Vessel Wall MRI Enhancement in Noninflammatory Cerebral Amyloid Angiopathy

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ABSTRACT

SUMMARY: Cerebral amyloid angiopathy is characterized by deposition of amyloid-β fibrils in the walls of small-to-medium-sized blood vessels. In this retrospective review of 5 patients with histologically confirmed noninflammatory cerebral amyloid angiopathy, high-resolution vessel wall MRI showed arterial wall enhancement in 2 patients (40%). Despite common consensus of equating vessel wall enhancement with inflammation, this report demonstrates that β-amyloid accumulation alone without inflammation can be associated with arterial wall enhancement in a subset of patients.

ABBREVIATIONS: CAA = cerebral amyloid angiopathy; VWMRI = vessel wall MR imaging

Cerebral amyloid angiopathy (CAA) is characterized by deposition of amyloid-β fibrils in the vessel walls of leptomeningeal and cortical arteries, arterioles, and, less often, capillaries and veins of the central nervous system. Histologically, Puchtler alkaline Congo Red stain remains the standard diagnostic method for staining amyloid deposition, which can be seen in the adventitia, tunica media, or all layers of vessel wall.

Along the spectrum of amyloid-deposition-related angiopathies, there are 3 entities: 1) CAA (generally there is an absence of wall inflammation); 2) inflammatory CAA (varying amounts of perivascular inflammation are present, including multinucleated giant cells); and 3) amyloid-β-related angiitis (in which there is destructive vasculitis). There is often overlap in the clinical presentations of these entities, which may include focal neurologic deficits, cognitive impairment, or seizure related to hemorrhagic or ischemic changes. The clinical diagnosis of CAA relies on clinical history and imaging evidence of microbleeds or lobar hemorrhage that are not explained by other etiologies.

High-resolution vessel wall MR imaging (VWMRI) is an emerging technique that is used to assess various vasculopathies, such as atherosclerotic disease, vasculitis, Moyamoya disease, reversible cerebral vasoconstriction syndrome, and dissection. While vessel wall enhancement using VWMRI has been reported in the inflammatory variety of amyloid-related angiopathies, very little is known about the VWMRI findings in patients with noninflammatory CAA. We report the VWMRI findings in a patient with histologically confirmed CAA.

Brief Report

A 75-year-old patient with a history of hypertension, diabetes, progressive memory loss, and behavior changes during 6 months presented with worsening of mental status followed by seizures and status epilepticus. The initial neurologic examination revealed impaired attention and language; however, cranial nerves, motor strength, and gait were unremarkable. The patient had fluctuating mental status during the first 3 days and developed a clinical seizure on the fourth day with subsequent nonconvulsive status epilepticus, which was revealed on video-electroencephalography and required multiple antiepileptic agents. MR imaging showed no acute infarction but a few chronic microhemorrhages (Fig 1A) and some white matter changes. Due to the presence of chronic microhemorrhages, the differential diagnosis of vasculitis or amyloid angiopathy was entertained. Conventional angiography showed normal intracranial arteries. VWMRI (3D T1-weighted fast-spin-echo sequence with variable refocusing flip angle acquisition with inherent black-blood effect [sampling perfection with application-optimized contrasts by...
Arterial wall enhancement is associated with amyloid accumulation within the vessel wall. A, Gradient-echo image shows microhemorrhages in the left temporal lobe (red arrowheads). B and C, coronal and sagittal views of postcontrast T1-weighted VWMRI show enhancement in the wall of cortical branches of the left middle cerebral artery in the parietal and temporal lobes (red arrows). D, Hematoxylin-eosin stain of a left temporal lobe sample shows thickened, hyalinized blood vessels containing amorphous eosinophilic material (blue arrows) in small- and medium-sized arteries within the leptomeninges and superficial cortical gray matter. E, Immunostain for β-amyloid shows amyloid accumulation within the vessel wall (blue arrowhead). No inflammatory cells were observed surrounding the vessels.

Using different flip angle evolution, SPACE sequence; Siemens, Erlangen, Germany: FOV, 230 mm; TR/TE, 700/18 ms; matrix, 256 mm²; 144 sections × 0.9-mm thick; voxels, 0.9 × 0.9 × 0.9 mm³) on the eighth day of admission showed arterial wall enhancement along the cortical branches of the left middle cerebral artery in the left parietal and temporal lobes (Fig 1B, -C). Findings of clinical and laboratory work-ups were negative for CNS infection, systemic vasculitis, or metabolic abnormalities except for elevated blood glucose (340 mg/dL on admission). Biopsy revealed β-amyloid accumulation in the vessel wall without associated inflammation (Fig 1D, -E).

Following institutional review board approval, we subsequently reviewed digital charts of patients with a histologically proved diagnosis of noninflammatory CAA between August 2010 and February 2019. From a total of 10 patients identified, only 4 patients had adequate imaging for inclusion in this study (no MR imaging, n = 2; inadequate MR imaging image quality to assess vessel wall enhancement, n = 4). Of these 4 patients, 1 demonstrated arterial wall enhancement similar to that in our report. Clinical and demographic data, imaging findings including VWMRI and the number of microhemorrhages, and histopathologic findings are summarized in the Online Table.

**DISCUSSION**

Intracranial vessel wall enhancement detected by high-resolution MR imaging has been considered a sign of inflammation and/or vasa vasorum in-growth in patients with atherosclerosis or vasculitis. Although vessel wall enhancement can be expected with vascular inflammatory processes such as primary angiitis of the central nervous system or an inflammatory type of amyloid such as amyloid β-related angiitis, in our report, we show that vessel wall enhancement can also be seen in a subset of patients with noninflammatory CAA (40%). This suggests that β-amyloid accumulation by itself and without inflammation can also result in vessel wall enhancement. This report, to our knowledge, is the first on VWMRI findings in patients with histopathologically proved noninflammatory CAA. An earlier study by De Havenon et al showed vessel wall enhancement in about 50% of patients with CAA; however, in this study, there was no discrimination between inflammatory and noninflammatory CAA because the patients did not have histopathology to confirm the diagnosis.

One hypothesis or explanation for our findings is potential change and disruption of the blood-brain barrier in CAA. CAA-positive vessels often exhibit morphologic changes such as loss of smooth-muscle cells and vessel wall thickening, which, in turn, can result in increased leakiness and an increased incidence of microbleeds. It is likely that this increased permeability is responsible for wall enhancement in a subset of CAA (despite the absence of inflammation). Most interesting, our patients who exhibited vessel wall enhancement also had numerous microhemorrhages, while those without vessel wall enhancement had none or just a few.

There are several limitations to our study. First, our sample size is very small, precluding any meaningful statistical analysis. Second, there is potential selection bias in this retrospectively collected series. Third, due to the retrospective nature of our study, there may have been variability in our VWMRI sequence parameters. Also, we did not use delay alternating with nutation for tailored excitation (DANTE) preparation to refine our blood suppression. However, we used the best option available to us from our vendors to perform these clinical studies using FDA-approved sequences in a clinical environment. Future prospective radiologic-histologic studies using dedicated VWMRI are needed to better characterize the arterial wall changes in CAA with and
without inflammation, and their role in CAA-related cerebral pathologies.

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