD is <0.5%. The efficacy of PCI in SCAD in comparison to medical therapy and CABG has been the subject of extensive evaluation.

#### 8.1.1 Type of stent and dual antiplatelet therapy

Bare metal stents (BMS) are associated with a 20–30% rate of recurrence of angiographic stenosis within 6–9 months after implantation. Drug-eluting stents (DES) reduce the incidence of angiographic restenosis and ischaemia-driven repeat revascularization. For the first

### Table 29: Treatment in patients with microvascular angina

<table>
<thead>
<tr>
<th>Recommendations</th>
<th>Class</th>
<th>Level</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>It is recommended that all patients receive secondary prevention medications including aspirin and statins.</td>
<td>I</td>
<td>B</td>
<td>371</td>
</tr>
<tr>
<td>β-blockers are recommended as a first line treatment.</td>
<td>I</td>
<td>B</td>
<td>372</td>
</tr>
<tr>
<td>Calcium antagonists are recommended if β-blockers do not achieve sufficient symptomatic benefit or are not tolerated.</td>
<td>I</td>
<td>B</td>
<td>367</td>
</tr>
<tr>
<td>ACE inhibitors or nicorandil may be considered in patients with refractory symptoms.</td>
<td>IIIb</td>
<td>B</td>
<td>368</td>
</tr>
<tr>
<td>Xanthine derivatives or non-pharmacological treatments such as neurostimulatory techniques may be considered in patients with symptoms refractory to the above listed drugs.</td>
<td>IIIb</td>
<td>B</td>
<td>373–375</td>
</tr>
</tbody>
</table>

ACE = angiotensin converting enzyme.

* Class of recommendation.

† Level of evidence.

§ Reference(s) supporting levels of evidence.
The most recent or ‘second- generation’ DES (with thinner struts and biodegradable or more biocompatible polymers) showed superior clinical outcomes for both efficacy and safety when compared with first-generation DES. Second-generation DES—preferring those tested in large all-comers trials and compared with other DES with proven outcome—are therefore the recommended option in SCAD patients with no contra-indication to DAPT (see Table 30).

A recent meta-analysis confirmed that clopidogrel pretreatment in stable patients undergoing elective PCI does not reduce mortality or major adverse cardiac events (MACE), as compared with clopidogrel administration in the catheterization laboratory. On the basis of several randomized trials and this meta-analysis—and contrary to a diffuse common practice—SCAD patients who undergo diagnostic coronary angiography with the possibility of undergoing ad-hoc PCI (revascularization within the same procedure) should not be treated with clopidogrel before the coronary anatomy is known. The bleeding risk of routine DAPT, administered before catheterization in patients who do not require stenting (no significant CAD or CAD requiring CABG surgery), is not balanced by a detectable benefit in terms of ischaemic events in those undergoing PCI. Despite the overwhelming advantages shown in ACS patients—and especially diabetic patients—in the absence of randomized clinical trials, the use of prasugrel or ticagrelor cannot be recommended in SCAD patients undergoing elective PCI. An off-label use of these drugs is, however, common practice in some high-risk patients, especially in cases of documented stent thrombosis. After stenting, premature discontinuation of antiplatelet therapy is a major risk factor for stent thrombosis and should be avoided.

Current guidelines recommended 6–12 months of DAPT after first-generation stents. New-generation DES have been associated with lower rates of stent thrombosis and recent data from registries and randomized controlled trials suggested that a shorter duration of DAPT might be sufficient in stable coronary patients. Considering the risk–benefit ratio of DAPT beyond 6 months—and while waiting for more information from the ongoing studies exploring various durations of treatment including more than a year—we endorsed the current ESC recommendation of 6–12 months of DAPT in SCAD patients undergoing PCI revascularization with a latest-generation DES (see section 9.5 for more details and for recommendations). Shorter durations (1–3 months) are reasonable in patients with high bleeding risk or undergoing undeferrable surgery or on concomitant anticoagulant treatment for which the use of clopidogrel only has shown significant advantages in a single small-scale trial (What is the Optimal antiplatelet and anticoagulant therapy in patients with oral anticoagulation and coronary StenTIng (WOEST)).

### Table 30 Stenting and peri-procedural antiplatelet strategies in stable coronary artery disease patients

<table>
<thead>
<tr>
<th>Recommendations</th>
<th>Class</th>
<th>Level</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>DES is recommended in SCAD patients undergoing stenting if there is no contra-indication to prolonged DAPT.</td>
<td>I</td>
<td>A</td>
<td>172</td>
</tr>
<tr>
<td>Aspirin is recommended for elective stenting.</td>
<td>I</td>
<td>A</td>
<td>172</td>
</tr>
<tr>
<td>Clopidogrel is recommended for elective stenting.</td>
<td>I</td>
<td>B</td>
<td>172</td>
</tr>
<tr>
<td>Prasugrel or ticagrelor should be considered in patients with stent thrombosis on clopidogrel without treatment interruption.</td>
<td>IIa</td>
<td>C</td>
<td>-</td>
</tr>
<tr>
<td>GP IIbIIIa antagonists should be considered for bailout situation only.</td>
<td>IIa</td>
<td>C</td>
<td>172</td>
</tr>
<tr>
<td>Platelet function testing or genetic testing may be considered in specific or high risk situations (e.g. prior history of stent thrombosis; compliance issue; suspicion of resistance; high bleeding risk) if results may change the treatment strategy.</td>
<td>IIb</td>
<td>C</td>
<td>-</td>
</tr>
<tr>
<td>Prasugrel or ticagrelor may be considered in specific high risk situations of elective stenting (e.g. left main stenting; high risk of stent thrombosis; diabetes).</td>
<td>IIb</td>
<td>C</td>
<td>-</td>
</tr>
<tr>
<td>Pretreatment with clopidogrel (when coronary anatomy is not known) is not recommended.</td>
<td>III</td>
<td>A</td>
<td>386, 388, 387</td>
</tr>
<tr>
<td>Routine platelet function testing (clopidogrel and aspirin) to adjust antiplatelet therapy before or after elective stenting is not recommended.</td>
<td>III</td>
<td>A</td>
<td>347, 398</td>
</tr>
<tr>
<td>Prasugrel or ticagrelor is not recommended in low risk elective stenting.</td>
<td>III</td>
<td>C</td>
<td>-</td>
</tr>
</tbody>
</table>

DATP = Dual antiplatelet therapy; SCAD = stable coronary artery disease.

* Class of recommendation.

* Level of evidence.

* Reference(s) supporting levels of evidence.

### 8.1.2 Intracoronary assessment of stenosis severity

- Fractional flow reserve
- Intravascular ultrasound
- Optical coherence tomography

(see web addenda)
infusion is particularly helpful in identify haemodynamically or functionally significant stenosis, inducing ischaemia, justifying revascularization (see Table 31). In patients with FFR > 0.80, studies in the BMS era have demonstrated that medical treatment provides better outcomes than immediate revascularization.110,172,399 Accordingly, a patient with a stenosis and an FFR > 0.80 (two measurements or during adenosine infusion) should not be revascularized. The recent Fractional Flow Reserve vs. Angiography for Multivessel Evaluation (FAME-2) study confirmed that SCAD patients with stenoses having FFR < 0.80 gain a benefit from PCI revascularization in addition to OMT, a benefit driven only by the reduced need for urgent revascularization (the study was stopped prematurely by the DSMB for that reason). Patients without ischaemia have excellent outcomes on medical therapy alone.405 Although the study suffers from significant limitations (the open nature of the trial may have affected the decision on ‘urgent’ revascularization; low-risk population), FFR can guide PCI in a clinically-effective way.

Fractional flow reserve, although in general not useful in very high grade lesions (angiographically > 90%), which practically always have an FFR ≤ 0.8, may help the decision on when to revascularize in many uncertain clinical conditions. One such condition is ‘multivessel disease’, which occurs in a very heterogeneous population. In these patients, FFR measurement may change the strategy of revascularization (PCI vs. CABG) and the extent of revascularization according to the functional assessment of stenoses in critical coronary locations. Another critical situation for revascularization is LM stenosis, a lesion site that is of major prognostic importance and often determines the type of treatment. A recent single-centre study showed that angiography is not always reliable in the determination of the severity of LM stenoses and that deferred revascularization, if FFR > 0.80, may be a safe approach.401

Another situation relevant to these Guidelines, where FFR may be useful, is in ‘post-ACS’ patients. Once the culprit lesion has been treated, the patient can be considered as a stable or stabilized CAD patient. Non-invasive stress testing/imaging immediately after the acute phase may be impossible, contra-indicated or hazardous. Non-culprit stenoses in patients with recent ACS can be evaluated by FFR, either during the index procedure or in a staged procedure.399,402

The use of intravascular ultrasound (IVUS) has been broadly investigated in SCAD with many different subsets of lesions (see Table 31). Unlike FFR, IVUS is an imaging diagnostic tool and does not provide assessment of the functional severity of a stenosis. Previously accepted cut-off limits of 3.5 or 4.0 mm² for major epicardial artery stenosis and 6.0 mm² for left main stenosis403 have been shown to be unreliable and poorly correlated with FFR, with somewhat better results when the absolute IVUS measurements are corrected for the reference vessel size. Once the indication to treatment is established, when more information is needed, IVUS is far superior to FFR because it provides an anatomical characterization of the lesion in terms of vessel size and plaque composition and can control stent expansion and strut apposition. More recently, optical coherence tomography (OCT) has been developed as a new intracoronary imaging tool with superior resolution (<10 μm) able to offer detailed assessment of superficial components including measurements of the thickness of the fibrous cap of lipidic plaques.404 The usefulness of OCT in SCAD patients with vulnerable plaques has not been well established,404 and certainly treatment of non-functionally critical lesions based only on the presence of elements of instability is not recommended. The facilitated technique of image acquisition allows optimization of stent expansion and apposition, and long-term assessment of stent healing.404

### 8.2 Coronary artery bypass surgery

#### 8.2.1 Arterial vs. venous grafts

For the last 25 years the principle technique underpinning CABG has been the use of an internal mammary artery (IMA) to the LAD coronary artery with supplemental vein grafts as required. This followed a seminal publication from the Cleveland Clinic in 1986, showing that an IMA to the LAD improved survival and reduced the subsequent incidence of MI, recurrent angina and the need for repeat revascularization.407

Since then, several angiographic studies have confirmed the superior patency of both IMA grafts in comparison to vein grafts both early and late after CABG.408,409 Most importantly, this superior graft patency appears to translate into a survival benefit. In 2001 a systematic review comparing single (SIMA) and bilateral (BIMA) IMA grafting reported a significant survival benefit with BIMA grafts with a hazard ratio for death of 0.81.410 Recent studies have reported that a survival
The benefit of BIMA grafts extends to the second and third decades of follow-up, and especially in patients with diabetes.

Previous concerns that the use of BIMA grafting may increase early postoperative mortality and/or morbidity have been dispelled by the Arterial Revascularization Trial (ART) which, in one of the largest trials ever conducted in cardiac surgery, randomized 3102 patients to SIMA or BIMA, with supplemental grafts as necessary. Whilst the primary outcome of this trial is 10-year survival, an interim analysis of safety at 1 year showed similar mortality of around 2% in both groups, with no difference in the incidence of MI, death or stroke but a slight increase in the incidence of sternal wound reconstruction in the BIMA group (1.9 vs. 0.6%). The data are currently being analysed to determine the key patient and operating factors that predispose to sternal dehiscence.

The radial artery has also been proposed as a second arterial graft, rather than a second IMA graft. In two randomized trials, the radial artery patency at 1 year was variously reported to be ‘superior’ and ‘equivalent’ to that of vein grafts. In an additional small, randomized trial the 5-year patency of the radial artery was significantly superior to that of vein grafts when placed to the circumflex coronary system.

Nevertheless, despite angiographic and clinical evidence of the potential superiority of arterial grafts, the reality is that the vast majority of bypass grafts—with the exception of the IMA to the LAD—are performed with saphenous vein grafts. Best current evidence suggests that the patency rate of saphenous vein grafts is slightly lower than open techniques.

The patency rate of saphenous vein grafts is slightly lower in off-pump surgery and when harvested using endoscopic, rather than open techniques.

8.2.2 On-pump vs. off-pump surgery (see web addenda)

Off-pump surgery was initially proposed almost three decades ago. Numerous randomized trials and meta-analyses have shown no significant beneficial effect on mortality, but there have been reductions in stroke, transfusion, re-operation for peri-operative bleeding and postoperative complications, possibly at the cost of an excess of repeat revascularization with off-pump CABG. The two largest randomized trials, the Veterans Affairs (VA) Randomized On/Off Bypass (ROOBY) (n = 2203) and The CABG Off or On Pump Revascularization Study (CORONARY) (n = 4752) both reported no difference in the primary composite endpoint at 30 days. ROOBY reported a poorer outcome (death or complication) in the off-pump composite endpoint at one year (9.9 vs. 7.4%) while CORONARY has still to report at the time of writing. In contrast to the randomized trials, several large propensity-matched registries, which generally include higher-risk patients, have reported a reduction in mortality in patients undergoing off-pump CABG, although off-pump surgery is still performed in a minority of centres.

8.3 Revascularization vs. medical therapy

8.3.1 General rules for revascularization (see web addenda)

The decision to revascularize a patient should be based on the presence of significant obstructive coronary artery stenosis, the amount of related ischaemia and the expected benefit to prognosis and/or symptoms (Figure 5). There are many clinical, anatomical, technical and environmental factors that may be discussed before the benefit of revascularization can be anticipated (Table 32, Figure 5). The vast number of possible combinations makes absolute recommendations difficult to mandate in every situation. In this regard, for a given patient in a given hospital, clinical judgement with consensual rather than individual decision-making, with a Heart Team discussion, should prevail, although this has to be individualized since, in many patients, the preferred approach is often quite clear-cut.

When technically feasible, with an acceptable level of risk and a good life expectancy, revascularization is indicated in chronic angina refractory to OMT. It can also be considered as first-line treatment in the situations discussed below.

8.3.1.1 Post-myocardial infarction

The Swiss Interventional Study on Silent Ischemia Type II (SWISSII trial), involving 201 patients with a recent ST-segment elevation MI or non-ST segment elevation MI, investigated whether revascularization with PCI was better than drug therapy in stable patients with silent myocardial ischaemia (see description below).

During a lengthy 10-year follow-up period, which was survival free of cardiac death, non-fatal MI or revascularization, was significantly better in the PCI group.

PCI also significantly reduced the rates of cardiac death and all-cause mortality or MI. In addition, objective evidence of ischaemia was reduced in the revascularization group.

The Danish trial in Acute Myocardial Infarction (DANAMI) compared a deferred invasive strategy of PCI or CABG with a conservative strategy in 503 patients with inducible myocardial ischaemia, who had received thrombolysis for a first MI. Stress testing was performed at discharge and in patients randomized to the invasive strategy, angiography was performed within two weeks of stress testing. Patients with unstable angina were excluded.

Angina plus ischaemia was present in 25%, angina alone in 16% and 57% had silent ischaemia on stress testing. At 2.5 years follow-up, the invasive strategy was associated with a reduction in the incidence of re-infarction and less frequent angina. This was noted in patients with both symptomatic and asymptomatic ischaemia.

In contrast, the Occluded Artery Trial (OAT), following a strategy of routine PCI 3 to 28 days after acute MI, found no discernible benefit in terms of death, re-infarction, or heart failure at 4-year follow-up among asymptomatic or minimally symptomatic patients with occlusion of the infarct-related artery. The findings of the OAT study, however, should not be interpreted as applying to all patients experiencing ST-elevation MI, but just to those with a late occluded artery and no- or minimal angina. Two smaller studies (The Open Artery Trial (TOAT) and Desobstruction Coronario En Post-Infarto (DECOPI)) dealt with similar situations of stable patients after a Q-wave MI, without residual ischaemia and a persistent total occlusion of the infarct-related artery, and these studies did not show any clinical benefit of stenting over medical therapy.

Post-lytic revascularization studies. Old trials (not discussed here) comparing an invasive- with a conservative approach after fibrinolytic therapy did not show any differences in patient outcomes, but these studies antedated the use of stenting and modern antiplatelet therapies. In contrast, more recent randomized studies, comparing systematic early PCI with a conservative ischaemia guided strategy, have demonstrated favourable trends with early PCI and a significant reduction of death or MI in a meta-analysis.
8.3.1.2 Left ventricular dysfunction
In general, revascularization improves survival in ‘sicker’ patients, especially in the presence of LV dysfunction. From the early days of coronary angiography, it has been well recognized that LV dysfunction is one of the most powerful indicators of an adverse prognosis. As techniques of revascularization improved, LV dysfunction has become a prime target—as opposed to a contra-indication—for coronary revascularization. Several older studies, including a meta-analysis, suggested that survival was improved by CABG over medical therapy in patients with mild-to-moderate systolic dysfunction. The CASS randomized trial of bypass surgery vs. medical therapy demonstrated no overall differences in survival, except in the subset of patients with an ejection fraction (EF) of 0.35–0.49, in association with triple vessel disease. The more contemporary Surgical Treatment for Ischemic Heart Failure (STICH) trial of subjects with more severe impairment of LV function (EF <0.35) demonstrated no survival difference at 5 years between CABG and OMT, although CV mortality was reduced, as were hospitalization rates for major CV causes, in the CABG group. Moreover, if the data are analysed by treatment received and as per protocol, due to the large number of crossovers to both therapies, the differences in all-cause mortality reached statistical significance in favour of the CABG group; in this respect the trial can be considered to have demonstrated a modestly positive result in favour of surgery, with potentially important clinical implications. The subset analysed by viability testing is inconclusive.

8.3.1.3 Multivessel disease and/or large ischaemic territory
Observational studies from the CASS registry and the meta-analysis of seven randomized trials—comprising a total of 2649 patients—of CABG vs. medical therapy suggested a survival advantage of surgery in patients with three-vessel disease (or LM disease), but no difference in patients with one- or two-vessel disease, except in patients with involvement of the proximal LAD plus one other major coronary artery. In addition, these studies demonstrated a greater efficacy from CABG over medical therapy in respect of symptom relief, bearing in mind the caveat that, in these trials, the methods of medical therapy and secondary prevention were obsolete by today’s standards. The more contemporary Medical, Angioplasty, or Surgery Study (MASS II) trial of CABG, PCI and medical therapy, patients treated with CABG enjoyed a better survival and lower rates of subsequent MI and need for additional revascularization procedures over a follow-up period of 10 years. The importance of the severity of symptoms was emphasized by two studies from the CASS registry, which demonstrated that, in patients with mild angina pectoris and triple vessel disease, survival advantage was confined to those with mild-to-moderate LV dysfunction. On the other hand, among patients with severe symptoms, survival was

Figure 5 Global strategy of intervention in stable coronary artery disease (SCAD) patients with demonstrated ischaemia. CABG = coronary artery bypass graft; CAD = coronary artery disease; LAD = left anterior descending; LV = left ventricular; OMT = optimal medical treatment; PCI = percutaneous coronary intervention.

Indication of revascularization for prognosis or symptoms (see Table 32).
Not suitable for revascularization due to anatomy or clinical conditions.
See section 9.
improved irrespective of LV function. In addition, the greater the number of proximal stenosis, the greater the surgical benefit.460,460 Observational studies also support a survival advantage for CABG in patients with double-vessel disease in the presence of severe or extensive ischaemia or severe angina.197,461 – 464 The concept of a revascularization benefit in patients with extended ischaemia is currently being tested in the ongoing International Study of Comparative Health Effectiveness with Medical and Invasive Approaches (ISCHEMIA).197,214,465

8.3.1.4 Left main coronary artery disease
The survival advantages for bypass surgery in patients with 50% or greater stenosis of the LM coronary artery was established by the striking differences noted in the Veterans Administration Cooperative Study in a subgroup of 113 patients,466,467 and confirmed in a subsequent meta-analysis,368 and in studies from the CASS registry.69,470 The data now need to be re-interpreted in the light of more recent data evaluating the functional severity of LM stenoses and the possibility that revascularization can be safely deferred if FFR > 0.80.461

Irrespective, LM CAD (stenosis 50% or greater) continues to be a Class 1 indication for revascularization.172,471 No further randomized, controlled trials of bypass surgery or PCI vs. medical therapy are likely to be undertaken in patients with LM CAD.

### Table 32  Indications for revascularization of stable coronary artery disease patients on optimal medical therapy (adapted from ESC/EACTS 2010 Guidelines)172

<table>
<thead>
<tr>
<th>Indication a</th>
<th>To improve prognosis:</th>
<th>To improve symptoms persistent on OMT:</th>
</tr>
</thead>
<tbody>
<tr>
<td>A Heart Team approach to revascularization is recommended in patients with unprotected left main, 2–3 vessel disease, diabetes or comorbidities.</td>
<td>Class d Level e</td>
<td>Class d Level e</td>
</tr>
<tr>
<td>Left main &gt;50% diameter stenosis.</td>
<td>I C</td>
<td>I C</td>
</tr>
<tr>
<td>Any proximal LAD &gt;50% diameter stenosis.</td>
<td>I A</td>
<td>I A</td>
</tr>
<tr>
<td>2–3 vessel disease with impaired LV function / CHF</td>
<td>I B</td>
<td>IIa B</td>
</tr>
<tr>
<td>Single remaining vessel (&gt;50% diameter stenosis).</td>
<td>I C</td>
<td>I A</td>
</tr>
<tr>
<td>Proven large area of ischaemia (&gt;10% LV)</td>
<td>I B</td>
<td>I B</td>
</tr>
<tr>
<td>Any significant stenosis with limiting symptoms or symptoms non responsive/intolerant to OMT.</td>
<td>NA NA</td>
<td>I A</td>
</tr>
<tr>
<td>Dyspnoea/ cardiac heart failure with &gt;10% ischaemia/viability supplied by stenosis &gt;50%.</td>
<td>IIb B</td>
<td>469, 470</td>
</tr>
<tr>
<td>No limiting symptoms with OMT in vessel other than left main or proximal LAD or single remaining vessel or vessel subtending area of ischaemia &lt;10% of myocardium or with FFR ≥ 0.80.</td>
<td>III A</td>
<td>III C</td>
</tr>
</tbody>
</table>

References attached to these recommendations can be found in Table 8 of the original ESC guidelines for myocardial revascularization.172

CCS = Canadian Cardiovascular Society; CHF: congestive heart failure; FFR = fractional flow reserve; LAD = left anterior descending; LV = left ventricle; NA: not available; OMT = optimal medical treatment; SCAD = stable coronary artery disease.

a In asymptomatic patients, the decision will be guided by the extent of ischaemia on stress testing.

b With documented ischaemia or FFR < 0.80 for angiographic diameter stenoses 50–90%.

c As assessed by non-invasive test (SPECT, MRI, stress echocardiography).

d Class of recommendation.

e Level of evidence.

f Reference(s) supporting levels of evidence.

### 8.3.2 Revascularization in lower-risk populations

#### 8.3.2.1 The randomized studies (see web addenda)
The older randomized studies that investigated revascularization vs. OMT are selectively reviewed in the web addenda.26,41,461,472,473,459,474 – 477 The three most recent studies are also the largest and most informative studies for this comparison of revascularization with OMT.

The Clinical Outcomes Utilizing Revascularization and Aggressive Drug Evaluation (COURAGE) trial (n = 2287) compared PCI + OMT with OMT only, in patients with SCAD or ischaemia and coronary lesions suitable for PCI. The target study population for the COURAGE trial were patients with chronic angina pectoris Canadian Cardiovascular Society (CCS) Class I–III, stable post-MI patients and asymptomatic patients with objective evidence of myocardial ischaemia. All patients had angiographically defined CAD, with at least one vessel meeting AHA/American College of Cardiology (ACC) Class I or II indications for PCI. Patients with a prior CABG were accepted. Patients with stenosis >80% in one or more vessels, subtending a large area of myocardium, could be enrolled even in the absence of objective ischaemia. The primary end-point of all-cause death or non-fatal MI did not differ between the two groups during a mean follow-up of 4.6 years.23,478 However, in patients who were invasively treated, freedom from angina was significantly better up to 3 years of follow-up. In a sub-study, patients with >10% ischaemia on stress myocardial perfusion scintigraphy...
had a higher rate of death or MI. More PCI + OMT patients exhibited significant ischaemia reduction (33 vs. 19%; \( P = 0.0004 \)). Patients with ischaemia reduction had lower unadjusted risk for death or MI, particularly if baseline ischaemia was moderate to severe.214

The Clinical Outcomes Utilizing Revascularization and Aggressive Drug Evaluation (BARI-2D) trial (\( n = 2368 \)) evaluated whether PCI or CABG (choice left to the discretion of the treating physician), combined with OMT, would be better than OMT alone in patients with SCAD and type 2 diabetes mellitus.25 The target study population were patients with a diagnosis of type 2 diabetes and angiographically documented CAD for which revascularization was not required for prompt control of severe or unstable angina. Patients with stenosis >70% presenting with angina symptoms were eligible for randomization, even without documented ischaemia. In contrast, approximately 30% of patients were asymptomatic with a positive stress test. The primary endpoint of all-cause mortality at 5 years follow-up did not differ between the two treatment strategies, nor did rates of MI or stroke. The patients with most severe disease were selected for CABG rather than PCI and were a higher-risk group that drew a greater benefit from early revascularization (reduction of MI compared with OMT).

In the Fractional Flow Reserve vs. Angiography for Multivessel Evaluation (FAME-2) trial, 888 SCAD patients with functionally significant stenosis (\( \text{FFR} \leq 0.80 \)) were randomly assigned to FFR-guided PCI plus OMT, or to OMT alone.400 The target study population were patients who had at least one functionally significant stenosis and, on average, large areas of ischaemic myocardium (mean FFR value of 0.68) while the low-risk patients with non-ischaemic FFR values were not randomized but followed in a separate registry. The study was stopped prematurely by the Data Safety Monitoring Board, due to a highly significant reduction in hospital re-admission and urgent revascularization in the FFR \(< 0.80\)-PCI group, compared with the FFR \(< 0.80\)-OMT group. There was no difference in rates of death or MI between the two strategies. In patients without ischaemia (registry), the outcome appeared to be favourable with OMT only.

Altogether, seven major (\( n \) of 200 or more) randomized trials of revascularization vs. medical therapy in chronic SCAD have been published over the past 10 years (Table 33). Typically, populations of these studies were selected after an angiogram, had demonstrated at least one significant stenosis of an epicardial coronary artery in patients with typical or suspected angina—with or without documented myocardial ischaemia—with, in general, good LV function, no comorbidities and excluding patients at high angiographic risk, patients with LM coronary disease, CABG, multivessel disease, or lesions deemed to be treated with revascularization without further discussion for OMT only.

The results of these studies comparing myocardial revascularization with OMT have been somewhat consistent in confirming that, except for better symptom relief and lesser frequency of urgent revascularization, there is no advantage of revascularization over OMT alone to reduce mortality in angiographically selected patients.

### Table 33 Characteristics of the seven most recent randomized trials

<table>
<thead>
<tr>
<th>Study</th>
<th>Recruitment (years)</th>
<th>Study size (n)</th>
<th>Mean age (years)</th>
<th>Angina CCS</th>
<th>Stress ischaemia (% of patients)</th>
<th>Prior MI (% of patients)</th>
<th>Mean LVEF (%)</th>
<th>Angiographic selection</th>
<th>Mandatory documented ischaemia</th>
<th>Revascularization</th>
<th>Primary Endpoint (PEP)</th>
<th>Revascularization better on PEP</th>
</tr>
</thead>
<tbody>
<tr>
<td>TIME475</td>
<td>1996–2000</td>
<td>301</td>
<td>80</td>
<td>II–IV</td>
<td>69</td>
<td>47</td>
<td>52</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>MASS II479</td>
<td>1995–2000</td>
<td>611</td>
<td>60</td>
<td>II–III</td>
<td>NA</td>
<td>44</td>
<td>67</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SWISSI II481</td>
<td>1991–97</td>
<td>201</td>
<td>55</td>
<td>0</td>
<td>NA</td>
<td>100</td>
<td>57</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>COURAGE22</td>
<td>1999–2004</td>
<td>2287</td>
<td>61</td>
<td>0–III</td>
<td>NA</td>
<td>39</td>
<td>62</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>BARI–2D25</td>
<td>2001–2005</td>
<td>2368</td>
<td>62</td>
<td>0–II</td>
<td>NA</td>
<td>38</td>
<td>NA</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>JSAP477</td>
<td>2002–2004</td>
<td>384</td>
<td>11–II</td>
<td>NA</td>
<td>NA</td>
<td>15</td>
<td>38</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FAME–2400</td>
<td>2010–2012</td>
<td>888</td>
<td>11–IV</td>
<td>NA</td>
<td>NA</td>
<td>37</td>
<td>65</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Notes:**
- CABG = coronary artery bypass graft; CCS = Canadian Cardiovascular Society; LVEF = left ventricular ejection fraction; MI = myocardial infarction; PCI = percutaneous coronary intervention.
- ESC Guidelines 2990

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presenting with SCAD, acknowledging the possibility of crossover from medical therapy to intervention during follow-up. Although interventional and surgical techniques have improved in the past two decades, medical therapy has also improved over the same period.

As a result, OMT may substantially improve long-term outcomes of patients treated conservatively but also of patients undergoing revascularization, reducing the impact of revascularization itself on survival in non-ACS patients.

8.3.2.2 Limitations of the randomized studies (see web addenda)
All of these studies have their limitations, which may limit their general applicability. These limitations are discussed in the additional web addenda. In brief:

- Some patient subsets that are commonly encountered in clinical practice were under-represented and the amount of evidence may appear insufficient, or even contradictory to other studies.
- Patients were considered for inclusion only after coronary angiography was performed: therefore conclusions from these trials cannot be extended to patients with unknown coronary anatomy.
- Crossover rates from OMT to revascularization were high and much higher than initially expected, suggesting that revascularization was merely deferred in 33–42% of patients randomized to a conservative approach.
- Documented ischaemia was not mandatory for enrolment in COURAGE or in BARI 2D. Many patients with severe ischaemia—and, as such, considered to be at higher risk—were not randomized in the study.
- The rapid evolution of revascularization techniques (e.g. DES for PCI and arterial grafts for CABG) and antiplatelet, anticoagulant, lipid-lowering and anti-ischaemic drugs render many of the studies obsolete by contemporary standards or difficult to interpret (e.g. stents used were mostly BMS).
- OMT was particularly well carried out in these trials (not reflecting current practice), which emphasizes the need to educate physicians in clinical practice about the necessity and the scope of OMT.
- COURAGE and BARI 2D failed to meet the statistical endpoint of superiority and, as such, were neutral trials demonstrating that an initial approach of intervention was neutral upon death or MI.
- Several meta-analyses of randomized studies have shown divergent results on hard outcomes, as have registries with propensity analyses.

8.3.2.3 Overall interpretation
In low-risk, stable CAD patients, after an ischaemic documentation and a careful clinical and angiographic selection, the strategy of initial OMT is safe and should be the default approach. When a period of OMT has not been adequately conducted, cardiologists and surgeons should be more conservative when making a decision over revascularization, especially in case of high-risk comorbidities, difficult anatomies, mildly symptomatic patients or in patients without extensive provokable ischaemia. The trials have shown that, despite frequent crossovers to revascularization, the majority of patients remain on OMT alone for the duration of the trial.

When initial OMT has failed and patients remain symptomatic, or when the ischaemic risk appears important, the various options need to be discussed (OMT reinforcement or revascularization). The advantages, limitations and advice from the Heart Team must be fully presented in the discussion with the patient.

The early hazards of revascularization are well known: early procedural MI, stent thrombosis or late restenosis (much reduced now by second-generation DES) after PCI, peri-operative MI, stroke, cognitive dysfunction, surgical wound infection, prolonged hospital stay and rehabilitation after CABG. Potential advantages of an initial revascularization strategy (PCI or CABG) include better relief of symptoms and no significant excess of mortality, fewer drugs, fewer hospital visits and less revascularization within the first year with globally improved QoL. The advantage of revascularization over OMT on symptom relief is, however, blunted over time. OMT is safer in the short term, and as safe as revascularization for mortality up to 5 years in patients meeting the low-risk inclusion criteria of these trials. However, OMT requires larger doses and numbers of medications that may have a direct impact on adherence to treatment, side-effects, drug interactions, QoL and long-term cost to the patient and third party payers.

8.3.2.4 Ongoing studies for management of stable coronary artery disease patients with documented ischaemia
Several studies have suggested that patients with more extensive ischaemia benefit from revascularization—and that this benefit could translate into a long-term survival benefit if ischaemia is severe and reduction of ischaemia is significant. This hypothesis has been poorly investigated prospectively, although the positive randomized trials ‘Asymptomatic Cardiac Ischaemia Pilot’ (ACIP) and SWISSI II—both with subsets analyses of the CABG population in BARI 2D, plus the results of PCI in FAME 2—strongly suggest that ischaemia plays a key role in the benefit afforded by revascularization.

The hypothesis of deciding upon an invasive approach prior to angiography and not after (as in COURAGE and BARI 2D)—on the basis of documented clinically meaningful ischaemia during stress testing or haemodynamic assessment of stenoses identified at the time of angiography—certainly needs re-evaluation. This hypothesis is currently being evaluated in randomized trials: in the ongoing ISCHEMIA trial, patients are randomized—before coronary angiography—to a conservative OMT strategy or to an invasive strategy when they have documented myocardial ischaemia, the primary endpoint being death or MI.

8.4 Percutaneous coronary intervention vs. coronary artery bypass graft (see web addenda)

8.4.1 Recent data and recommendations
The relative indications for PCI and CABG in SCAD patients have been clearly defined by the recent recommendations. There has been an increasing recognition of the value of the Heart Team in reaching consensus over if, when, and how to revascularize patients. Figures 6 and 7 show suggested algorithms to help simplify the decision-making process and to possibly avoid the need for systematic discussion of every patient with locally agreed protocols (refer to specific ESC Guidelines on myocardial revascularization for class and LOE concerning the respective indications of PCI and CABG). The Guidelines emphasize the importance of
OMT in all patients and for both procedures, and the pivotal role of the heart team in most decisions over revascularization in patients with multivessel or left main disease. This is particularly true for patients with three-vessel disease when they present with a syntax score ≥ 22 or when complete revascularization is not achievable by one technique of revascularization, or when they have diabetes. For these patients, CABG should most often be the preferred option.

In SYNergy between percutaneous coronary intervention with TAXus and cardiac surgery (SYNTAX), 1800 patients with three-vessel or left main coronary artery disease were randomized to undergo CABG or PCI. Publication of the 5-year results of the SYNTAX trial have confirmed the initial findings, with a higher rate of major adverse cardiac or cerebrovascular events at 12 months in the PCI group, in large part because of an increased rate of repeat revascularization. At 5 years, all-cause death was 13.9% with PCI, against 11.4% with CABG (P = 0.10) and cardiac death was 9.0 vs. 5.3% (P = 0.003) in favour of CABG. MACE was significantly reduced with CABG also.

Interestingly, this benefit was driven by the upper two tertiles of the SYNTAX score; although PCI and CABG performed as well on all endpoints for SYNTAX scores of 22 or less, there was a clear benefit with CABG at 5 years, especially in patients with scores of 33 or more. In patients with intermediate or high SYNTAX scores, MACE was significantly increased with PCI (intermediate score, 25.8% of the CABG group vs. 36.0% of the PCI group; P = 0.008; high score, 26.8 vs. 44.0%; P < 0.0001).

These findings are consistent with the survival benefit of CABG reported in several large propensity-matched registries comparing outcome of PCI and CABG. Indeed, in a recently published study of 7235 pairs of patients, matched for numerous baseline characteristics, the overall 8-year survival rates were 78.0% for CABG and 71.2% for stenting (HR 0.68; 95% CI 0.64–0.74; P < 0.001). For anatomic groups, the HRs ranged from 0.53 (P < 0.001) for patients with three-vessel disease involving the proximal LAD to 0.78 (P = 0.05) for patients with two-vessel disease but no disease in the LAD artery. A lower risk of death after CABG was observed in all subgroups stratified by a number of baseline risk factors. Most recently, the Asymptomatic atrial fibrillation and Stroke Evaluation in pacemaker patients and the atrial fibrillation Reduction atrial pacing Trial (ASSERT) reported survival in 86 244 CABG and 103 549 PCI propensity-matched patients with two- or three-vessel CAD. At 4-year follow-up there was increased mortality with PCI, compared with CABG. Despite statistical adjustment, this huge registry cannot eliminate confounding variables and the fact that sicker patients may have been assigned to PCI.

In SYNTAX, the results for 705 patients with LMS disease differ from the remaining patients with three-vessel CAD. For these patients, there was no overall difference between CABG and PCI in terms of death (8.4% CABG vs. 7.3% PCI; P = 0.64) or MI (4.1 vs. 6.9%; P = 0.14) but a higher incidence of stroke with CABG (4 vs. 1.2%; P = 0.02). The advantage of CABG was reduced repeat revascularization at 12% vs. 20% for stents (P = 0.004).
‘Premier of Randomized Comparison of Bypass Surgery vs. Angioplasty Using Sirolimus-Eluting Stent in Patients with Left Main Coronary Artery Disease’ (PRECOMBAT), another randomized trial of 600 patients with left main stem (LMS) disease, reported a composite endpoint of death, cerebrovascular accident and MI as 4.7% for CABG and 4.4% for PCI. Furthermore, the incidence of stroke was substantially lower than in SYNTAX and similar for PCI (0.4%) and CABG (0.7%). It should be acknowledged that Left Main SYNTAX was a subgroup analysis and PRECOMBAT was not powered to detect a difference in hard clinical endpoint. Accordingly, further large randomized controlled trials are needed to establish the optimal mode of LM revascularization with this degree of complexity [e.g. the Evaluation of XIENCE PRIME or XIENCE V vs. Coronary Artery Bypass Surgery for Effectiveness of Left Main Revascularization (EXCEL)].

Meanwhile, angiographic characteristics of the LM disease are key in selection between PCI and CABG (calcifications, ostial/mid/distal, LM size, distal lesions, etc.) and, for at least lower severity of LMS disease, PCI produces at least equivalent—if not superior—outcomes to CABG.

The Design of the Future Revascularization Evaluation in patients with Diabetes mellitus: Optimal management of Multivessel disease (FREEDOM) trial (see chapter 9.2 on diabetes for details)—which demonstrated a significant reduction on the primary ischaemic outcome at 5 years in patients treated with CABG vs. PCI—taking into consideration previous trials, suggests that there is a significant mortality benefit from bypass surgery vs. PCI in diabetic patients with multivessel disease when both options of revascularization are technically feasible, albeit at the price of an increased risk of non-fatal stroke.

The role of hybrid coronary revascularization (Figure 5) combining Left Internal Mammary Artery (LIMA)-to-LAD artery grafting and PCI of at least one non-LAD artery is evolving and is an option in patients with multivessel disease and technical issues over one of the two techniques of revascularization, comorbidities, prior history of revascularization with limitations in graft conduits or access for PCI (e.g. occlusion). It allows complete revascularization with the advantages of both modes of revascularization. No randomized study, and only small series of patients have so far been published, precluding any firm recommendation.

**8.4.2 Target populations of the randomized studies (see web addenda)**

Over the last two decades there have been approximately 20 trials of PCI vs. CABG, which have consistently reported no overall difference in survival between the two interventional techniques, possibly related to the low risk of the populations studied.

In contrast, several propensity-matched registries have consistently demonstrated a survival benefit of CABG after intervention, accompanied by a marked reduction in the need for repeat intervention, although it may still be susceptible to confounding factors.

**8.5 Scores and decisions (see web addenda)**

**8.5.1 Scores (see web addenda)**

Interventional and surgical scores have been developed to evaluate the risk of the different revascularization strategies.
we lack prospective validation of these scores in randomized studies comparing CABG to PCI, the recommendations—and now the practice—are heavily based on these scores as tools for decision-making in individual patients.

8.5.2 Appropriate utilization of revascularization (see web addenda)
Appropriateness criteria are based upon expert consensus as to when a procedure is appropriate. This is, however, an important and complex area of concern as the cost of imaging and revascularization comes under increasing but appropriate scrutiny.

9. Special groups or considerations

9.1 Women (see web addenda)
Coronary artery disease develops 5–10 years later in women than in men. Recent studies indicate that the decline in mortality from CAD does not extend to younger women, in whom it has remained constant. CVD guidelines in general are based on research conducted primarily in men, the mean percentage of women enrolled in clinical trials since 2006 being 30%. CVD risk factors in women and men are the same, although their distribution differs over time and between regions. Stable angina is the most common initial presentation of CAD in women. There is a widespread understanding that women with CAD present with symptoms that are different from those in men. Some of this is due to women presenting at older ages and symptoms becoming less specific with advancing age. Several studies have indicated gender-related differences in the care of both acute and chronic CAD, in part related to differences in presentation and pathophysiology. Compared with men, women have higher rates of procedural complication, including mortality, stroke and vascular complications. Women also have higher complication rates following CABG but, although the numbers of women included in trials are limited, results do not indicate gender-related differences in outcome. Nonetheless, it may be prudent to adopt a more conservative approach in undertaking PCI and CABG in women.

Probably the most important difference between CAD in men and women is that women, presenting with MI and angina twice as often as men, have no significant obstructive CAD. (see section 6.7.1 on microvascular angina). However, the notion that these women have ‘normal’ coronary arteries should be reconsidered in light of the IVUS sub-study within the WOmen’s Ischemia Syndrome Evaluation (WISE) showing that, among a sample of 100 such women, ~80% had definite coronary atherosclerosis that was concealed by positive remodelling. Until sufficient trial-based evidence is available, women with chest pain and no obstructive coronary disease should be screened for CVD risk factors and treated according to risk stratification, as described in CVD prevention Guidelines, supplemented by individualized symptomatic treatment for angina (see sections 7.5.1 and 7.5.2 on treatment of microvascular and vasospastic angina). At present HRT is not recommended for primary or secondary prevention of CVD.

9.2 Patients with diabetes (see web addenda)
Mortality due to CVD is increased three-fold in diabetic men and two- to five-fold in diabetic women, compared with age-and sex-matched non-diabetic persons. A target HbA1c <7% (<53 mmol/mol) and target blood pressure <140/85 mmHg are recommended in recent European Guidelines on CVD prevention. The high prevalence of significant CAD and prohibitively high cardiovascular mortality may suggest the usefulness of routine screening extended to asymptomatic patients. In the absence of outcome trials confirming a clinical benefit, this strategy is not recommended. Coronary artery revascularization of diabetes remains a challenge. The decision to use either PCI or CABG as preferred mode of revascularization should be based on anatomical factors, together with clinical factors and other logistical or local factors (see chapter 8 and Figure 6). As a rule, PCI is recommended in diabetic patient with single-vessel disease. Conversely, given the results of the FREEDOM trial, CABG is recommended in diabetic patients with multivessel disease after discussion in a Heart Team meeting.

9.3 Patients with chronic kidney disease (see web addenda)
Chronic kidney disease is a risk factor for— and strongly associated with—CAD and has a major impact on outcomes and therapeutic decisions. The use of drugs and iodinated contrast agents is exposes patients to more complications. This is also a group of patients poorly explored in clinical trials, with limited strong evidence based medicine.

9.4 Elderly patients (see web addenda)
This population is specific in many ways:

(1) Higher prevalence of comorbidities.
(2) Population is usually undertreated and under-represented in clinical trials.
(3) Difficult diagnosis due to atypical symptoms and difficulties in performing stress testing.
(4) Patients are more often referred to PCI than CABG but age should not be the sole criterion for the choice of type of revascularization.
(5) Higher risk of complications during and after coronary revascularization.

9.5 The patient after revascularization (see web addenda)
Therapy and secondary prevention should be initiated during hospitalization, when patients are highly motivated. Follow-up strategies should focus on the assessment of the patient’s symptoms, functional status and secondary prevention, and not only on the detection of stenosis or graft occlusion. Recommendations are given below in Table 34.
9.6 Repeat revascularization of the patient with prior coronary artery bypass graft revascularization (see web addenda)

Repeat revascularization in the patient who has undergone prior CABG poses a clinical challenge. Considerations in determining the preferred modality of revascularization include the age of patients, co-morbidities and diffuseness of coronary disease, as well as the potential for damage to patent grafts, intraluminal embolization in saphenous vein grafts, lack of suitable arterial and venous conduits and instability of a graft-independent circulation. PCI may be preferred in patients with discrete lesions in grafts and preserved LV function or accessible native vessel disease. Repeat bypass surgery may be preferred when the vessels are unsuitable for PCI and when there are good distal vessel targets for bypass graft placement.

The use of distal embolic protection devices is recommended in saphenous vein graft interventions. Any revascularization strategy needs to be accompanied by optimizing medical therapy with anti-anginal drugs and risk factor reduction.

9.7 Chronic total occlusions (see web addenda)

Chronic total occlusions (CTO) are identified in 15–30% of all patients referred for coronary angiography. A worse prognosis has been attached to chronic total occlusions. Revascularization needs to be discussed in patients with symptoms of occlusion or large ischaemic areas. Percutaneous coronary intervention (PCI) of CTOs is technically challenging and requires familiarity with advanced techniques and specialized equipment. Surgical treatment, with the implantation of a distal bypass graft, is also a valid option for discussion.

9.8 Refractory angina (see web addenda)

The term ‘refractory angina’ is defined as “a chronic condition caused by clinically established reversible myocardial ischaemia in the presence of CAD, which cannot be adequately controlled by a combination of medical therapy, angioplasty or coronary artery bypass graft”. For this patient group, a number of treatment options has
9.9 Primary care (see web addenda)

Primary care physicians have an important role in the identification and management of patients with SCAD. In particular:

- identifying those patients presenting with symptoms of possible SCAD that requires further evaluation and investigation
- identifying those at increased risk of developing SCAD and ensuring that modifiable risk factors are actively managed, with lifestyle and therapeutic interventions, in order to reduce their future risk
- ensuring that those with SCAD are aware of the benefits, both in respect of symptom control and prognosis, of optimal medical therapy and, in appropriate cases, the benefits of percutaneous intervention or surgery
- establishing a systematic approach to the follow-up of patients with SCAD, at appropriate intervals, for the primary care physician to re-appraise the patient’s clinical symptoms, medication and risk factors.

9.10 Gaps in evidence (see web addenda)

These guidelines suffer from limitations inherent in the evidence available, uncertainties on the best imaging modalities, on what is the best modern pharmacologic approach and on what is the real benefit from myocardial revascularization.

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**Table 35** Treatment options in refractory angina

<table>
<thead>
<tr>
<th>Recommendations</th>
<th>Class a</th>
<th>Level b</th>
<th>Ref. c</th>
</tr>
</thead>
<tbody>
<tr>
<td>EECP should be considered for symptom relief in patients with invalidating angina refractory to optimal medical and revascularization strategies.</td>
<td>IIa</td>
<td>B</td>
<td>509, 510</td>
</tr>
<tr>
<td>TENS may be considered to ameliorate symptoms of invalidating angina refractory to optimal medical and revascularization strategies.</td>
<td>IIb</td>
<td>C</td>
<td>-</td>
</tr>
<tr>
<td>SCS may be considered to ameliorate symptoms and quality of life in patients with invalidating angina refractory to optimal medical and revascularization strategies.</td>
<td>IIb</td>
<td>B</td>
<td>511</td>
</tr>
<tr>
<td>TMR is not indicated in patients with invalidating angina refractory to optimal medical and revascularization strategies.</td>
<td>III</td>
<td>A</td>
<td>514</td>
</tr>
</tbody>
</table>

EECP = enhanced external counterpulsation; TENS = transcutaneous electrical nerve stimulation; TMR = transmyocardial revascularization; SC = spinal cord stimulation.

a Class of recommendation.

b Level of evidence.
c Reference(s) supporting levels of evidence.

Among non-pharmacological treatments, enhanced external counterpulsation therapy and neurostimulatory techniques have shown that they can ameliorate symptoms and improve quality of life, although convincing evidence regarding reduction in both ischaemia burden and mortality is still lacking. Conversely, transmyocardial or percutaneous myocardial revascularization have been abandoned because they are ineffective.


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