Role of transcranial Doppler ultrasonography in stroke
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Transcranial Doppler sonography is a non-invasive, non-ionising, inexpensive, portable and safe technique that uses a pulsed Doppler transducer for assessment of intracerebral blood flow. This article deals with the principles and technique of transcranial Doppler sonography. It gives a brief overview of its use in evaluation of intracranial steno-occlusive disease, subarachnoid haemorrhage, and extracranial diseases (including carotid artery disease and subclavian steal syndrome). The role of transcranial Doppler in detection of microembolic signals and evaluation of right to left shunts is also dealt with. Finally, its use in acute stroke is briefly outlined.

Ultrasound has been used for the evaluation of cerebrovascular disease for over a decade and has made considerable progress. Transcranial Doppler sonography is a non-invasive, non-ionising, inexpensive, portable and safe technique that uses a pulsed Doppler transducer for assessment of intracerebral blood flow.

With the advent of thrombolytic treatment for acute ischaemic stroke, the internist would probably benefit from having a knowledge of transcranial Doppler ultrasound (TCD), which is a useful tool for the detection of occlusion of intracranial vasculature. In addition, success of thrombolytic treatment can also be assessed by TCD.

This review article aims to provide a basic understanding about the use of TCD in clinical practice. A brief outline is provided of the principles and techniques of TCD and its role in acute ischaemic stroke, including abnormalities affecting both intracranial and extracranial parts of vessels supplying the brain. We then explore the role of TCD in the detection of microembolic signals, which help in stratification of risk of recurrence of stroke or transient ischaemic attack (TIA), and its role of in the detection and quantification of right-to-left shunts. We also outline the possible role of TCD in subarachnoid haemorrhage and subclavian steal syndrome. Finally, the role of TCD during carotid endarterectomy is discussed (box 1).

PRINCIPLES

The “Doppler effect”, which was enunciated by Christian Andreas Doppler in 1842, forms the basis of TCD. In 1982, Aaslid and colleagues introduced a 2 Mhz Doppler which allowed adequate penetration through the intact skull. A pulsed wave transducer emits (insonant) waves, and then receives their reflections off the surfaces of the red blood cells within the intracranial vasculature. This information is analysed by a computer to give us both numerical and visual output, which is useful for inferring the flow characteristics within a blood vessel.

A particular vessel may be traced superficially or deep (that is, proximally and distally)—for example, the middle cerebral artery may be traced at a depth of between 30–60 mm via the transtemporal window. Transcranial Doppler spectra obtained may be single-gated TCD or multigated TCD (M mode Doppler).

The peak systolic and diastolic velocities, mean flow velocity and Gosling pulsatility index (PI) are routinely calculated and displayed. Transcranial Doppler diagnoses are based on the detection of altered blood flow velocity, absence of blood flow, changes in the spectral waveform, or changes in pulsatility in a specific vessel.

TECHNIQUE

TCD is performed at the bedside. The first step is to localise a cranial “window” where the ultrasound beam can penetrate without being excessively dampened. The three main windows for accessing the intracranial arteries are listed below and shown in fig 1:

- **Transtemporal window**—found between the angle of the eye and the pinna above the zygomatic ridge and is the major route for insonating the anterior, middle and posterior cerebral arteries
- **Transorbital window**—through the eye for insonation of the ophthalmic artery and the siphon of the internal carotid artery
- **Transforaminal window**—through the foramen magnum insonated from the top of the neck below the occiput for the basilar artery and the intracranial segments of the vertebral arteries

TCD is a blind technique. The various blood vessels are identified from the window used, the depth of insonation, the direction of blood flow with respect to the probe, and characteristics of the TCD waveform.

The middle cerebral artery (MCA) is identified through the transtemporal window, with the flow direction normally towards the probe, about 30–60 mm from the skull surface. At about 60 mm,
the internal carotid artery (ICA) divides into the MCA and anterior cerebral artery (ACA), which flows away from the probe, and this bifurcation is one of the most important reference points for TCD. ACA is insonated between 65–80 mm with the probe directed anterosuperiorly through the transtemporal window. The posterior cerebral artery (PCA) is insonated at 55–70 mm with the probe directed posteroinferiorly through the transtemporal window. At times, provocative manoeuvres are used to confirm the identity of the vessels. Anomalies of the circle of Willis are common and a thorough knowledge of the anatomy of the blood vessels is an essential requisite for accurate interpretation. The technique is highly operator dependent and requires considerable skill and experience (figs 2–4).

TCD can be performed at the bedside and repeated as needed (even for continuous monitoring). It is non-invasive and does not need a contrast agent. Its chief limitations are that it is an operator dependent technique and attenuation of ultrasound occurs through the skull and soft tissues. The mean loss of power through the skull can be up to 80%. The transtemporal window can be absent in up to 5–20% of the patients, and the vessels cannot be properly insonated due to inadequacy of the acoustic windows.

However, with skill and experience in detecting accurate signals, TCD can be used at the bedside to help with stroke management. The pros and cons of TCD and its use in clinical practice are summarised in box 2.

**Box 1: Use of transcranial Doppler ultrasound**

- Detection of site/degree of stenosis/occlusion of cerebral vasculature
- Assessment of recanalisation following occlusion (with/without thrombolytic treatment)
- Assessment of collateral flow in intracranial vasculature in cases of critical carotid artery stenosis (extracranial)
- Detection of microemboli: stratification of risk of recurrence of stroke/TIA
- Detection and quantification of right to left shunts
- Detection of degree of vasospasm following subarachnoid haemorrhage
- Complementary to duplex carotid scan in diagnosis of subclavian steal syndrome
- Intraoperative monitoring of carotid endarterectomy

**Figure 1** The three main windows for accessing the intracranial arteries.

**ACUTE ISCHAEMIC STROKE**

Cerebral angiography shows acute occlusion in 76% of acute MCA territory infarcts within 6 h of stroke onset. Follow up studies show spontaneous recanalisation in the majority by the end of 48 h and in up to 86% by 2 weeks. TCD can detect these angiographic occlusions with high sensitivity and specificity, and has a high positive predictive value.

TCD has a specificity of 90% in demonstrating MCA occlusions in patients with acute MCA stroke within 5 h. Alexandrov et al have shown major arterial occlusions in 69% of patients with acute hemispheric stroke, who may be eligible for thrombolytic treatment. Recanalisation can be inferred by TCD by the appearance of flow in the vessel or an improvement in...
the flow, with or without reduction in the PI in the proximal segments of the vessel. Thus, in the setting of acute ischaemic stroke, TCD can reveal the presence of arterial occlusion and it can also show whether recanalisation has occurred following intravenous thrombolysis.

TCD is also useful in prognostication of stroke.

- A normal TCD at 6 h post-ischaemic stroke is an independent predictor of early improvement.
- TCD may predict occurrence of spontaneous haemorrhagic transformation in ischaemic infarcts. MCA occlusions within 6 h of stroke onset may be an independent predictor of spontaneous haemorrhagic transformation with a positive predictive value of 72%.
- In acute MCA stroke, blood flow velocity on TCD of <30 cm/s within 12 h after stroke correlated with poor recovery.
- Microemboli are an important independent predictor of early ischaemic recurrence in patients with stroke or TIA of arterial origin (see later).

**USE OF TCD IN EVALUATING INTRACRANIAL ARTERIAL DISEASE**

**Intracranial steno-occlusive disease**

Intracranial atherosclerosis is responsible for up to 10% of TIA and strokes. Sensitivity, specificity, and positive and negative predictive value of TCD are generally higher in detecting abnormalities of the anterior circulation than in the vertebrobasilar circulation, as the latter has more anatomical variations and can be difficult to localise for TCD insonation.

TCD can ascertain higher grades of stenosis (based upon elevated blood flow velocities) fairly accurately. However, its

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**Figure 3** Middle cerebral artery (MCA)–anterior cerebral artery (ACA) tracing at the division of the internal carotid artery into the MCA (tracing above baseline) and the ACA (below baseline).

**Figure 4** Normal anterior cerebral artery (ACA) tracing below the baseline reflecting flow away from the probe at a depth of 66 mm (transtemporal window).
role in diagnosing milder grades of stenosis is still uncertain. The primary sign of stenosis is a focal increase in mean flow velocity at the site of luminal narrowing. Secondary TCD signs of stenosis include decreased velocity and increased pulsatility upstream from the lesion and abnormal flow immediately downstream from the lesion (box 3).

With a tight narrowing (>50% reduction in lumen diameter), the velocity increases dramatically at the site of narrowing (mean flow velocity ≥80 cm/s and a velocity difference of ≥30% compared to the control side). In addition, there is lowering of signal intensity because of compromised blood flow due to narrowing. MCA stenosis has been most widely studied and can be diagnosed with a sensitivity of 86% and a specificity of 99% using TCD.

Distal MCA stenosis, however, can be missed with TCD, as the distal branches are not well visualised. Stenosis-like waveforms also occur in other conditions like arterial spasm, as in subarachnoid haemorrhage (SAH) or intracerebral haemorrhage, or due to increased collateral flow through hypoplastic communicators. In the case of spasm the changes tend to occur over a longer segment and/or involve multiple arteries.

Intracranial arterial occlusions have been diagnosed using TCD with even greater accuracy and Demchuk et al have noted detailed diagnostic criteria for occlusion of large arteries. They have used the TIBI (Thrombolysis In Brain Ischaemia) classification to describe changes in the waveform found in association with recanalisation during recombinant tissue plasminogen activator treatment at or just beyond the site of occlusion.

In general, the criteria for diagnosing an occluded artery based on TCD include: (1) absence of signal from the artery, taking care that a good acoustic window has been obtained by confirming flow in other vessels from the same window; and (2) sonographic evidence of collateral flow.

In general TCD has a high specificity, though sensitivity of TCD is lower in detecting intracranial stenosis and occlusions.

**USE OF TCD IN EVALUATION OF EXTRACRANIAL ABNORMALITIES**

**Extracranial carotid lesions**

Extracranial ICA disease is a significant cause of neurologic deficits ranging from TIAs to progressive ischaemic eye disease and cerebral infarction. The risk rises as the degree of stenosis increases. Carotid duplex sonography is a sensitive and specific technique for demonstrating the presence and degree of narrowing of the proximal ICA. TCD is a useful adjunct to this investigation in evaluating its haemodynamic consequences on the intracranial circulation. In patients with haemodynamically significant extracranial ICA disease TCD may demonstrate: (1) a decrease in mean blood flow velocity, with diminished pulsatility in the ipsilateral MCA together with normal flow in the contralateral MCA; (2) diminished flow acceleration in the ipsilateral MCA; (3) increased blood flow in potential collateral routes of circulation, typically the contralateral ACA, PCA, anterior communicating (Acom) and posterior communicating (Pcom) arteries; and (4) reversed direction of flow in the ipsilateral ACA and PCA and ophthalmic arteries (fig 5).

**TCD and transient microembolic signals**

Microemboli in the cerebral circulation can originate from atherothrombotic lesions of the carotid arteries and cardiac sources. Interventional procedures such as cerebral angiography, carotid angioplasty, carotid endarterectomy and cardio-pulmonary bypass may also give rise to microemboli. In addition, in right to left shunts microemboli may occur.

Gaseous or solid microemboli within the MCA can be detected by TCD as high intensity transient signals (HITS), also called microembolic signals (MES). They are characterised by: (1) a duration <300 ms; (2) an amplitude that is 3 dB higher than the background blood flow signal; (3) are typically unidirectional and occur randomly within the cardiac cycle; and (4) produce a characteristic sound like a “moan” or “chirp” on audio signal (fig 6). The vessels have to be monitored for at least 30–60 min for detection of MES. An individual MES does not cause neurological symptoms but may represent an early warning sign for greater risk of future TIA/stroke. Regarding characterisation of the nature of emboli, it has been shown that gaseous emboli have signals of higher amplitude and intensity compared to formed solid particles.

MES correlate well with prior ischaemic events and may represent a higher risk for future recurrence ischaemic events. Higher prevalence rates of MES have been reported in strokes caused by large vessel disease and in cardioembolic strokes as compared to lacunar strokes. In asymptomatic patients with a critical ICA stenosis, when the microembolic signal rate is >2/hour in the ipsilateral MCA, there is an associated increased risk of development of ischaemia. MES arising from cardiac sources have higher total signal power, longer signal duration,
and higher percentage of signals occurring in diastole as compared to those seen in carotid artery disease.\textsuperscript{29}

TCD in ischaemic strokes due to right-to-left shunts

In young patients with cryptogenic stroke, right-to-left shunts (including patent foramen ovale (PFO)) are thought to be a risk factor for ischaemic stroke. However, whether it is only a chance association or a risk factor of stroke remains to be determined. Until recently, transoesophageal echocardiography (TOE) with a contrast agent was the only means of diagnosing right-to-left intracardiac shunts. It had many limitations—being a semi-invasive technique it relied on the compliance of the patient to a great extent and also could not always detect latent shunts. TCD with a gaseous contrast medium is now used for the diagnosis of right-to-left shunts.

When a gaseous contrast agent is injected into the peripheral vein of a patient with shunts, microbubbles pass from the right to the left circulation during cardiac cycle, and enter the systemic circulation; TCD picks up the microbubbles as

Figure 5  Effect of extracranial internal carotid artery (ICA) stenosis on cerebral haemodynamics (patient had high grade stenosis of proximal right ICA). (A) Decrease in mean flow velocity of right (ipsilateral) middle cerebral artery (MCA). (B) Normal flow in left (contralateral) MCA with increased flow in left (contralateral) anterior cerebral artery (ACA) (due to collateral flow). (C) Reversal of flow in right (ipsilateral) ACA. (D) Increased flow in left (contralateral) ACA.

Figure 6  Transcranial Doppler waveform demonstrating an emboli.
Box 4 Quantification of right-to-left shunt

- Test negative: no microbubble
- Low grade shunt: 1–10 microbubbles
- Medium grade shunt: >10 microbubbles but without “curtain effect”
- High grade shunt: curtain effect, seen when the microbubbles are so numerous as to be no longer distinguishable separately

TCD in subarachnoid haemorrhage

Focal or diffuse cerebral vasospasm may follow in about 30% of patients having SAH due to rupture of an intracranial aneurysm or other pathologic conditions. The temporal course of vasospasm varies. In most cases it develops on the third or fourth day and gradually increases for a week, and then peaks between the 11th and 17th day, before gradually abating.10–17 TCD monitoring enables identification of patients at particular risk of developing ischaemic events secondary to vasospasm. This could be avoided by adequate early medical treatment.18

TCD is a sensitive and specific technique for the detection of vasospasm. There is an increase in blood flow velocity in the MCA.19

Intraoperative TCD involves monitoring the velocity of flow of blood in the MCA during carotid endarterectomy. It assesses the adequacy of cerebral blood flow while the carotid artery is cross-clamped during carotid endarterectomy.44–45 Ischaemia during cross clamping is a classic complication and is considered severe if reduction in flow velocity is >85%, mild to moderate if reduction is between 60–85%, and absent if there is <60% reduction.

Hyperaemic phenomena may occur and result in a sudden increase in blood flow velocity in the MCA. Microemboli (MES/HITS) caused by operative manoeuvres such as shunt insertion or removal can be documented in real time with TCD.

Postoperatively TCD may help in diagnosing hyperperfusion syndromes caused by deficient autoregulation.46

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