Guidelines for the management of acute coronary syndromes 2006

Heart Foundation
National Heart Foundation of Australia

The Cardiac Society of Australia and New Zealand
Guidelines for the management of acute coronary syndromes 2006

These guidelines were developed by means of a consensus approach which involved an independent assessment of key Australian and international evidence-based clinical guidelines, scientific articles and trial data, which are incomplete in some areas.

Recommendations are not necessarily congruent with current Pharmaceutical Benefits Scheme criteria for eligibility for subsidy in all areas.

The guidelines provide a general framework for appropriate practice, to be followed subject to the practitioner's judgement in each individual case. All treatments should be individualised according to the patient's comorbidities, drug tolerance, lifestyle and living circumstances, and wishes.

For all medications, observe usual contraindications, be mindful of the potential for significant and possibly adverse drug interactions and allergies, and monitor and review patients carefully and regularly.

Where drug therapy is recommended for indefinite use, these recommendations have been based on the extrapolated findings of clinical trials which are by their nature of limited duration.

These guidelines were published in April 2006 and are based on the literature up to September 2005. Please check Heartsite regularly for updates and amendments: <http://www.heartfoundation.com.au>.
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GUIDELINES FOR THE MANAGEMENT OF ACUTE CORONARY SYNDROMES 2006

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Summary of key recommendations

Systems of care for patients with acute coronary syndromes
- Effective systems of care are required to deliver optimal care for patients with acute coronary syndromes (ACS), particularly in rural and remote areas.
- Systems of care should be regionally based, and have formal links with specialist centres for consultation and acute interhospital transfer.
- Systems should include appropriate monitoring, feedback and quality improvement components.
- Clinical decisions about care and transfer should take into account patients’ cultural and personal beliefs and wishes.

New acute coronary syndrome terminology and implications for diagnosis
- It is important to establish an initial working diagnosis to guide clinical decision making.
- New definitions of myocardial infarction, based heavily on the presence of cardiac biomarkers, have implications for coding and epidemiological studies. However, clinically they do not influence the indications for ongoing prevention therapies.
- Use of the ACS Dataset (part of the National Health Data Dictionary) can facilitate the collection of data relating to the presentation and management of ACS that can be compared and collated within and between health care providers.

Acute management of chest pain
- People experiencing symptoms of an ACS should seek help promptly and activate emergency medical services.
- The most important initial need is access to a defibrillator to avoid early cardiac death resulting from reversible arrhythmias.
- Aspirin should be given early (ie, by emergency or ambulance personnel) unless already taken or contraindicated.
- Oxygen should be given, as well as glyceryl trinitrate and intravenous morphine as required.
- As a minimum, medical facilities receiving patients should be given warning of incoming patients in whom there is a high suspicion of an ACS — particularly ST-segment-elevation myocardial infarction (STEMI) — or whose condition is unstable.
- Where appropriate, a 12-lead electrocardiogram (ECG) should be taken en route and transmitted to a medical facility.
- Where formal protocols are in place, prehospital treatment (including fibrinolysis in appropriate cases) should be facilitated.

Investigations
- The ECG is the sole test required to select patients for emergency reperfusion (fibrinolytic therapy or direct percutaneous coronary intervention [PCI]).
- Patients with STEMI who present within 12 hours of the onset of ischaemic symptoms should have a reperfusion strategy implemented promptly.
- Patients with a suspected ACS without ST-segment elevation on ECG should undergo further observation and investigation to rule out other diagnoses, enable risk stratification and determine the most appropriate treatment strategy.
- Patients whose ECG and cardiac marker levels are normal after a suitable period of observation should, where practicable, undergo provocative testing (eg, stress test) before discharge.

Management of patients with ST-segment-elevation myocardial infarction

Adjuvant therapy in association with reperfusion
- All patients undergoing reperfusion therapy for STEMI (PCI or fibrinolysis) should be given aspirin and clopidogrel unless these are contraindicated.
- Antithrombin therapy should be given in combination with PCI or fibrinolytic therapy with fibrin-specific fibrinolytic agents, but antithrombin therapy in conjunction with streptokinase is optional.
- It is reasonable to use abciximab with primary PCI, but glycoprotein (GP) IIb/IIIa inhibitors should generally be avoided with full or reduced doses of fibrinolytic therapy.

Choice of reperfusion strategy
- Time delay (both to first medical contact and potential PCI or fibrinolysis) plays a major role in determining best management of STEMI.
- In general, PCI is the treatment of choice, providing it can be performed promptly by a qualified interventional cardiologist in an appropriate facility.
- In general, the maximum acceptable delay from presentation to balloon inflation is:
  - 60 minutes if a patient presents within 1 hour of symptom onset; or
  - 90 minutes if a patient presents later.
- Note: for patients who present late (between 3 and 12 hours after symptom onset) to a facility without PCI capability, it is appropriate to consider transfer for primary PCI if balloon inflation can be achieved within 2 hours (including transport time).
- All PCI facilities should be able to perform angioplasty within 90 minutes of patient presentation.
- Fibrinolysis should be considered early if PCI is not readily available, particularly in rural and remote areas.
- When there are major delays to hospitalisation (ie, more than 30 minutes), prehospital fibrinolysis should be considered.
- Reperfusion is not routinely recommended in patients who present more than 12 hours after symptom onset and who are asymptomatic and haemodynamically stable.

Choice of fibrinolytic agent
- Second-generation fibrin-specific fibrinolytic agents that are available as a bolus (ie, reteplase, tenecteplase) are the fibrinolytics of choice.
- These agents should be available at all centres where fibrinolysis may be required.
- Streptokinase is an inappropriate choice in Aboriginal and Torres Strait Islander patients, or in patients with previous exposure to the drug.
Transfer after STEMI
- Patients who have had STEMI should be considered for early transfer to a tertiary cardiac centre with PCI facilities and links to cardiac surgical facilities.
- If immediate transfer is not possible, patients should be transferred or referred as soon as is practicable for assessment of need for revascularisation (through PCI or coronary artery bypass grafting).

Management of patients with non-ST-segment-elevation acute coronary syndromes
- All patients with non-ST-segment-elevation acute coronary syndromes (NSTEACS) should have their risk stratified to direct management decisions (see page 20 for stratification criteria).
- All patients with NSTEACS should be given aspirin, unless contraindicated.
- High-risk patients with NSTEACS should be treated with aggressive medical management (including aspirin, clopidogrel, unfractionated heparin or subcutaneous enoxaparin, intravenous tirofiban or eptifibatide and a β-blocker), and arrangements should be made for coronary angiography and revascularisation, except in those with severe comorbidities.
- Intermediate-risk patients with NSTEACS should undergo an accelerated diagnostic evaluation and further assessment to allow reclassification as low or high risk.
- Low-risk patients with NSTEACS, after an appropriate period of observation and assessment, may be discharged on upgraded medical therapy for outpatient follow up.

Long-term management after control of myocardial ischaemia
- Before discharge, patients with an ACS should be initiated on a medication regimen, including antiplatelet agent(s), β-blocker, angiotensin-converting enzyme inhibitor, statin and other therapies as appropriate.
- Implantable cardiac defibrillators should be considered in some patients who, despite optimal medical therapy, have persistently depressed left ventricular function more than 6 weeks after STEMI.
- Patients should be given advice on lifestyle changes that will reduce the risk of further coronary heart disease (CHD) events, including smoking cessation, nutrition, alcohol, physical activity and weight management as relevant.
- All patients should have access to, and be actively referred to, comprehensive ongoing prevention and cardiac rehabilitation services.
- All patients should be provided with a written action plan for chest pain.
- Depression and CHD frequently coexist, and in patients with CHD, the presence of depression is more likely to lead to poorer outcomes. Social isolation and lack of social support are also associated with worse outcomes. All patients with CHD should be assessed for depression and level of social support.

Levels of evidence and grades of recommendation
The levels of evidence and grades of recommendations used in these guidelines are adapted from the National Health and Medical Research Council (NHMRC) levels of evidence for clinical interventions and the US National Institutes of Health clinical guidelines. These classifications allow the ability to differentiate between strengths of recommendations and the levels of evidence on which these are based, and allow a classification for recommendations based on panel consensus judgement.

<table>
<thead>
<tr>
<th>Level of evidence*</th>
<th>Study design</th>
<th>Grade of recommendation†</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>Evidence obtained from a systematic review of all relevant randomised controlled trials.</td>
<td>A</td>
</tr>
<tr>
<td>II</td>
<td>Evidence obtained from at least one properly designed randomised controlled trial.</td>
<td>B</td>
</tr>
<tr>
<td>III-I</td>
<td>Evidence obtained from well designed pseudo-randomised controlled trials (alternate allocation or some other method);</td>
<td>B</td>
</tr>
<tr>
<td>III-2</td>
<td>Evidence obtained from comparative studies with concurrent controls and allocation not randomised (cohort studies), case–control studies, or interrupted time series without a control group.</td>
<td>B</td>
</tr>
<tr>
<td>III-3</td>
<td>Evidence obtained from comparative studies with historical control, two or more single-arm studies, or interrupted time series with a parallel control group.</td>
<td>C</td>
</tr>
<tr>
<td>IV</td>
<td>Evidence obtained from case series, either post-test or pre-test and post-test.</td>
<td>C</td>
</tr>
</tbody>
</table>

Rich body of high-quality RCT data

Limited body of RCT data or high-quality non-RCT data

Limited evidence

No evidence available — panel consensus judgement

RCT = randomised controlled trial.

Acute coronary syndromes (ACS) include “a broad spectrum of clinical presentations, spanning ST-segment-elevation myocardial infarction, through to an accelerated pattern of angina without evidence of myonecrosis”. Collectively, they represent one of the most common causes of acute medical admissions to Australian hospitals.

The current guidelines for the management of both ST-segment-elevation ACS and non-ST-segment-elevation ACS have been developed by a joint working party of the National Heart Foundation of Australia (NHFA) and the Cardiac Society of Australia and New Zealand (CSANZ).

The aim of these guidelines is to incorporate contemporary information on the diagnosis and management of ACS into a set of recommendations that defines the boundaries of highest quality care. The guidelines expand on previous guidelines by consolidating recommendations for the management of ST-segment-elevation myocardial infarction (STEMI), non-ST-segment-elevation myocardial infarction and unstable angina, as well as incorporating the newer developments that have arisen since the previous guidelines, Management of unstable angina — 2000 (and addenda, available at: http://www.heartfoundation.com.au) and Reperfusion therapy for acute myocardial infarction (2002).

These new guidelines provide a general framework for appropriate practice, to be followed subject to clinical judgement in each individual patient. They are primarily for doctors in a hospital environment (emergency physicians, general physicians, rural doctors and cardiologists) who manage patients with ACS, but they also contain information relevant to general practitioners and others, including ambulance personnel. The guidelines are designed to provide information to assist decision making, and are based on the best information available up to September 2005. It should be understood that the context in which clinical trials are performed and the local environment in which practice is undertaken must always be considered when assessing the evidence base for guidelines and, at times, their local implementation.

These new guidelines represent a local synthesis of the most recent evidence including recent international guidelines. Where relevant, the evidence has been interpreted with regard to the Australian context in which the guidelines will be implemented.

Key recommendations are summarised at the beginning of these guidelines.

### Systems of care for patients with acute coronary syndromes

The ability to implement best-practice guidelines for the management of ACS will depend on local resources and systems of care. The following guidance is offered to assist practitioners and organisations in facilitating the most effective systems of care for the communities they serve.

Effective management of ACS requires collaborative systems of care to ensure that patients have access to the services that they need in a timeframe commensurate with their clinical condition and the potential benefit of treatments available in larger or specialised centres. The guiding principles for developing these systems are equity of access, equity of care and evidence-based care, taking into account patients’ preferences.

The systems of care should be regionally based, formal rather than ad-hoc, and should cover the continuum of care from the first point of presentation to a health professional to definitive care and rehabilitation. Responsibility for establishing these systems should be at board or executive level within health services.

The systems of care should address:

- clinical issues such as consultation, treatment and acute interhospital transfer protocols (note that systems should enhance options for patients without disempowering decision making by appropriate local clinicians);
- education; and
- quality monitoring, such as time to specific treatments and outcomes.

The structure of these systems will vary depending on the features of the region in which they are placed. In a metropolitan setting, a hospital without percutaneous coronary intervention (PCI) capabilities may have arrangements with a local PCI-capable facility for timely transfer of selected patients. In a rural or remote setting, the system is usually considerably more complex and involves general practitioners or community health centres, prehospital care providers, retrieval services (such as Careflight, Victorian Adult Emergency Retrieval and Coordination Service, Royal Flying Doctor Service), and regional and metropolitan referral hospitals. The systems should be tailored to a region’s needs.

The key elements of successful systems include:

- clear lines of communication (e.g., single points of contact for consultation or referral and coordination of acute interhospital transfers; the consultation component is particularly important as the benefits of some treatments for ACS are time-dependent, so early decision making is vital);
- clear triage protocols where appropriate, recognising the fact that the closest hospital may not be the most suitable in all cases (algorithms can be developed to guide decisions about the best primary destination for patients);
- effective and timely feedback (this should be two-way, and should address ways to improve the process as well as collecting outcomes information; the latter should be both specific for the patient referred and pooled so that trends in outcomes and issues for improvement can be identified);
- agreed treatment protocols, with processes to facilitate drug availability if required;
- agreed acute interhospital transfer protocols and processes;
- program quality monitoring, including analysis of adverse events and system breakdowns;
- identified leaders (these may be drawn from across the system, but leaders should jointly accept responsibility for monitoring the system, providing education and feedback, developing improvements to the system if required, facilitating arrangements with relevant extra-regional organisations and acting as public spokespeople for the system); and
- ownership of the established systems at a senior level within hospital or health service management.
Taking patient preferences into account

On occasion, the pathway of care may recommend that patients be transferred from their local community or region to a distant centre. There may be strong personal or cultural reasons that make this difficult or unacceptable for some patients. Every effort should be made to overcome these barriers by appropriate explanation and discussion, involvement of family and community members and preferential transfer to centres that have specific programs and resources for relevant cultural groups (e.g., Aboriginal Liaison Officers). If the barriers to transfer cannot be overcome and the patient asserts his or her right to be treated more locally, the patient should have the best care that can be delivered in that setting. This includes consultation with specialists.

Aboriginal and Torres Strait Islander peoples

Aboriginal and Torres Strait Islander peoples have a high rate of ACS, and lower intervention rates and poorer outcomes than non-Indigenous people.4 The reasons for this are complex and include barriers to health care access and language and cultural differences. To optimise outcomes for Indigenous people, systems of care that recognise these factors are needed in both metropolitan and rural and remote areas. These might include:

- providing culturally appropriate education and information to Indigenous patients and their families through Aboriginal Health Workers and Hospital Liaison Officers; and
- facilitating interhospital transfer arrangements by involving the local Aboriginal health sector and metropolitan hospital Aboriginal Liaison Officers.

New acute coronary syndromes terminology and implications for diagnosis

The terminology used to describe ACS continues to evolve, with the emergence of the term “non-ST-segment-elevation acute coronary syndrome” (NSTEMI). This reflects a shift away from establishing a definitive diagnosis at presentation, and towards a more clinically appropriate strategy of forming a rapid working diagnosis with its implications for initial clinical decision making.

At presentation, the initial diagnostic nomenclature focuses on risk stratification to direct treatment strategies. Establishing a definitive diagnosis often requires time, particularly for evidence of myocardial necrosis to emerge, and has important implications pertaining to prognosis, diagnostic coding, and social issues such as insurance and licensure. See Box 1 for a representation of diagnosis over time, from presentation to final diagnosis.

Initial working diagnosis

The initial working diagnosis is based on the clinical presentation and the initial electrocardiogram (ECG) findings and, in particular, the presence or absence of ST-segment elevation. As the vast majority of patients who present with initial ST-segment elevation develop biochemical evidence of myocardial necrosis, the term “ST-segment-elevation myocardial infarction” (STEMI) is often used from the outset in these patients.

ACS without ST-segment elevation on the presenting ECG represent a broad spectrum of risk, but are collectively referred to as NSTEMI. This grouping is useful because emergency reperfusion therapy is not indicated (unless ST-segment elevation develops later), and further investigation is required to classify the patient's risk and determine the most suitable treatment (see sections on Investigations [page 12] and Management of patients with STEMI [page 13] for further details).

Final diagnosis

The final diagnostic attribution (i.e., clinical label) has important and persisting implications for the patient, both prognostically and socially. Current international criteria for the diagnosis of myocardial infarction have a strong emphasis on biomarkers, specifically troponin, given its high sensitivity and, in particular, specificity for myocardial necrosis.3 The diagnostic criteria for acute, evolving or recent myocardial infarction are defined as:

- Typical rise in the serum level of troponin or a more rapid rise in the serum level of the MB isoenzyme of creatine kinase (CK-MB) with at least one of the following:
  - Ischaemic symptoms;
  - Development of pathological Q waves on the ECG;
  - ECG changes indicative of ischaemia (ST-segment elevation or depression); and
  - Coronary artery intervention (e.g., coronary angioplasty or coronary bypass surgery);

or

- Pathological findings of an acute myocardial infarction.

This definition requires a temporal appreciation of the cardiac markers, and therefore differentiation between non-ST-segment-elevation myocardial infarction and unstable angina (without evidence of myocardial necrosis) must be delayed.

While this differentiation is necessary for patient education and licensure (e.g., permission to drive, particularly commercial vehicles), it does not change the indication for ongoing prevention strategies.

Standardised terminology and data collection

In 2004, the NHFA and the CSANZ developed a National ACS Dataset, which has now been included in the National Health Data
1 Defining acute coronary syndromes over time: presentation to final diagnosis

- **Presentation** (clinical presentation, initial ECG)
  - **Working diagnosis**
    - ST-segment-elevation myocardial infarction
    - Non-ST-segment-elevation acute coronary syndrome
  - **Time**
    - Evolution of ECG and biomarkers
      - Myonecrosis confirmed
      - Myonecrosis not confirmed
  - **Final diagnosis**
    - ST-segment-elevation myocardial infarction
    - Non-ST-segment-elevation myocardial infarction
    - Unstable angina

**ECG** = electrocardiogram.

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**Acute management of chest pain**

**Getting to hospital**

Chest discomfort at rest or for a prolonged period (more than 10 minutes, not relieved by sublingual nitrates), recurrent chest discomfort, or discomfort associated with syncope or acute heart failure are considered medical emergencies. Other presentations of ACS may include back, neck, arm or epigastric pain, chest tightness, dyspnoea, diaphoresis, nausea and vomiting. Very atypical pain, including sharp and pleuritic pain, is more common in women, people with diabetes and older people.3,7,8

People experiencing such symptoms should seek help promptly and activate emergency medical services to enable transport to the nearest appropriate health care facility for urgent assessment (grade D recommendation). Ideally, transport should be by ambulance. However, where ambulance response times are long, alternatives may need to be considered. Patients should be strongly discouraged from driving themselves because of the risk to other road users.

The most important initial requirement is access to a defibrillator to avoid early cardiac death from reversible arrhythmias. All Australian ambulances now carry defibrillators, and there is promise in further exploring public access defibrillation opportunities. In the case of cardiac arrest occurring in a setting where a defibrillator is not immediately available, cardiopulmonary resuscitation should be commenced immediately.

**Actions in transit**

Aspirin (300mg) should be given unless already taken or contraindicated (grade A recommendation), and should preferably be given early (eg, by emergency or ambulance personnel) (grade D recommendation). Oxygen should also be given (grade D recommendation).

Glyceryl trinitrate and intravenous morphine should be given as required (grade D recommendation).

Where appropriate, a 12-lead ECG should be taken en route and transmitted to a medical facility (grade B recommendation).

Receiving medical facilities should be given warning of incoming patients in whom there is a high suspicion of ACS, particularly STEMI, or those whose condition is unstable (grade B recommendation).

Where formal protocols are in place, prehospital treatment should be given, including fibrinolysis in appropriate cases (grade A recommendation). See section on management of patients with STEMI (page 13) for further discussion of prehospital fibrinolysis.

**On arrival**

All patients presenting with suspected ACS should be subject to ongoing surveillance and have an ECG completed within 5 minutes of arrival at the medical facility (grade A recommendation). The ECG should be assessed promptly by an appropriately qualified person (grade D recommendation).

Oxygen and pain control should be given as required (grade D recommendation).

**Key messages**

- People experiencing symptoms of ACS should seek help promptly and activate emergency medical services.
- The most important initial requirement is access to a defibrillator to avoid early cardiac death from reversible arrhythmias.
- Aspirin should be given early (eg, by emergency or ambulance personnel) unless already taken or contraindicated.
- Oxygen should be given, as well as glyceryl trinitrate and intravenous morphine as required.
- As a minimum, receiving medical facilities should be given warning of incoming patients in whom there is a high suspicion of ACS, particularly STEMI, or whose condition is unstable.
- Where appropriate, a 12-lead ECG should be taken en route and transmitted to a medical facility.
- Where formal protocols are in place, prehospital treatment (including fibrinolysis in appropriate cases) should be facilitated.

Initial investigations

Patients presenting with a suspected ACS should undergo immediate electrocardiography. Further investigations may be necessary, but should not delay treatment.

While other serious diagnoses can present similarly to ACS (eg, pulmonary embolism, aortic dissection, pericarditis), once these have been excluded and ACS is considered the most likely diagnosis further delay in treatment is unnecessary and inappropriate.

Investigations and invasive vascular access techniques should not delay reperfusion therapy if indicated on the basis of ST-segment elevation on the ECG.

Patients whose condition is unstable should have early consultation with a cardiologist.

Cardiac biomarkers are becoming increasingly important to the diagnosis of myocardial infarction. See Box 2 for recommendations and rationale regarding their measurement.

Electrocardiography

Electrocardiography is necessary to detect ischaemic changes or arrhythmias. It should be noted that the initial ECG has a low sensitivity for ACS, and a normal ECG does not rule out ACS. However, the ECG is the sole test required to select patients for emergency reperfusion (fibrinolytic therapy or direct PCI). Patients with STEMI who present within 12 hours of the onset of ischaemic symptoms should have a reperfusion strategy implemented promptly (grade A recommendation) — see the section on management of patients with STEMI (page 13) for recommendations.

Blood tests

Measurements should include:

- Serum troponin I or T levels (or CK-MB if troponin is not available).
- Full blood count.
- Serum creatinine and electrolyte levels, particularly potassium concentration, as hypokalaemia is associated with an increased risk of arrhythmias, especially ventricular fibrillation (grade B recommendation). Knowledge of kidney function (expressed as estimated glomerular filtration rate) is strongly encouraged (grade B recommendation) given the association between renal impairment and adverse outcomes (evidence level III). 11
- Serum creatine kinase (CK) level.
- Serum lipid levels (fasting levels of total cholesterol, low-density-lipoprotein cholesterol, high-density-lipoprotein cholesterol and triglycerides) within 24 hours.
- Blood glucose level.

### 2 Recommendations and rationale for measuring cardiac biomarker levels

<table>
<thead>
<tr>
<th>Cardiac biomarker</th>
<th>Recommendation</th>
<th>Rationale</th>
</tr>
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<tbody>
<tr>
<td>Troponin level</td>
<td>On arrival</td>
<td>Troponin rise indicates myonecrosis, and is a high-risk feature in NSTEACS. Troponin is the preferred marker because about a third of patients with elevated troponin, but normal CK and CK-MB levels, will develop an adverse outcome. 9</td>
</tr>
<tr>
<td></td>
<td>Not repeated if positive</td>
<td>Troponin remains elevated for 5–14 days, and therefore may not be useful for identifying early re-infarction.</td>
</tr>
<tr>
<td></td>
<td>Repeated &gt; 8 hours after last episode of pain or other symptoms of coronary</td>
<td>Troponin elevation is often delayed by 4–6 hours. Therefore, repeat troponin testing is necessary to identify patients at high risk who may benefit from aggressive therapy and an early invasive strategy.</td>
</tr>
<tr>
<td></td>
<td>insufficiency if initially negative</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Serial troponin measurements in patients with NSTEACS suspected to be at high risk</td>
<td>Appearance of typical rise of troponin indicates high risk NSTEACS and may be an indication for more aggressive therapy.</td>
</tr>
<tr>
<td>Total CK level</td>
<td>Serial measurements performed for 48 hours in patients with myocardial</td>
<td>Can be remeasured to confirm a second event if re-infarction is suspected later.</td>
</tr>
<tr>
<td></td>
<td>infarction</td>
<td></td>
</tr>
<tr>
<td>CK-MB level</td>
<td>Should be measured in all patients with an ACS if troponin assay unavailable</td>
<td>While troponin is the preferred marker of myocardial damage, if it is unavailable CK-MB is more specific than CK for myocardial injury. CK-MB may also be used to confirm a re-infarction.</td>
</tr>
</tbody>
</table>

Chest x-ray
A chest x-ray is useful for assessing cardiac size, evidence of heart failure and other abnormalities (grade D recommendation), but should not delay reperfusion treatment where indicated.

Further investigations
Patients without ST-segment elevation on the initial ECG should be further observed and investigated to promptly identify patients suitable for an emergency reperfusion strategy (based on ECG changes) and/or determine the best management protocol for NSTEACS based on risk stratification (see section on Management of patients with NSTEACS [page 20]).

Continuing investigations include:
• continuous ECG monitoring of heart rhythm (ST-segment monitoring is desirable, if available; grade D recommendation);
and
• serial ECGs should be performed in patients with NSTEACS who have high and intermediate risk features (see Management of patients with NSTEACS [page 20]; grade B recommendation). The frequency of ECGs will depend on clinical features (eg, every 10–15 minutes during ongoing symptoms, immediately if symptoms change while the patient is under observation, or at the same intervals as biomarker measurements if the patient is asymptomatic).

Ongoing discomfort requires frequent follow-up 12-lead ECGs (15 minutes apart if continuous ST-segment monitoring is unavailable) to rapidly detect ST-segment elevation and diagnose eligibility for a reperfusion strategy (grade D recommendation). A repeat serum troponin measurement (unless already positive) should be performed at least 8 hours after the last episode of pain or any other symptoms of coronary insufficiency. This has a high sensitivity for detecting acute myocardial infarction, but levels may be normal in other presentations of ACS (grade A recommendation).12

Patients with normal ECG and cardiac markers after an appropriate period of observation should, where practicable, undergo provocative testing (eg, stress test) before discharge. If not immediately available, provocative testing should be arranged at the earliest opportunity, optimally within 72 hours of the index episode (grade C recommendation).13

During ongoing care
In patients with myocardial infarction, serial measurements of total CK should be performed for 48 hours, so that if re-infarction is suspected later, total CK can be remeasured to confirm a second event. A specific marker such as CK-MB may also be used for the diagnosis of re-infarction. Troponins are not useful for diagnosing early re-infarction, as they remain elevated for 5–14 days.12

KEY MESSAGES
• The ECG is the sole test required to select patients for emergency reperfusion (fibrinolytic therapy or direct PCI).
• Patients with STEMI who present within 12 hours of the onset of ischaemic symptoms should have a reperfusion strategy implemented promptly.
• Patients with a suspected ACS without ST-segment elevation on the ECG should undergo further observation and investigation to rule out other diagnoses, enable risk stratification and determine the most appropriate treatment strategy.
• Patients with a normal ECG and cardiac biomarker levels after an appropriate period of observation should, where practicable, undergo provocative testing (eg, stress test) before discharge.

Management of patients with ST-segment-elevation myocardial infarction
STEMI is defined as presentation with clinical symptoms consistent with an acute coronary syndrome with ECG features including any of:
• Persistent ST-segment elevation of ≥ 1 mm in two contiguous limb leads;
• ST-segment elevation of ≥ 2 mm in two contiguous chest leads; or
• New left bundle branch block (LBBB) pattern.6
(Note that LBBB is presumed new unless there is evidence otherwise; echocardiography may be useful to detect regional wall contraction abnormalities.)

Patients with STEMI usually have a completely occluded coronary artery with thrombus at the site of a ruptured plaque. Restoring coronary patency as promptly as possible is a key determinant of short-term and long-term outcomes (level 1 evidence).14,18

Patients with STEMI who present within 12 hours of the onset of ischaemic symptoms should have a reperfusion strategy implemented promptly (grade A recommendation).

Reperfusion therapy
Reperfusion may be obtained with fibrinolytic therapy or PCI. A combination of fibrinolysis and PCI may also be used (facilitated or rescue PCI). Coronary artery bypass graft (CABG) surgery may occasionally be more appropriate — particularly in patients who have suitable anatomy and are not candidates for fibrinolysis or PCI. CABG surgery may also be considered in patients with cardiogenic shock19 or in association with mechanical repair.12

Adjuvant therapy associated with reperfusion
See Box 3 for a summary for recommended adjuvant therapy.

Antiplatelet therapy
Aspirin (300 mg) should be given to all patients with STEMI unless contraindicated and, in the absence of significant side effects, low-dose therapy should be continued in the long term (grade A recommendation).16,20
3 Summary of adjuvant therapy associated with reperfusion

<table>
<thead>
<tr>
<th>Medication</th>
<th>Primary percutaneous coronary intervention</th>
<th>Fibrin-specific fibrinolytic</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aspirin</td>
<td>Yes</td>
<td>Yes (unless the need for acute CABG is likely)</td>
</tr>
<tr>
<td>Clopidogrel</td>
<td>Yes (unless the need for acute CABG is likely)</td>
<td>Yes (unless the need for acute CABG is likely)</td>
</tr>
<tr>
<td>Heparin</td>
<td>Unfractionated heparin (ACT 200–300 s if using glycoprotein IIb/IIIa inhibitors, 300–350 s if not) or Enoxaparin*</td>
<td>Unfractionated heparin (APTT 1.5–2 times control [approx 50–70 s]) or Enoxaparin*</td>
</tr>
<tr>
<td>Glycoprotein IIb/IIIa inhibitors</td>
<td>Abciximab optional</td>
<td>No</td>
</tr>
</tbody>
</table>

* Care should be taken in patients aged over 75 years, or those who have significant renal dysfunction — dose adjustment is required.

CABG = coronary artery bypass graft. ACT = activated clotting time. APTT = activated partial thromboplastin time.

There is evidence that clopidogrel (300–600 mg loading dose) should be prescribed in addition to aspirin for patients undergoing PCI with a stent.21-23 In patients selected for fibrinolytic therapy, clopidogrel (300 mg) should be given in addition to aspirin, unless contraindicated (grade B recommendation).24 Note, however, that if it is thought that the patient is likely to require CABG acutely, clopidogrel should be withheld.

Clopidogrel (75 mg daily) should be continued for at least a month after fibrinolytic therapy, and for up to 12 months after stent implantation, depending on the type of stent and circumstances of implantation (level II evidence; grade B recommendation).25

Antithrombin therapy

**With PCI:** Antithrombin therapy should be used in conjunction with PCI (grade A recommendation). The dose of unfractionated heparin therapy will depend on concomitant use of glycoprotein (GP) IIb/IIIa inhibitors. The aim should be to obtain an activated clotting time (ACT) between 200 and 300 seconds if using GP IIb/IIIa inhibitors, or between 300 and 350 seconds if these drugs are not used (grade B recommendation). It may be advisable to give a bolus of heparin while the patient is in transit to the catheterisation laboratory (grade D recommendation). The role of enoxaparin in acute STEMI in conjunction with PCI remains to be determined, but it appears to be safe and effective at a dose of 0.75 mg/kg (grade D recommendation).

**With fibrinolysis:** Antithrombin therapy should be used with fibrin-specific fibrinolytic agents (grade A recommendation).26,27 Unfractionated heparin should be given as an initial bolus dose of 60 units per kilogram of body weight (with a maximum dose of 4000 units) followed by an initial infusion of 12 units per kilogram per hour (maximum units 1000 per hour), adjusted to attain the activated partial thromboplastin time (APTT) at 1.5 to 2 times control (about 50–70 seconds; grade B recommendation).12 Enoxaparin may be used in conjunction with fibrin-specific fibrinolytic agents in patients under the age of 75 years, provided they do not have significant renal dysfunction. An intravenous bolus dose of 30 mg followed by a 1 mg/kg subcutaneous injection every 12 hours in combination with tenecteplase is the most comprehensively studied therapy.12 Care should be taken in patients who are aged over 75 years, or who have renal dysfunction, as dose adjustment is required.12 The use of antithrombin therapy in conjunction with streptokinase therapy is optional.28

**Glycoprotein IIb/IIIa inhibitors**

It is reasonable to use abciximab with primary PCI, although there are conflicting data (grade B recommendation). It appears the earlier it is used, the greater the advantage.29 When used in patients with STEMI undergoing primary PCI, the timing of administration of abciximab is a matter of clinical judgement.30,31 Full-dose GP IIb/IIIa inhibitors should be avoided with fibrinolytic therapy (grade B recommendation) as there is evidence of excessive bleeding (including intracranial haemorrhage) with this combination.32 It is unclear how early full-dose GP IIb/IIIa inhibitors can be safely given after fibrinolysis, but it is probably at least 4 hours after administration of fibrin-specific fibrinolytic agents and 24 hours after administration of streptokinase.32 The combination of GP IIb/IIIa inhibitors with reduced doses of fibrinolytic therapy is not recommended. There is no significant advantage over full-dose fibrinolytic therapy alone, and the risk of bleeding is increased, particularly in the elderly.32 This combination has been used for facilitated PCI.32

**KEY MESSAGES**

- All patients undergoing reperfusion therapy (PCI or fibrinolysis) for STEMI should be given aspirin and clopidogrel unless contraindicated.
- Antithrombin therapy should be given in combination with PCI or fibrinolytic therapy with fibrin-specific fibrinolytic agents, but its use in conjunction with streptokinase is optional.
- It is reasonable to use abciximab with primary PCI, but GP IIb/IIIa inhibitors should generally be avoided with full or reduced doses of fibrinolytic therapy.

**Choice of reperfusion therapy**

The choice of reperfusion therapy is usually between PCI and fibrinolytic therapy.

PCI is the best available treatment if provided promptly by a qualified interventional cardiologist in an appropriate facility (level I evidence; grade A recommendation).

PCI will improve both short-term and long-term outcomes (reduced deaths, myocardial infarctions and strokes) in patients with STEMI presenting within 12 hours compared with fibrinolytic therapy (level I evidence).33,34 However, this benefit may only occur if the additional time delay associated with PCI — over and above that associated with giving fibrinolysis — is less than 1 hour (level IV evidence).35 See below for further details.
Where PCI is not available or is delayed, reperfusion with fibrinolytic therapy should occur unless contraindicated (level I evidence; grade A recommendation).

Cardiologists performing primary PCI should have significant expertise in both coronary angioplasty and management of patients with acute myocardial infarction (level III evidence; grade B recommendation). The cardiologist and the unit should fulfill the minimum requirements defined by the CSANZ for competency in angioplasty.36 The unit should also perform a sufficient volume of primary PCIs — international experience suggests this might be more than 36 per unit per year (level III evidence; grade B recommendation).37

On-site surgical backup is not a requirement for primary angioplasty (level III evidence; grade B recommendation); however, established networks for urgent referral should be in place.

The choice of reperfusion therapy will depend on a number of factors, including:
- time delay to PCI;
- time from symptom onset to first medical contact;
- time to hospital fibrinolysis;
- contraindications to fibrinolytic therapy;
- location and size of infarction;
- presence of cardiogenic shock; and
- special circumstances.

The major factor determining the choice of reperfusion strategy is time, including time since symptom onset, time delay for transportation, and time delay for PCI.

### Time delay to percutaneous coronary intervention

The acceptable delay to PCI will vary with time from symptom onset to presentation. Time to PCI in this context relates to time from presentation to balloon inflation, not arrival at a PCI-capable hospital or even at the catheter laboratory.

In general (see below for exceptions), a time delay of 90 minutes from first medical contact to balloon inflation is the maximum desirable, otherwise fibrinolysis should be used (level I evidence; grade A recommendation). This time is arrived at by presuming a delay of 30 minutes from presentation to delivery of fibrinolysis and recognising that PCI is of benefit if performed within 60 minutes of potential fibrinolysis.35 All PCI facilities should be able to perform angioplasty within 90 minutes of patient presentation.

In circumstances where the delay to hospital for fibrinolytic therapy is significant (more than 30 minutes), prehospital fibrinolysis should be considered (level II evidence; grade B recommendation) — see below.

**Transfer for primary PCI versus immediate fibrinolysis:** For patients presenting with STEMI at a facility without PCI facilities, transfer to a PCI-capable facility (rather than immediate fibrinolysis) may provide benefits in terms of risk of death, stroke and re-infarction, provided PCI can be performed promptly (level II evidence).38,39 If PCI can be performed in an appropriate time frame (see Box 4), the patient should be transferred to a PCI centre (grade B recommendation). Otherwise, fibrinolysis should be given.

Warning the cardiac catheter laboratory that the patient is being transferred is vital for implementation of this strategy.39

**Direct transport to a PCI centre:** Significant improvements in delay to PCI may be made by directly transporting patients to a PCI centre rather than transporting them to the nearest hospital if interhospital transfer will subsequently be required to obtain primary PCI (level II evidence; grade B recommendation).40 However, to translate such a protocol into benefit for patients, processes must be in place to identify appropriate patients accurately and minimise delay to PCI at the receiving hospital.

**Transfer for PCI following fibrinolytic therapy — facilitated PCI:** Facilitated PCI refers to a strategy of planned immediate PCI after an initial thrombolytic or fibrinolytic regimen, regardless of reperfusion status.

---

### Hospital management of ST-segment-elevation myocardial infarction*

<table>
<thead>
<tr>
<th>Symptom onset</th>
<th>Percutaneous coronary intervention available</th>
<th>Fibrinolysis†</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;1 hour</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>1–3 hours</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>3–12 hours</td>
<td>Yes</td>
<td>No</td>
</tr>
</tbody>
</table>

* Assuming no contraindications to fibrinolytic therapy — see Box 5. † Time delay refers to time from first medical contact to balloon. ‡ Patients with ongoing symptoms or instability should be transferred for PCI.

Note: Reperfusion after 12 hours is indicated for cardiogenic shock, ongoing pain or haemodynamic instability (see text).
5 Contraindications and cautions for fibrinolysis use in ST-segment-elevation myocardial infarction* 

Absolute contraindications

Risk of bleeding
- Active bleeding or bleeding diathesis (excluding menses)
- Significant closed head or facial trauma within 3 months
- Suspected aortic dissection (including new neurological symptoms)50
- Risk of intracranial haemorrhage
  - Any prior intracranial haemorrhage
  - Ischaemic stroke within 3 months
  - Known structural cerebral vascular lesion (eg, arteriovenous malformation)
  - Known malignant intracranial neoplasm (primary or metastatic)

Relative contraindications

Risk of bleeding
- Current use of anticoagulants: the higher the international normalised ratio (INR), the higher the risk of bleeding
- Non-compressible vascular punctures
- Recent major surgery (< 3 weeks)
- Traumatic or prolonged (> 10 minutes) cardiopulmonary resuscitation
- Recent (within 4 weeks) internal bleeding (eg, gastrointestinal or urinary tract haemorrhage)
- Active peptic ulcer
- Risk of intracranial haemorrhage
  - History of chronic, severe, poorly controlled hypertension
  - Severe uncontrolled hypertension on presentation (> 180 mmHg systolic or > 110 mmHg diastolic)
  - Ischaemic stroke more than 3 months ago, dementia, or known intracranial abnormality not covered in contraindications
- Other
  - Pregnancy

*Modified with permission of the American College of Cardiology and the American Heart Association.12

The possible drug regimen associated with facilitated PCI includes:
- full-dose fibrinolytic; and
- half-dose fibrinolytic and GP IIb/IIIa inhibitor.

This strategy is theoretically attractive, but has not been proven. A recent trial with full-dose tenecteplase followed by immediate angioplasty (within a median time of 104 minutes) showed inferior outcomes compared with direct PCI.41 While other strategies continue to be tested in ongoing trials, immediate PCI after full dose fibrinolysis cannot be recommended at this time (grade D recommendation).

Rescue PCI (where reperfusion has not occurred) may be of benefit, and is discussed below.

Time from symptom onset to first medical contact

Early presentation (< 1 hour after symptom onset): Reperfusion treatment has a much greater benefit in patients who present very early (level I evidence). Fibrinolytic therapy given early after symptom onset can result in a reduction in death of up to 50% (level I evidence),42 and, in patients who present very early (ie, within 1 hour), the delays to PCI become even more important. In this situation, fibrinolytic therapy should be considered unless PCI is available within 1 hour (level III evidence, grade B recommendation).

Presentation 1–3 hours after symptom onset: Both primary PCI and fibrinolysis are effective for treating STEMI within 1–3 hours of symptom onset (level I evidence).38 PCI is preferable if it can be performed in a timely manner (balloon inflation within 90 minutes of first medical contact or 60 minutes of potential fibrinolysis), or if fibrinolysis is contraindicated (grade A recommendation). Otherwise, fibrinolysis should be used (grade A recommendation).

Late presentation (3–12 hours after symptom onset): For patients who present more than 3 hours after symptom onset, reperfusion with PCI is superior to fibrinolytic therapy (level II evidence, grade B recommendation).36 and the window of efficacy is wider. Balloon inflation should be achieved within 90 minutes of presentation to a PCI-capable facility. However, for facilities without PCI capability, consideration of transfer for primary PCI is appropriate if balloon inflation can be achieved within 2 hours (including transport time). If PCI is not available within this time frame, fibrinolysis should be given (grade A recommendation).

Very late presentation (> 12 hours after symptom onset): Reperfusion therapy with either PCI or fibrinolysis is not routinely recommended in patients who are asymptomatic and haemodynamically stable, and who present more than 12 hours after symptom onset (grade B recommendation).43 Preliminary evidence, however, suggests that patients may benefit from PCI performed between 12 and 24 hours after symptom onset.44 If the patient has ongoing symptoms or is haemodynamically unstable, either reperfusion strategy can be considered (grade B recommendation).

Cardiogenic shock is covered below.

Time to hospital — prehospital fibrinolysis

In some cases, the delay between a patient’s first presentation (to emergency medical services, general practitioner or health clinic) and either PCI or hospital-based fibrinolysis may be considerable. In such cases, prehospital fibrinolysis should be considered (grade B recommendation).

In the Australian context, prehospital fibrinolysis (by general practitioners, ambulance paramedics, nurses or other qualified staff in a variety of prehospital settings) needs to be considered:
- when the delay to PCI is outside acceptable limits defined above; and
- when transport delay to a hospital for fibrinolysis exceeds 30 minutes (grade B recommendation).

These circumstances will usually apply to patients from rural and remote areas, and some fringe or commuter areas around major cities.45 The above recommendation is based on data from Victoria showing a greater relative risk of death if fibrinolysis is delayed beyond this time46 and by a meta-analysis comparing in-hospital and prehospital fibrinolysis.47 There is evidence that, among patients receiving fibrinolysis in the Northern Territory, Indigenous patients are more likely to receive prehospital fibrinolysis than non-Indigenous patients.48

Prehospital fibrinolysis should be considered as a component of the system of care. It requires established linkages for patient transfer for further care; drug and transfer protocols; processes for consultation, training and quality assurance; and processes to facilitate access to appropriate fibrinolytic agents. These system issues should be addressed on a regional basis (grade D recommendation).
Contraindications to fibrinolytic therapy

Patients with a contraindication to fibrinolytic therapy benefit from early interventional therapy and should be considered for early transfer if invasive facilities are not available (grade B recommendation). The delay to PCI can be longer than discussed above, as it is important that an attempt to reperfuse is made even if there is a long delay of up to 12 hours from symptom onset (level III evidence, grade B recommendation). See Box 5 for absolute and relative contraindications to fibrinolytic therapy.

Location and size of infarction

In patients with acute myocardial infarction involving a large area of risk, successful reperfusion is even more important. The area of risk can be defined by the extent of ECG changes. High-risk patients include those with anterior ST-segment elevation (with the more chest leads involved, the higher the risk), inferior infarctions with significant anterior ST-segment depression, signs of right ventricular infarction or left bundle branch block (level I evidence).

In situations where resources do not allow PCI for all patients, its selective use for patients with large infarctions is advisable (grade A recommendation).

Presence of cardiogenic shock

PCI is the preferred strategy in patients aged under 75 years with cardiogenic shock, provided it can be performed promptly. The aim should be balloon inflation within 90 minutes of first medical contact (level II evidence, grade B recommendation). In patients aged over 75 years with significant comorbidity, initial conservative treatment may be preferable.

While fibrinolytic therapy does not provide a major benefit in cardiogenic shock, it should be considered in patients with ST-segment elevation and cardiogenic shock if PCI is not a realistic option (grade D recommendation).

CABG surgery should be considered in patients with cardiogenic shock and appropriate coronary artery anatomy (level II evidence, grade B recommendation).

Special circumstances

Diagnosis of STEMI is in doubt: In selected high-risk patients who present with what appear to be ischaemic symptoms, but no clear evidence of ST-segment elevation on the ECG (as can occur particularly with occlusions of the circumflex artery), coronary angiography with possible angioplasty may be preferable to treatment with fibrinolytic therapy (grade D recommendation). If this is not available, repeated ECGs or echocardiography (to detect regional wall contraction abnormalities) may be useful in identifying candidates suitable for reperfusion therapy, and the opinion of an experienced cardiologist should be sought urgently.

Heart failure: PCI is the preferred strategy for patients with severe heart failure (Killip class, ≥ 3) (level III evidence, grade B recommendation).

### 6 Prehospital management of ST-segment-elevation myocardial infarction (STEMI)

<table>
<thead>
<tr>
<th>STEMI confirmed by 12-lead electrocardiogram with expert interpretation</th>
<th>No contraindications to fibrinolytic therapy*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Time delay to percutaneous coronary intervention acceptable (see below)</td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Time delay to hospital for fibrinolysis &lt; 30 minutes</td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Prehospital fibrinolysis</td>
<td></td>
</tr>
</tbody>
</table>

Direct to percutaneous coronary intervention-capable hospital

Direct to hospital for fibrinolytic therapy

Hospital

<table>
<thead>
<tr>
<th>Time since onset of symptoms</th>
<th>&lt; 1 hour</th>
<th>1–3 hours</th>
<th>3–12 hours</th>
<th>&gt; 12 hours</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acceptable delay to percutaneous coronary intervention (from first medical contact to balloon inflation)</td>
<td>60 minutes</td>
<td>90 minutes</td>
<td>120 minutes</td>
<td>Not routinely recommended (see text)</td>
</tr>
</tbody>
</table>

*If fibrinolysis is contraindicated, it is important that an attempt to reperfuse is made, even if there is a long delay (up to 12 hours).
Summary

In general, PCI is the treatment of choice (grade A recommendation), but only if:
• it can be performed promptly; and
• it is performed by appropriately qualified interventional cardiologists in an appropriate facility.

When PCI is not promptly available, fibrinolytic therapy should be used.

See Boxes 4 and 6 for simplified algorithms to direct decision making based on these parameters in the hospital and prehospital settings, respectively.

**Key messages**

- Choice of reperfusion strategy depends on a number of factors, with time delay (both to presentation and potential PCI or fibrinolytic therapy) playing a major role in determining best management.
- In general, PCI is the treatment of choice, provided it can be performed promptly by a qualified interventional cardiologist in an appropriate facility.
- In general, the maximum acceptable delay from presentation to balloon inflation is:
  - 60 minutes if a patient presents within 1 hour of symptom onset; and
  - 90 minutes if a patient presents later.

**Note:** for patients who present late (3–12 hours after symptom onset) to a facility without PCI capability, it is appropriate to consider transfer for primary PCI if balloon inflation can be achieved within 2 hours (including transport time).
- All PCI facilities should be able to perform angioplasty within 90 minutes of patient presentation.
- Fibrinolysis should be considered early if PCI is not readily available, particularly in rural and remote areas.
- When there are major delays to hospitalisation (more than 30 minutes), prehospital fibrinolysis should be considered.
- Reperfusion is not routinely recommended in patients who present more than 12 hours after symptom onset and who are asymptomatic and haemodynamically stable.

### Choice of fibrinolytic agent

There are four fibrinolytic agents currently available in Australia; streptokinase and the tissue fibrin-specific fibrinolytic agents alteplase, reteplase and tenecteplase. The properties of these agents are summarised in Box 7.

Fibrin-specific fibrinolytic agents have been shown to reduce mortality compared with streptokinase in patients with STEMI who present within 6 hours of symptom onset (level I evidence). Streptokinase may be associated with a lower incidence of intracranial haemorrhage, particularly in older people (level I evidence), but the overall mortality is still lower with the use of fibrin-specific fibrinolytic agents (level II evidence). Tenecteplase is associated with a lower rate of bleeding than alteplase (level II evidence).

Second-generation fibrin-specific fibrinolytic agents can be given as either single or double bolus injections, which makes them significantly easier to use than streptokinase.

In combination therapy, PCI combined with fibrin-specific fibrinolytic agents appears to have greater efficacy and results in fewer complications than PCI combined with streptokinase (level III evidence, grade B recommendation).

Streptokinase should not be given to patients with previous exposure (more than 5 days ago) to the drug (grade B recommendation). There is also evidence that streptokinase may be less effective in Aboriginal and Torres Strait Islander peoples because of the high levels of skin infection (and thus streptococcal antibodies), particularly in remote populations. Streptokinase should be available at all centres where fibrinolysis may be required (grade D recommendation).

### 7 Fibrinolytic agents currently available in Australia — prescribing information and properties

<table>
<thead>
<tr>
<th>Details</th>
<th>Tenecteplase (TNK)</th>
<th>Reteplase (r-PA)</th>
<th>Alteplase (rt-PA)</th>
<th>Streptokinase</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Dose</strong> (also see product information)</td>
<td>Up to 10 000 international units (50 mg) on basis of body weight:</td>
<td>10 units x 2, 30 minutes apart (give each bolus slowly over no more than 2 minutes)</td>
<td>For patients &gt; 65 kg: 15 mg bolus; then 50 mg over 30 minutes and 35 mg over the next 60 minutes</td>
<td>1.5 million international units over 30–60 minutes</td>
</tr>
<tr>
<td>&lt; 60 kg — 6000 units</td>
<td></td>
<td></td>
<td>For patients ≤ 65 kg: 15 mg bolus; then 0.75 mg/kg over 30 minutes and 0.5 mg/kg over the next 60 minutes.</td>
<td></td>
</tr>
<tr>
<td>60–70 kg — 7000 units</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>70–80 kg — 8000 units</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>80–90 kg — 9000 units</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&gt; 90 kg — 10 000 units</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Bolus administration</strong> Yes</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td><strong>Antigenic</strong> No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td><strong>Systemic fibrinogen depletion</strong> Minimal</td>
<td>Moderate</td>
<td>Mild</td>
<td>Marked</td>
<td></td>
</tr>
<tr>
<td><strong>Lives saved per 1000 patients treated</strong> (approximate number at 30 days) 3553</td>
<td>3554</td>
<td>3555</td>
<td>2514</td>
<td></td>
</tr>
</tbody>
</table>
**Failed reperfusion**

After reperfusion therapy, patients should be monitored for symptoms and changes in ST-segment elevation (grade D recommendation). Non-invasive findings suggestive of successful reperfusion include relief of symptoms, restoration of haemodynamic or electrical stability, and reduction by 50% of the initial ST-segment elevation within 60–90 minutes of initiation of therapy.12

In patients in whom fibrinolysis fails, rescue PCI should be considered (grade B recommendation). Ideally, patients who receive fibrinolysis at a facility not capable of PCI should be transferred rapidly so that rescue PCI can occur no later than 90 minutes after fibrinolysis if necessary (grade C recommendation). It is recognised that this may have systems implications that will need to be explored.

Patients in whom reperfusion fails, or in whom re-occlusion occurs, in a setting where rescue PCI can not be performed within a reasonable time should be considered for further medical reperfusion.

There are few data for patients in whom reperfusion fails, but an additional dose of fibrinolytic therapy has been given (grade D recommendation). The balance between the risk of the myocardial infarction and the risk of bleeding must be considered.

While GP IIb/IIIa inhibitors are not generally recommended at the time of fibrinolytic therapy, preliminary data suggest that administering intravenous tirofiban to patients with failed fibrinolysis or recurrent ST-segment elevation results in a superior outcome compared with historical controls (level III evidence).60 In patients who re-occlude after initial fibrinolytic therapy, a further dose of fibrinolytic can be given safely, as long as the effects of the first dose have abated (6 hours after fibrin-specific fibrinolytic agents, 24 hours after streptokinase).

**Rescue percutaneous coronary intervention**

Rescue PCI may be beneficial in patients with:

- haemodynamic and electrical instability (level II evidence);
- persistent ischaemic symptoms (level III evidence);
- cardiogenic shock (level II evidence).

Rescue PCI refers to PCI after failed fibrinolysis for patients with continuing or recurrent myocardial ischaemia. PCI should be considered in patients who develop cardiogenic shock within 36 hours after a myocardial infarction, or those who develop severe haemodynamic instability (grade B recommendation).51,61

The data for rescue PCI are still evolving, but general consensus is that obtaining normal coronary flow has long-term beneficial effects (grade D recommendation).

---

**Cardiac surgery**

Emergency or urgent bypass surgery should be considered in patients with STEMI and:

- failed PCI with persistent pain or haemodynamic instability and coronary anatomy suitable for surgery (grade B recommendation); or
- persistent or recurrent ischaemia refractory to medical therapy and suitable anatomy (grade B recommendation).

**Transfer to a tertiary cardiac centre after STEMI**

**Patients who have had STEMI should be considered for transfer to a tertiary cardiac centre** with PCI facilities and links to cardiac surgical facilities (grade B recommendation). There is a significant incidence of re-occlusion and re-infarction after reperfusion therapy.31,35 Many patients will develop symptomatic angina or require hospital admission later.26 There is evidence that transfer for further assessment and appropriate revascularisation will reduce symptomatic angina and re-admission,62 but few data suggest improved survival or reduction in recurrent myocardial infarction.

The timing of the transfer will depend on the success of reperfusion, haemodynamic and electrical stability, and the availability of transfer. Decisions about transfer should be made within the local context.

Early transfer should be considered in all patients, but particularly those with any of the following:

- ongoing pain;
- a large area of myocardium at risk (including those with anterior ST-segment elevation, inferior infarctions with significant anterior ST-segment depression, signs of right ventricular infarction or LBBB);
- known poor left ventricular function; and
- renal impairment.

If immediate transfer is not possible, all patients should be transferred or referred as soon as is practicable for coronary angiography and assessment of the need for revascularisation (by PCI or CABG, grade D recommendation).

**Routine revascularisation after STEMI**

For patients with objective evidence of recurrent myocardial infarction in whom there is spontaneous or inducible ischaemia or haemodynamic instability, coronary angiography with a view to PCI or coronary surgery, if appropriate, should be performed (grade B recommendation).

CABG may be considered in patients with poor ventricular function and appropriate coronary anatomy, patients with left main disease and patients with severe triple vessel disease (grade B recommendation).63,64

**Key messages**

- Patients who have had STEMI should be considered for early transfer to a tertiary cardiac centre with PCI facilities and links to cardiac surgical facilities.
- If early transfer is not possible, all patients should be transferred or referred as soon as is practicable for assessment of the need for revascularisation through PCI or CABG.
Management of patients with non-ST-segment-elevation acute coronary syndromes

Risk stratification

The initial objective of evaluation is to define the likelihood of an ACS as the cause of a patient’s presentation. Most patients will present with prolonged or recurrent central chest discomfort but others, particularly the elderly, people with diabetes and women, may present with atypical symptoms. These include neck, jaw, back or epigastric discomfort or dyspnoea, diaphoresis, nausea and vomiting. Age is an important risk factor, and the presence (or absence) of coronary risk factors adds little to the accuracy of the diagnosis in middle-aged or elderly patients, but is more useful in making a diagnosis in younger patients. A history of physical or emotional stress before symptom onset increases the likelihood of an ACS. Most patients with NSTEACS are normal on physical examination. An abnormal ECG, particularly dynamic ST-segment deviation (≥ 0.5 mm) or new T-wave inversion (≥ 2 mm) will confirm the diagnosis, but the ECG may be normal or show minor changes in up to 50% of cases.

The second objective of evaluation is to determine the risk of short-term adverse outcomes, which will direct the management strategy. Box 8 provides a paradigm for the risk stratification of patients presenting with suspected NSTEACS, and a simplified risk assessment algorithm is shown in Box 9. Most patients admitted to hospital with possible NSTEACS will have intermediate-risk or high-risk features (Box 8), and these patients are best managed with a structured clinical pathway (see Investigations section [page 12]). Patients with clinical features consistent with NSTEACS and high-risk features are best managed with aggressive medical and invasive therapy (detailed later). Patients with diabetes or chronic kidney disease with typical symptoms of ACS would be considered to be at high risk, but those with atypical symptoms and normal ECGs and cardiac biomarker levels may initially be considered at intermediate risk until a diagnosis is made. Patients with low-risk unstable angina may be managed with upgraded medical therapy and outpatient cardiac referral.

Emerging risk factors

In patients with NSTEACS, diabetes has emerged as an independent risk factor for adverse cardiac events (level I evidence), and should be regarded as a high-risk feature in patients who present with typical symptoms of ACS. People with diabetes have an increased risk similar to that of patients with an elevated troponin level or ST-segment deviation.65,66 It should be noted that there is likely to be a high rate of undiagnosed diabetes among people presenting with acute coronary syndromes.65

Chronic kidney disease (CKD): Substantial clinical data from registries and clinical trials document the excess risk of mortality, recurrent cardiac events and bleeding events associated with reduced renal function among patients with an ACS.11 While the relationship between reduced renal function and clinical events is proportional, this evidence supports a threshold level of renal impairment of a glomerular filtration rate less than 60 mL/minute as having significant negative prognostic impact.68 Despite this, few studies have formally sought strategies for reducing risk in this specific population. Although there are limited data on the use of invasive strategies among patients with CKD who present with an ACS, there appears to be a trend towards benefit in the use of early invasive strategies in these patients.69,70 There is emerging evi-

8 Features associated with high-risk, intermediate-risk and low-risk non-ST-segment-elevation acute coronary syndromes (NSTEACS)

<table>
<thead>
<tr>
<th>High-risk features</th>
<th>Intermediate-risk features</th>
<th>Low-risk features</th>
</tr>
</thead>
<tbody>
<tr>
<td>Presentation with clinical features consistent with acute coronary syndromes (ACS) and any of the following high-risk features:</td>
<td>Presentation with clinical features consistent with ACS and any of the following intermediate risk features AND NOT meeting the criteria for high-risk ACS:</td>
<td>Presentation with clinical features consistent with an acute coronary syndrome without intermediate-risk or high-risk features. This includes onset of anginal symptoms within the last month, or worsening in severity or frequency of angina, or lowering of anginal threshold.</td>
</tr>
<tr>
<td>• Repetitive or prolonged (&gt; 10 minutes) ongoing chest pain or discomfort;</td>
<td>• Chest pain or discomfort within the past 48 hours that occurred at rest, or was repetitive or prolonged (but currently resolved);</td>
<td></td>
</tr>
<tr>
<td>• Elevated level of at least one cardiac biomarker (troponin or creatine kinase-MB isoenzyme);</td>
<td>• Age &gt;65 years;</td>
<td></td>
</tr>
<tr>
<td>• Persistent or dynamic electrocardiographic changes of ST-segment depression ≥ 0.5 mm or new T-wave inversion ≥ 2 mm;</td>
<td>• Known coronary heart disease — prior myocardial infarction with left ventricular ejection fraction &lt; 0.40, or known coronary lesion more than 50% stenosed;</td>
<td></td>
</tr>
<tr>
<td>• Transient ST-segment elevation (≥ 0.5 mm) in more than two contiguous leads;</td>
<td>• No high-risk changes on electrocardiography (see above);</td>
<td></td>
</tr>
<tr>
<td>• Haemodynamic compromise — systolic blood pressure &lt; 90 mmHg, cool peripheries, diaphoresis, Killip Class &gt; I, and/or new-onset mitral regurgitation;</td>
<td>• Two or more of the following risk factors: known hypertension, family history, active smoking or hyperlipidaemia;</td>
<td></td>
</tr>
<tr>
<td>• Sustained ventricular tachycardia;</td>
<td>• Presence of known diabetes (with typical symptoms of ACS);</td>
<td></td>
</tr>
<tr>
<td>• Syncope;</td>
<td>• Chronic kidney disease (estimated glomerular filtration rate &lt; 60 mL/minute) (with typical symptoms of ACS);</td>
<td></td>
</tr>
<tr>
<td>• Left ventricular systolic dysfunction (left ventricular ejection fraction &lt; 0.40);</td>
<td>• Chronic kidney disease (estimated glomerular filtration rate &lt; 60 mL/minute) (with atypical symptoms of ACS); or</td>
<td></td>
</tr>
<tr>
<td>• Prior percutaneous coronary intervention within 6 months or prior coronary artery bypass surgery;</td>
<td>• Prior aspirin use.</td>
<td></td>
</tr>
<tr>
<td>• Presence of known diabetes (with typical symptoms of ACS); or</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Chronic kidney disease (estimated glomerular filtration rate &lt; 60 mL/minute) (with typical symptoms of ACS).</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

S20 MJA • Volume 184 Number 8 • 17 April 2006
dence to suggest that the benefits of conventional medical therapies used for ACS confer similar if not greater benefit on patients with CKD than those without the disease,\textsuperscript{71-75} and there is no significant evidence to suggest that patients with CKD are at increased risk of drug-related toxicities with the use of aspirin, β-blockers, statins, or angiotensin-converting enzyme inhibitors.\textsuperscript{71-74,76} Thus, CKD should not discourage the use of either an early invasive strategy or established pharmacotherapies in the management of ACS (grade B recommendation). However, where relevant (eg, with β-blockers), titration of such agents should be performed cautiously to avoid drug accumulation in the context of renal clearance.\textsuperscript{77,78}

Other markers of risk: C-reactive protein and brain natriuretic peptide are under active investigation as markers of risk in ACS. Currently there are insufficient data available to support their routine use.

Treatment of NSTEACS

Aspirin is recommended (unless contraindicated) in all low-risk, intermediate-risk and high-risk patients (grade A recommendation). \textbf{High-risk patients} should be treated with aggressive medical management (level I evidence, grade A recommendation) (see below) and arrangements should be made for coronary angiography and revascularisation (level I evidence, grade A recommendation), except in those with severe comorbidities, including general frailty (grade A recommendation). Age alone should not be a barrier to aggressive therapy.

Patients at intermediate risk should be observed by staff trained in cardiac care practice and should undergo an accelerated diagnostic evaluation and further risk stratification (level III evidence, grade B recommendation). Accurate assessment can be improved by the use of structured forms for admission and continuing evaluation.\textsuperscript{79-82} During the evaluation process, intermediate-risk patients are observed as described in the Investigations section (page 12), with frequent electrocardiography (with or without continuous ST-segment monitoring), repeat troponin testing and provocative testing if a repeat troponin assay is negative.

\textbf{Low-risk patients} may be discharged on upgraded medical therapy after an appropriate period of observation and assessment (see Investigations section). These patients (including those manifesting anginal symptoms for the first time within the previous month or with a change in the tempo of their anginal) are considered unstable, as some will have atherothrombotic disease with a definite risk of progression to myocardial infarction. These patients should be treated with β-blockers and aspirin, and cardiac assessment should be obtained urgently.

Treatment of patients with NSTEACS on the basis of risk is summarised in Box 10.

\textbf{Medical management of high-risk patients}

\textit{Antiplatelet therapy:} Early treatment should be initiated with aspirin\textsuperscript{83-86} (grade A recommendation) and clopidogrel (300 mg loading dose and 75 mg daily)\textsuperscript{87} (grade B recommendation), with the following considerations:

- Clopidogrel should be avoided in patients likely to require emergency coronary bypass surgery (those with severe widespread ST-segment depression or haemodynamic instability);
- If possible, clopidogrel should be discontinued 5 days before coronary bypass surgery;
- Clopidogrel should be given (preferably more than 6 hours) before planned percutaneous coronary intervention (level I evidence, grade A recommendation),\textsuperscript{21,22} but may be omitted if coronary angiography is planned immediately;
- If relevant, warfarin therapy should be discontinued and heparin given along with the recommended antiplatelet therapy (grade D recommendation).

\textbf{10 Treatment strategies for patients with non-ST-segment-elevation acute coronary syndromes (NSTEACS), based on risk stratification}

\begin{tabular}{|c|c|c|}
\hline
\textbf{Symptom} & \textbf{Low (< 2%) risk*} & \textbf{Intermediate (2%–10%) risk*} & \textbf{High (> 10%) risk*} \\
\hline
Any pain & Yes & Yes & Yes \\
Pain at rest, repetitive or prolonged pain & No & Yes & Yes \\
Changes on electrocardiogram or elevated troponin level & No & No & Yes \\
\hline
\end{tabular}

* Risk categories are based on the presence of clinical factors known to increase rates of myocardial infarction and death within 6 months.

\textbf{9 Simplified risk assessment algorithm}

6-month risk of death or myocardial infarction

- **Any pain:** Yes
- **Pain at rest, repetitive or prolonged pain:** No
- **Changes on electrocardiogram or elevated troponin level:** No

\textbf{10 Treatment strategies for patients with non-ST-segment-elevation acute coronary syndromes (NSTEACS), based on risk stratification}

\begin{itemize}
\item **High-risk NSTEACS:** Aggressive medical management and coronary angiography and revascularisation
\item **Intermediate-risk NSTEACS:** Further observation and risk stratification (see text) → Reclassification into either high risk or low risk
\item **Low-risk NSTEACS:** Discharge on upgraded medical therapy with urgent cardiac follow-up
\end{itemize}
Antithrombin therapy: Unfractionated heparin or subcutaneous enoxaparin should be given until angiography or for 48–72 hours (level I evidence, grade A recommendation). The enoxaparin dose must be reduced in patients with impaired renal function.

GP IIb/IIIa inhibitors: Intravenous tirofiban or eptifibatide is particularly recommended in high-risk patients in whom an invasive strategy is planned (level I evidence, grade A recommendation). Administration should commence as soon as a high-risk feature is identified. Intravenous tirofiban or eptifibatide are also recommended if patients continue to have ischaemia while receiving enoxaparin or unfractionated heparin (level III evidence, grade B recommendation).

Concomitant tirofiban is particularly beneficial and recommended in patients with diabetes (level I evidence, grade A recommendation).

Other: A β-blocker should be given unless contraindicated (level I evidence, grade A recommendation). Intravenous glyceryl trinitrate can be given for refractory pain (grade D recommendation).

In patients with diabetes, good glycaemic control should be targeted in hospital and after discharge. This may require considering an insulin-based regimen in hospital and for 3 or more months after discharge in selected patients (grade B recommendation). The TIMI risk score has been validated as a valuable measure of early risk in NSTEMI. It uses a seven-point score derived from:
- age greater than or equal to 65 years;
- more than three coronary risk factors;
- prior angiographic coronary obstruction;
- ST-segment deviation;
- more than two angina events within 24 hours;
- use of aspirin within 7 days; and
- elevated levels of cardiac biomarkers.

Additional risk stratification on the basis of a TIMI risk score of greater than three for deciding which patients might be transferred for early invasive management may be considered where funding is constrained, but it must be remembered that 14-day cardiac event rates are still considerable, even for those with low scores (see Box 12). Appropriate patients should be transferred for angiography within 48 hours, and aggressive medical therapy with initial stabilisation of symptoms does not mitigate the need for early angiography.

### Key Messages
- All patients with NSTEMIs should have their risk stratified to direct management decisions.
- All patients with NSTEMIs should be given aspirin unless contraindicated.
- Patients with high-risk NSTEMIs should be treated with aggressive medical management (including aspirin and clopidogrel, unfractionated heparin or subcutaneous enoxaparin, intravenous tirofiban or eptifibatide, and a β-blocker), and arrangements should be made for coronary angiography and revascularisation, except in those with severe comorbidities.
- Patients with intermediate-risk NSTEMIs should undergo an accelerated diagnostic evaluation and further assessment to allow reclassification into low-risk or high-risk categories.
- Patients with low-risk NSTEMIs, after an appropriate period of observation and assessment, may be discharged on upgraded medical therapy for urgent outpatient cardiac follow-up.
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Long-term management after control of myocardial ischaemia

Considerations before discharge

Initiating long-term therapy with a number of medications should be considered before discharge for all patients who have had an ACS (see Box 13). Other predischarge and longer term considerations are summarised in Box 14.

13 Recommended discharge medications

<table>
<thead>
<tr>
<th>Medication</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aspirin</td>
<td>All patients should take 75–150 mg daily unless contraindicated (level I evidence, grade A recommendation).</td>
</tr>
<tr>
<td>Clopidogrel</td>
<td>There is evidence that clopidogrel should be prescribed for up to 12 months after an acute coronary syndrome, in particular after stent implantation, with the duration of therapy depending on the particular type of stent and circumstances of implantation (level II evidence). Clopidogrel may also be prescribed as an alternative when aspirin is contraindicated, or in addition to aspirin, particularly in patients with unstable angina or recurrent cardiac events (level I evidence, grade A recommendation).</td>
</tr>
<tr>
<td>β-blocker</td>
<td>Should be prescribed for most patients after a myocardial infarction unless contraindicated, and continued indefinitely, especially in high-risk patients (level I evidence, grade A recommendation). Carvedilol, bisoprolol or metoprolol (extended release) should be used in patients with heart failure.</td>
</tr>
<tr>
<td>Angiotensin-converting enzyme inhibitor</td>
<td>Should be given early after an acute coronary syndrome, and its use reviewed later (level II evidence, grade B recommendation).</td>
</tr>
<tr>
<td>Statin</td>
<td>Statin therapy should be initiated in hospital for all patients with coronary heart disease (level II evidence, grade B recommendation).</td>
</tr>
<tr>
<td>Warfarin</td>
<td>Recommended after myocardial infarction for those at high risk of systemic thromboembolism because of atrial fibrillation (level I evidence, grade A recommendation), mural thrombus, congestive heart failure or previous embolisation (level III evidence, grade B recommendation). Warfarin may sometimes be combined with aspirin, but in this circumstance patients should be observed closely for signs of bleeding (grade D recommendation).</td>
</tr>
<tr>
<td>Nitrates</td>
<td>All patients should be prescribed a short-acting nitrate (unless contraindicated) and provided with a written action plan for chest pain (level III evidence, grade C recommendation) — see Box 14.</td>
</tr>
<tr>
<td>Insulin/oral hypoglycaemics</td>
<td>Good glycaemic control should be obtained and continued in patients who have had acute coronary syndromes and who have diabetes (level II evidence, grade B recommendation).</td>
</tr>
<tr>
<td>Aldosterone antagonists</td>
<td>Initiation of eplerenone therapy should be considered early after myocardial infarction in those with left-ventricular systolic dysfunction and symptoms of heart failure (level II evidence, grade B recommendation).</td>
</tr>
</tbody>
</table>

14 Other discharge and longer term considerations

<table>
<thead>
<tr>
<th>Factor</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lifestyle advice</td>
<td>All patients should be given advice on lifestyle changes that will reduce the risk of further cardiac events, including smoking cessation, good nutrition, moderate alcohol intake, regular physical activity and weight management as appropriate.</td>
</tr>
<tr>
<td>Ongoing prevention and cardiac rehabilitation programs</td>
<td>Cardiac rehabilitation is a proven effective intervention. All patients with cardiovascular disease should have access, and be actively referred, to comprehensive ongoing prevention and cardiac rehabilitation services (grade A recommendation). Specific guidelines are available for Indigenous populations.</td>
</tr>
<tr>
<td>Chest pain action plan</td>
<td>All patients should be provided with a written action plan for chest pain which includes:</td>
</tr>
<tr>
<td></td>
<td>• rest and self-administration of short-acting nitrates;</td>
</tr>
<tr>
<td></td>
<td>• self-administration of aspirin unless contraindicated (most patients should already be taking aspirin);</td>
</tr>
<tr>
<td></td>
<td>• calling an ambulance (dialling 000) if chest pain or discomfort is not completely relieved within 10 minutes; and</td>
</tr>
<tr>
<td></td>
<td>• individualised clinician notification and action plan for those living in areas where an ambulance is not readily available.</td>
</tr>
<tr>
<td>Fish oil</td>
<td>A diet high in omega-3 fatty acids from fish and the use of fish oil tablets is recommended (grade B recommendation).</td>
</tr>
<tr>
<td>Psychosocial factors</td>
<td>Depression, social isolation and lack of quality social support are likely to lead to significantly worse outcomes in those with coronary heart disease. All patients with coronary heart disease should be assessed for comorbid depression and level of social support.</td>
</tr>
<tr>
<td>Diabetes</td>
<td>An early glucose tolerance test should be considered in those without diagnosed diabetes.</td>
</tr>
<tr>
<td>Implantable cardiac defibrillators (ICDs)</td>
<td>ICDs should be considered in some patients who, despite optimal medical therapy, have persistently depressed left ventricular function after ST-segment-elevation myocardial infarction (level I evidence). See Appendix 1 for a suggested management algorithm.</td>
</tr>
</tbody>
</table>
KEY MESSAGES

- Before discharge of patients who have had an ACS, therapy with an appropriate medication regimen should be initiated, including antiplatelet agent(s), β-blocker, angiotensin-converting enzyme inhibitor, statin and other therapies as appropriate.
- Implantable cardiac defibrillators should be considered in some patients who, despite optimal medical therapy, have persistently depressed left ventricular function more than 6 weeks after STEMI.
- Patients should be given advice on lifestyle changes that will reduce the risk of further coronary heart disease events, including smoking cessation, good nutrition, moderate alcohol intake, regular physical activity and weight management, as appropriate.
- All patients should have access, and be actively referred, to comprehensive ongoing prevention and cardiac rehabilitation services.
- All patients should be provided with a written action plan for chest pain.
- Depression and coronary heart disease frequently coexist, and in patients with heart disease, depression, social isolation and lack of social support are more likely to lead to poorer outcomes. All patients with coronary heart disease should be assessed for depression and level of social support.

Conclusion

A better understanding of the pathophysiology of the acute coronary syndromes has developed, along with more accurate diagnostic tools, better risk stratification and improved medical and invasive treatments. However, these advances have led to an increase in the complexity of possible treatment strategies.

These recommendations will be regularly updated as required to provide a continuing resource to health providers. Check http://www.heartfoundation.com.au regularly for updates.

Interested individuals are invited to register with Heartline, the Heart Foundation’s national telephone information service, to receive notification of updates to these guidelines and related activities. In addition, we may contact you to invite your participation in implementation and evaluation activities. If you would like to register, please email your name to heartline@heartfoundation.com.au with “ACS” in the subject line.

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Appendices

Appendix 1: Implantable cardiac defibrillator (ICD) implantation after ST-segment-elevation myocardial infarction (STEMI): proposed management algorithm

<table>
<thead>
<tr>
<th>Path A</th>
<th>Path B*</th>
<th>Path C*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ejection fraction ≤ 30% or left ventricular ejection fraction ≤ 35% and New York Heart Association Class II or III heart failure</td>
<td>Asymptomatic patients with ejection fraction 30%–40% &gt; 30 days after STEMI</td>
<td>Ejection fraction &gt; 40%</td>
</tr>
<tr>
<td>&gt; 6 weeks after STEMI</td>
<td>Monitor</td>
<td>No ICD</td>
</tr>
<tr>
<td>(provided the patient has been appropriately revascularised)</td>
<td>Non-sustained ventricular tachycardia</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Electrophysiological studies</td>
<td>+</td>
</tr>
<tr>
<td>ICD</td>
<td>−</td>
<td></td>
</tr>
</tbody>
</table>

This algorithm is for suggested management in an area which is still evolving. There may be considerable resource issues that will need to be explored, and cost-effectiveness data are currently lacking. Other factors such as comorbidities and conditions that significantly shorten life expectancy and reduce quality of life should be considered before ICD implantation. The evidence for benefit is strongest in patients with a left-ventricular ejection fraction ≤ 30% and New York Heart Association Class II or III heart failure.

* Patients with sustained ventricular tachyarrhythmias or unexplained syncope after STEMI and an ejection fraction > 35% should also be considered for electrophysiological evaluation.

Appendix 2: Development process

The guidelines were developed on a foundation of evidence-based criteria, using a consensus approach. They are the outcome of a review of recent evidence, representations of key expert groups and stakeholders, and many meetings of writing group members during 2004 and 2005. Broad consultation was undertaken to finalise the content of these guidelines, and they have been endorsed by:

- Australasian College for Emergency Medicine
- Australian Cardiac Rehabilitation Association
- Australian Indigenous Doctors’ Association
- Australian Resuscitation Council
- Council of Ambulance Authorities
- Council of Remote Area Nurses of Australia Inc
- Internal Medicine Society of Australia and New Zealand
- Kidney Health Australia
- National Aboriginal Community Controlled Health Organisation
- Royal Australian College of General Practitioners
- Royal College of Nursing Australia
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Competing interests

The following working group members are consultants, advisory committee members, or receive honoraria, fees for service, or travel assistance (independent of research related meetings) from, or have research or other associations with the organisations listed: Roger Allan — Merck Sharp & Dohme, Sanofi, Con Aroney — CSL, Merck Sharp & Dohme, Sanofi-aventis, Phil Aylward — Sanofi-aventis, Pfizer, Merck, Bristol-Myers Squibb, Boehringer Ingelheim, AstraZeneca, Procter & Gamble, Eli Lilly, The Medicines Co, Servier, CSL, Schering Plough; David Brierre — Aventis, Sanofi, Boehringer Ingelheim, Merck Sharp & Dohme, Alex Brown — National Heart Foundation of Australia, Australian Indigenous Doctors’ Association, Alice Springs Hospital Management Board, Bristol-Myers Squibb, Pfizer, Gerard Carroll — Aventis, Bristol-Myers Squibb, AstraZeneca, Merck Sharp & Dohme, Servier, Solvay, Roche; Derek Chew — Merck Sharp & Dohme, Sanofi, Pfizer; Ian Jacobs — St John Ambulance, Australian Government Department of Health and Ageing, Convention of Ambulance Authorities Australia, National Health and Medical Research Council, Laerdal Foundation, National Heart Foundation of Australia, Health Department of Western Australia; Anne-Maree Kelly — Proctor & Gamble/Alexion, Boehringer Ingelheim; Shiong Tan — Health Department of Western Australia (Office of Safety & Quality and Sentinel event review group), Royal Australian College of General Practitioners (Quality Care National Standing Committee), National Prescribing Service (Director), Royal Australian College of General Practitioners (WA) Faculty (Director), Andrew Tonkin — AstraZeneca, Bristol-Myers Squibb, Pfizer, Sankyo, Fournier, Servier, Merck Sharp & Dohme; Warren Walsh — Roche; Harvey White — The Medicines Company, AstraZeneca, Aventis, Bayer, Boehringer Ingelheim, Eli Lilly, Merck Sharp & Dohme, Novartis, Pfizer, Roche, Servier, Wyeth Ayerst.


43 Late Assessment of Thrombolytic Efficacy (LATE) study with alteplase 6–24 hours after onset of acute myocardial infarction. Lancet 1993; 342: 759-766.


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endorsed by