Accurate classification of stroke has significant impact on patient care and conduction of stroke clinical trials. The current systems such as TOAST, SSS-TOAST, Korean TOAST, and A–S–C–O have limitations. With the advent of new imaging technology, there is a need to have a more accurate stroke subclassification system. Chinese ischemic stroke subclassification (CISS) system is a new two step system aims at the etiology and then underlying mechanism of a stroke. The first step classify stroke into five categories: large artery atherosclerosis (LAA), including atherosclerosis of aortic arch and intra-/extracranial large arteries, cardiogenic stroke, penetrating artery disease, other etiology, and undetermined etiology. The second step is to further classify the underlying mechanism of ischemic stroke from the intracranial and extracranial LAA into the parent artery (plaque or thrombosis) occluding penetrating artery, artery-to-artery embolism, hypoperfusion/impaired emboli clearance, and multiple mechanisms. Although clinical validation of CISS is being planned, CISS is an innovative system that offers much more detailed information on the pathophysiology of a stroke.

**Keywords:** ischemic stroke, subclassification, etiology, mechanism, Chinese
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of stenosis in the parent artery (detected by TCD, MRA, CTA, or DSA); (3) No evidence of potential cardiac-origin embolic cause; (4) Other possible causes have also been excluded.

**Cardiogenic stroke**

(1) Acute multiple infarcts, especially involving bilateral anterior and/or anterior and posterior circulations (including cortical infarcts) that have occurred closely in time; (2) No evidence of atherosclerosis on relevant intracranial or extracranial large arteries (vulnerable plaques or stenosis ≥50% or occlusion); (3) No evidence of other etiologies that can cause multifocal acute ischemic infarcts such as vasculitides, hemostatic disturbances, and tumorous embolism; (4) Evidence of cardiac disease that has a potential for embolism; (5) If the possibility of aortic arch atherosclerosis has been excluded, CS is definite. Otherwise, the category should be possible CS.

The potential lesions included under this category are: mitral stenosis, prosthetic heart valve, myocardial infarction within the past 4 weeks, mural thrombus in the left cavities, left ventricular aneurysm, any documented history of permanent or transient atrial fibrillation or flutter with or without spontaneous echo contrast or left atrial thrombus, sick sinus syndrome, dilated cardiomyopathy, ejection fraction <35%, endocarditis, intracardiac mass, PFO plus *in situ* thrombosis, PFO plus concomitant PE, or DVT preceding the brain infarction (Amarenco et al., 2009).

**Penetrating artery disease**

Acute isolated infarct in the territory of one penetrating artery caused by atherosclerosis at the proximal segment of the penetrating arteries or lipohyalinotic degeneration of arterioles is called penetrating artery disease (PAD).

**2. Intra- and extracranial large arteries atherosclerosis**

(1) Any distribution of acute infarcts (except isolated infarct in the territory of one penetrating artery), with evidence of atherosclerosis involving intracranial or extracranial large arteries (vulnerable plaques or stenosis ≥50%) that supply the area of infarction; (2) Concerning isolated penetrating artery territory infarct, the following circumstance should also be included in LAA: with evidence of atherosclerotic plaque (detected by HR-MRI) or any degree of significant aortic arch atherosclerosis (aortic plaques >4 mm and/or aortic thrombi, detected by HR-MRI/MRA and/or TEE; Reynolds et al., 2003; Harloff et al., 2008).

Figure 1 | Illustration of large artery atherosclerosis, cardiogenic stroke, and penetrating artery disease.

Figure 2 | Detailed illustration for intracranial large artery atherosclerosis and penetrating artery disease.
1. Parent artery (plaque or thrombus) occluding penetrating artery: Isolated acute infarct in penetrating artery territory, the parent artery has evidence of plaque or any degree of stenosis. For example, an isolated acute infarct located in the basal ganglia, and no other acute infarct in the territory supplied by the ipsilateral MCA; or an isolated acute infarct located in the pons, and no other acute infarct in the territory supplied by the basilar artery. The acute isolated infarct is posited to be caused by the plaque of the parent artery protruding and subsequently occluding blood flow to the penetrating artery.

2. Artery-to-artery embolism: Imaging shows small cortical infarcts or a single territory infarct in the area supplied by the relevant intracranial or extracranial artery atherosclerosis. No borderzone infarct related to this diseased artery. This diagnosis is confirmed if the infarcts are multiple, or the single infarct is accompanied by TCD–MES on TCD. However, a single cortical or territorial infarct without obtaining MES can also be diagnosed as artery-to-artery embolism.

3. Hypoperfusion/impaired emboli clearance: the acute infarcts occur solely in the borderzone area. No acute cortical or territorial infarct related to this diseased artery. The degree of stenosis of the clinically relevant intracranial or extracranial artery is usually >70% with or without evidence of hypoperfusion or poor collateral compensation in the related region of perfusion.

4. Multiple mechanism: two or more underlying mechanisms mentioned above combined together.

**Other etiologies**
Evidence of other specific diseases (e.g., vascular related disease, infective disorder, inherited disease, hematological system disorder, vasculitis), that are relevant to the index stroke and can be demonstrated by blood tests, cerebrospinal fluid (CSF) tests, and vascular imaging. The possibility of LAA or CS has been excluded.

**Undetermined etiology**
No evidence of any specific potential etiology that is clinically relevant to the index stroke.

*Multiple*: Evidence of more than one potential cause, but difficult to determine which was the relevant cause of the index stroke.

*Unknown*: No determined cause is responsible for the index stroke unless more investigations would be performed.

*Inadequate evaluation*: Routine assessments of intracranial and extracranial arteries or heart are not completed, which makes the etiology undetermined.

**THE SECOND STEP, DEFINING THE UNDERLYING MECHANISM FOR ISCHEMIC STROKE OF INTRACRANIAL OR EXTRACRANIAL LARGE ARTERYATHEROSCLEROSIS (FIGURE 3)**

In the CISS system, the underlying mechanisms of ischemic stroke caused by intracranial or extracranial LAA are further defined as the parent artery (plaque or thrombus) occluding penetrating artery, artery-to-artery embolism, hypoperfusion/impaired emboli clearance, and multiple mechanism.

**ILLUSTRATIONS OF CISS**
Comparing to other classification systems, CISS has the following major differences: (1) aortic arch atherosclerosis belongs to LAA. Given that the intrinsic lesion is atherosclerosis, it is more reasonable to classify aortic arch atherosclerosis into LAA (Amarenco et al., 2009). (2) In acute isolated penetrating artery territory infarcts, they would be classified into LAA if there was atherosclerotic plaque or stenosis of the parent artery regardless of the presence of any plaques or stenosis. (3) PAD is a new concept. Excluding other diseases, acute isolated penetrating artery territory infarcts are...
considered to be caused by the penetrating artery lesions. Pathology has shown that atheromatous disease at the proximal segment of the penetrating arteries mainly lead to symptomatic infarcts, while fibrohyalnosis of arterioles is mainly associated with asymptomatic lacunes or diffuse white matter hyperintensities (Fisher, 1969, 1979; Fisher and Caplan, 1977).

Caplan (1989) noted that intracranial branch atheromatous disease was a neglected, understudied, and underused concept. Unfortunately, in the past 20 years, it has been rare to find vascular pathology studies that trace the entire penetrating arteries and their parent arteries. Researchers either studied the parent artery (leading to the finding of atherosclerosis of parent artery) or more distal segment of the penetrating artery and the arteriole (leading to the finding of arterial fibrohyalnosis; Lammie et al., 1979; Chen et al., 2008; Klein et al., 2010). Fibrohyalnosis of small arteries and arterioles causes symptomatic acute penetrating artery infarcts less often than atheromatous branch disease. Therefore, the current opinion that symptomatic penetrating artery territory lesions are equivalent to intrinsic small vessel disease is inappropriate.

Even if atherosclerotic lesions involving the proximal segment of the penetrating arteries was not considered the main cause of symptomatic penetrating artery territory infarcts, it should be taken into account as well as arteriolar fibrohyalinosis. Diffuse white matter hyperintensities can reflect arteriolar fibrohyalinosis, but atherosclerotic lesions and fibrohyalinosis often coexist (Fisher, 1965), and current imaging technology can not show the penetrating artery wall directly. Therefore, clinically it is difficult to distinguish between the two. With further improvements in imaging, there could develop further PAD subclassifications. In addition, the introduction of the concept of PAD not only avoids confusion with intrinsic small vessel disease, but also avoids confusion with the "lacunar" concept.

In LAA, several outcomes of atherosclerotic stenosis or occlusion of the carotid artery may occur: (1) There is no infarct or ischemic symptoms because the fragments of plaques or thrombi do not dislodge and there is a well-compensated Willis’ circle. (2) The fragments of plaques or thrombi do not dislodge, but borderzone infarction may occur if there is a poorly compensated Willis’ circle. Cerebral perfusion may be compromised due to hypoperfusion events such as sudden drop in blood pressure or in other systemic hypoperfusion events (Momjian-Mayor and Baron, 2005). (3) If the fragments of plaques or thrombi dislodge and travel to distal branches, this may lead to a mechanism of artery-to-artery embolism or impaired emboli clearance depending on the location of the infarcts (Fisher and Caplan, 1977; Derdeyn et al., 2001; Chaves et al., 2003). The mechanism of vertebral artery infarction is similar to that of the extracranial carotid artery.

For intracranial large arteries (MCAs), thrombosis on atheroma will make the stenosis more severe and may lead to complete occlusion of the artery. There are several processes and outcomes of atherothrombotic stenosis or occlusion of the MCA (Wong et al., 2002; Bang et al., 2004; Lee et al., 2005; Caplan et al., 2006): (1) The fragments of plaques or thrombi do not dislodge, the penetrating artery is not blocked by plaques or thrombi, there is a well-compensated pia mater collateral circulation and the newly generated collateral arteries can supply the territory of the penetrating arteries. This MCA territory can endure long-term ischemia and there will be no infarcts in this region even after the MCA is completely occluded. (2) The fragments of plaques or thrombi do not dislodge, the penetrating artery is not blocked by atherosclerosis and there is poorly established collateral circulation. Borderzone infarction may occur if cerebral perfusion is compromised due to hypoperfusion. This mechanism is called hypoperfusion. (3) If the orifice of one or more penetrating arteries are blocked by plaque or thrombi, infarction will occur in the regions of the penetrating arteries. This mechanism is called parent artery occluding penetrating artery. (4) If the fragments of plaques or thrombi are dislodged and travel to distal branches, this mechanism is labeled artery-to-artery embolism or impaired emboli clearance based on the location of the infarcts. The underlying mechanism of infarction related to the basilar artery is similar to that of MCA.

The mechanism of hypoperfusion refers to borderzone infarction from compromised hemodynamics. The mechanism of impaired emboli clearance means that the borderzone infarct is posited to result from impaired clearance of emboli due to both hypoperfusion and embolism. Actually, it is difficult to completely separate these two mechanisms. For a patient with ICA or MCA stenosis (>70%), develop a borderzone infarct and TCD also detects MES, then the underlying mechanism would be impaired emboli clearance. On the other hand, we cannot exclude the existence of impaired emboli clearance mechanism even though there is no MES detected by TCD. Therefore, it would be more appropriate to combine the two mechanisms together.

CONCLUSION

Chinese ischemic stroke subclassification introduces a new way of subclassifying acute ischemic stroke that takes into consideration of both etiological and pathophysiological information. In CISS, aortic arch atherosclerosis is classified into LAA, and a new subclassification of PAD has been created. The underlying mechanism of ischemic stroke from the intracranial and extracranial LAA has been subclassified into four categories: the parent artery (plaque or thrombus) occluding penetrating artery, artery-to-artery embolism, hypoperfusion/impaired emboli clearance, and multiple mechanisms. CISS is an improved and more rational way of stroke subtyping. It has deepened our understanding into the pathophysiology of stroke.

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