Cerebral Small Vessel Disease and Arterial Stiffness: Tsunami Effect in the Brain?

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Key Words
Ankle-brachial index · Arterial stiffness · Cerebral small vessel disease · Cognitive impairment · Pulse wave velocity

Abstract
Background: Cerebral small vessel diseases, including silent lacunar infarcts, white matter hyperintensities, and microbleeds, pose a risk for cerebrovascular disease, cognitive impairment, and the geriatric syndrome via effects on arterial stiffness. However, the vascular, physiological, and metabolic roles of arterial stiffness in cerebral small vessel diseases remain unclear. Summary: Arterial stiffness can be assessed using various indicators such as the ankle-brachial index, pulse wave velocity, cardio-ankle vascular index, and augmentation index. Arterial stiffness is independently associated with all components of cerebral small vessel disease including silent lacunar infarcts, white matter hyperintensities, and microbleeds, although there are some methodological differences between the various surrogate markers. Evidence of arterial stiffness indicates microvessel arteriosclerosis presenting with vascular endothelial dysfunction. Further, vascular narrowing due to atherosclerosis and vascular stiffness due to lipohyalinosis can accelerate the pulse waves. This hemodynamic stress, pulsatile pressure, or blood pressure variability can cause a ‘tsunami effect’ towards the cerebral parenchyma and lead to cerebral small vessel disease. Previous studies have shown that silent lacunar infarcts and white matter hyperintensities are strongly associated with arterial stiffness. However, the association between microbleeds and arterial stiffness remains controversial, as there are two vessel mechanisms related to microbleeds: cerebral amyloid angiopathy and hypertensive small vessel disease. Key Messages: Cerebral small vessel disease with as-
associated arterial stiffness is a risk factor for silent cerebral lesions, stroke, and cognitive impairment. Improvement of the living environment, management of risk factors, and innovation and development of novel drugs that improve arterial stiffness may suppress the progression of cerebral small vessel disease, and may reduce the risk for stroke and dementia.

**Introduction**

Cerebral small vessel disease (SVD) is the most common vascular cause of dementia, a major contributor to mixed dementia, and the cause of approximately one fifth of all strokes worldwide [1, 2]. Recently, the neuroimaging features of SVD on magnetic resonance imaging (MRI) have been reclassified into five categories: (1) recent small subcortical infarcts (formerly categorized as acute lacunar stroke); (2) white matter hyperintensity (WMH); (3) lacunae [formerly defined as silent lacunar infarcts (SLI)]; (4) microbleeds (MB), and (5) perivascular space [1].

Cerebral SVD is of clinical importance as a risk factor for cerebrovascular diseases and cognitive impairment [1, 2]. There is also widespread acceptance of the existence of vascular cognitive impairment. Additionally, the continuity between cerebrovascular diseases and cognitive impairment is of increasing interest. Cerebral SVD poses a risk for both cerebrovascular disease and cognitive impairment via effects on arterial stiffness, presenting as microvessel arteriosclerosis with vascular endothelial dysfunction [3]. Recent studies also suggest an association between arterial stiffness and the geriatric syndrome [4, 5].

Stroke, dementia, and the geriatric syndrome are important in our aging society, as such diseases increase caregiver burden [6]. The increased number and proportion of older adults presenting with these diseases in the Japanese population and worldwide is heightening the financial pressures on health care systems. Here, we review the characteristics of cerebral SVD from the viewpoint of silent cerebral lesions, stroke, and dementia, with a particular focus on the association with arterial stiffness.

**Arterial Stiffness**

**Outline**

Arterial stiffness is a known predictor of mortality, cardiovascular diseases, and cerebrovascular diseases [7–9]. Recently, numerous studies have also shown novel associations between cognitive impairment and arterial stiffness [10–15]. Methods for assessing arterial stiffness are noninvasive and allow simple assessment and data evaluation. Surrogate markers to assess arterial stiffness include the ankle-brachial index (ABI), pulse wave velocity (PWV; e.g., aortic, brachial-ankle, and carotid-femoral PWV), cardio-ankle vascular index (CAVI), and augmentation index. A decreased ABI indicates atherosclerotic changes in large arteries [9]. By contrast, an increased ABI indicates progression of arterial stiffness [13]. Therefore, the ABI is considered a component of arterial stiffness. Although there are some methodological differences between these markers, their overall utility is well established.

**Measurement**

Patients were placed in the supine position and rested for 5 min before the measurements of arterial stiffness commenced. Electrocardiographic electrodes were placed on both wrists, cuffs were wrapped around both ankles and the brachial regions of both forearms, and the heart sounds were monitored. Firstly, ABI was calculated by the ratio of the systolic blood
pressure at the ankle to the systolic blood pressure at the arm. Secondly, brachial-ankle PWV was automatically calculated according to the height of the patients, the distance between the brachium and the ankle, and the time intervals between the wave at the right brachium and those at both ankles [16]. Thirdly, CAVI was calculated independent of blood pressure using the stiffness parameter $\beta$ in the Bramwell-Hill equation [17]. The stiffness parameter $\beta$ indicates patient-specific, blood pressure-independent vascular stiffness as measured by arterial ultrasound. The detailed formula of CAVI is shown elsewhere [17, 18].

### Cerebral SVD

**Outline**

Cerebral SVD such as WMH, SLI, and MB are considered to involve silent cerebral changes. Firstly, WMH characterized by bilateral, mostly symmetrical hyperintensities on T2-weighted MRI are common in older individuals. WMH is strongly associated with cerebrovascular disease, vascular risk factors, and covert neurological and cognitive symptoms and physical difficulties such as gait disturbance [1]. Secondly, SLI are frequently seen on imaging in older patients with no symptoms and are associated with an increased risk of stroke, gait impairment, and dementia [1]. Thirdly, MB are small hypointense lesions that are visible on T2*-weighted gradient-recalled echo or susceptibility-weighted sequences, and are most commonly located in the cortico-subcortical junction and deep gray or white matter in the cerebral hemispheres, brainstem, and cerebellum [1]. MB are associated with cognitive impairment such as Alzheimer’s disease.

**Assessment and Clinical Meaning**

We defined WMH as an irregular periventricular hyperintensity (Fazekas grade $\geq 3$) and/or early confluent or confluent separate deep white matter hyperintense lesions (Fazekas grade $\geq 2$) on FLAIR images [19]. We defined SLI as a focal lesion of $\geq 3$ mm in diameter, with hyperintensity on T2-weighted MRI and hypointensity on FLAIR [20]. We defined MB as a focal area of signal loss in the brain parenchyma $<5$ mm in size on T2* scans [21].

Cerebral SVD is the most common vascular cause of stroke and dementia, and is a major contributor to mixed dementia [1]. In fact, these lesions are not ‘silent’, but rather they slowly increase the risk of stroke and dementia in the brain. For example, as the numbers of SLI and MB increase, the risk of cerebrovascular diseases and cognitive impairment increases. Likewise, as the area of WMH enlarges, the risks of disease increase. Age and hypertension are well-known risk factors for cerebral SVD [22]. Pulse pressure and blood pressure variability may also contribute to the risk of cerebral SVD and cardiovascular and cerebrovascular diseases [23, 24].

**Association with Arterial Stiffness**

Arterial stiffness is independently associated with cerebral SVD including WMH [19], SLI [20, 25], and MB [26]. The mechanism of cerebral SVD is considered to involve vascular endothelial dysfunction, which may be associated with arterial stiffness [27]. Further, blood-brain barrier failure or progression of arteriosclerotic changes may lead to cerebral parenchymal damage [3]. As there are vessel similarities between the kidney and the brain, the presence of chronic kidney disease may also be associated with cerebral SVD based on similar mechanisms [21, 28].

Taken together, there is a robust association between cerebral SVD and arterial stiffness, although there are some methodological differences between the surrogate markers of arterial stiffness. Both WMH and SLI are independently associated with increased PWV [25].
By contrast, the independent association between CAVI and WMH remains controversial, although CAVI is independently associated with SLI [20]. This discrepancy suggests that (1) microvessel arteriosclerosis such as lipohyalinosis is a major cause of SLI and (2) vessel stiffness, as well as other factors such as pulsatile pressure, may affect the etiology of WMH. The associations between PWV, CAVI, and MB are also controversial, which may be related to the different etiology of MB: the differences between deep and/or infratentorial MB and lobar MB. Deep MB located in the basal ganglia are mainly caused by hypertensive SVD, while lobar MB are caused by cerebral amyloid angiopathy [21]. Thus, both positive and negative associations between MB and arterial stiffness are expected, and are dependent on the differences in participating subjects.

**Stroke**

Stroke is a major cause of long-term care in Japan [6]. Lacunar infarcts are a common subtype of brain infarction [22]. Acute lacunar infarcts are associated with WMH [22] and can induce progressive neurological deficit presenting with infarct enlargement [3]. Arterial stiffness is a risk factor for progressive neurological deficit [3]. The major mechanism of lacunar infarcts is thought to involve traditional risk factors such as hypertension and diabetes mellitus [22], although cerebral SVD associated with arterial stiffness may also be an important contributor.

There are some clinical differences between the various surrogate markers of arterial stiffness used in patients with ischemic stroke: (1) a low ABI indicates advanced atherosclerosis, (2) an increased PWV strongly indicates the presence of cerebral SVD, and (3) an increased CAVI may indicate vessel stiffness due to arteriosclerosis [18]. Cerebral SVD may be associated with any of these surrogate markers. Such clinical differences and associations should be clarified in detail in future studies.

**Cognitive Impairment**

Cerebral SVD such as SLI, WMH, and MB are well-known risk factors for cognitive impairment [1, 2]. Cognitive impairment including dementia is also associated with arterial stiffness [8–12]. An ABI >1.30 and increased blood pressure variability are important predictors of cognitive impairment among patients without vascular diseases [13]. Thus, an increased ABI indicates arterial calcification that leads to vessel sclerosis (advanced vessel stiffness). Increased PWV is also associated with cognitive impairment [12, 14], an independent predictor of cognitive decline [10, 11], and β-amyloid (Aβ) deposition in the brains of older patients [15]. More specifically, Hughes et al. [15] showed the associations between arterial stiffness and Aβ deposition in the brain independent of blood pressure. Increased arterial stiffness has a direct impact on penetrating arteries of the brain, leading to an altered structure and function, with subsequent effects on perivascular amyloid clearance from the brain via the cerebrospinal fluid drainage along the perivascular space. This may contribute to a disruption of vascular dynamics and complicate perivascular flow of Aβ, thus indirectly causing decreased Aβ clearance leading to plaque formation. Therefore, arterial stiffness may contribute to the pathogenesis of cognitive impairment through its potential role in the effects of hypertension on the brain, cerebrovascular disease, vascular dementia, and Alzheimer’s disease pathology [24].

There are two vessel-related mechanisms thought to be involved in cognitive impairment: cerebral amyloid angiopathy and hypertensive SVD. The etiology of MB is associated with both cerebral amyloid angiopathy and hypertensive SVD [29], and both may be associated with arterial stiffness. Therefore, MB are a surrogate marker of both vascular cognitive impairment and stroke. Collectively, the risk of cognitive impairment depends on the presence of any components of cerebral SVD including WMH, SLI, and MB, all of which are associated with arterial stiffness.
The geriatric syndrome is defined as having clinical symptoms characteristic of older adults, typically including cognitive functional impairment, falls, foreign body aspiration, impaired appetite of unknown cause, frailty, and sarcopenia [30]. This syndrome can be associated with cerebral SVD, as shown above. Recently, novel associations between the geriatric syndrome and arterial stiffness have been reported. Frailty syndrome in older adults is associated with subclinical peripheral arterial disease indicated by ABI [4]. Increased brachial-ankle PWV is also associated with an increased risk for mortality and the onset of impairment in activities of daily living [5, 7]. These factors may pose risks for dementia; this is an issue that should be investigated in more detail in the future.

**Mechanism**

Numerous studies have reported associations between arterial stiffness and various patterns of cerebral damage, as well as injury to other organs such as the retina, heart, and kidney. These organs have a common vascular mechanism of injury consisting of perforating arteries. This mechanism has been indicated as a water hammer effect [31]. In detail, Fazekas et al. [31] suggested the pathological findings of WMH as follows: arteries with walls thickened by hyalinosis and fibrosis, large perivascular spaces bordered by atrophic neuropil, and extensive arterial pulsations which could be implicated in their pathogenesis by initiating a

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**Fig. 1.** Overview of the tsunami wave model. a The heights of tsunami waves increase as the waves move closer to the land because of the geographical features of the bottom of the sea and/or river and those of the seashore. This mechanism is due to the depths (b) and the widths (c) of the bay and/or river. Words in parentheses are added by the authors to suggest how this tsunami wave model may explain the mechanism of cerebral damage in cerebral SVD, cognitive impairment, and the geriatric syndrome due to increased PWV. This figure is cited from [33]. b Vertical section of the wave. The heights of the tsunami waves increase (H₁ → H₂) as the waves move closer to the land because the depths of the sea and/or river (D₁ → D₂) become shallower. c Horizontal section of the wave. The heights of tsunami waves increase (H₁ → H₂) as the waves move closer to the land because the widths of the sea and/or river (W₁ → W₂) become narrower.
water hammer effect on the surrounding tissue. Recently, some researchers have suggested physiological considerations [27, 32]. Van Sloten et al. [27] showed that increased pulsatile pressure and flow might directly load to cerebral microcirculatory damage and induce a microvascular remodeling response. O'Rourke and Hashimoto [32] also showed in detail that repetitive pulsations cause fatigue and fracture of elastin lamellae of central arteries, causing them to stiffen, so that reflections return earlier to the heart; in consequence, aortic systolic pressure rises, diastolic pressure falls, and pulsations of flow extend further into cerebral small arteries, which leads to damage to the cerebral parenchyma. Stiffening also increases stresses on cerebral small arteries.

Based on our findings and previous studies, we propose a hypothesis using the ‘tsunami wave model’ to improve our understanding of the risk and the importance of arterial stiffness (fig. 1) [33]. The tsunami wave model shows that wave damage to the land increases with river narrowing (fig. 1a). The mechanism involves an increase in tsunami wave height as the wave moves closer to the land, owing to the geographical features of the bottom of the sea and/or the width of the river (fig. 1b, c). The power of the tsunami wave depends on the depth and width of the bay and/or river. This tsunami wave model could explain the physiological action of increased arterial stiffness. Although both a water hammer model and this tsunami wave model might indicate a similar physiological mechanism [32], this mechanism could also contribute to an improved understanding of the association between cerebral SVD and increased arterial stiffness.

Vascular narrowing due to atherosclerosis and vascular stiffness due to lipohyalinosis may accelerate the pulse waves. Based on this mechanism, we also propose a hypothesis regarding the connection of vascular risk factors, cerebral SVD, and stroke with cognitive impairment via arterial stiffness (fig. 2) [33]. This hypothesis will be investigated in more detail to clarify the mechanism in future studies.

**Conclusion**

Cerebral SVD is a well-known and important visual finding on brain MRI in older patients with and without risk factors for cerebrovascular diseases and cognitive impairment. The vascular, physiological, and metabolic roles of arterial stiffness require detailed clarification.
in future studies. Further, improvement of the living environment, management of risk factors, and innovation and development of novel drugs may help decrease the progression of cerebral SVD and reduce the risk of stroke and dementia.

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