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Campbell, BCV

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**Authors:**

	Title	First name	Mid inits	Last name	Postnom (eg, PhD) [3 only for publication]	Position1	Address1	Position2	Address2	Tel	Email
1	Prof.	Bruce	CV	Campbell	MBBS, BMedSc, PhD, FRACP	bruce.campbell@mh.org.au	1				bruce.campbell@mh.org.au
2											
3											
4											
5											

Number of corresponding author:	1
Number of alternative corresponding author:	

**Addresses:**

	Institution	City	State	Post Code	
1	Royal Melbourne Hospital, University of Melbourne	Melbourne	VIC	3050	
2					
3					
4					
5					

Postal address of first corresponding author (if different from the institutional address given above)	Department of Neurology, Melbourne Brain Centre at the Royal Melbourne Hospital, 300 Grattan St, Parkville VIC 3050
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
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# Advances in stroke medicine

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## Summary

- In recent years, reperfusion therapies such as intravenous thrombolysis and endovascular thrombectomy for ischaemic stroke have dramatically reduced disability and revolutionised stroke management.
- Thrombolysis with alteplase is effective when administered to patients with potentially disabling stroke, who are not at high risk of bleeding, within 4.5 hours of the time the patient was last known to be well. Emerging evidence suggests that other thrombolytics such as tenecteplase may be even more effective. Treatment may be possible beyond 4.5 hours in patients selected using brain imaging.
- Endovascular thrombectomy (via angiography) effectively reduces risk of death or dependency in patients with large vessel occlusion (internal carotid, proximal middle cerebral and basilar arteries) if applied within 6 hours of the time they were last known to be well.
- Endovascular thrombectomy is also beneficial 6–24 hours from the last known well time in selected patients with favourable brain imaging. Thus, some patients with wake-up stroke are now treatable, and protocols for stroke need to include computed tomography (CT) perfusion scan and CT angiography as routine, in addition to the non-contrast CT brain scan.
- Optimised pre-hospital and emergency department systems (eg, code stroke response teams, pre-notification by ambulance, direct transport from triage to CT scanner) are essential to maximise the benefit of these strongly time-dependent therapies. Telemedicine is increasingly providing specialist guidance for these more complex treatment decisions in rural areas.
- Important developments in secondary stroke prevention include the use of direct oral anticoagulants or left atrial appendage occlusion for atrial fibrillation, and endovascular closure of patent foramen ovale.

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Acute ischaemic stroke caused by occlusion of a cerebral artery is a leading cause of disability globally. In Australia, there are over 56 000 stroke cases annually.<sup>1</sup> Stroke care has been revolutionised in recent years with the advent of effective reperfusion therapies. It is now possible for some patients with severe stroke to recover and be discharged home in a matter of days. However, major challenges remain in the prevention and treatment of stroke. This review outlines the current state of evidence and practice in Australia.

The literature review was performed searching the PubMed online database, including articles published between 2010 and 2018, and the Australian stroke guidelines ([www.informme.org.au/guidelines](http://www.informme.org.au/guidelines)). Randomised controlled trials in the areas of acute stroke treatment and secondary prevention were selected based on personal assessment of importance and relevance.

## Evidence-based therapies for stroke

The basis of all stroke care is management in a geographically defined, specialist-led stroke unit (Box 1). This “black box” of skilled personnel reduces disability and death for all stroke subtypes and severities through a combination of acute treatment, prevention of complications and early rehabilitation.<sup>2</sup> Aspirin remains an important, generalisable and inexpensive therapy to reduce early ischaemic stroke recurrence, with a modest number needed to treat.<sup>3,4</sup> In contrast, hemispherectomy is a treatment for patients selected on a case-by-case basis who have large hemispheric infarction to

prevent mass effect causing tonsillar herniation. The number needed to treat to reduce disability is 4.6, and 2 to save a life, but relatively few patients require the procedure.<sup>5</sup> Thrombolysis for ischaemic stroke was first shown to be effective in 1995<sup>6</sup> but remains underutilised globally. Recent randomised trials suggest that selection by tissue status using advanced brain imaging rather than time may allow additional patients to receive treatment.<sup>7,8</sup> Endovascular thrombectomy came of age in 2015 with the publication of five randomised controlled trials demonstrating treatment efficacy within 6 hours of stroke onset.<sup>9-13</sup> In 2018, that time window was extended to 24 hours in patients selected based on advanced brain imaging to identify salvageable ischaemic penumbra.<sup>14,15</sup> Box 2 summarises the indications and evidence of benefit for each therapy.

## Reperfusion therapies for ischaemic stroke

### Intravenous thrombolysis

Intravenous alteplase was initially established as effective within 3 hours of symptom onset<sup>6</sup> and subsequently extended to 4.5 hours.<sup>20</sup> An individual patient data meta-analysis of all randomised trials of alteplase clarified the relationship between longer time to treatment and poorer outcome.<sup>16</sup>

Thrombolysis with 0.9 mg/kg alteplase within 4.5 hours of stroke onset is now recommended in stroke guidelines globally.<sup>21-24</sup> The benefits are generalisable across all ages and clinical severities for patients with a potentially disabling stroke.<sup>16</sup> Thrombolysis beyond 4.5 hours is not recommended in current guidelines. However, the recent WAKE-UP randomised trial used magnetic resonance imaging (MRI) to identify patients with a diffusion lesion that was not yet visible on fluid-attenuated inversion recovery imaging as a marker of recent onset in patients with stroke of unknown onset and it demonstrated improved outcomes with alteplase versus placebo.<sup>7</sup> Urgent MRI is not practical in most Australian centres. Fortunately, the Australian-led EXTEND randomised trial, which selected patients based on the presence of salvageable brain tissue predominantly using computed tomography (CT) perfusion also showed improved functional outcomes with alteplase in patients within 9 hours of the last known well time or from the midpoint of going to bed and waking with stroke symptoms.<sup>8</sup> [Author, we can include the *NEJM* reference that will be published on 8 May and add “in press”. Please kindly provide all the details for this reference and we’ll replace it in the reference list. - done] These results will likely be incorporated into treatment guidelines and allow the treatment of additional patients. However, the imperative to minimise treatment delays is not reduced by this expansion in treatment time window as neuronal death continues rapidly and the prevalence of favourable brain imaging decreases as time passes.

About 20% of all patients with ischaemic stroke are eligible for thrombolysis based on the 4.5-hour time window in data from the most active centres, but the current Australia-wide rate is 13%.<sup>25</sup> Thrombolysis reduces disability, with a number needed to treat to achieve an additional excellent patient outcome (ie, return to all their usual activities, modified Rankin Scale score 0–1) of 4.5 when treated within 90 minutes, 9 when treated within 90–180 minutes and 14 when treated within 180–270 minutes.<sup>26</sup> Clearly, the earlier the treatment, the greater the benefit. Unfortunately, only 30% of Australian patients with stroke treated with thrombolysis receive it within 60 minutes of hospital arrival.<sup>25</sup> This rate compares with approximately 60% of thrombolysis patients treated within 60 minutes in the United States and the United Kingdom.<sup>25</sup>

### Risks

The main risk of alteplase is symptomatic intracerebral haemorrhage. Early trial definitions of symptomatic haemorrhage included patients with any bleeding within a large infarct, in whom the bleeding was clearly not the cause of neurological deterioration. Indeed, haemorrhagic transformation is part of the natural history of large infarcts. Current definitions of intracerebral haemorrhage require that the bleeding occupy at least 30% of the infarct volume

and cause mass effect.<sup>27</sup> Using that definition, current real-world registry data indicate a 1.7% risk of symptomatic intracerebral haemorrhage.<sup>27</sup>

Serious systemic bleeding is rare in appropriately screened patients, and endovascular thrombectomy is an option for patients with large vessel occlusion (internal carotid, proximal middle cerebral and basilar arteries) and systemic bleeding risk. In some patients with post-operative stroke, thrombolysis can be considered, in discussion with the surgeon, based on careful consideration of individual risk and benefit. Orolingual angioedema (usually affecting unilateral tongue and palate on the paretic side) occurs in about 1% of patients (5% if taking angiotensin-converting enzyme inhibitors) and is mediated by bradykinin.<sup>28</sup> Icatibant (a bradykinin receptor antagonist) may be used for severe cases to prevent the need for intubation.<sup>29</sup>

### ***Mild stroke with non-disabling symptoms***

In patients with mild stroke with non-disabling symptoms, there is a particularly difficult balance of risk and benefit, and 30% of patients deemed “too mild to treat” subsequently deteriorate (beyond the time frame for treatment) and become disabled.<sup>30</sup> The risk of symptomatic haemorrhage is lower in patients with smaller strokes but the natural history is also better. The recent PRISMS (Potential of rtPA for Ischemic Strokes With Mild Symptoms) randomised controlled trial in patients with minor, non-disabling stroke (eg, isolated sensory loss, facial droop or dysarthria) showed that thrombolysis with alteplase was not beneficial.<sup>31</sup> However, PRISMS did not use brain imaging to demonstrate a target for thrombolysis. As patients with arterial occlusion are most at risk of deterioration despite initially mild symptoms,<sup>32</sup> trials of thrombolysis in mild stroke that use imaging to select patients with arterial occlusion are in progress. Importantly, this uncertainty does not apply to patients with isolated motor weakness, aphasia or hemianopia that would be regarded as potentially disabling. These patients should be considered candidates for thrombolysis.

Ongoing research is investigating whether intravenous thrombolysis efficacy can be improved. Tenecteplase has shown superior outcomes compared with alteplase in two trials that enrolled patients with vessel occlusion.<sup>33,34</sup> Other trials that did not select patients with vessel occlusion showed similar results to alteplase.<sup>35,36</sup> This is consistent with the minimal benefit of thrombolysis in patients without demonstrable vessel occlusion seen in multiple studies.<sup>37</sup>

### **Endovascular thrombectomy**

The concept of endovascular treatment of stroke via angiography preceded intravenous thrombolysis but, despite a positive trial in 1999,<sup>38</sup> it did not enter mainstream practice. Initially, treatment was provided with intra-arterial thrombolytics, but mechanical clot retrieval devices appeared in the first decade of the 21st century. Three neutral randomised trials published in 2013<sup>39-41</sup> dampened the enthusiasm but also reset clinician equipoise and facilitated recruitment into the next generation of trials. A new generation of devices was developed: retrievable stents that are deployed within the clot and then retrieved under negative pressure aspiration (Box 3). In 2015, five positive trials of endovascular thrombectomy using predominantly stent-retriever devices essentially doubled the rate of successful reperfusion compared with previous generation devices, and also incorporated a variety of important imaging selection and streamlined workflow approaches to reduce treatment delays.<sup>9-13</sup>

A subsequent individual patient data meta-analysis has demonstrated remarkable consistency of treatment benefit across the spectrum of age, clinical severity, and imaging characteristics of participants.<sup>17</sup> As in most conditions, increasing age and more severe clinical presentations were associated with worse outcomes. However, older and severely affected patients benefitted from endovascular thrombectomy compared with medical therapy, with effect sizes similar to other subgroups. There were trends to greater mortality reduction in older patients. Using general anaesthesia was associated with worse outcomes than performing the procedure with the patient awake,<sup>42</sup> in contrast to single centre randomised trials that showed little difference.<sup>43-45</sup> However, the randomised trials were performed by specialist

neuroanaesthetic teams with strict protocols to maintain physiological parameters and minimal delay (approximately 9 minutes) to achieve general anaesthesia. These practices are not routine in most centres.

Brain imaging is key to stroke diagnosis but also provides important prognostic information regarding the extent of irreversible injury and tissue at risk. Individual patient data meta-analyses of centrally reviewed imaging data from all the endovascular thrombectomy trials have examined the effect of non-contrast CT hypodensity, collateral blood flow quality,<sup>46</sup> CT perfusion and MRI<sup>47</sup> parameters on prognosis and benefit from endovascular thrombectomy versus medical therapy. Even patients with extensive injury on pre-treatment imaging can still benefit from thrombectomy. Further randomised trials are specifically addressing the treatment benefit of endovascular thrombectomy in patients with a large irreversibly injured ischaemic core, but at present there is no reason to exclude a patient with large vessel occlusion and onset within 6 hours from endovascular therapy based purely on imaging appearances.

Time to reperfusion remains a crucial determinant of outcome, with every 4-minute delay after reaching the emergency department associated with one in 100 patients having a worse outcome.<sup>48</sup> Despite this time-sensitivity, individual patients can have prolonged survival of salvageable brain due to good collateral blood flow. These patients can be identified using CT perfusion or perfusion-diffusion MRI (Box 4). CT perfusion dynamically tracks the passage of an intravenous bolus of contrast, identifying regions of the brain with delayed contrast arrival due to arterial occlusion with retrograde filling via collateral pathways. Significant delay indicates tissue at risk of infarction, and regions with severely reduced flow are likely to already be irreversibly injured. The use of automated processing software to apply delay and blood flow thresholds to CT perfusion allows reasonably accurate determination of tissue viability in individual patients.

The 2018 DAWN<sup>14</sup> and DEFUSE 3<sup>15</sup> randomised controlled trials showed large absolute benefits of thrombectomy in patients up to 24 hours after the time they were last seen well if they had favourable advanced brain imaging. Recovery to independence occurred in 47% of patients with thrombectomy compared with 15% of controls, and there was significant improvement across the spectrum of disability<sup>14,15</sup> Having broader, simpler eligibility criteria, essentially requiring < 70 mL irreversible injured brain, DEFUSE 3 allowed inclusion of additional patients, who experienced comparable treatment benefit.<sup>15</sup> This has led to a major expansion in access to endovascular thrombectomy for patients with wake-up onset or delayed presentation, including rural patients with stroke. However, the proportion of patients with favourable imaging declines rapidly, so the imperative to treat as fast as possible remains.

### **Risks**

The risks of endovascular thrombectomy are relatively low when performed by expert neurointerventionalists. Arterial injury or perforation occurs in less than 2% of patients.<sup>42</sup> Symptomatic intracerebral haemorrhage occurred in 4.3% of patients in the HERMES meta-analysis, similar to the control group.<sup>17</sup> Intravenous thrombolysis was administered to all eligible patients in the trials. Ongoing studies are testing whether intravenous thrombolysis is beneficial in patients who have immediate access to on-site thrombectomy. However, available observational data suggest improved recanalisation and outcomes in patients pre-treated with thrombolysis, without increased risk of symptomatic intracerebral haemorrhage.<sup>49</sup> Most Australian patients with stroke currently do not present directly to an endovascular-capable hospital; therefore, “bridging” thrombolysis preceding thrombectomy remains standard — analogous to the pharmaco-invasive strategy for ST-elevation myocardial infarction when a delay of more than 90 minutes from first medical contact to reperfusion is expected.<sup>50</sup> Stroke thrombectomy has also not yet achieved the over 95% success rate of percutaneous coronary intervention for ST-elevation myocardial infarction, with about 75% of patients achieving more than 50% reperfusion, but only 32% of patients achieving more than 90% reperfusion in the positive trials.<sup>17</sup> Therefore, intravenous thrombolysis may provide some benefit for patients in whom thrombectomy fails to fully reperfuse the brain.

## **Systems of care**

The evidence from randomised trials has indicated a strong time dependence of reperfusion therapies. It therefore follows that the greatest value for money in maximising benefits for patients comes from delivering these proven therapies faster and to a larger proportion of appropriate patients.

### **Ambulance assessment and pre-notification**

Ambulance paramedics should use validated stroke screening tools to identify suspected stroke.<sup>21</sup> Patients within potential treatment time frames (now 24 hours) should be transported to the nearest stroke-capable hospital and pre-notified to the receiving emergency department so that the stroke team can prepare in advance of the patient's arrival and clear the CT scanner. Pre-notification should include clinical details, name and date of birth to allow record searching and pre-ordering of the CT scan. On arrival, after a brief check to ensure the patient is stable from a cardiorespiratory viewpoint, the patient should be transported directly to the CT scanner for imaging, as this is the rate-limiting step in determining treatment. Clinical examination can occur en route to the scanner. These simple workflow changes, rather than taking the patient to an emergency cubicle, offloading from the ambulance stretcher and assessing the patient extensively before the CT scan, save significant time and translate to improved patient outcomes.<sup>51,52</sup>

With the advent of endovascular thrombectomy and the potential for treatment beyond 6 hours, routine acquisition of CT perfusion and CT angiography of the aortic arch to the cerebral vertex to identify a therapeutic target is advisable, in addition to non-contrast CT brain scan (Box 4).<sup>53</sup>

### **Pre-hospital triage**

As in-hospital workflow improves, the bulk of the onset to treatment time for patients with stroke now elapses pre-hospital. Community awareness is a prerequisite and, ideally, everyone would know the “facial droop, arm weakness, speech disturbance, time to call 000” (FAST) stroke recognition message and dial “000” immediately. The standard pre-hospital approach to patients with suspected stroke is to transport them directly to the nearest thrombolysis-equipped stroke unit if that is possible within a reasonable time frame (often < 60-minute drive time). States such as Victoria are well served with a network of such hospitals linked by telemedicine, and 99% of the population are within a 60-minute drive time of a stroke centre. However, other regions face greater geographical challenges. Even in metropolitan areas, endovascular thrombectomy is only available at selected centres. If a patient with large vessel occlusion first presents to a primary (thrombolysis-only) stroke centre, the process of assessment and transfer to a comprehensive (endovascular-capable) stroke centre introduces a delay of about 2 hours, which has major consequences for patient outcomes.<sup>54</sup> For this reason, various clinical triage tools have been introduced to assist paramedics in identifying likely large vessel occlusion stroke so that they can bypass directly to an endovascular-capable hospital. In Western Australia, the Rapid Arterial Occlusion Evaluation (RACE) scale (developed in Catalonia, Spain)<sup>55</sup> has been successfully implemented. In Victoria, a simpler algorithmic approach called Ambulance Clinical Triage for Acute Stroke Treatment (ACT-FAST) is currently being trialled (Box 5).<sup>56</sup>

### **Mobile stroke units**

The high technology solution to reducing treatment times is to bring the traditionally hospital-based elements of care to the pre-hospital environment. Mobile stroke units are ambulances equipped with CT scanners and clinicians specialised in stroke.<sup>57</sup> The CT brain scan allows definitive diagnosis of intracerebral haemorrhage versus ischaemic stroke, permitting pre-hospital thrombolysis. CT angiography allows definitive diagnosis of large vessel occlusion requiring thrombectomy, permitting bypass directly to an endovascular-capable hospital. Blood pressure management, reversal of anticoagulants and treatment of seizures are also possible. Pre-hospital thrombolysis and triage to a thrombectomy-

capable hospital cuts treatment delays substantially, and trials to evaluate the clinical effectiveness of such interventions are ongoing.

### **Intracerebral haemorrhage**

About 15% of strokes in Australia are due to intracerebral haemorrhage but these contribute to 50% of the mortality from stroke.<sup>58</sup> In other regions (eg, western China) intracerebral haemorrhage causes up to 40% of all stroke cases.<sup>59</sup> Stroke unit care benefits all stroke subtypes, including intracerebral haemorrhage. The Australian-led INTERACT-2 randomised controlled trial showed a modest 3% absolute reduction in death and disability, with no safety concerns when the systolic blood pressure was lowered to 140 mmHg.<sup>19</sup> The guidelines recommend lowering blood pressure to this level but not substantially below,<sup>21</sup> as results of the more intensive ATACH-2 randomised controlled trial did not show benefit of targeting a systolic blood pressure of 120 mmHg, which was associated with increased renal adverse events.<sup>60</sup> Ongoing trials of haemostatic agents (eg, tranexamic acid, recombinant activated factor VII) and minimally invasive surgery provide hope for future effective treatments of this major cause of disability.

### **Cytoprotection and neurorecovery**

As yet, there are no proven strategies to protect the brain from the effects of stroke other than reperfusion. Multiple trials of neuroprotective agents have been unsuccessful. However, with the advent of effective reperfusion with endovascular thrombectomy, human stroke treatment more closely resembles the temporary middle cerebral artery occlusion model used in animal studies, and a new wave of trials are underway to re-examine whether cytoprotection is feasible in humans.

### **Secondary prevention**

There is a strong evidence base for secondary stroke prevention, which relies on accurate determination of stroke mechanism. The guidelines now recommend the imaging of the vessels from aortic arch to cerebral vertex with CT angiography as the preferred vascular imaging approach to identify atherosclerosis, dissection and other arterial lesions.<sup>21</sup> Cardiac investigations search for a cardiac source of embolism. In particular, prolonged monitoring for paroxysmal atrial fibrillation is increasingly used, as the traditional 24-hour Holter monitoring has very limited sensitivity for this high risk cause of stroke that warrants anticoagulation. Implantable loop recorders can provide 3 years of continuous cardiac monitoring with an atrial fibrillation yield of 30% in patients with stroke without clear cause for their stroke, compared with 3% using standard clinical surveillance.<sup>61</sup> Consumer electronics have increasingly sophisticated electrocardiogram monitoring capabilities and may play an important role in atrial fibrillation detection.

High blood pressure is a risk factor for cardiovascular disease in general, but particularly for intracerebral haemorrhage and ischaemic stroke. Most patients who have had a stroke should take blood pressure-lowering medication unless contraindicated by symptomatic hypotension.<sup>21</sup> The SPRINT trial, which showed that a systolic blood pressure target of 120 mmHg improved outcome versus a 140 mmHg target, specifically excluded patients with stroke, but did reduce the incidence of stroke.<sup>62</sup> Post-hoc analysis of the PROGRESS trial of perindopril–indapamide for secondary stroke prevention also suggested the benefit of lowering the blood pressure, regardless of the baseline blood pressure.<sup>63</sup>

Cholesterol-lowering also reduces the risk of ischaemic stroke. Atorvastatin 80 mg showed significant benefits<sup>64</sup> and current guidelines recommend the use of a high dose, high potency statin.<sup>21</sup> The role of proprotein convertase subtilisin/kexin type 9 (PCSK9) inhibitors is evolving but may certainly benefit patients with stroke, particularly those who are intolerant of statins.

Antithrombotic therapy is essential after ischaemic stroke. Recent publications have re-emphasised the value of early aspirin commencement.<sup>65</sup> Clopidogrel or aspirin–dipyridamole provide slightly greater efficacy than aspirin alone.<sup>21</sup> Combined aspirin and clopidogrel for about 3 weeks reduces the risk of recurrent stroke,<sup>66,67</sup> but longer term dual antiplatelet therapy is not used in stroke prevention due to increased bleeding and minimal benefit.<sup>21</sup>

For patients with atrial fibrillation, aspirin has no role in stroke prevention.<sup>21</sup> Anticoagulation for atrial fibrillation remains underutilised in Australia and globally. The guidelines recommend first-line direct oral anticoagulants for non-valvular atrial fibrillation (ie, no mechanical prosthetic heart valve or moderate to severe mitral stenosis), provided the creatinine clearance rate is adequate (> 30 mL/min for dabigatran and rivaroxaban; > 25 mL/min for apixaban using the Cockcroft–Gault formula).<sup>21</sup> Warfarin (target international normalised ratio, 2.0–3.0) is still initiated in patients who are unable to take direct oral anticoagulants. Common problems with anticoagulation management leading to preventable strokes are the overestimation of the risk of bleeding (eg, falls) relative to the risk of stroke, leading to underutilisation of anticoagulation, and the use of lower doses than recommended in the applicable product information. For patients with genuine contraindications to anticoagulation, left atrial appendage occlusion is emerging as a relevant treatment option.<sup>68</sup> Anticoagulation in suspected embolic stroke without proving the diagnosis of atrial fibrillation has not been supported by recent randomised trials.<sup>69</sup>

Patent foramen ovale has been a controversial potential cause of stroke. However, new trials published in 2017 using improved patient selection to thoroughly exclude alternative stroke mechanisms have demonstrated the unequivocal benefit of percutaneous device closure of patent foramen ovale in highly selected patients.<sup>70–72</sup> The absolute benefit is about 1% per annum. However, unlike front-loaded atherosclerotic risk that decreases over time, risk of stroke related to patent foramen ovale appears to accumulate at a constant rate, implying a major lifetime risk reduction for young patients.

## Conclusion

Stroke medicine is an exciting field with major advances in recent years. In particular, reperfusion therapies for ischaemic stroke have transformed the prognosis for long term disability. However, optimal implementation to maximise patient benefit requires increased community recognition, immediate “000” activation, faster emergency treatment systems and innovative therapeutic developments to further improve outcomes.

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## Author details

Bruce CV Campbell

Royal Melbourne Hospital, University of Melbourne, Melbourne, VIC.

**Bruce.Campbell@mh.org.au**

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[Insert boxes]

[Box 1; cam\_mja18.00888\_gr1]

## 1 Ischaemic stroke pathophysiology\*

[Box 1 foot]

ACA = anterior cerebral artery. PCA = posterior cerebral artery. \* Most ischaemic stroke is thromboembolic: a thrombus forms in the heart or large arteries and then migrates to the intracranial circulation. When this occurs, symptoms develop immediately as neurons lose the oxygen and glucose required for electrical function. However, collateral vessels are often able to sustain metabolic viability in the affected brain for some time. This hibernating but salvageable brain is termed the ischaemic penumbra and is the reason that rapid reperfusion can dramatically alter the natural history of an otherwise disabling stroke. Collateral circulation is highly variable between individuals and the genetic and environmental determinants are poorly understood. Over time, collateral circulation tends to fail leading to growth in the irreversibly injured ischaemic core. The patient's clinical deficit generally remains stable but it has now become irreversible. One of the important insights from recent endovascular reperfusion trials in patients 6–24 hours after stroke onset is that good collaterals and ischaemic penumbra can persist for many hours in a subgroup of patients. However, despite this initial resilience, without treatment the outcome is very poor with recovery to independence in about 15% of patients.

[Box 2]

## 2 Evidence-based acute therapies for stroke

Intervention	Indication	Outcome*	Relative risk (95% CI)	NNT
All stroke types and severities				
Stroke unit care <sup>2</sup>	All admitted patients with stroke	Death/dependency	0.94 (0.88–0.99)	28
Ischaemic stroke				
IV alteplase 0–4.5 h <sup>16</sup> (thrombolysis)	Potentially disabling stroke without excessive bleeding risk	Return to all usual activities (mRS, 0–1)	Treatment within 0–3 h	1.42 (1.21–1.68) 10
			Treatment within 3–4.5 h	1.17 (1.05–1.31) 19
Endovascular thrombectomy 0–6 h <sup>17</sup>	Large vessel occlusion (internal carotid or proximal middle cerebral artery) <sup>†</sup>	Death/dependency	0.74 (0.67–0.80)	5
Endovascular thrombectomy 6–24 h <sup>14,15</sup>	Large vessel occlusion and favourable CT perfusion/MRI (core < 70 mL)	Death/dependency	0.62 (0.53–0.71)	3
Aspirin 0–48 h <sup>3,4,18</sup>	All ischaemic stroke (unless atrial fibrillation or allergy)	Recurrent stroke/death	0.90 (0.84–0.96)	109
Hemicraniectomy 0–48 h <sup>5</sup>	Malignant MCA infarction (mass effect, reduced conscious state), age < 60 years	Death/severe disability <sup>‡</sup>	0.72 (0.54–0.96)	4.6
Intracerebral haemorrhage				
BP-lowering in ICH (target 140 mmHg) <sup>19</sup>	BP > 150 mmHg (maintain at 140 mmHg for the first 7 days)	Death/dependency	0.87 (0.75–1.01)	28
		Disability severity <sup>§</sup>	0.87 (0.77–1.00)	na

BP = blood pressure. CT = computed tomography. ICH = intracerebral haemorrhage. IV = intravenous. MCA = middle cerebral artery. MRI = magnetic resonance imaging. mRS = modified Rankin Scale. na = not applicable. NNT = number needed to treat. \* Outcomes and NNT assessed at 3 months, except hemicraniectomy (12 months). † Basilar artery occlusion excluded from these trials. ‡ mRS 4–6: mRS 6 = death, mRS 5 = nursing home, mRS 4 = need for assistance with personal activities of daily living. § Change by at least one level on the mRS.

[Box 3; cam\_mja18.00888\_gr2]

## 3 Thrombus retrieved by endovascular thrombectomy with a stent retriever device

[Box 4; cam\_mja18.00888\_gr3]

#### 4 Imaging in a patient\* who underwent endovascular thrombectomy†

[Box 4 foot]

\* An older patient living independently at home presented one hour after onset of left hemiparesis, dysarthria, hemianopia and inattention.  
† Multimodal computed tomography (CT) including (A) non-contrast scan to exclude haemorrhage and established infarction, CT perfusion confirming diagnosis of ischaemic stroke with (B) delayed blood flow arrival in right middle cerebral artery territory (ie, territory at risk), and (C) preserved cerebral blood volume suggesting salvageable tissue. (D) CT angiography for confirmation of right middle cerebral artery occlusion (arrow) and thrombectomy planning. (E) Pre-thrombectomy and (F) post-thrombectomy angiography showing complete reperfusion 2 hours after onset. (G) Follow-up diffusion magnetic resonance imaging scan showing successful tissue salvage. The patient was discharged directly home on Day 3.

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[Box 5]

5 Pre-hospital identification of suspected large vessel occlusion

RACE* score	Points	ACT-FAST* algorithm
Facial droop		<b>Step 1:</b> arm drift to stretcher within 10 seconds
Absent	0	▪ No = ACT-FAST negative
Mild	1	▪ Yes = proceed to Step 2
Severe	2	
Arm drift		<b>Step 2:</b>
Absent/mild	0	▪ If right arm weak = assess language:
Drifts to stretcher	1	▶ no deficit = ACT-FAST negative;
Cannot get antigravity	2	▶ significant deficit = proceed to Step 3
Head and gaze deviation		▪ If left arm weak = assess eye deviation and shoulder tap:
Absent	0	▶ no eye deviation and patient appropriately localises to a tap on left shoulder and calling their name = ACT-FAST negative;
Present	1	▶ significant deficit = proceed to Step 3
Aphasia (if right arm weak)†		
Both tasks performed correctly	0	
One error	1	
Two errors	2	
Agnosia (if left arm weak)‡		<b>Step 3:</b> eligibility and mimic exclusion (all criteria needs to be fulfilled)
Both tasks performed correctly	0	▪ Stroke onset (last known to be well) < 24 hours
One error	1	▪ Pre-morbid independent function
Two errors	2	▪ Not comatose, no seizure at onset, no known malignant brain tumour, glucose level > 2.8 mmol/L
RACE score ≥ 5 indicates high probability of large vessel occlusion		All 3 ACT-FAST steps positive indicates high probability of large vessel occlusion

ACT-FAST = Ambulance Clinical Triage for Acute Stroke Treatment. RACE = Rapid Arterial Occlusion Evaluation. \* Both scales test arm drift by positioning both arms at 90 degrees and observing for unilateral drift over 10 seconds. † RACE tests aphasia by asking the patient "close your eyes" and "make a fist" and evaluating if the patient obeys. ‡ RACE tests agnosia by asking the patient, while showing them the paretic arm, "whose arm is this" and evaluating if the patient recognises their own arm; and by asking the patient "can you lift both arms and clap" and evaluating if the patient recognises their functional impairment.