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Reprint Collection

Update on stroke

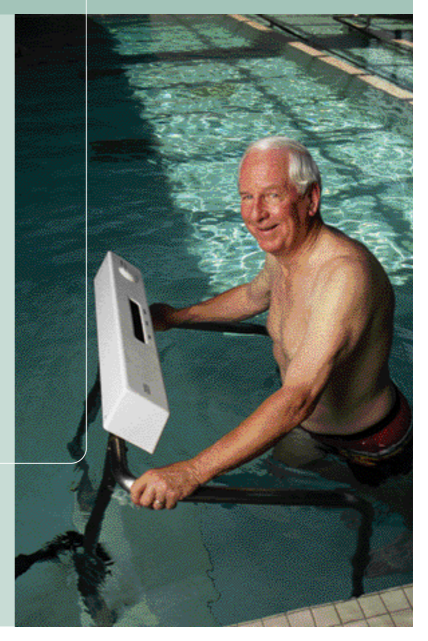
Atherothrombosis, antiplatelet therapy and ischaemic stroke prevention

Current management of atrial fibrillation

Diagnosing and evaluating patients with a transient ischaemic attack

TIA and stroke: a management guide for GPs

Managing stroke survivors in the community



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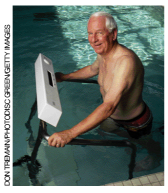
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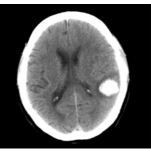
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Current management of atrial fibrillation

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A stroke can transform a person physically, socially and psychologically. The community care of stroke survivors includes promoting independent living, preventing further strokes and caring for the carers.

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Atherothrombosis, antiplatelet therapy and ischaemic stroke prevention

Strategies to prevent the development of atherosclerosis and mitigate the formation of surface thrombus on atherosclerotic plaque can potentially translate into substantial reductions in the risk of ischaemic stroke.

CHRISTOPHER R. LEVI

BMedSc, MB BS, FRACP

Dr Levi is Director of Acute Stroke Services, John Hunter Hospital, New Lambton Heights, Newcastle, and Head – Stroke Research Group, Hunter Medical Research Institute, Newcastle, NSW. Stroke is the third most common cause of death in Australia after heart disease and all cancers combined. It is the most common cause of long term disability in the adult community, and costs the nation approximately \$1.7 billion annually.

Ischaemic stroke is the major stroke subtype, comprising 85% of all strokes. The dominant underlying pathophysiology in ischaemic stroke is brain embolism originating from either atherosclerosis of the large extracranial or intracranial arteries (large artery stroke) or embolism originating from the heart (cardioembolic stroke). Atherosclerosis with thrombosis producing vessel occlusion *in situ*, particularly in the small intracranial vessels, is also common; small vessel occlusion of the

penetrating arteries supplying the basal ganglia and pons is classified as lacunar stroke. Overall, the process of atherosclerosis with thrombosis involving either large or small cerebral vessels is responsible for approximately 45% of all strokes (Figure 1).

Atherosclerosis

Atherosclerotic large vessel disease has a predilection to develop at arterial branch points where haemodynamic factors and vessel wall shear stresses predispose to intimal injury, one factor involved in the initiation of the atherosclerotic process (Figures 2a to d). With the development of regions of atherosclerosis, the resulting plaque and secondary vessel stenosis produce increasing

I SUMMARY

- Strategies to prevent the development of atherosclerosis and to mitigate the formation
 of surface thrombus on atherosclerotic plaque can potentially translate into substantial
 reductions in the risk of ischaemic stroke.
- Antiplatelet therapies, blood pressure lowering therapies, smoking cessation, cholesterol lowering therapies and carotid endarterectomy each reduce the absolute risk for stroke.
- Enhanced benefit over aspirin use can accrue from the judicious use of the newer antiplatelet agents, either alone or in combination with aspirin, both as first line and as second line therapy options.
- Clopidogrel is indicated in patients with aspirin intolerance or in whom aspirin is
 contraindicated. Cerebrovascular disease patients who have peripheral arterial disease,
 have had previous coronary artery surgery or have diabetes gain greater relative benefit
 from clopidogrel than aspirin.
- The convincing additional benefit seen with combined extended release dipyridamole
 plus aspirin over aspirin alone supports the use of this combination as first line
 antiplatelet therapy in patients who have had an ischaemic stroke or a transient
 ischaemic attack.

of flow disturbance. Disturbed flow is associated with relative platelet activation.

Many aspects of the pathology of atherosclerosis are inflammatory, with subsequent processes within the plaque leading over time to intimal damage, rupture of the luminal surface, exposure of subintimal plaque components to circulating blood and development of plaque surface thrombosis. This produces further vessel narrowing and often an intraluminal thrombus, carrying a high risk of downstream embolisation. The breakdown of plaque surface and secondary thrombosis is a dynamic process with recurrent episodes of destabilisation, intimal healing and subsequent stabilisation of plaque over time.

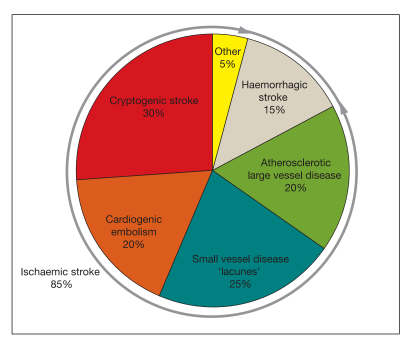
The dynamic nature of the process of embolism from carotid plaque has been demonstrated in patients with recently symptomatic carotid artery atherosclerosis, where frequent showers of cerebral microemboli can be detected with transcranial Doppler ultrasound. In asymptomatic carotid stenosis, in contrast, microemboli are detected much less often (Figure 3).2

Common sites for the development of atherothrombotic cerebrovascular disease are the extracranial carotid bifurcation, carotid siphon, vertebral artery origin, distal intracranial vertebral and proximal basilar artery.

Risk factors for atherosclerosis

Atherosclerosis is an inflammatory disease involving the intima and subintimal arterial wall. It is a common but complex disorder, showing increasing incidence with age under the influence of both genetic and environmental factors. The most important recognised modifiable environmental risk factors for atherosclerosis are those with high population prevalences, including hypertension, hyperlipidaemia, diabetes and cigarette smoking.

Risk factors for atherosclerosis induce intimal injury by many different pathways. They allow lipid deposition to occur in the subintimal zone, which stimulates secondary inflammatory effects and eventual intimal damage.3 The early intimal changes of atherosclerosis, measured as increasing intimal-media vessel wall thickness on B mode ultrasonography of the common carotid artery, can be minimised or reversed in some circumstances with appropriate management of vascular risk factors.



Plaque surface thrombosis

When atherosclerotic plaque destabilises there is exposure of subintimal components, particularly the potent prothrombotic agent, tissue factor. Tissue factor activates the extrinsic coagulation cascade through interaction with factor VII. The tissue factor:activated factor VII complex leads to the generation of thrombin, recruitment of platelets and conversion of fibrinogen to fibrin, forming a platelet rich fibrin thrombus at the plaque surface (Figure 4). This thrombus can then embolise and occlude a more distal vessel (as typically seen with atherothrombosis of the larger extracranial vessels) or result in in situ thrombotic occlusion (as typically seen in small vessel stroke syndromes).

Management of atherothrombosis

Strategies to prevent the development of atherosclerosis and to mitigate the formation of surface thrombus on atherosclerotic plaque can potentially translate into substantial reductions in the risk of ischaemic stroke. The use of antiplatelet therapy produces a 1 to 2% absolute risk reduction, thereby avoiding about 10 to 20 strokes per 1000 patients treated every year. Blood pressure lowering therapies, smoking cessation and cholesterol lowering therapies can each produce an approximate 2% absolute risk reduction, resulting in 20 to 25 strokes avoided per 1000 patients

Figure 1. Proportions of the various stroke subtypes. Atherothrombotic stroke comprises 45% of all cases of stroke, consisting of both atherosclerotic large vessel and small vessel disease.

Vessel wall changes in the development of atherosclerosis and plaque surface thrombosis*

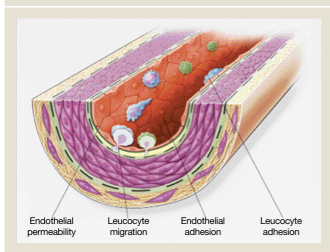


Figure 2a. Endothelial dysfunction in atherosclerosis. The earliest changes that precede the formation of lesions of atherosclerosis take place in the endothelium. These changes include increased endothelial permeability to lipoproteins and other plasma constituents, up-regulation of leucocyte adhesion molecules and endothelial adhesion molecules, and migration of leucocytes into the artery wall.

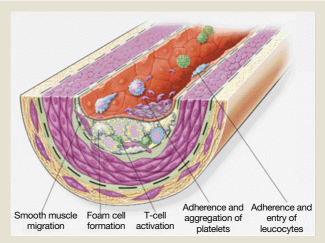


Figure 2b. Fatty streak formation in atherosclerosis. Fatty streaks initially consist of lipid laden monocytes and macrophages (foam cells) together with T-lymphocytes. Later they are joined by various numbers of smooth muscle cells. The steps involved in this process include smooth muscle migration, T-cell activation, foam cell formation and platelet adherence and aggregation.

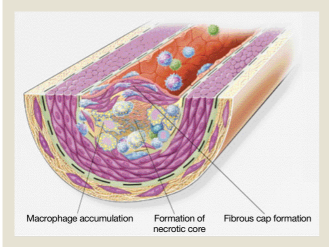


Figure 2c. Formation of an advanced, complicated lesion of atherosclerosis. As fatty streaks progress to intermediate and advanced lesions, they tend to form a fibrous cap that walls off the lesion from the lumen. This represents a type of healing or fibrous response to the injury. The fibrous cap covers a mixture of leucocytes, lipid and debris, which may form a necrotic core. These lesions expand at their shoulders by means of continued leucocyte adhesion and entry.

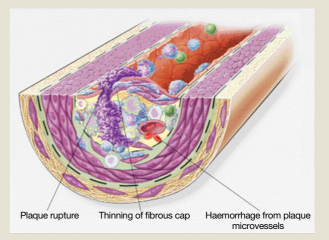


Figure 2d. Unstable fibrous plaques in atherosclerosis. Rupture of the fibrous cap or ulceration of the fibrous plaque can rapidly lead to thrombosis and usually occurs at sites of thinning of the fibrous cap that covers the advanced lesion. Thinning of the fibrous cap is apparently due to the continuing influx and activation of macrophages, which release metalloproteinases and other proteolytic enzymes at these sites. These enzymes cause degradation of the matrix, which can lead to haemorrhage from the vasa vasorum or from the lumen of the artery and can result in thrombus formation and occlusion of the artery.

^{*} FIGURES REPRODUCED COURTESY OF THE MASSACHUSETTS MEDICAL SOCIETY FROM: ROSS R. ATHEROSCLEROSIS – AN INFLAMMATORY DISEASE. N ENGL J MED 1999; 340(2): 115-126. © 1999 MASSACHUSETTS MEDICAL SOCIETY. ALL RIGHTS RESERVED.

treated each year for each intervention. Carotid endarterectomy for recently symptomatic high grade carotid artery stenosis results in a 4% absolute risk reduction, avoiding up to 40 strokes per 1000 patients treated each year.4 The benefits of these preventive therapy approaches are likely to be incremental, with relative risk reductions of up to 40% potentially achievable with combination antiatherothombotic therapies. (See the box on this page for an explanatory note on absolute and relative risk reductions.)

Antiplatelet therapies and ischaemic stroke prevention

The Antithrombotic Trialists' Collaboration meta-analysis of the use of any antiplatelet therapy following any ischaemic stroke or transient ischaemic attack (TIA) showed that antiplatelet therapy (all varieties combined) reduced the odds of subsequent adverse vascular events by 22%, with an absolute reduction of 36 major vascular events per 1000 patients treated for two years.5 The events prevented over that time frame were predominantly recurrent nonfatal strokes (25 per 1000 patients treated); however, antiplatelet therapy also prevented six nonfatal myocardial infarcts and seven vascular deaths per 1000 patients treated over the two-year timeframe.5

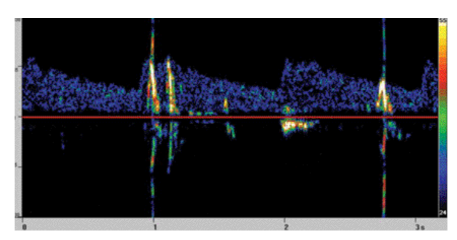


Figure 3. Asymptomatic cerebral microemboli detected as high intensity transient Doppler signals downstream from symptomatic high grade internal carotid artery stenosis.

The potential benefit of antiplatelet therapies far outweighs their risk; however, the various medications in the metaanalysis were associated with an increased risk of haemorrhagic stroke of 0.8 cases per 1000 patients treated over two years. The risk of serious extracranial bleeding (both fatal and nonfatal) was estimated at five per 1000 patients treated over two years.

With aspirin therapy per se, a more modest preventive benefit for stroke events in patients presenting with ischaemic cerebrovascular disease is evident from the meta-analysis of aspirin trials in stroke patients (a 13% relative risk reduction).6 No significant difference in net benefit was

seen in a comparison of low dose (75 mg or less) and higher doses of aspirin.5

Enhanced antiplatelet therapy in stroke prevention

Over the past decade, the following five major clinical trials have compared the use of clopidogrel (Plavix, Iscover) or extended release dipyridamole (either with aspirin [Asasantin SR] or without aspirin [Persantin SR]) with that of aspirin:

- CAPRIE Clopidogrel versus Aspirin in Patients at Risk of Ischaemic Events
- ESPS-2 the second European Stroke Prevention Study

A note on absolute and relative risk reductions

The absolute risk reduction is the difference in the outcome event rates between treatment and control groups. So, if in a study the risk (or rate) of a negative event was 10% in the control group and 8% in the treatment group, the absolute risk reduction associated with treatment would be 2%.

The relative risk reduction is the proportional difference in the outcome event rates between treatment and control groups. So, in the above example in which the control group's risk (or rate) of an event was 10% and the treatment group's risk was 8%, the relative risk reduction would be 20%. That is, the risk itself (10%) was reduced proportionally by 20% in the treatment group.

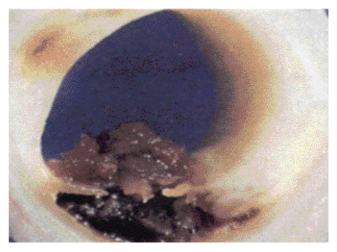


Figure 4. Plaque surface thrombosis.

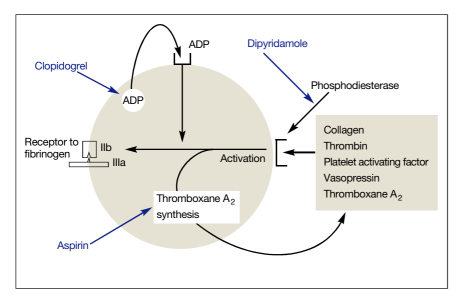


Figure 5. Pharmacological sites of action in the platelet of the antiplatelet agents approved for use in stroke prevention. Abbreviation: ADP = adenosine diphosphate.

REPRODUCED COURTESY OF PROFESSOR GEOFFREY DONNAN, NATIONAL STROKE RESEARCH INSTITUTE

- MATCH Management of ATherothrombosis with Clopidogrel in High-risk patients with recent TIA or ischaemic stroke
- CHARISMA Clopidogrel for High Atherothrombotic Risk and Ischemic Stabilization, Management and Avoidance
- ESPRIT European/Australasian Stroke Prevention in Reversible Ischaemia Trial.

The principle message from these various trials is that enhanced benefit can accrue from the judicious use of the newer antiplatelet agents, either alone or in combination with aspirin, both as first line and as second line therapy options.

The pharmacological modes of action of the newer antiplatelet agents are illustrated in Figure 5. The distinct mechanisms of action suggest that combination therapy may result in additive benefit; however, the outcomes from combination therapy trials have been variable.

Clopidogrel

In the CAPRIE trial the risks and benefits of clopidogrel 75 mg were compared head

to head with those of aspirin 325 mg daily in 19,185 patients with symptomatic atherothrombosis involving the coronary, cerebral and peripheral arterial circulations.7 Clopidogrel produced a statistically significant relative risk reduction against the comparator aspirin for the composite primary endpoint of ischaemic stroke, myocardial infarction or vascular death (8.7% relative risk reduction, p=0.043). Although the study was not powered to study the individual vascular bed outcomes within the composite endpoint, a 7.3% relative risk reduction (p=0.26) was seen for stroke alone. This trial led to the PBS approval for the use of clopidogrel in stroke in patients who are either intolerant of aspirin or have recurrent vascular events despite aspirin therapy.

A more substantial benefit on the primary end-point of the CAPRIE trial was seen in the entry subgroup with peripheral arterial disease (23.8% relative risk reduction; p=0.003). In addition, post-hoc subgroup analyses of the CAPRIE trial data suggest incremental benefit in patients with a history of diabetes and

previous coronary artery bypass grafting.

The combination of clopidogrel plus aspirin was subsequently studied in both the MATCH and CHARISMA trials. In MATCH, clopidogrel 75 mg alone was compared with aspirin 75 mg added to baseline clopidogrel within three months of stroke or TIA onset in 7599 patients with a least one major vascular risk factor.8 The primary outcome event for this trial was the composite of ischaemic stroke, myocardial infarction, vascular death or rehospitalisation for an ischaemic event. The addition of aspirin to clopidogrel resulted in a 6.4% relative risk reduction (nonsignificant; p=0.24) for the primary outcome but was associated with a significant increase in life threatening bleeding (2.6% for combination therapy versus 1.3% for clopidogrel alone; p<0.0001). The increased rates of intracerebral bleeding rates on combination therapy began to occur approximately three months after the addition of aspirin to baseline clopidogrel.

The addition of clopidogrel to baseline aspirin therapy was subsequently studied in the CHARISMA trial. This trial divided 15,603 trial entrants into an established vascular disease ('symptomatic') subgroup (12,153 patients) and a multiple risk factor ('asymptomatic') subgroup at trial entry (3284 patients). In the 'symptomatic' group, the patients with cerebrovascular disease were eligible if they had suffered a TIA or ischaemic stroke over the preceding five years. The primary endpoint was defined as myocardial infarction, stroke or vascular death.

In the overall trial population, the combination of aspirin plus clopidogrel produced a 7% relative risk reduction for the primary outcome (nonsignificant, p=0.22). Interestingly, there was significant heterogeneity between the two entry subgroups with the 'symptomatic' patients showing a 12% relative risk reduction with combination therapy (p=0.046) compared with the 'asymptomatic' multiple risk factor group showing

a 20% relative increase in the occurrence of the primary endpoint (p=0.20). A trend towards higher rates of severe bleeding was seen in the combination therapy arm of CHARISMA; however, in contradistinction to the MATCH trial, there was no significant increase in the rates of fatal and intracranial bleeding.

The data from both MATCH and CHARISMA indicate that combining clopidogrel and aspirin in patients with symptomatic vascular disease can produce an incremental benefit; however, this benefit is modest (nonsignificant in MATCH, borderline significant in CHARISMA). In addition, more prolonged combination therapy (exceeding three months in cerebrovascular patients) can be hazardous because of the significant risk of serious intracranial and extracranial bleeding. Based on the data from CHARISMA, the use of combination aspirin plus clopidogrel in patients with vascular risk factors but no symptomatic atherothrombotic disease cannot be supported.

In patients presenting with cerebrovascular disease, clopidogrel has a modest additional protective benefit over and above aspirin in the prevention of adverse vascular events. Clopidogrel is clearly indicated in patients with aspirin intolerance or in whom aspirin is contraindicated. In addition, cerebrovascular patients with peripheral arterial disease, previous coronary artery surgery and diabetes gain greater relative benefit from clopidogrel than from aspirin.

Dipyridamole extended release

The combination of extended release (ER) dipyridamole plus aspirin was compared with aspirin in ESPS-2.10 This trial used a factorial design to examine aspirin 50 mg daily, dipyridamole ER 400 mg daily, and combination therapy against single agents and also against placebo in 6602 patients who had had an ischaemic stroke or transient ischaemic attack within the preceding three months. The primary outcome for the study was the individual outcomes

of all stroke (fatal and nonfatal), all-cause death and stroke and/or death.

The combination of aspirin plus dipyridamole ER showed a 23% relative risk reduction over aspirin alone for the endpoint of stroke (p=0.006) and a 13% relative risk reduction for the endpoint of stroke and/or death. Dipyridamole ER alone was approximately equivalent to aspirin 50 mg, with a 16% relative risk reduction for stroke (p=0.04) compared with placebo. Headache and gastrointestinal symptoms were the most common adverse effects associated with dipyridamole ER requiring cessation of trial medication, occurring in 8% and 6% of patients, respectively.

The second large trial testing the combination of dipyridamole ER plus aspirin against aspirin, ESPRIT, randomised 2739 patients within six months of a transient ischaemic attack or minor ischaemic stroke of presumed arterial origin to receive either aspirin 30 to 325 mg or aspirin 30 to 325 mg plus dipyridamole ER 400 mg daily.11 The primary outcome was the composite of all vascular death, nonfatal stroke, nonfatal myocardial infarction or major bleeding complication.

Combination therapy with aspirin (mean dose 75 mg) plus dipyridamole ER produced a 20% hazard reduction (95% confidence interval [CI] 2% to 34%) for the primary composite endpoint compared with aspirin alone (mean dose 75 mg). A nonsignificant trend towards less major bleeding was seen in the combination therapy arm. Discontinuation of trial therapy occurred in 34% of patients allocated to the combination of dipyridamole ER plus aspirin, with 26% who discontinued combination therapy reporting headache as at least one reason for this.

The convincing additional benefit seen with combined dipyridamole ER plus aspirin over aspirin alone supports the use of this combination as first line antiplatelet therapy in patients with ischaemic stroke or TIA.

Summary

Antiplatelet therapy is effective in reducing the risk of thrombosis and subsequent embolism arising from atherosclerotic plaque affecting the cerebral vasculature. It forms an essential part of a package of therapy, which includes blood pressure and cholesterol lowering strategies, aimed at preventing plaque surface thrombosis, reducing plaque activity and progression and preventing rupture.

Aspirin treatment alone results in an approximate 1% absolute stroke risk reduction for each year of therapy. Enhanced benefits over aspirin alone are achieved with the use of clopidogrel in selected patients, particularly those with a history of peripheral arterial disease, and with the use of dipyridamole ER plus aspirin in those suitable for aspirin therapy. Clopidogrel produces a worthwhile additional 0.5% absolute stroke risk reduction per year of treatment (absolute stroke risk of 5.83% for aspirin v. 5.32% for clopidogrel per annum in CAPRIE). In ESPRIT, dipyridamole ER plus aspirin produced an approximate 1% per annum absolute stroke risk reduction compared with aspirin alone.

Although the absolute risk reductions seen with clopidogrel and dipyridamole ER over and above that seen with aspirin alone are relatively modest, the use of these agents does translate into worthwhile benefits given the disease burden of stroke and its financial implications. The PBS listed base dispensed prices for 12 months' therapy with these agents at recommended doses are A\$983 for clopidogrel and A\$401 for the combination of aspirin plus dipyridamole ER.

No head-to-head comparisons are presently available for clopidogrel (with or without aspirin) versus dipyridamole ER plus aspirin. However, the PROFESS trial, which is currently in follow up phase, will examine the comparative safety and efficacy of these two agents tested head-to-head in a cerebrovascular disease population.

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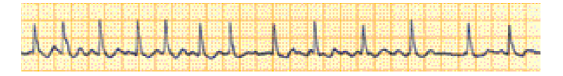
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Current management of atrial fibrillation



Rate control plus anticoagulation is the usual therapy for atrial fibrillation. A rhythm control strategy has not been shown to have prognostic benefits over rate control with anticoagulation but is useful for symptom management.

DAVID GALLAGHER

MB BS, FRACP

Dr Gallagher is a Cardiologist in private practice, Liverpool, and Visiting Medical Officer, Liverpool Hospital, Liverpool, NSW.

Atrial fibrillation is the most common arrhythmia encountered in clinical practice. Historically associated with rheumatic valvular heart disease, the condition is now associated with ageing, and hypertension and is becoming more prevalent in Western society. Current prevalence ranges from less than 1% in people under 60 years of age to over 6% in those over 80 years. Morbidity relates to reduced left ventricular function and thromboembolic complications.

Treatment strategies have focused on minimising cardiac embolism complications and preserving left ventricular function. GPs are involved in the initial management of patients presenting with atrial fibrillation, and in the ongoing management after specialist review.

Aetiology

Patterns of atrial activation in atrial fibrillation are thought to be due to multiple wavelet re-entry or increased focal activity. In some cases, the re-entrant circuit may arise from the region of pulmonary vein insertion. Anatomical and physiological processes that can promote electrophysiological changes favouring atrial fibrillation include atrial dilatation, ischaemia, fibrosis and altered autonomic innervation.

Given the variety of anatomical and physiological changes promoting the condition, it is not surprising that atrial fibrillation is associated with a large number of conditions and may be precipitated by others. These conditions are listed in Table 1.

- Most patients with atrial fibrillation are elderly or have hypertension and are at high risk
- The goals of management include control of symptoms, preservation of left ventricular function and prevention of cardiac embolism.
- Rate control and anticoagulant therapy can be initiated in stable patients by the GP while awaiting specialist review.
- Anticoagulation is underutilised in high risk groups.
- A patient's therapy should be re-evaluated over time because both the risk of embolism and the incidence of conduction disease increase with age.

Table 1. Conditions related to atrial fibrillation

Cardiac causes

Hypertension

Valvular heart disease

Coronary artery disease

Chronic heart failure

Cardiomyopathy

Pericardial disease

Cardiac tumours

Sinus node disease

Familial

Cardiothoracic surgery

Congenital heart disease

Noncardiac causes and precipitants

Autonomic dysfunction

(sympathetic/parasympathetic)

Toxins (e.g. alcohol)

Thyrotoxicosis

Pulmonary disease, hypoxia

Neurological disorders

Idiopathic atrial fibrillation

Sepsis

Exertion

Table 2. Classification of atrial fibrillation for management guidance*

Acute atrial fibrillation

- Reversible extrinsic cause
- First episode

Chronic atrial fibrillation

- Paroxysmal (self-terminating within 7 days, most cases within 24 hours)
- Sustained (lasting more than 7 days):
 - persistent (successful cardioversion)
 - permanent (unsuccessful or not attempted cardioversion)
- * Patients may move between categories with treatment and passage of time.

Classification

A descriptive clinical classification has been proposed by the American Heart Association and the European Society of Cardiology. There appears to be value in determining whether an episode of atrial fibrillation is first detected or recurrent (more than one episode) and furthermore

Initial approach to atrial fibrillation

History

- · New, recurrent, longstanding
- Symptoms
- · Reversible conditions, precipitants
- Past response to treatment

Examination

- Blood pressure
- Cardiac failure

ECG

- · Confirm diagnosis
- Assess for pre-excitation, ventricular response, QT interval, LVH, ischaemic changes

Holter monitor

 May be needed to confirm diagnosis in paroxysmal atrial fibrillation

Blood tests

- Electrolytes (including potassium), urea, creatinine, magnesium
- Thyroid stimulating hormone (TSH)

Chest x-ray

 To assess cardiac failure and pulmonary disease

Echocardiogram

 To assess valvular or myopathic disease and LV function

Further investigations

• As required

Initial management

- · Referral: urgent or elective
- Rate control: digitalis, β-blockers, calcium channel blockers
- Anticoagulation: warfarin

whether the episode is paroxysmal, persistent or permanent. These latter definitions are essentially based on time. Paroxysmal episodes resolve within seven days, most within 24 hours; persistent episodes last more than seven days and generally require cardioversion to return to sinus rhythm. Patients classified as having permanent atrial fibrillation have had the arrhythmia for over one year; this includes people who have failed cardioversion and those in whom cardioversion has not been attempted.

Atrial fibrillation may present as an acute episode or as chronic condition (Table 2). The acute episode may be a transient occurrence related to a reversible extrinsic cause or a first episode of atrial fibrillation. Chronic atrial fibrillation may be subclassified into paroxysmal (not sustained) or sustained forms. Sustained atrial fibrillation may be further divided into persistent (successfully cardioverted) or permanent (unsuccessful or not attempted cardioversion).

These classifications are used to guide management and are not static in that patients may move between categories depending on response to treatment and passage of time.

Clinical presentation

Patients with atrial fibrillation usually present with symptoms directly related to the arrhythmia, such as palpitations, fatigue and dyspnoea. Less commonly, the arrhythmia may exacerbate the symptoms of underlying conditions such as congestive cardiac failure and ischaemic heart disease, and the presenting symptoms may be suggestive of heart failure and angina. Syncope is a less common presentation that may be associated with sinus node disease (sick sinus syndrome) and structural conditions such as severe aortic stenosis or hypertrophic obstructive cardiomyopathy. Another presentation may be that of a thromboembolic episode (stroke or systemic emboli).

Not infrequently atrial fibrillation is





Figure 1 (left). Rhythm strip of atrial flutter with 4:1 block. Flutter rate is 300 per minute with ventricular rate 75 beats per minute. Atrial flutter with block at this rate is usually variable.

Figure 2 (above). Rhythm strip of polymorphic ventricular tachycardia. Proarrhythmia is a potential complication of antiarrhythmic therapy.

identified incidentally in asymptomatic patients.

Diagnosis and assessment History

The history should attempt to identify symptoms associated with the arrhythmia, the pattern and duration of the arrhythmia, and any precipitating factors, in particular alcohol or events that increase adrenergic or vagal tone (such as exercise and a vasovagal episode, respectively).

The presence of reversible conditions (such as hyperthyroidism) should be identified. The presence of underlying heart disease (valvular, ischaemic or myopathic) should also be identified, as should associated conditions such as hypertension, diabetes and previous cerebrovascular or systemic embolic episodes. A checklist for GPs is given in the box on page 10.

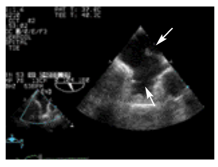
Physical examination

Physical examination includes assessment of ventricular response (apical and radial), blood pressure and signs of heart failure, valvular heart disease, chronic airways disease and thyrotoxicosis.

When to refer

Urgent referral for emergency treatment, including cardioversion, should be considered in patients presenting with atrial fibrillation and recent or ongoing symptoms of collapse, chest pain and dyspnoea





or signs of cardiac failure. A patient with a more distant history of similar symptoms and paroxysmal arrhythmia should be referred for urgent specialist review.

Stable patients with atrial fibrillation may be initiated on treatment (rate control and anticoagulant therapy) while awaiting specialist review.

Investigations

Initial investigations include a 12-lead ECG to confirm arrhythmia and assess for left ventricular hypertrophy (LVH), conduction disease or prior myocardial infarction, and to serve as a baseline



Figure 3 (above left). Transthoracic echocardiogram (TTE) showing the biatrial dilatation (arrows) seen in established atrial fibrillation.

Figure 4 (above). TTE (apical four-chamber view) showing concentric left ventricular hypertrophy (arrow) in a hypertensive patient. Hypertension is commonly associated with atrial fibrillation.

Figure 5 (left). Transoesophageal echocardiogram (TOE) showing a left atrial thrombus (arrow at 1 o'clock) protruding over the left apical appendage (arrow at 5 o'clock).

before antiarrhythmic therapy (RR and QT intervals) (Figures 1 and 2). Holter monitoring may be used to confirm the diagnosis in patients with a history of paroxysmal symptoms.

A chest x-ray may be performed to assess for pulmonary disease or heart failure if clinically suspected.

An echocardiogram should be performed to identify valvular heart disease, left ventricular function, LVH, atrial

dimensions and pericardial disease (Figures 3 to 5). Transthoracic echocardiograms have a low sensitivity in detecting left atrial thrombus and should not be relied upon for that purpose.

Thyroid function studies should also be performed, as should baseline electrolytes, urea and creatinine before commencing antiarrhythmic drug therapy or digitalis.

Further investigations that may be carried out include stress testing and electrophysiological studies.

Treatment

The goals of therapy are to minimise symptoms of atrial fibrillation, maintain exercise capacity, prevent thromboembolic episodes and preserve left ventricular function. The two treatment strategies available are:

- return to and maintenance of sinus rhythm, using cardioversion or antiarrhythmic drugs, or both
- rate control therapy with anticoagulation, in individuals at high risk of thromboembolic complications.

The choice of strategy depends on several factors. It can usually be based on classification of the arrhythmia according to pattern (e.g. acute or chronic). However, in each individual case, the symptoms associated with the atrial fibrillation, the patient's age and activity level, and the underlying conditions need to be considered. A general strategy according to classification is as follows:

- acute atrial fibrillation identify any transient or reversible causes and treat these; if no reversible or transient cause can be found the patient should be cardioverted
- chronic paroxysmal atrial fibrillation

 prevent paroxysmal episodes and maintain sinus rhythm, or minimise symptoms during episodes with rate control therapy
- chronic sustained atrial fibrillation either restore and maintain sinus rhythm or control the rate and

anticoagulate; in patients with longstanding atrial fibrillation the chance of successful cardioversion and rhythm maintenance diminish with duration of atrial fibrillation.

One goal common to all groups is the minimisation of thromboembolic events. Patients with paroxysmal or persistent atrial fibrillation should be assessed for thromboembolic risk and anticoagulated with warfarin (Coumadin, Marevan) if appropriate.

Rate control

The primary aims of rate control in atrial fibrillation are to improve symptoms and effort tolerance and maintain or improve left ventricular function. Treatment targets are a resting heart rate of 60 to 80 beats per minute and an exercise heart rate of below 120 beats per minute during moderate exercise.

Pharmacological options for achieving rate control include:

- digoxin (Lanoxin, Sigmaxin)
- nondihydropyridine calcium channel antagonists (class IV antiarrhythmic agents) – diltiazem, verapamil
- β-blockers (class II antiarrhythmic agents) esmolol (Brevibloc), metoprolol, pindolol (Barbloc, Visken), propranolol (Deralin, Inderal).

The choice of rate control agent depends on comorbidities and the clinical setting, particularly the age and activity level of the patient and the presence or absence of conditions such as coronary artery disease, heart failure, hypertension, asthma and renal impairment.

Amiodarone (a class III antiarrhythmic agent) may also be used for rate control but concerns over long term toxicity tend to limit its use for long term rate control. Digoxin, traditionally used for the control of ventricular response, is relatively ineffective at controlling ventricular response associated with increased adrenergic activity (such as exercise) and, therefore, a β -blocker may be more effective for younger, more active subjects.

Therapy should be titrated to achieve symptom control, and Holter monitoring, exercise stress testing or both may be used to assess the effectiveness of treatment. Often combinations of treatment are required for control, such as β -blockers plus digitalis, β -blockers plus calcium channel antagonists, and calcium channel antagonists plus digoxin. Exercise stress testing and Holter monitoring can identify excessive slowing of ventricular response.

In patients intolerant of pharmacotherapy or those in whom ventricular rate is inadequately controlled despite maximal pharmacotherapy, catheter ablation of the His bundle and pacemaker insertion may be required. Pacemaker insertion may also be required for patients with coexistent conduction disease and episodes of symptomatic bradycardia.

Anticoagulation

All patients presenting with atrial fibrillation should be assessed as to their risk of systemic embolism and cardioembolic stroke. Clinical criteria derived from anticoagulation and epidemiological studies are used to estimate an individual's stroke risk (Table 3). The yearly risk of stroke in a patient with nonvalvular atrial fibrillation is classified as low (1%), moderate (2 to 5%) or high (more than 5%). Anticoagulation therapy with warfarin is recommended in patients with moderate or high risk of stroke and reduces the stroke rate to 1 to 2%.

The decision to anticoagulate involves a balance between the risk of stroke and the risk of complications from anticoagulation. Unfortunately, many patients at high risk from cardioembolic stroke are also at higher risk of complications from warfarin

The risk of cardioembolic stroke varies inversely with the international normalised ratio (INR) level, with the lowest stroke rates recorded in trials treating to INR 2.5 to 3.0. Lower INR values have been proposed for elderly patients (INR,

2.0) in an attempt to reduce warfarin complications. Higher stroke rates have been reported in these trials, probably reflecting periods of subtherapeutic anticoagulation when the lower target is used.

Aspirin provides partial reduction in stroke risk in patients with atrial fibrillation, probably due to reduction in atherothrombotic stroke. It is not considered an alternative to warfarin in moderate and high risk groups. It may be considered in low risk groups or in intermediate risk groups who are noncompliant or who have a major contraindication to warfarin.

Nonpharmacological techniques aimed at reducing embolism risk, including left atrial appendage exclusion or amputation via surgery or percutaneously implanted device (PLAATO) and the Maze procedure, are currently being explored. The efficacy of these procedures is not yet determined.

Rhythm control

Maintenance of sinus rhythm has been advocated as a means of preserving left ventricular function, reducing the need for anticoagulation and thereby reducing morbidity. A fairly aggressive approach is taken, several cardioversions and a variety of antiarrhythmic agents often being required, and frequently there is no ultimate success. Against the putative benefits of maintaining sinus rhythm weigh the real risks of morbidity related to use of antiarrhythmic agents. In addition, there remains a risk of cardiac thromboembolism from paroxysmal atrial fibrillation that may be asymptomatic.

Two recent trials have looked at rhythm maintenance versus rate control and anticoagulation strategies (the Atrial Fibrillation Follow-up Investigation of Rhythm Management [AFFIRM] and the Rate Control versus Electrical Cardioversion for Persistent Atrial Fibrillation [RACE] studies).2,3 These trials enrolled elderly cohorts with chronic atrial fibrillation. Both showed no improvement in morbidity using rhythm control over rate control, with the AFFIRM study suggesting a trend towards increased mortality in the rhythm control group. Stroke rates were similar in both cohorts (1%), and most strokes occurred in patients who were not on warfarin or were subtherapeutically anticoagulated.

These studies, therefore, did not support the use of rhythm control as a means of reducing morbidity and cerebrovascular events in this group, which form the bulk of patients with atrial fibrillation. However, a rhythm maintenance strategy may still be valid for younger patients and those who are symptomatic from atrial fibrillation.

Cardioversion

Cardioversion to sinus rhythm may be attempted electrically, using synchronised direct current shock, or pharmacologically. The most commonly used pharmacological agents are the class III antiarrhythmics sotalol and amiodarone, although the former is not thought to be particularly effective in achieving cardioversion but is thought effective in preventing recurrence of atrial fibrillation. The fairly high rate of spontaneous reversion confounds studies of drug effectiveness in acute reversion. Class 1a

and 1c agents (such as procainamide [Pronestyl] and flecainide [Flecatab, Tambocor], respectively) are also effective but are less commonly used. Pharmacological agents and electrical cardioversion may be used together to increase initial success rates and maintenance rates, particularly in patients who have previously failed cardioversion.

The success of cardioversion, both initially and in the long term, is inversely proportional to the preceding duration of atrial fibrillation. Increased left atrial size (greater than 50 mm diameter) is associated with poor outcome, as is underlying valvular disease or left ventricular dysfunction. Up to 50% of cases of paroxysmal atrial fibrillation revert spontaneously within 72 hours, and success at maintaining sinus rhythm following cardioversion in patients with persistent atrial fibrillation is less than 50%.

The timing of cardioversion is based on clinical stability and requirement for cardioversion. Emergency cardioversion is required in patients with haemodynamic instability, particularly in the setting of Wolff-Parkinson-White syndrome. In this condition, atrial impulses are conducted anterogradely down the accessory pathway, resulting in very rapid

Table 3. Stroke risk and anticoagulation in atrial fibrillation

Patient age group	Stroke risk factor status*	Risk of stroke	Anticoagulation therapy
Less than 65 years	Positive	High (>5%)	Warfarin
	Negative	Low (<1%)	?Aspirin
65 to 75 years	Positive	High	Warfarin
	Negative	Moderate (2 to 5%)	Warfarin (?aspirin)
More than 75 years	Positive	High	Warfarin
	Negative	Moderate to high	Warfarin

^{*} The presence of any one of the following stroke risk factors confers a positive stroke risk factor status: previous cerebrovascular accident, left ventricular dysfunction, hypertension, diabetes mellitus, coronary artery disease, thyrotoxicosis.

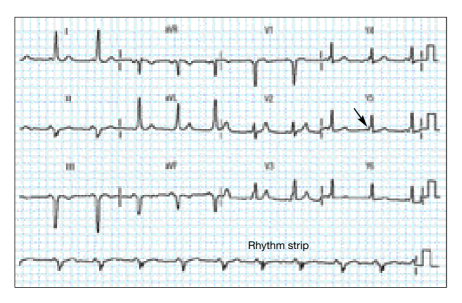


Figure 6. 12-lead ECG showing pre-excitation, seen as short PR intervals and delta waves (marked by arrow in lead where they are most clearly shown), in a patient with Wolff-Parkinson-White syndrome.

Table 4. Anticoagulation in cardioversion of atrial fibrillation

Before cardioversion

- Patients with arrhythmia duration less than 48 hours:
 - anticoagulation generally not required but heparin may be used if necessary
- Patients with arrhythmia duration more than 48 hours:
 - anticoagulation with warfarin (INR, 2 to 3) for three to four weeks
 - prolonged preceding anticoagulation not necessary if transoesophageal echocardiogram negative for intra-atrial thrombus (cardioversion is performed while therapeutically anticoagulated with either heparin or warfarin)

After successful cardioversion

- Patients with arrhythmia duration less than 48 hours:
 - anticoagulation generally not required
- Patients with arrhythmia duration more than 48 hours (with or without left atrial thrombus):
 - anticoagulation mandated, warfarin (INR, 2 to 3) for at least four weeks

ventricular response and possible ventricular fibrillation. Atrioventricular (AV) node blocking agents (digoxin and calcium channel antagonists) are contraindicated in this setting because the increased AV block enhances conduction down the accessory pathway. When in sinus rhythm, pre-excitation is often evident on the patient's ECG (Figure 6).

Anticoagulation before and after cardioversion

Anticoagulation is thought not to be required for patients with atrial fibrillation of less than 48 hours' duration (Table 4). Patients with atrial fibrillation of less than 48 hours' duration appear to have a low risk of atrial stunning (the delayed return of atrial contractile function following

atrial fibrillation), and prolonged anticoagulation after cardioversion appears not to be necessary in this group.

Anticoagulation is, however, required to minimise thromboembolic events before cardioversion in patients with structurally normal hearts and atrial fibrillation of more than 48 hours' duration. This anticoagulation should be for three to four weeks. Cardioversion may be performed earlier if transoesophageal echocardiogram (TOE) excludes intraatrial thrombus. Both early (TOE-guided) and delayed cardioversion have similar complication rates, and the choice of approach is determined by the patient's symptoms and the available resources. Heparin is used as the anticoagulant in early TOE-guided cardioversion.

Anticoagulation should be continued for at least four weeks following cardioversion of patients with atrial fibrillation of more than 48 hours' duration as the thromboembolic risk remains high in this period (thought to be because of atrial stunning). Embolism can occur after successful cardioversion in patients without thrombus on TOE before cardioversion (presumably due to stunning) and, therefore, anticoagulation is indicated in these patients following cardioversion. One month of anticoagulation following cardioversion should be considered a minimum, and anticoagulation may be continued indefinitely in patients at high risk of cardiac embolism from recurrent atrial fibrillation.

Maintenance of sinus rhythm

Pharmacological agents used for maintenance of rhythm control include sotalol, flecainide and amiodarone. These agents have varying side effect profiles and incidences of proarrhythmia. The choice of agent depends on the patient's age, associated cardiac disease and comorbidities.

Monitoring patients on long term antiarrhythmics

Sotalol has β-blockade activity and while

useful in patients with coronary disease may exacerbate conduction disease or airways disease. Patients may also complain of fatigue and exertional dyspnoea. Sotalol has a small incidence of proarrhythmia (prolonged QT syndrome), and should, therefore, be administered incrementally at low dose, with ECGs at baseline and several days after each dose increase. It is renally excreted. Drug levels and renal function should be monitored and it is best not used in patients with renal impairment.

Flecainide, although effective at reverting and maintaining sinus rhythm, is also associated with proarrhythmia and is contraindicated in patients with left ventricular dysfunction or ischaemic heart disease.

Amiodarone has a lower proarrhythmic profile than sotalol and flecainide and is generally well tolerated. Over the longer term, however, organ toxicity is significant (thyroid, liver, lung, brain). Although lower doses tend to minimise this, the effects of long term (more than five years) administration are uncertain and these agents tend not to be used long term in younger patients. Regular monitoring is required, including thyroid function studies, liver function studies (six-monthly) and chest x-ray (yearly).

Curative catheter ablation

Nonpharmacological procedures for rhythm maintenance include surgical atrial interruption procedures such as the Maze procedure, and radiofrequency catheter ablation techniques directed to pulmonary vein isolation. These latter procedures appear to have a particular role in patients with idiopathic paroxysmal atrial fibrillation. The role of these procedures in reducing systemic embolism risk has not been determined.

Summary

Atrial fibrillation is increasing in prevalence in the community and is associated with ageing, hypertension and left ventricular dysfunction. Most patients presenting with this condition will require a rate control and anticoagulation strategy. At present a rhythm control strategy has not proven superior to rate control and anticoagulation with respect to morbidity relating to reduced left ventricular function and thromboembolic complications, but is still indicated for symptomatic management of the condition.

Ongoing morbidity and inconvenience associated with warfarin has stimulated the development of newer antithrombin agents. Curative catheter based strategies for atrial fibrillation are evolving and may have applicability in the subgroup of patients with idiopathic paroxysmal atrial fibrillation.

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DECLARATION OF INTEREST: None.



Diagnosing and evaluating patients with a transient ischaemic attack

In this series, we present authoritative advice on the investigation of a common clinical problem, specially commissioned for family doctors by the Board of Continuing Medical Education of the Royal Australasian College of Physicians.

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Series Editor: Dr Pokorny is a member of the Board of Continuing Education, Royal Australasian College of Physicians, and a Gastroenterologist in private practice, Sydney, NSW. A transient ischaemic attack (TIA) is not a benign event and should under some circumstances be treated as a medical emergency. Failure to adequately recognise and evaluate this warning sign could mean missing an opportunity to prevent permanent disability or death. The risk of stroke after a TIA is up to 10% at seven days and 15% at 30 days, with 50% of strokes occurring in the first 48 hours. Peccent evidence shows that by using a simple scoring scheme (the ABCD² score), those patients at high risk of stroke can be identified and should be referred for emergency assessment.

A case study

A 69-year-old man, Mr L, consults his GP one hour after recovering from a 45-minute episode of difficulty speaking and weakness of the right side of his face and right arm. He has a history of hypertension and hypercholesterolaemia.

What is the appropriate management of this patient?

Definition and clinical presentation

A TIA is a clinical syndrome characterised by the sudden onset of focal neurological symptoms and/or signs that last less than 24 hours. These symptoms or signs are presumed to have been caused by focal cerebral or retinal ischaemia as a result of either arterial thrombosis or embolism associated with arterial, cardiac or haematological disease.

The vast majority of TIAs are brief, with symptoms lasting less than one hour. Focal neurological and ocular symptoms and signs common in TIA are outlined in Table 1. In general, a TIA presents as a syndrome rather than any one sign or symptom.

About 15 to 20% of patients who have a stroke report a preceding TIA, and a similar proportion

N SUMMARY

- A transient ischaemic attack (TIA) is a clinical syndrome characterised by the sudden onset of focal neurological symptoms and/or signs that last less than 24 hours.
- A focused and prompt history and physical examination are important for all patients with acute transient neurological symptoms to confirm the diagnosis of the cerebral ischaemic syndrome and to localise the vascular territory affected.
- All patients with an ABCD² score of 4 or greater should be sent to an emergency department for urgent evaluation; those with a score of less than 4 can complete their investigations as outpatients.
- Failure to adequately recognise and evaluate a TIA could mean missing an opportunity to prevent permanent disability or death.

have a preceding minor stroke.1 These 'warning' events provide an opportunity for therapeutic intervention. A recent study shows that urgent assessment of patients in a rapid response TIA clinic and early initiation of a combination of existing preventative treatments can reduce the risk of early recurrent stroke by about 80%.4

Primary care (pre-emergency department)

For all patients with acute transient neurological symptoms it is important to take a focused and prompt history and perform a physical examination to confirm the diagnosis of the cerebral ischaemic syndrome and localise the vascular territory affected. This can be important in directing investigations and determining the potential cause of the TIA.

A TIA affecting the carotid artery territory usually results in contralateral motor, sensory and higher cortical symptoms. Amaurosis fugax is strongly associated with internal carotid artery stenosis, because the ophthalmic artery is the first branch of the internal carotid artery. A TIA affecting the posterior circulation can manifest in a multitude of symptoms and signs, including vertigo, imbalance, ataxia and cranial nerve dysfunction (see Table 1).

All patients with ongoing symptoms suggestive of cerebral ischaemia should be immediately sent to a hospital emergency department via ambulance since there is no reliable way to determine if the neurological deficits represent ischaemia without subsequent brain damage (TIA) or if ischaemia will result in permanent damage to the brain (stroke). Patients presenting to the emergency department within three hours of onset of symptoms may be considered for thrombolytic therapy in centres capable of administering this treatment. More than 50% of patients with symptoms lasting longer than one hour but less than 24 hours (i.e. clinically classified as a TIA) will have imaging evidence of infarction if acute MRI scanning is performed.

Differential diagnosis

The diagnosis of a TIA can be problematic because it is often based on clinical history alone and specifically on the recollection of the patient who was neurologically impaired during the event. The

Investigating transient ischaemic attacks



The short-term risk of stroke in patients after a transient ischaemic attack (TIA) is high and relatively predictable. Identification of TIAs thus provides an important opportunity for the clinician to intervene and prevent strokes. For all patients who have had acute transient neurological symptoms, a focused and prompt history should be taken and physical examination performed to confirm the diagnosis of cerebral ischaemic syndrome and localise the vascular territory affected.

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differential diagnosis of TIA is wide and includes metabolic, structural and vascular disorders.

The most common mimics of TIA-like symptoms are:

- seizure with Todd's paralysis
- migraine headache with aura
- glucose derangements
- global cerebral ischaemia from hypotension

Table 1. Focal neurological and ocular symptoms and signs common in TIAs				
Affected function	Symptoms and signs	Localisation		
Motor function	Weakness or paralysis of one side of the body, in whole or in part (hemiparesis) Simultaneous bilateral weakness Difficulty swallowing (dysphagia)	Carotid artery territory, posterior circulation Brainstem Brainstem		
Co-ordination	Imbalance (ataxia) with standing or walking Clumsy arms or legs	Cerebellum, brainstem Cerebellum, brainstem		
Sensory function	Altered feeling on one side of the body, in whole or in part (hemisensory disturbance)	Carotid artery territory, posterior circulation		
Vestibular function	Spinning sensation (vertigo). Isolated vertigo is more likely to be due to a peripheral (labyrinthine) cause	Posterior circulation		
Vision	Visual loss in one (amaurosis fugax) or both eyes. Bilateral loss may indicate onset of basilar ischaemia Loss of vision in the left or the right half or quarter of the visual field (hemianopia, quadrantanopia) Double vision (diplopia)	Retinal artery Carotid artery territory, posterior circulation Brainstem		
Speech/language	Difficulty comprehending, pronouncing or 'finding' words (aphasia), difficulty reading (alexia) and writing (agraphia) Slurred speech (dysarthria)	Left carotid artery territory Brainstem, carotid artery territory		
Higher cortical function	Difficulty dressing or using tools – e.g. combing hair, cleaning teeth (apraxia) Visual–spatial–perceptual dysfunction Inattention to surrounding environment on one side (hemineglect)	Carotid artery territory Carotid artery territory Carotid artery territory (usually right)		

• structural intracranial abnormalities (especially with acute haemorrhage).

A TIA is typically rapid in onset, and maximum intensity is usually reached within one minute. Distinguishing a TIA from migraine aura can be difficult. In general, migraine aura tends to have a more gradual onset and resolution and a marching quality. For example, symptoms such as tingling may progress from the finger to the forearm to the face within minutes. Associated headache, nausea and photophobia with a previous history of migraine are more suggestive of migraine than TIA.

In the absence of other focal neurological symptoms, loss of consciousness, fatigue or light-headedness is unlikely to represent TIA. Syncope and seizures should be considered in patients with loss of consciousness.

TIA referral guidelines using the ABCD² score

If the clinical features of a TIA have resolved by the time the patient is first seen, the seven-point ABCD² score is a validated model to predict the early risk of stroke after this event (Table 2).³ This can help the front-line physician identify which patients with a suspected TIA should be referred for emergency assessment.

In a cohort of 4799 patients with TIA assessed with the ABCD² scoring system, the two-day stroke risk was 1% in patients who had a score of 0 to 3 (low risk), 4.1% in those with a score of 4 or 5 (moderate risk) and 8.1% in those with

a score of 6 or 7 (high risk).3

Patients with a score of 4 or greater should be sent to an emergency department or to a rapid referral TIA clinic for urgent assessment. Those classified as being at high risk are likely to benefit from urgent inpatient evaluation, treatment and observation, particularly in view of the high social and economic costs of stroke. On the other hand, most patients with a score of less than 4 will not need hospital observation. Ideally, these patients should be assessed at a routine stroke/TIA clinic or by a stroke physician, preferably within a week.

Initial investigations

Initial investigations are often carried out at the emergency department.

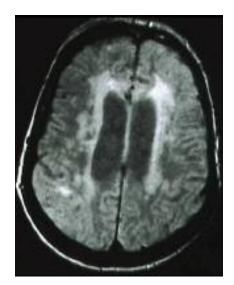


Figure. FLAIR MRI image showing white matter changes with acute cerebral infarction in the right posterior parietal lobe.

Laboratory tests

A complete blood count with platelet count should be obtained to rule out polycythaemia, thrombocytopenia, and thrombocytosis.

It is important to determine the prothrombin time (PT), activated partial thromboplastin time (aPTT) and international normalised ratio (INR) before antiplatelet or anticoagulation therapy is administered. The PT and aPTT can be prolonged and INR elevated in hypercoagulable states.

The glucose level should be determined to rule out hypo- or hyperglycaemia.

Creatinine levels are relevant, because poor renal function may prohibit the use of contrast media in imaging.

An elevated erythrocyte sedimentation rate may indicate rare causes of cerebral ischaemia such as bacterial endocarditis or temporal arteritis.

Electrocardiography and echocardiography

A 12-lead electrocardiogram (ECG) is important for detecting atrial fibrillation as well as concurrent cardiovascular disease. If the ECG is unrevealing, cardiac

	Risk	Score
Α	Age 60 years or greater	1
В	Blood pressure: systolic greater than 140 mmHg and/or 1 diastolic 90 mmHg or greater	
С	Clinical features:	
	Unilateral weakness	2
	Speech disturbance without weakness	1
	• Other	0
D	Duration of symptoms:	
	• 60 minutes or more	2
	• 10 to 59 minutes	1
	Less than 10 minutes	0
D	Diabetes mellitus	1

Patients who score 4 or greater should be sent immediately to the emergency department; those scoring less than 4 should be assessed, preferably within a week, at a routine stroke/TIA clinic or by a stroke physician.

monitoring may help diagnose paroxysmal atrial fibrillation. Echocardiography may identify a thromboembolic source or left ventricular systolic dysfunction, both of which are predictors of ischaemic stroke.

Transoesophageal echocardiography is superior to transthoracic echocardiography for evaluating dysfunction of the left atrium, intracardiac thrombus, patent foramen ovale and atrial septal defects, and aortic arch plaque. This is usually performed in an elective setting.

Brain imaging

All patients with a suspected TIA should undergo cranial imaging by CT or MRI on the same day as their presentation to rule out structural causes, such as intracerebral haemorrhage, subdural haematoma or brain tumour (Figure). Urgent identification of these conditions is critical because special management or neurosurgical intervention may be required. A head CT scan can also identify signs of early infarction or evidence of previous strokes. MRI is more sensitive than CT for detec-

tion of early ischaemia, especially in the brain stem because of increased bony artefact in the posterior fossa affecting CT.

Neurovascular imaging

Imaging of the extracranial circulation is an important component of the diagnostic workup. Patients with anterior circulation ischaemia should have carotid imaging, usually ultrasonography, within 24 hours. The presence or absence of a neck bruit on clinical examination should not alter the decision to image the carotid bifurcation, since this finding is neither sensitive nor specific.

The discovery of severe carotid stenosis (greater than 70%) in patients who present after a TIA is an important finding because there is good evidence that endarterectomy is superior to medical management in this group of patients. Endarterectomy is most beneficial if performed within two weeks of the patient's last symptoms.⁵

Carotid ultrasonography is a good screening tool, but may not provide enough information on its own for surgical decision making.⁶

Often CT angiography or MR angiography and occasionally conventional catheter angiography are used to confirm the ultrasound findings in patients with estimated stenosis of more than 70% before proceeding to surgery. CT angiography or MR angiography of the vertebral and intracranial arteries may reveal stenosis or dissection. Evaluation of the intracranial cerebral circulation may be important in selected patients, but in general it is unlikely to alter management decisions for the patient presenting with a TIA. Catheter angiography should be reserved for patients being considered for surgery or in whom diagnostic uncertainty remains after noninvasive imaging.

Case continued

Mr L had an ABCD² score of 5 and was referred to the emergency department by his GP. He was found have normal sinus rhythm, a blood pressure of 150/85 mmHg, normal routine blood tests and a normal head CT scan. Carotid duplex ultrasono graphy revealed a 90% stenosis of the left internal carotid artery.

Aspirin therapy was commenced, and an uncomplicated endarterectomy was performed three days later.

At follow-up four weeks later he was found to be hypertensive, and his LDLcholesterol level was high. He was started on an ACE inhibitor with a diuretic and a cholesterol-lowering agent, and aspirin was changed to combination aspirin/ dipyridamole therapy.

Conclusion

The short-term risk of stroke in patients after a TIA is high and relatively predictable. Emergency evaluation and treatment in patients such as the one described in the clinical vignette is justified.

TIA provides an important opportunity for the clinician to intervene and prevent stroke. Effective interventions for secondary stroke prevention exist but need to be implemented rapidly to be effective. The most important step in the prevention of disability from stroke is the early recognition of TIA by patients and their physicians. All patients with an ABCD2 score of 4 or greater should be sent to an emergency department for urgent evaluation, whereas most patients with a score of less than 4 can complete their workup as outpatients.

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TIA and stroke a management guide for GPs

The incidence of stroke in Australia is about 46,000 per year. It is the third most common cause of death in this country (about 15,000 per year) and a major cause of disability. All GPs need to be aware of rapid changes occurring in management.

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Dr Ma is Clinical Research Fellow, National Stroke Research Institute; Dr Ly is Clinical Research Fellow, National Stroke Research Institute; Professor Donnan is Professor of Neurology, University of Melbourne, and Director, National Stroke Research Institute, Heidelberg, Vic. For stroke, perhaps above all other medical conditions, there is a need for a seamless continuum of care from the acute presentation, through management in hospital or other healthcare facility, to integration back into the community. With the availability of level I evidence to show that the burden of stroke may be reduced by specific strategies for the acute phase as well as effective forms of secondary prevention, management is changing rapidly. This article discusses current approaches to management, and highlights the specific roles that GPs play – these are summarised in the box on page 24. The evidence underlying changes in management approaches is also explained (see the box on page 25). 1-10

There are now three specific strategies in acute stroke management that have been proven to improve outcomes (that is, are supported by level I evidence). Specifically:

 administration of intravenous tissue plasminogen activator (tPA; alteplase [Actilyse]) within three hours of stroke onset may increase the number of patients improving to

- virtually no clinical deficit by about 30%1,2
- administration of aspirin within 48 hours of ischaemic stroke onset may reduce death and nonfatal stroke in about nine patients per 1000 treated^{3,4}
- management in a stroke care unit may reduce mortality and the need for institutionalised care by about 20%.^{5,11}

Pre-hospital management

For the stroke patient, time is brain. The GP's initial tasks are to recognise the symptoms and signs of an acute transient ischaemic attack (TIA) or stroke (Table 1), and to facilitate rapid transfer to a healthcare facility where the strategies listed above can be put into place with minimal delay. If there is any doubt about the diagnosis, the patient should be transferred to hospital for further investigation. Differential diagnoses are listed in Table 2.

A number of factors have been identified that may prolong hospital arrival times. These include the following:

• lack of recognition of cause of symptoms

N SUMMARY

- . Stroke is one of the major causes of death and disability in Australia.
- All patients with acute stroke should be referred directly to a hospital with imaging facilities (CT and/or MRI), and preferably a stroke unit that provides thrombolytic therapy.
- Management in a stroke unit can reduce long term outcomes of death and dependency.
- · Risk factor identification and management are important.
- Early rehabilitation of patients in a multidisciplinary team environment is ideal.
- Rural GPs are in a unique position to take on multiple roles in stroke management (physician, service co-ordinator, staff and family educator, and support service).

Table 1. Symptoms and signs of TIA and stroke

- · Motor symptoms: weakness or clumsiness (unilateral or bilateral)
- Sensory symptoms: altered feeling on one side of the body
- Difficulty swallowing
- Speech or language disturbance: slurred speech, difficulty with reading or understanding
- Vestibular dysfunction
- Unsteady gait/cerebellar features
- Visual symptoms: loss of vision in one eye, loss of visual field, diplopia
- Nonfocal symptoms: generalised weakness, incontinence, imbalance, altered state of consciousness, dizziness

Table 2. Differential diagnoses

- Migraine
- Syncope
- Epilepsy (Todd's palsy)
- Intracranial structural lesions
- Encephalopathy
- Encephalitis
- Multiple sclerosis
- Transient global amnesia
- Peripheral neuropathy or myopathy
- Metabolic derangement e.g. hypoglycaemia
- Peripheral causes of vertigo, such as benign postural vertigo
- nonambulance transport
- GP contact
- less severe stroke
- ischaemic stroke
- living alone
- being asleep at stroke onset.

The fact that 'GP contact' delays transfer to hospital may seem paradoxical, but it is understandable given that GPs cannot always immediately attend the home to establish the diagnosis.12 Hence, it is crucial that they, having being contacted by a patient or carer by telephone, act to facilitate rapid ambulance (ideally) or other mode of transport to

TIA and stroke



Stroke is a common cause of death and permanent disability in adults. With the results of key trials and meta-analyses that have recently become available, management is undergoing rapid change. Increasing emphasis is being placed on the value of specific treatment strategies in the acute phase and effective forms of secondary prevention.

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hospital if there is reasonable suspicion that stroke has occurred.

Hospital care

After transfer to hospital has been effected, time is still the most important factor. The GP has an important role in a number of the steps involved.

Clinical confirmation of diagnosis

Rapid triage by a nurse trained to recognise the symptoms and signs of stroke is helpful, and expeditious clinical evaluation should establish the diagnosis more clearly. The presentation of the stroke syndromes can often be divided according to the vascular territories using, for example, the Oxford classification of cerebral infarction, which may assist in management. It is also important to identify risk factors such as hypertension, atrial fibrillation, diabetes, cigarette smoking, hyperlipidaemia, a past history of stroke and previous ischaemic heart disease.

Brain imaging

Computed tomography (CT) remains the workhorse of acute stroke management, and has the important advantage over magnetic resonance imaging (MRI) of immediately distinguishing between haemorrhage (increased signal) and infarction (usually little change within the first few hours). Examples are shown in Figures 1a and b.

MRI is becoming more readily available – diffusion-weighted MRI (DWI) has the great advantage of detecting acute ischaemia almost immediately after stroke onset when a CT scan would still be negative. In fact, about one-third

The GP's role in managing patients with stroke

Pre-hospital management

- Recognising the symptoms and signs of acute transient ischaemic attack (TIA) or stroke
- Facilitating rapid transfer to a healthcare facility where acute stroke management strategies can be put into place with minimal delay

Hospital care

- Assisting with medical management (in some hospitals)
- Providing additional medical and historical information about patients to hospital doctors
- Preparing family and carers for the patient's discharge
- Liaising with 'hospital in the home' or 'rehabilitation in the home' programs
- Care-planning with hospital team for discharge

Community care

- Implementing secondary prevention strategies
- Re-integrating stroke patients into the community

of clinically proven TIAs have MRI ischaemic changes, particularly in DWI, that correspond with the clinical presentation. This subset of patients may have a different clinical course compared with patients with TIAs but without MRI (DWI) changes. 13,14

MRI also provides better resolution than CT for imaging the posterior fossa and brainstem.

Acute management with tPA and/or aspirin

Therapy with intravenous tPA may be considered if:

- the length of time from symptom onset is less than three hours
- CT scans are essentially normal
- blood pressure is less than 185/110 mmHg
- the coagulation profile is normal, and
- there are no other contraindications to thrombolysis.

For every 100 patients treated within three hours, 12 patients will have little or no residual clinical deficit (i.e. the number needed to treat to benefit one individual is about eight). tPA was licensed for use in managing stroke in appropriate centres in August 2003.

Oral aspirin (150 to 300 mg) should be administered as early as possible, preferably within 48 hours of stroke onset. Note, however, that tPA and aspirin are never given before brain imaging.

Ultrasound of carotid or vertebrobasilar arteries

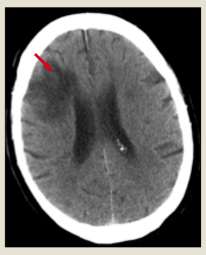
Carotid and transcranial Doppler ultrasound may show focal narrowing in the internal carotid artery and the posterior circulation, respectively. Each patient's suitability for surgical intervention (carotid endarterectomy) should be considered before he or she is subjected to the test. Significant symptomatic stenosis in the internal carotid artery is defined as 70% or more.

Management in a stroke care unit

Level I evidence is now available to show that managing acute stroke patients in a

CT imaging in acute stroke management





Figures 1a and b. CT scans are mandatory for all stroke patients and crucial in making decisions about therapy. a (left). An area of haemorrhage appears white (arrow). b (right). Cerebral infarcts (arrow) appear darker than normal brain.

stroke care unit reduces the risk of death and dependency at one year by about 20% compared with management in a general ward,5 and, therefore, the former approach should now be mandatory. There has been a general recommendation that stroke care units be established Australia-wide,15 and the initiative is being taken up by a number of States and Territories. At present, however, only 23% of Australians have access to stroke care unit facilities.

Minimum investigation set

To define the type of stroke and associated comorbidities, a minimum set of investigations is recommended. This

comprises the following:

- brain CT
- chest x-ray
- full blood examination
- erythrocyte sedimentation rate
- random blood sugar
- urea, creatinine and electrolytes
- ECG.

Newer strategies for reducing the burden of stroke: a look at the evidence

Acute management

There is now level I evidence for three acute management strategies in reducing the burden of stroke: tissue plasminogen activator (tPA), aspirin and care in a stroke care unit. This is summarised in Table A.

Table A. Benefits (avoidance of death or disability) of acute interventions

Strategy	Absolute risk reduction	Number needed to treat*	Potential benefit per 1000 cases of stroke in Australia [†]
Tissue plasminogen activator (tPA) 0-3 hours ^{1,2}	11.0	9	11'
Aspirin ^{3,4}	1.2	83	6 [†]
Stroke care unit ^{5,6}	5.6	18	46 [†]

^{*} The number of patients needed to be treated in one year to avoid one death or disability per year.

The evidence relies on a number of pivotal trials and meta-analyses that have been conducted over the last 10 years. Of these strategies, the number of patients needed to treat to obtain benefit for one individual is most favourable for tPA. It should be noted that although tPA does increase the risk of symptomatic intracranial haemorrhage (about 7% of patients), it does not appear to increase mortality.^{1,2} tPA is now licensed in Australia for management of stroke in appropriate centres.

Aspirin is less effective than tPA, but it is cheaper and safer. There is no evidence that intravenous or subcutaneous heparin is of benefit.3

Management of patients in a stroke care unit is effective and safe, and should not be particularly expensive. Even in smaller hospitals, a stroke care unit can be established and operated by reorganising existing resources.

Secondary prevention

Level I evidence now exists for at least four early strategies for secondary prevention in improving long term outcomes after TIA or stroke, including use of antiplatelet agents, blood pressure lowering, warfarin and carotid endarterectomy. This is summarised in Table B.

Table B. Benefits of secondary prevention strategies

Strategy	Relative risk reduction	Absolute risk reduction	Number needed to treat*
Antiplatelet agent⁴	23	1.0	100
Blood pressure lowering ⁷	28	2.2	45
Warfarin ⁸	67	8.0	12
Carotid endarterectomy ^{7,9,10}	44	3.8	26

^{*} The number of patients needed to be treated in one year to avoid one recurrent stroke per year.

[†] Case reduction of death and dependency from an estimated number of stroke cases in Australia in 1998 sourced from several Australian studies.

Special issues for rural GPs

- Access to CT and duplex ultrasound.
 Can this be arranged urgently? If not, the patient should be transferred.
- More complete medical
 management. In many instances
 the GP may be the sole medical
 manager, hence a knowledge of
 level I evidence for acute care (tPA,
 aspirin, stroke unit management)
 as well as secondary prevention
 strategies (antiplatelet agents,
 warfarin, blood pressure lowering,
 referral for carotid endarterectomy
 or lipid lowering) is essential. If not,
 the patient should be referred or
 transferred.
- Facilitating the continuum of care from acute presentation through to rehabilitation and community adjustment. Rural GPs are probably even more involved in this than urban GPs.
- Family and emotional support, which are crucial in the recovery process.
 Depression is not uncommon among stroke patients.

Transoesophageal echocardiography is an additional investigation that may be used to detect aortic arch atheroma as an embolic source if no obvious cardiac or large artery sources have been identified. In patients with atrial fibrillation or recent acute myocardial infarction, this investigation is not usually indicated unless more information about cardiac function is required.

Management of medical complications

Each complication should be managed on its merits. These include:

- infections (e.g. pneumonia, urinary tract infections)
- deep venous thrombosis (DVT) or pulmonary embolism

- cardiac failure, fluid imbalance
- hypertension
- hyperglycaemia
- cardiac arrhythmias/cardiac infarction
- pressure sores
- urinary and bowel problems
- seizures
- · falls.

Significant hypertension is not an uncommon finding in the acute stroke setting, and generally there is no need for intervention in ischaemic stroke unless the systolic pressure is above 220 mmHg or the diastolic pressure is above 120 mmHg. Fortunately, elevated blood pressure at admission usually settles within hours, or within a few days at most.

Pressure stockings are essential for preventing DVT; subcutaneous heparin is often used in patients who cannot be mobilised or with a history of DVT or pulmonary embolus.¹⁶

Speech assessment is important for patients with speech impairment, and appropriate dietary advice is essential for those with swallowing difficulties.

Early rehabilitation

Although there is rapidly accumulating evidence that rehabilitation improves clinical outcomes, it is uncertain how early this should occur. Nevertheless, it is generally considered that rehabilitation should usually be commenced as soon as practicable, and this is the practice in most stroke care units.

Early secondary prevention

Based on their risk factor profile, people who have experienced a TIA may have a risk of stroke of up to 31% within the next seven days. ¹⁷ Conversely, in patients who have experienced an ischaemic stroke preceded by a TIA, 43% of these TIAs will have taken place within seven days before the ischaemic stroke. ¹⁸ Thus it is crucial to instigate appropriate investigations and secondary prevention as early as possible.

There is now level I evidence that at least four early strategies for secondary prevention may improve long term outcomes after TIA or stroke. Some of these may be initiated in hospital, but most often they are commenced by the GP when the patient has returned to the community. These strategies are discussed in the section 'Secondary prevention' below.

Discharge planning

Of the patients with stroke who enter hospital, about 20% will die during their stay and the remainder will be discharged:

- directly home (33%)
- to rehabilitation centres (33%)
- to nursing homes (only about 10%).

For the two-thirds who go either home or to rehabilitation, early planning is essential. Increasingly, patients are being discharged early with the support of 'hospital in the home' programs, in which ongoing medical management of stroke comorbidities is undertaken. The use of 'rehabilitation in the home' programs is an increasingly common trend.

Community care

The GP, who may have been involved during the early phases of stroke onset and hospital care, assumes an even more important role in caring for a patient after discharge from hospital. The main management issues for GPs include use of secondary prevention strategies, and it is helpful to have a checklist to ensure that all appropriate steps are taken. GPs are also involved in reintegrating the patient into the community.

Secondary prevention

The evidence-based secondary stroke prevention strategies that need to be implemented where appropriate are as follows:

- · antiplatelet agents
 - aspirin4
 - clopidogrel (Iscover, Plavix)¹⁹
 - aspirin with dipyridamole (Asasantin SR)^{20,21}

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- blood pressure lowering agents
 - using perindopril (Coversyl) alone or with indapamide²² (a combined preparation [Coversyl Plus] is available)
 - for patients with a previous history of stroke there is some evidence that ramipril (Ramace, Tritace) may be effective²³
- warfarin (Coumadin, Marevan), in patients in atrial fibrillation^{8,24}
- carotid endarterectomy, in symptomatic patients with carotid stenosis of 70% or more 9,10
- lipid lowering agent (atorvastatin [Lipitor]) in ischaemic stroke.²⁵

The box on page 29 summarises the benefits of the first four of these secondary prevention strategies.

Compared with the use of aspirin alone, clopidogrel has been shown to further reduce the relative risk of vascular events, including ischaemic stroke, by about 8.7% (CAPRIE).¹⁹ The major use of clopidogrel is in the setting of further cerebrovascular events despite the use of aspirin or in the patient who cannot tolerate aspirin due to its side effects.

Compared with the use of aspirin alone, the combination of aspirin and dipyridamole also shows a relative reduction of risk of stroke or death by a further 13% (ESPS-2).20 This has recently been confirmed by another trial (ESPRIT),²¹ and makes this combination a suitable alternative to aspirin as first line antiplatelet therapy in secondary prevention of ischaemic stroke. This trial, which compared aspirin with combination therapy of aspirin plus dipyridamole in patients with TIA or minor ischaemic stroke, showed a relative risk reduction of 20% in reaching the primary end point (composite of death from all vascular causes, nonfatal stroke, nonfatal myocardial infarction or major bleeding complication) in the combination group compared with the aspirin only group.²¹ The other major use of the combination of aspirin and dipyridamole is in the setting of further cerebrovascular events despite the use of aspirin.

For patients with nonvalvular atrial fibrillation one should aim for an INR of 2.5 (range, 2.0 to 3.0).²⁴ Heparin is still occasionally used as a prelude to warfarin, but evidence for this is lacking.³

A recent prospective randomised controlled study has shown that in patients with recent ischaemic stroke or TIA and serum LDL cholesterol levels between 2.6 and 4.9 mmol/L, atorvastatin 80 mg/day reduced the five-year absolute risk of fatal or nonfatal stroke by 2.2% compared with placebo (SPARCL).25 The five-year risk of major cardiovascular events was also reduced by 3.5%. This means that 46 patients will need to be treated over a period of about five years to prevent one stroke. However, there was a slight increase in the incidence of haemorrhagic stroke (adjusted hazard ratio 1.66) in the treatment group. Hence atorvastatin should be used only in patients with ischaemic stroke.

Risk factor modification measures are also important. These include cessation of cigarette smoking and avoidance of heavy alcohol consumption. In addition, blood glucose in patients with diabetes should be strictly controlled (although there is no level I evidence to support this recommendation).

Re-integration

Adjustments to the patient's environment may be required because he or she may need to be re-introduced to work or home activities gradually. Increased stress on carers is likely, sometimes with attendant illnesses. In some instances, respite care may need to be arranged.

Role of the rural GP

Rural GPs are faced with additional challenges for managing patients with stroke or TIA – these are described in the box on page 26. Therefore, awareness of the evidence base for management strategies is most important – particularly if

they undertake more complete medical management. Since access to specialist services is more limited in rural areas, GPs usually assume greater responsibility, thus making it even more important that they are aware of the need to refer to rural based specialists when appropriate (for example, for carotid endarterectomy).

Resources

The National Stroke Foundation (NSF) produced a set of clinical guidelines for acute stroke management in 2003.

Final comments

The management of stroke is changing quite rapidly with the introduction of the evidence based strategies discussed in this article. The role of the GP remains central.

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Managing stroke survivors in the community

Your patient has survived a stroke and presents at your surgery a week after discharge.

Where do you start?



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A stroke can transform a person's physical, psychological and social functioning. Stroke survivors are usually managed for the first seven or so days in an acute stroke unit, and rehabilitation promoting independent movement begins during this time. About one-third of these patients, depending on each individual's needs, then undergo further rehabilitation in specialised inpatient rehabilitation units. About 45% of stroke survivors die within five years of their initial stroke and 20 to 25% become significantly

Stroke rehabilitation helps patients maximise their potential for recovery and provides practical ways of dealing with ongoing disability. Family members and other carers are also involved in rehabilitation in that they are trained in how to care for stroke survivors. A stroke rehabilitation program involves a co-ordinated program of exercises (physiotherapy), the practice of daily activities such as dressing (occupational therapy), the optimising of speech and swallowing (speech therapy) and the mapping of memory and cognitive losses (neuropsychology). This is done within a framework of support for the patient and family (social work) and culminates in the stroke survivor's return to the community to live with a disability and draw out the best quality of life possible.

This article provides a practical guide to helping stroke survivors who have a persistent disability maintain and enhance the gains made in rehabilitation. The article is divided into three sections: what to do once the patient has been discharged from hospital, secondary prevention and how to approach general post-stroke deterioration.

Discharge summary: stroke

The initial post-stroke consultation probably occurs a week after discharge when the stroke

- To optimise the quality of life of a stroke survivor it is important both to become re-acquainted with the patient and to be aware of all aspects of his or her new care infrastructure.
- As stroke survivors have a 6 to 10% yearly risk of recurrence, secondary prevention is essential. Tight control of blood pressure and diabetes, the use of antiplatelet agents (or warfarin in those with atrial fibrillation), cholesterol level reduction, smoking cessation, a healthy diet and avoiding excessive alcohol intake are all important.
- Daily exercise is central to maintaining mobility and quality of life in stroke survivors.
- General deterioration in a stroke survivor is not always 'the final curtain' once acute causes have been excluded; quite often some simple rehabilitation is all that is required.
- Stroke rehabilitation helps patients maximise their potential for recovery and provides practical ways of dealing with ongoing disability. It involves an intense multidisciplinary program that can be undertaken in the community or as an inpatient.

Trivia points

- · Stroke was first likened to the 'stroke of God's hand' in 1599. The Greek word plesso means to 'stroke, hit or beat' and gives us the word plegia, as in hemiplegia.2
- · Babinski didn't give his name to a reflex but described a loss of plantar response now known as the Babinski sign. There are over 20 different ways to elicit plantar/extensor responses.
- The League of Nations (the precursor to the United Nations) was formed following World War I essentially because US President Woodrow Wilson suffered post-stroke vascular dementia and world leaders felt sympathy for him. His wife hid him away from the press.

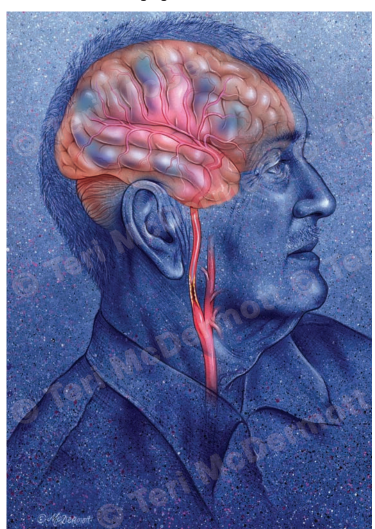
survivor's supply of hospital medications is about to run out. This will be your first opportunity to assess your patient's condition and the presence and level of disabilities. Be aware that changes are not only physical but also psychological and social, and that communication may be affected (for example, the patient may be aphasic).

Ideally, GPs should be involved in the discharge case conference while the patient is in the inpatient rehabilitation unit so that the handover to community care can be streamlined.

The post-stroke follow up appointment

A comprehensive assessment of the patient is essential to establish a new baseline of physical and psychological function. If time does not permit this at the patient's first presentation after discharge, organise a follow up appointment. From the discharge summary determine whether the stroke was a hemisphere stroke (physical and cognitive effects) or a brainstem stroke (usually physical affects only), and the mechanism involved (thrombus 40%, embolus 30%, lacunar 20% or haemorrhage 10%). This differentiation will have determined the medical and/or surgical treatments used (such as warfarin, aspirin or statin, and carotid endarterectomy or carotid stenting, respectively).

Managing stroke survivors



The community care of stroke survivors includes promoting independent living, preventing further strokes and caring for the carers. These patients can benefit greatly from rehabilitation both in the first few weeks after their stroke and also later on when carers report that the patient has deteriorated but no acute cause can be found.

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A checklist for assessing a stroke survivor is given in Table 1. Your focus will necessarily start with ascertaining the patient's level of communication (Table 2) to determine if the history should be taken from the stroke survivor or the carer. You can then explore the patient's new

functional baseline and document the level of dependence in self-care, homecare and work and leisure skills – i.e. the activities of daily living (ADLs; Table 3). Ensure that the patient undergoes a Mini Mental State Examination (MMSE) early in the assessment as post-stroke dementia occurs in 12 to 22% (and possibly more) of stroke survivors.³

The patient may still be in stroke rehabilitation, either as an outpatient or in the community. He or she should have a home-based exercise program prescribed as part of the rehabilitation, and compliance in minutes per day should be documented.

If swallowing is a problem, the consistency of the food that is eaten (pureed food, normal food but thickened fluids or normal diet and thin fluids) can be recorded.

Complaints of fatigue (which occur in

10 to 39% of patients) or pain can be documented using visual analogue scales (for example: 'If 0 is no pain [or fatigue] and 10 is the worst you can imagine, what level of pain [or fatigue] are you feeling right now?').4

Ask also about mood, as up to 23% of stroke patients living in the community have reported significantly depressed mood.⁵

Examining a stroke survivor

Attention should be paid to blood pressure, which should be tightly controlled and as low as possible without there being symptoms of hypotension (at least below 135 mmHg systolic and 85 mmHg diastolic), and to heart rate. Careful examination of respiratory status, cardiac status, visual loss, visuospatial neglect or inattention, and weakness are important. Testing muscle tone is helpful because a

change in tone is often a harbinger of generalised illness or infection, or an indicator of a hidden site of pain.

Observe whether the patient is able to stand from a seated position, walk independently and get on and off the examination couch without assistance. Examine the hemiplegic shoulder with care because there is a high incidence of hemiplegic shoulder pain (from a combination of rotator cuff tears, adhesive capsulitis and complex regional pain syndrome).

Secondary prevention

You are likely to see a stroke survivor about every six months for repeat prescriptions and this provides an opportunity to review secondary prevention measures. This population has a risk of recurrent stroke per year of 6 to 10%, which can be reduced by up to 13% with

Table 1. Assessing a stroke survivor: baseline history and examination

First step

- Assess communication (see Table 2)
- Perform a Mini Mental State Examination

History

Ask about:

- problems since discharge from hospital
- activities of daily living (ADLs; see Table 3)
- mobility and falls
- continence
- pain shoulder, hemisensory, arthritic
- mood sleep disorders, social withdrawal, appetite, interests and hobbies, suicidal thoughts
- devices and equipment used gastrostomy feeding tube, suprapubic catheter, walking stick, wheelchair, commode, dressing stick, personal medical monitoring system (e.g. VitalCall)
- food consistency and diet pureed food, normal food but thickened fluids or normal diet and thin fluids
- medications and side effects
- carer stress ask carers how they are managing and whether they have any health problems of their own

Examination

Assess:

- blood pressure should be as low as possible without symptoms of hypotension (at least <135/85 mmHg)⁶
- cardiovascular status atrial fibrillation, cardiac failure or murmurs, carotid bruits
- respiratory status swallow deficit poses risk of aspiration
- transfers chair to standing, standing to examination couch
- mobility stick use, hemiplegic gait, stumbling, hemiplegic foot clearance (foot drop), arm swing (abnormal posturing during walking)
- vision hemianopic, diplopia, eye movement abnormalities
- speech and swallow dysarthria, dysphonia
- muscle tone hypertonicity, clonus, spasticity; plus any associated pain
- muscle weakness in face, arms, legs test reflexes
- hemiplegic shoulder pain, contracture, poor axillary hygiene
- sensory loss visuospatial neglect or inattention, hemisensory loss of pain and temperature (risk of burns), loss of proprioception in leg (risk of falls)

Table 2. Assessing a stroke survivor: communication skills

- · Check the patient's ability to hear and see
- Establish how many steps the patient can comprehend by asking him or her to:
 - close your eyes (one-step command)
 - close your eyes and open your mouth (two steps)
 - close your eyes, open your mouth and poke out your tongue (three steps)
- · Establish naming ability by asking the patient to name three items of increasing complexity. For example:
 - show a watch
 - show the band of the watch
 - show the clasp of the watch
- Establish repetition ability by asking the patient to repeat a number sequence such as 1,2,3,4 and the phrase 'No ifs, ands or buts'
- Establish alexia and agraphia by asking the patient to read and write

antiplatelet therapy and by up to 60% with the use of warfarin (Coumadin, Marevan) in those subjects with atrial fibrillation.7,8

Studies have shown hypertension to be the strongest modifiable risk factor for stroke and the lower the blood pressure is kept then the better the prevention. Tight asymptomatic lowering of the blood pressure by as little as 3 to 6 mmHg has been shown to decrease the risk of recurrent stroke by up to 32%.9 While reducing the blood pressure seems more important than the type of agent chosen, ACE inhibitors such as perindopril and ramipril have been shown to reduce recurrent stroke even in normo tensive patients.

Tight diabetic control is extremely important, and needs to be reviewed regularly. Also, a healthy diet, smoking cessation and avoiding excessive alcohol should all be encouraged.

Many stroke survivors (up to 35%) have concurrent coronary and carotid artery disease, and cholesterol reduction is important in these patients. Although the relation of statin use to stroke prevention may be much looser than to ischaemic heart disease prevention, it is likely that more than 22% of stroke

survivors will die from a cardiovascular event.10

Exercise is required on a daily basis to minimise muscle contractures, assist in blood pressure control and improve endurance and spasticity. Stroke survivors might need to attend outpatient rehabilitation centres where specialised home exercise programs can be developed. Carers are often trained to help supervise these exercise programs and physiotherapists may monitor an individual's program once or twice over a six-month period.

A medication review is needed, includ ing of antiplatelet therapy (or warfarin use in those with atrial fibrillation). If a patient is on an antiepileptic medication that is metabolised by the liver (such as phenytoin [Dilantin, Phenytoin Injection], carbamazepine [Carbamazepine-Sandoz, Carbamazepine-BC, Tegretol, Teril] or sodium valproate [Epilim, Valpro]), it may be wise to check liver function tests every six months. If levels are elevated, increasing or associated with symptoms, referral to a neurologist or a rehabilitation physician should be considered. A one to two-year period free of fits should encourage the gradual withdrawal of the medication, particularly in stroke

Table 3. Assessing a stroke survivor: activities of daily living (ADLs)

ADL functioning levels

- Independent no help needed
- Supervision required someone standing by just in case
- Assistance required (i.e. dependent) hands-on help needed

Types of ADLs* Personal ADLs (PADLs or 'paddles')

- Feeding
- Grooming
- Dressina
- Showering
- Toileting
- Transfers

Domestic (DADLs or 'daddles')

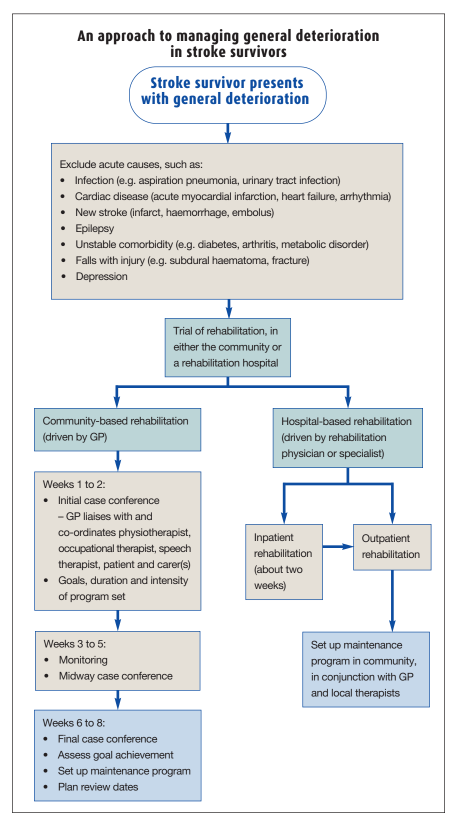
- Cooking
- Cleaning
- Laundry

Community (CADLs or 'caddles')

- Driving
- Shopping
- Gardening
- Telephone use
- Money management
- Medication management
- * Domestic and community ADLs also fall in the category of instrumental ADLs (i.e. activities enabling community living).

survivors with cognitive deficits that might be worsened by the sedative side effects of the antiepileptic agent.

Carer stress cannot be overemphasised. The carer should be seen separately or spoken to directly during the patient's consultation. Enquiring about his or her health, cardiovascular risk factors and the number of hours per week he or she has for recreation time, as well as discussing respite services in the community, is vital for preserving the care infrastructure needed by the stroke survivor.11 In



extreme situations, such as family breakdown or acute illness in the carer, inpatient respite care (in a hostel or nursing home) may need to be organised for the stroke survivor.

Post-stroke deterioration

Deterioration after a stroke – 'He or she is just not him or herself' – can be difficult to diagnose and manage. An approach is presented in the flowchart on this page.

It is essential to determine whether there are focal or general symptoms (such as an overall increase in tone), and acute causes of deterioration should be excluded (Table 4). Acute focal neurological symptoms or signs should be treated as an emergency. The most common infections in people who have had a stroke are chest infections (particularly in those with dysphagia), urinary tract infections and cellulitis. Motor or psychomotor seizures are reported by carers in 5 to 10% of patients, and a work up (including EEG, blood tests [sodium and antiepileptic medication levels] and a brain scan) is necessary in these patients to exclude causes of seizures. Stroke extension, haemor rhage, or fitting despite good antiepileptic cover usually requires referral to a neurologist, physician or rehabilitation physician.

Cardiac disease is common in stroke survivors and they may develop arrhythmias (up to 50% of patients), myocardial infarcts or heart failure. These patients need to be treated urgently because cardiac reserve, related to endurance, is often reduced in patients who have motor deficits.

Falls need attention as sensation can be abnormal and pain may not be reported.

Finally, continence changes can be due to infection, constipation or outflow obstruction (for example, prostate disease), as well as failing mobility.

Post-stroke depression

Post-stroke depression can occur as late as two years after the stroke and is often difficult to diagnose. Lack of participation in usual activities and decreasing levels of functioning in ADLs are good indicators, as are suicidal thoughts, sleep disorder and social withdrawal.

Personality change, blunted affect, crying, cognitive decline, changes in appetite and weight, and anhedonia can often be confused with the neurological manifestations of stroke (such as cognitive impairment, emotional lability, facial paralysis, dysphagia and immobility).

It is reasonable to give the stroke survivor with symptoms of depression a trial of a selective serotonin reuptake inhibitor (SSRI), choosing one likely to have the least adverse effects in that patient. Citalopram or sertraline are often appropriate. Tricyclic antidepressants can worsen swallow, cognition and urinary outflow and diminish ADL functioning; these drugs should probably be avoided.12

Deterioration in the absence of obvious cause

If there is no obvious cause of a stroke survivor's deterioration, it might be that deconditioning is involved. After the patient's return from rehabilitation he or she may have been allowed to rest excessively and have everything done for him or her because of circumstances such as environmental change, lack of a support framework, poor motivation or inadequate carer education. Such a 'killing with kindness' scenario may lead to muscle weakness and poor mobility, followed by the development of contractures (usually at the knee and hip but also at the hemiparetic shoulder), falls or even urge incontinence (because it takes too long to walk to the toilet). Recent evidence demonstrates that stroke survivors improve in walking speed and cardiovascular fitness following a six-week burst of rehabilitation, no matter how long after their stroke this occurs.

A burst of multidisciplinary rehabilitation can be undertaken either in the community with GPs co-ordinating the treatment or as an inpatient or outpatient of a rehabilitation medicine unit. (The website of the Australasian Faculty of Rehabilitation Medicine, www.afrm. racp.edu.au [then click regional, NSW, rehabsearch], has a search engine that allows an entered postcode to bring up all the local rehabilitation facilities and information on outpatient and inpatient programs.) A two-week burst of inpatient rehabilitation has the added benefit of giving the carer a break.

If the rehabilitation is to be undertaken in the community, physiotherapy practices accredited by the Australian Physiotherapy Association should be able to put you in touch with local private occupational therapists and speech therapists if needed. Therapists can also be found through OT Australia and Speech Pathology Australia, respectively (www. ausot.com.au and www.speech pathologyaustralia.org.au). The initial case conference, a phone hook-up or brief meeting with the therapists, carers and possibly the patient, can set the goals. Possible goals could be the improving of the following:

- mobility and transfers
- education and carer training
- upper limb strength
- intelligibility of communication
- independence in ADLs.

The physiotherapist, occupational therapist and/or speech therapist will need to see the patient two to three times a week for six to eight weeks to achieve the set goals. Quite often this is a trial by fire, and lack of progress can point to a more serious or complex organic process such as the progression of vascular dementia. For this reason, a case conference needs to be held with therapists, patient and carers about halfway through the process. Another case conference is needed at the end of the process to determine whether goals have been achieved. A monitoring stage then follows, during which the GP reviews the patient every

Table 4. Post-stroke deterioration: acute causes

- Infection e.g. aspiration pneumonia, urinary tract infection
- Cardiac disease acute myocardial infarction, heart failure, arrhythmia
- New stroke infarct, haemorrhage, embolus
- Epilepsy
- Unstable comorbidity e.g. diabetes, arthritis, metabolic disorder
- Fall injury to subdural haematoma, fracture
- Depression

few months to ensure the goals are maintained.

There are Medicare arrangements in place to provide several allied health services; however, there can be significant transport issues for the patient and carer. Rehabilitation in the community also relies greatly on excellent communication between the various team members.

The rural setting

In rural settings, access to allied health personnel and transport are major issues. It may be necessary to admit the patient to the local district hospital for investigation of deterioration and for physiotherapy and occupational therapy services. Patients unable to return to work such as farming or driving because of deterioration qualify for therapy services provided by the Commonwealth Rehabilitation Service (a government agency with a charter to assist disabled people to return to the workplace).

Telemedicine facilities exist in some rural hospitals and can be used to contact hospitals in large cities for advice from rehabilitation physicians and allied health staff on approaches to post-stroke rehabilitation.

Trainers at local swimming pools, instructors at local gyms and even carers can be educated by physiotherapists to oversee the performing of specific exercise programs.

Conclusion

Stroke survivors with substantial disabilities are often put in the too hard basket and may be thought of as complex cases with poor outcomes. On the contrary, assisting a stroke survivor in maintaining his or her best quality of life is a great achievement and well appreciated. In the words of one of my patients, a retired doctor: 'I summit Mt Everest every day; thank God for the sherpas!'

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