The role of cross-sectional imaging of the extracranial and intracranial vasculature in embolic stroke of undetermined source

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Despite an extensive workup, nearly one third of ischemic strokes are defined as Embolic Stroke of Undetermined Source (ESUS), indicating that no clear etiologic cause has been identified. Since large vessel atherosclerotic disease is a major cause of ischemic stroke, we focus on imaging of large vessel atherosclerosis to identify further sources of potential emboli which may be contributing to ESUS. For a stroke to be considered ESUS, both the extracranial and intracranial vessels must have <50% stenosis. Given the recent paradigm shift in our understanding of the role of plaque vulnerability in ischemic stroke risk, we evaluate the role of imaging specific high-risk extracranial plaque features in non-stenosing plaque and their potential contributions to ESUS. Further, intracranial vessel-wall MR is another potential tool to identify non-stenosing atherosclerotic plaques which may also contribute to ESUS. In this review, we discuss the role of cross-sectional imaging of the extracranial and intracranial arteries and how imaging may potentially uncover high risk plaque features which may be contributing to ischemic strokes.

KEYWORDS: cerebrovascular disease/stroke, atherosclerosis, carotid artery stenosis, magnetic resonance angiography, carotid artery disease

Introduction

Despite an extensive workup, nearly one-third of ischemic strokes are defined as Embolic Stroke of Undetermined Source (ESUS) meaning that no definite cause of the stroke has been identified (1). ESUS has proven to be a difficult clinical entity to treat with an almost 5% per year stroke recurrence rate (1). Most non-lacunar ischemic strokes are embolic and can originate from cardiac sources, from more proximal arterial structures, such as the carotid arteries or aortic arch, or potentially from a venous source in the setting of paradoxical embolism.
Cardiac sources, specifically atrial fibrillation, were thought to play a major role in ESUS because occult atrial fibrillation was found in many patients with ESUS (2). However, two major randomized clinical trials comparing oral anticoagulants to aspirin in patients with ESUS had neutral results (3, 4), suggesting that other embolic causes for ESUS may play a larger role. A major contributor to ischemic stroke is large artery atherosclerotic disease accounting for approximately 25% of ischemic strokes and most commonly arising from the extracranial carotid artery. According to the most common methods for classifying stroke etiologies, in order to attribute an ischemic stroke to large artery atherosclerosis, there must be associated luminal stenosis of at least 50% (5). These criteria do not take into account the recent paradigm shift in our scientific understanding of the contribution of specific plaque features to ischemic stroke.

In this review article, we will review the role of cross-sectional imaging of the carotid arteries in patients presenting with ESUS. First, we will discuss the current standard of care and typical imaging workup to exclude carotid disease as a potential cause of stroke. We will then discuss the role of computed tomography angiography (CTA) and magnetic resonance angiography (MRA) in evaluating potential causes of ischemic stroke in the extracranial carotid artery. We will also review the role of intracranial vessel wall MR (VWMR) in assessing intracranial atherosclerosis, another potential contributor to ESUS.

Current paradigm/standard of care

Rather than being a diagnosis of exclusion, ESUS has a standardized, criteria-based definition requiring specific imaging and clinical workup. In order to meet criteria for a diagnosis of ESUS, an ischemic stroke must be a non-lacunar stroke detected on CT or MR imaging, the patient must have ≤50% luminal stenosis of the extracranial and intracranial vessels supplying the territory of the brain infarction, and have no major risk of a cardioembolic source or other specific identifiable cause of stroke, such as arteritis, dissection, migraine/vasospasm, or drug misuse. In order to make this diagnosis, suggested diagnostic assessment in evaluating those with ESUS is a brain CT or MR, 12-lead electrocardiogram, precordial echocardiography, cardiac monitoring for 24 h with automated rhythm detection, and imaging of both the extracranial and intracranial arteries supplying the area of brain ischemia with either digital subtraction angiography, MR or CT angiography, or cervical duplex and transcranial Doppler ultrasonography (1). These relatively recent guidelines have allowed for standardization in the identification of those with acute ischemic stroke and have made those with ESUS easier to identify. These more rigid definitions have led to more concentrated effort in mitigating stroke in this population and have paved the way for recent large randomized clinical trials (3, 4).

Limitations of current imaging techniques

While the current diagnostic criteria for ESUS require assessment of both the extracranial and intracranial arterial structures, the primary focus remains on the degree of luminal stenosis. For decades, the degree of stenosis has been the primary indicator of stroke risk in the extracranial and intracranial arteries. Carotid disease is thought to lead to ischemic stroke by two distinctive, but often synergistic factors: flow-limitation in the setting of stenosis leading to hypoperfusion and artery-to-artery embolism from plaque leading to thromboemboli (6). It is likely that hypoperfusion from flow limitation contributes to cerebral ischemia. Further, impaired perfusion in the setting of flow-limitation may lead to a potentially transient embolic event resulting in an infarct. While flow-limiting stenosis is clearly a risk factor for the development of ischemic stroke, there is mounting evidence that the plaque itself, regardless of the degree of accompanying stenosis is likely a contributor to ischemic strokes via artery-to-artery embolism (7).

Recent interest in the plaque components have furthered our understanding of the role of plaque features in contributing to embolic strokes. There is strong histopathologic evidence that plaque may have different features conferring higher risk for an embolic phenomenon. The American Heart Association plaque classification describes a spectrum of plaque with certain features, including intraplaque hemorrhage, lipid-rich necrotic core, and surface defects including fibrous cap rupture which are features indicative of more “vulnerable” plaque that is more likely to rupture and lead to emboli (8, 9). With the current recommendations for imaging, specific carotid plaque features are not always appropriately imaged or may not always be identified or treated as important drivers of ischemic stroke. Despite strong scientific evidence supporting the role of vulnerable plaque features in the development of ischemic stroke, specifically in those with non-stenotic carotid atherosclerosis, these plaque features are not always being routinely assessed using the current guidelines. There is strong evidence that specific plaque vulnerable features in non-stenosing plaque are more commonly seen ipsilateral to infarction in ESUS patients (10). By recognizing the importance of non-stenosing plaque in the extracranial and intracranial vasculature, we may potentially be able to reclassify patients originally thought to have ESUS (11).

Extracranial carotid plaque

Extracranial internal carotid artery atherosclerosis has traditionally been the most common source of large vessel
The extracranial carotid arteries are always imaged in the setting of acute ischemic stroke to evaluate for a source. The simplest method for imaging extracranial plaque is by duplex ultrasound (US) where the degree of stenosis based on flow measurements can be assessed. US can also evaluate various plaque features which are known to be higher risk, including echolucent plaque (12). Though US certainly plays a major role in the evaluation of stroke etiology and ESUS (13), we will focus on other cross-sectional imaging modalities which can more accurately assess specific plaque features as well as luminal stenosis for a more complete assessment of stroke risk.

When imaging the extracranial carotid artery in the setting of ischemic stroke, there are two major considerations. First, the degree of luminal stenosis must be assessed using a standardized system, most commonly using North American Symptomatic Carotid Endarterectomy Trial (NASCET) criteria (14). This can be accurately ascertained in any of the imaging modalities currently being utilized for the diagnosis of ESUS. In addition to the degree of luminal stenosis, the components of the plaque must also be assessed, even in those with non-stenosing plaque. All imaging modalities for the extracranial carotid arteries can assess plaque features to a certain degree, though some are more well-suited than others.

**MR imaging**

MR is the most studied method to visualize vulnerable plaque features with considerable evidence supporting its use to evaluate advanced atherosclerotic plaque. Intraplaque hemorrhage (IPH) is the most commonly encountered MR-detected plaque feature considered to be “high-risk” and is strongly associated with infarction (15–17). Other MR-detected vulnerable plaque features including lipid-rich necrotic core, ulceration, or fibrous cap rupture are also strongly associated with ischemic infarction. There is strong histopathologic evidence correlating MR imaging findings to known specific vulnerable plaque features (9, 18). In order to accurately image these plaque features, many utilize dedicated carotid coils or specific high-resolution MR sequences, including T1 and T2 weighted sequences, proton density, and time-of-flight sequences to evaluate flow (19). Contrast-enhanced MR sequences can also improve plaque characterization and allow for better characterization of plaque ulceration (20). While using dedicated carotid coils improves imaging by increasing signal-to-noise ratios, some studies have found that even simple MR sequences can accurately identify basic high-risk carotid plaque features (Figure 1) (21).

Multiple studies have specifically evaluated the role of MR imaging of plaque in the setting of ESUS. Several studies have evaluated individuals with ESUS and have found that these MR-detected vulnerable plaque features are more commonly seen ipsilateral to the side of stroke compared to the contralateral side (22–25). Recent prospective studies have confirmed these findings. The Plaque At RISK study showed that in patients with mild-moderate extracranial carotid stenosis, those with IPH and higher total plaque volume were more likely to experience recurrent ipsilateral ischemic stroke over a 5 years follow-up period, though plaque ulcerations and calcifications were not significantly associated (26). Another recent prospective study of patients with non-stenosing plaque found that patients with complicated plaque, such as IPH or surface defects, were much more likely to experience recurrent ipsilateral infarctions in the 30 months following an initial infarct (27). The findings from these studies suggest that even in the setting of non-stenosing plaque, certain higher-risk plaque features may be responsible for infarcts.

Though MR imaging of the extracranial carotid arteries can be helpful in identifying high risk plaque features, its widespread use is somewhat limited by availability, patient contraindications (e.g., implanted metal), and usually lengthy sequences making it a time-consuming imaging examination.

**CTA**

CTA is an increasingly commonly used imaging modality in the assessment of etiology of acute ischemic stroke. Because it is relatively cost-effective and quick to obtain, it is most often the first-line examination for those presenting emergently with acute stroke symptoms. While there is more prospective evidence that MR-assessed vulnerable plaque features contribute to future and recurrent ischemic stroke, many of these plaque features can also be assessed using CTA imaging (28). While MR is a superior imaging modality for differentiating histopathologic components of plaque, CTA is able to assess a few specific features which are known to increase risk of
emboli, including “soft” or predominantly non–calcified plaque which is thought to be a correlate of IPH or lipid-rich necrotic core, plaque ulceration, and plaque thickness (Figure 2). These features are readily visible on routine CTA imaging and are associated with increased likelihood of symptomaticity (28, 29). Similar to studies performed with MR-detected plaque features, several have found that non–stenotic plaques are more commonly seen ipsilateral to the infarcted cerebral hemisphere in patients with ESUS (30, 31). Specifically, several have found that having plaques >3 mm was more common ipsilateral to the side of stroke (30–32). Other studies have found that plaque with spotty calcification and a “rim sign” were also associated with cerebrovascular ischemic symptoms (33, 34). These studies indicate that though there is a paucity of prospective data evaluating the role of CT-plaque features in future ischemic stroke, certain imaging findings may be useful in identifying those at higher risk of ischemic stroke.

Other imaging techniques

Though not frequently used in everyday practice, there may be a role for more advanced imaging to evaluate for higher risk plaque. Positron Emission Tomography (PET) imaging has been studied as a method for assessing the vulnerability of carotid plaque. A recent systematic review and meta-analysis found that carotid arteries ipsilateral to recent ischemic events had more avid uptake of markers of inflammatory activity (e.g., 18-F fluorodeoxyglucose) than asymptomatic arteries (35). Other types of more advanced imaging has been studied to evaluate for plaque vulnerability, including dynamic contrast enhanced perfusion imaging (36). These and other findings point to a potential role for advanced imaging in evaluating plaque vulnerability in the future.

Intracranial atherosclerosis

Intracranial atherosclerosis leads to up to 9–15% of ischemic infarctions in the United States and up to 50% worldwide. Similar to extracranial atherosclerosis, intracranial atherosclerosis must result in at least 50% narrowing in order to be considered causative in the setting of ischemic stroke. Active atherosclerotic plaque can easily be overlooked when using conventional angiographic imaging because plaques do not always produce associated vessel narrowing. Because of this, intracranial vessel wall MR (VW-MR) can be used as an imaging assessment of atherosclerosis, particularly non-stenosing plaque.

Intracranial VW-MR

Intracranial VW-MR is a powerful tool to image beyond the vessel lumen and for evaluating non-stenosing plaque which may lead to ischemic stroke. In order to accurately assess the vessel wall, there are several critical components to intracranial VW-MR imaging (37). First, in order to highlight the wall itself and any potential plaque, it is essential to suppress flowing luminal blood and CSF, which can be done with a variety of different T1-weighted sequences. This is essential to increase conspicuity of any plaque features or enhancement. Further, high spatial resolution is needed in order to see the small vessel wall with most institutions performing approximately 0.5 mm voxels. Multiplanar acquisitions are also essential to assess the vessels en face because of the inherent tortuosity of the intracranial vessels. This is usually achieved by acquiring images using 3D techniques then creating reformats. Lastly, multiple tissue weightings are also performed to evaluate specific T1 and T2 characteristics in order to distinguish different plaque components.

Intracranial VW-MR can more easily detect smaller plaques or plaques with associated positive remodeling which may not produce narrowing on angiographic imaging but may still lead to ischemic stroke. Positive remodeling is an adaptive process where the outer wall of a vessel can outwardly bulge in the setting of an atherosclerotic plaque to preserve cerebral blood flow, leading to a normal, non-stenotic appearance on standard angiographic imaging techniques, including CTA, MRA, and DSA. Positive remodeling is commonly seen in the posterior circulation but can be seen in any intracranial arteries. Because of the common occurrence of positive remodeling, many patients presenting with acute ischemic stroke may have a normal appearing angiographic study without any suspicious findings for a contributing atherosclerotic lesion. When imaged using VW-MR, however, culprit atherosclerotic plaques may be
identified (Figure 3) and are generally assumed to be causative for ischemic stroke.

VW-MR uses a few specific imaging findings to identify active or culprit atherosclerotic plaques. The most important imaging finding for evaluating plaque in the setting of acute ischemic stroke is plaque enhancement. Several meta-analyses show that plaque/vessel wall enhancement is very strongly associated with culprit or symptomatic plaques (38–40). Plaque enhancement is readily assessable on post-contrast MR sequences and is a fundamental aspect of imaging with VW-MR techniques. In addition to plaque enhancement, positive remodeling is another important imaging finding that is strongly associated with symptomatic plaques (39, 40). This strong association of positive remodeling with symptomatic plaque highlights the importance of VW-MR in identifying potential culprit plaques. Similar to extracranial carotid plaque, discontinuities on the plaque surface indicative of fibrous cap rupture are also associated with ischemic stroke and symptomatic plaques (39, 40). Though intraplaque hemorrhage is a very strong marker of high risk plaque in the extracranial carotid artery, the association between IPH or intraplaque high T1 signal in the intracranial artery is not as strong, with a more modest association with ischemic stroke and symptomatic plaque, more commonly seen in the basilar artery (40, 41). Further studies evaluating the role of intracranial IPH in contributing to acute ischemic stroke are warranted.

When used in patients with ESUS, some studies have found that intracranial VW-MR can be a helpful tool. A study with over 240 patients with ESUS found that intracranial plaque was much more common ipsilateral to the side of stroke (42). They also found that there was increased wall remodeling in patients with ESUS, again highlighting the importance of non-stenosing plaque (42). A recent systematic review of 21 studies of patients with non–stenosing atherosclerosis found that intracranial plaque with higher risk features such as plaque enhancement and positive remodeling were more commonly seen in those with acute infarction, again indicating the role of specific plaque features (43). Another study found that using intracranial VW-MR could change the stroke etiology classification as it identified alternate causes of the ischemic stroke (44).

Intracranial VW-MR has become increasingly popular in evaluating ischemic stroke and ESUS patients, with a recent survey suggesting that more than 50% of neuroradiology practices routinely perform this type of study (45). Despite its increasing popularity, intracranial VW-MR imaging is limited by lengthy acquisitions, patient contraindications, and cost. Further, there has been limited histopathologic validation of MR signal characteristics of intracranial vessel wall pathology due to limitations in correlation with vessel samples (46, 47). This inherent limitation in our ability to correlate imaging findings with histopathologic components constrains our understanding of intracranial plaque characteristics.

Conclusion

Given recent randomized clinical trial findings that treating cardiac sources for ESUS may not be as beneficial as originally hoped, more attention is being placed on other potential embolic sources. Since the current ESUS definitions require <50% luminal narrowing, potential culprit plaques could be missed or inadequately treated because they are producing insignificant narrowing. In the extracranial carotid artery, both MR and CTA can be used to identify certain plaque features which indicate more plaque vulnerability including IPH on MR and increased soft plaque thickness on CTA. VW-MR can also be used as a powerful tool to identify non-stenosing but active atherosclerotic plaque in the intracranial arteries by identifying an enhancing plaque with positive remodeling. Though these studies can be helpful in determining the source of potential emboli, there are some further studies needed to validate these imaging techniques and pave a path for their routine use in ESUS.
Author contributions

HB, HK, and AG contributed to conception of the manuscript. HB wrote the first draft of the manuscript. HK and AG critically revised and wrote sections of the manuscript. All authors contributed to manuscript revision, read, and approved the submitted version.

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Conflict of interest

HK serves as a PI for the NIH-funded ARCADIA trial (NINDS U01NS095869), which receives in-kind study drug from the BMS-Pfizer Alliance for Eliquis® and ancillary study support from Roche Diagnostics; as Deputy Editor for JAMA Neurology; on clinical trial steering/executive committees for Medtronic, Janssen, and Javelin Medical; and on endpoint adjudication committees for AstraZeneca, Novo Nordisk, and Boehringer Ingelheim. He has an ownership interest in TETMedical, Inc.

The remaining authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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