High-resolution MRI of intracranial large artery diseases: how to use it in clinical practice?

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ABSTRACT

High-resolution MRI (HRMRI) has emerged as a useful tool for clinical research in recent years. Compared with traditional cranial and vessel imaging, HRMRI provides more additional valuable pathophysiology information that is helpful for the differential diagnosis of intracranial atherosclerosis, dissection and vasculitis. However, there are some points that a neurologist should keep in mind. First, although enhanced vessel wall imaging is widely applied for research purposes, it is not appropriate for routine clinical use. Any injury or inflammation within vessel wall can result in enhancement, which is unspecific for a diagnosis. Second, although plaque components identified on HRMRI arouse researchers’ interest, they may have limited positive predictive value for future stroke. Ruptured plaques may have higher prevalence in asymptomatic patients than expected. More prospective observational studies are required. Third, the vessel wall morphology features remain the useful and reliable clue for a diagnosis. It is true that eccentric vessel wall lesions most likely represent atherosclerosis if vessel dissection is easily excluded. For concentric wall lesions, however, the underlying pathophysiology is complicated, either atherosclerotic or non-atherosclerotic. Fourth, HRMRI can show luminal thrombus directly and provide valuable details. All in all, when HRMRI is used by a neurologist, it should not be viewed as the only key for a diagnosis. The diagnosis should be made based on patient history, lab works, other imaging technique and even genetic examinations.

In the recent years, high-resolution MRI (HRMRI) has emerged as a useful tool for clinical research. By HRMRI, intracranial vessel wall lesions can be observed in vivo. It has made it easier for the differential diagnosis of intracranial atherosclerosis (figure 1), dissection (figure 2), vasculitis (figure 3),1 and moyamoya disease (figure 4). HRMRI can also identify intracranial luminal thrombosis.2 Compared with traditional cranial and vessel imaging, HRMRI provides more additional valuable pathophysiology information. In 2017, HRMRI has been recommended for clinical practice to differentiate the causes of intracranial arterial narrowing and identify symptomatic, non-stenotic intracranial arterial diseases by American Society of Neuroradiology.1 However, there are some points that a neurologist should keep in mind when applying this technique.

VESSEL WALL ENHANCEMENT: NOT SPECIFIC

A lot of authors believe the enhanced intracranial vessel wall imaging has good image quality and is helpful for making a diagnosis. It was reported that enhanced eccentric plaque represents vulnerable atherosclerosis and enhanced concentric wall thickening represents vasculitis.3,4 In patients with moyamoya disease, enhanced vessel wall lesions seldom occur.5 However, all these opinions come from small sample size study. Any injury or inflammation within vessel wall can result in enhancement. Recent studies suggested even in healthy subjects, vessel wall can display enhancement to some extent.6 Vessel inflammation, either occurring in atherosclerosis, vasculitis or vasculopathy, has dynamic process that may have different imaging findings at different stages.

PLAQUE COMPONENTS: REMAIN AWAY FROM ROUTINE CLINICAL PRACTICE

Although fibrous caps have been observed in patients with intracranial atherosclerosis (figure 1),7 they are much thinner and have poorer image quality than those in carotid plaques. Intracranial fibrous cap fissure or rupture can be not sensitively identified by HRMRI. It is known from coronary atherosclerosis disease, ruptured plaques had higher prevalence in asymptomatic patients than expected.8 It could be assumed that the imaging findings of ruptured plaque, even identified, may have limited positive predictive value for stroke. With regard to intracranial intraplaque haemorrhage, the low prevalence suggested it is not useful marker for most patients.9

VESSEL WALL MORPHOLOGY: REMAINS USEFUL

When we observe intracranial vessel wall, the most important individual morphology
characteristic is eccentricity. It is true that eccentric vessel wall lesions most likely represent atherosclerosis. If a plaque develops in middle cerebral artery, it most likely locates at the ventral and inferior wall. Additional plaque components findings such as lipid core, fibrous caps and intraplaque haemorrhage may further strengthen the diagnosis. For a long stenosis, intracranial atherosclerosis may display concentric features on several image slices, but it always has at least one slice with eccentric lesion (figure 5). Different diagnosis may be required in patients with intracranial artery dissection. However, vessel wall haematoma at non-atherosclerosis sites is usually easier to be identified on HRMRI, as compared with other vessel imaging such as CT angiography and magnetic resonance angiography.  

For concentric wall lesions, the underlying pathophysiology is complicated. They can represent atherosclerosis (usually asymptomatic), vasculitis and other non-atherosclerotic vasculopathy. The evidence from histological study of intracranial vessel wall lesions is very lacking. In a young patient with an intracranial concentric stenosis, it is not easy to establish the diagnosis if no clue from labwork is available. The pathophysiology of early asymptomatic intracranial atherosclerosis in young patients with traditional risk factors is unknown, which need further studies.

**LUMINAL THROMBUS: A PROMISING TOOL**

Intracranial artery clot is the main cause and therapeutic target for patients who had acute ischaemic stroke. The conventional method to evaluate the presence of a clot is the occlusion of blood flow on MRA or CTA imaging. However, it is indirect and cannot provide more accurate and useful details of thrombus. Luminal clots can be identified as hyperintense or isointense signals on non-contrast T1-weighted HRMRI in extracranial and intracranial vessels (figure 6). A whole picture of clot burdens including number, location, length, components, associations with other lesions and other characteristics can be clearly displayed on HRMRI, which makes it possible to investigate stroke mechanisms further. Although there are a small number of researches using HRMRI to study luminal clot limited to imaging technology, it will be a promising tool and wildly used in clinical practice in the future.

**HOW SHOULD WE USE HRMRI?**

HRMRI makes it feasible for in vivo evaluating intracranial vessel wall lesions and luminal clot. It is one of the techniques that help neurologist observe intracranial diseases. For an individual patient, HRMRI is not the only ‘key’. The diagnosis should be made based on...
Magnetic resonance angiography shows basilar artery stenosis (white thick arrow (A)). On consecutive axial views of three-dimensional T1-weighted high-resolution MRI, image slices show a concentric stenosis (white thin arrow (B)) and an eccentric stenosis (white arrowhead (C)).

Figure 5

patient history, lab works, other imaging technique and even genetic examinations.\(^1\) It is not appropriate to give cortisol treatment simply because concentric enhanced vessel wall lesions are present on HRMRI. It is also not appropriate to give conservative treatment simply because non-enhanced eccentric asymptomatic atherosclerosis is observed. We are still far away from the ‘truth’ of intracranial vessel diseases even we have HRMRI now. In clinical practice, comprehensive diagnostic work should be done for making a final decision.

Magnetic resonance angiography shows right middle cerebral artery (MCA) occlusion (white thick arrow (A)). Isointense signals within the occluded MCA were noted on corresponding three-dimensional T1-weighted high-resolution MRI (3D T1 HRMRI) images suggesting luminal thrombosis (white arrowheads (B)). On consecutive sagittal views of 3D T1 HRMRI, image slices show a normal vessel lumen (white thin arrow (C)) and an occluded vessel lumen with clots (white thin arrow (D)).

Figure 6

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