The contribution of imaging in diagnosis, preoperative assessment, and follow-up of moyamoya disease

A review

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The aim of this review was to evaluate the imaging tools used in diagnosis and perioperative assessment of moyamoya disease, with particular attention to the last decade. (DOI: 10.3171.2009.01.FOCUS08296)

Key Words • moyamoya disease • diagnosis • neuroimaging modality • magnetic resonance imaging • computed tomography angiography • catheter angiography • cerebrovascular reactivity

This review will present different imaging techniques and their contribution to the evaluation of patients with moyamoya disease.

Baseline and Morphological Studies

The Angiography Method

The first description of a case of moyamoya disease was based on diagnostic catheter cerebral angiography.43,77,80 Catheter angiography still represents the gold standard for moyamoya diagnosis. Three features usually characterize the vascular picture on angiography studies: stenosis of the supraclinoid distal ICA; the development of dilated striate arteries (moyamoya collateral vessels with a characteristic blush); and sparing of the posterior fossa vessels (Fig. 1D, G, and H).

Stenoses involve the anterior circulation and start in the terminal, supraclinoid part (C1–C2 portion) of the ICA, and subsequently spread to the proximal part of the ACA and the MCA. It is only in severe cases and late-stage disease that the stenotic process involves the PCA. The formation of collateral pathways44 initially comprises dilation of intracranial collateral vessels (perforating lenticulostriate arteries, the thalamoperforating arteries, the anterior choroidal, and the posterior pericallosal or splenial arteries). The final stages are characterized principally by the development of collateral vessels from the external circulation (ethmoidal arteries and dural–pial supply). Some authors consider the dilation of the anterior choroidal and posterior communicating arteries predictive for hemorrhage.57

Suzuki and Takaku77 have described 6 stages of moyamoya progression: 1) narrowing of the CA termination; 2) dilation of the proximal portions of the ACA and MCA with initial basal moyamoya blush; 3) proximal portions of the ACA and MCA are no longer visualized; distal branches are still present due to collateral vessels from the PCA and intensification of the moyamoya blush; 4) minimization of the basal moyamoya network together with progressive occlusion of the ICA, which reaches the origin of the PCA; 5) further reduction of moyamoya vessels, with complete disappearance of the main arteries arising from the CA, continuous decrease of moyamoya collateral vessels that are more limited to the siphon area,
and increased collateral supply from the ECA; and 6) disappearance of the moyamoya blush together with the blood supply from the ICA. At this stage, only the ECA supplies the intracranial circulation.

Angiography might also reveal the presence of aneurysms that may be related to the increased flow load in relation to the normal vessel diameter in these patients. These aneurysms are found either at the periphery, on the collateral dilated vessels, or on the Circle of Willis. However, when evaluating the cause of SAH in these patients, one has to also consider hemorrhage from fragile transdural anastomotic vessels.

In extremely rare cases, other vascular anomalies (such as arteriovenous malformations) may be detected in these patients.

A further interesting, less investigated angiographic finding is the involvement of the ECA system: stenoses were found (and also histologically proven) on the STA and on the MMA. It was suggested that these findings are associated with lower treatment success rates.

Changes in the angiographic venous filling patterns parallel the arterial changes. In the compensated phases of moyamoya disease, a delayed but increased filling of intracerebral venous drainage is observed, together with an increased but not delayed filling of the external venous draining system. In particular, the dilation of deep intracranial draining vessels, such as the striate veins, reflects the different stages of moyamoya disease. The superior striate vessels dilate together with the increase in basal ganglia moyamoya vessels, and decrease in size in the phase of leptomeningeal compensation. These findings differ from those in atherosclerotic patients, in whom there is no delay in venous filling, but rather territorial rarefaction of venous filling.

**The CT Modality**

Usually, the first neuroimaging tool used in diagnosing an emergency patient is a CT scan. Besides evidence of SAH or intraventricular hemorrhage (Fig. 1A), common CT findings in patients with moyamoya disease are those of ischemic cortical stroke. In a series of 18 moyamoya patients, 14 presented with mono- or bilateral cortical or subcortical low-density lesions, consistent with en-
Cephalomalacic changes following previous infarctions; ventricular dilation was noted in 9 patients and cortical atrophy in 11. Only 1 patient in this series with presumable TIA had normal findings on a CT scan of the brain, even after the addition of contrast. The CT scans obtained after contrast administration revealed enhancing lesions in 3 of 16 patients, and basal ganglia enhancement in 4 of 4 patients undergoing high-load contrast injections. Other features described in the literature include low densities in the basal ganglia in 4 of 6 patients,25 and small hypodense subcortical hyperdensities, referable to calcium deposits, within old infarcted lesions.25

The CT Angiography Technique

The use of CT angiography to image the vessels of the head and neck has evolved rapidly since helical CT introduced the ability to scan quickly and to capture the arterial phase of enhancement. The recent introduction of multidetector row CT scanners such as the 16, 64, or 320 multidetector row scanners enables isotropic imaging, which provides high-resolution 3D reconstructions and a short acquisition time. This diagnostic tool is especially helpful in patients with suspected vascular abnormalities, particularly in cases of intracerebral hemorrhage.1 Stenoses, CA occlusion, moyamoya vessels, and areas of contrast enhancement can be seen.28 A CT angiography scan is also a useful device for preoperative planning10 and for postsurgical verification of patency of an EC-IC bypass (Fig. 1A and B, Fig. 2A and D).

The MR Imaging Modality

Types of MR imaging studies include standard acquisitions and MR angiography as well as functional and metabolic studies. Use of MR imaging offers high spatial and temporal resolution without radiation.

In conventional MR imaging, findings in patients with moyamoya disease have been described as primary and secondary. Primary findings include the loss of flow voids of the large arteries, attributable to occlusions of the circle of Willis, and numerous flow voids in the basal cistern and basal ganglia, referable to the augmented vascular network at those sites (Fig. 2E and F). These findings are best seen on T2-weighted images, in which the black vascular flow voids are beautifully contrasted against the bright CSF, a critical observation that, when present and not overlooked, is virtually diagnostic of moyamoya disease. Secondary findings concern signs of cerebral infarction, gliosis, atrophy, and hemorrhage.24

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Fig. 2. Neuroimages obtained in a 42-year-old patient presenting with multiple TIAs treated by bilateral EC-IC bypass. A: Coronal maximum intensity projection reconstruction of CT angiography studies, obtained after the first procedure, demonstrates a patent left STA-MCA bypass as it passes through the cranectomy and connects intracranially to a distal branch of the left MCA (arrowhead). B and C: Conventional cerebral angiogram of the left CCA in anteroposterior (B) and lateral (C) projections confirms the patency of this bypass with opacification of vessels of the left MCA territory. D: A 3D reconstruction of a CT angiography study, obtained after performing the second bypass, demonstrates that both STA-MCA anastomoses are patent (arrows). E: The MR imaging CVR studies obtained prior to treatment reveal reduced CVR in bilateral MCA and ACA territories (in blue) but preserved reactivity in the posterior circulation territory (in red and yellow). F: After the bypass procedures there is near normalization of CVR on the right side of the brain and significant improvement on the left. The ACA territory is not significantly changed because the bypass does not enhance perfusion of this region. Designators in this figure include the following: arrowheads denote the STA as it passes through the cranectomy (panels A–C); long arrows denote STA-MCA anastomosis (panels A, B, and D); and short arrows denote MCA branches (panels B–D).
The FLAIR images offer improved sensitivity to ischemic injury compared with T1- and T2-weighted images and are particularly helpful in the discrimination between subacute and chronic lesions, because the latter usually present with a lower or isointense signal. In contrast, diffusion-weighted imaging is the most sensitive sequence for identifying acute ischemic stroke, becoming abnormal before FLAIR and T2 images. This technique measures the microscopic motion of water protons and is quantified by measuring the apparent diffusion coefficient, which lumps the effects of random and restricted water motion in tissue. A reduced apparent diffusion coefficient is a sensitive marker of the failure of cellular energy metabolism in the ischemic brain. Furthermore, signal changes on diffusion-weighted imaging develop in areas of severely reduced perfusion, representing restricted diffusion that often follows cytotoxic edema.

Areas of severely reduced perfusion, representing restricted diffusion that often follows cytotoxic edema, can be identified on diffusion-weighted imaging. These lesions are classically present in the periventricular deep white matter, rather than at the corticocortical or at the basal ganglia/thalami as described in other diseases. The authors found nonsignificant differences in the incidence of microhemorrhages between patients with and without antiplatelet therapies. Usually antiplatelet agents are given to patients with moyamoya who have the occlusive ischemic, nonhemorrhagic disease type. Whether microhemorrhages represent a marker of intracranial hemorrhage has to be clarified.

A characteristic sign for moyamoya on MR imaging is leptomeningeal enhancement. It is related to the leptomeningeal flow engorgement, also known as the “ivy sign.” This sign is typically detected with T1-weighted contrast-enhanced and FLAIR images, and can be associated with poor visualization of the MCA. However, in patients with moyamoya who have large infarcts, poorer visualization of the MCA tree was not associated with leptomeningeal high signal intensity. The following differential diagnoses have to be kept in mind for the “ivy sign”: SAH, meningitis, brain tumor or meningeal carcinomatosis, subdural hematoma, CSF hypotension, and hyperbaric O2 therapy.

According to the guidelines for the diagnosis and treatment of spontaneous occlusion of the circle of Willis, angiography is not mandatory for the diagnosis of moyamoya disease, because MR imaging and MR angiography can show all the findings required for diagnosis. Nevertheless, these authors recommend diagnosis made with the aid of MR techniques alone only in childhood cases. The MR imaging should be performed on 1.5-T (or higher) systems, as suggested by the Research Committee for Diagnosis of moyamoya. Use of 3-T imaging provides higher resolution and can detect even smaller pathological vessels associated with moyamoya.

Different acquisition techniques are available for MR angiography, including 3D time-of-flight MR angiography, phase-contrast MR angiography, and contrast-enhanced MR angiography. Pulse sequence variations have been developed to improve these basic MR angiography methodologies, including turbo MR angiography, which represents 3D fast imaging with steady-state precession and magnetization transfer contrast, tilted optimized nonsaturated excitation, multislab techniques, and zero-filling interpolation, and has been described in a single report as a 98% accurate technique for diagnosis and assessment of moyamoya disease when compared with conventional angiography. These authors further analyzed the sensitivity, specificity, and accuracy of these MR angiography techniques regarding moyamoya features, including stenoocclusive lesions (100% for all 3 parameters); basilar cerebral moyamoya vessels (98, 100, and 98%, respectively); and leptomeningeal transdural collateral vessels (98, 100, and 98%, respectively) in a sample of 42 patients. Altogether this MR angiographic technique presents acceptable sensitivity (98%), specificity (100%), and accuracy (98%).

Postoperatively, MR angiography has been used for the assessment of bypass filling (with cine-phase MR angiography) and for the assessment of changes in the ECAs in cases of indirect revascularization (with time-of-flight). This method was used to evaluate the superficial as well as the deep temporal artery and the MMAs (with time-of-flight). These authors performed a very early postoperative follow-up evaluation and were able to describe dilatation of the superficial and of the deep temporal arteries in the first days after surgery.

From a temporal resolution point of view, MR DS angiography with high temporal resolution (< 1 second) provides images of cerebral hemodynamics that are almost equivalent to intraarterial DS angiography. However, spatial resolution is not as high. In the case of very small leptomeningeal anastomotic vessels in patients with moyamoya disease, these authors were able to recognize only an increased density of contrast rather than individual vessels themselves.

**Metabolic Studies**

Several metabolic studies can be used to assess the CBF reserve in patients with ischemia and to evaluate the need for revascularization.

Basically, 2 types of hemodynamic impairments can be assessed: 1) the response to a vasodilatory stimulus representing a reduced perfusion pressure, either through increased blood volume or through impaired blood flow response; and 2) noninvasive measurements of OEF that detect an increase in O2 extraction.

For the characterization of the level of stenosis in the main arteries, blood volume flow (in ml/minute), as used by one group, expresses the flow rate in the brain-supplying arteries and seems to be an indicator of vascular territory perfusion.

For the characterization of the parenchymal microcirculation, perfusion studies are performed. The CBF is measured, usually as the quantity of blood (in ml) perfusing a volume of brain (100 g) per unit of time (in minutes).

There are other parameters linked to CBF: the CBV is the amount of blood in a given volume of brain, whereas the MTT represents the time the blood takes to pass through the brain tissue. The 2 parameters are linked by the following relationship: CBF × MTT = CBV. The CBF and CBV values are tightly coupled.
Numerous factors affect CBF, including arterial perfusion pressure, intracranial pressure, blood viscosity, PaCO₂, pH, and O₂.

According to our current understanding of the brain microvasculature, there is autoregulation of CBF at the parenchymal level influenced by locoregional pH, nitric oxide, and other paracrine mediators through relaxation of the arteriolar smooth-muscle tone, permitting fine-tuning of blood flow. Assessment of autoregulation capacity presumes that blood flow within the microcirculation distal to stenotic large feeding vessels is maintained through the relaxation of arteriolar smooth muscle. In the case of patients with uncompensated moyamoya disease, once the maximum arteriolar dilation is reached, any further narrowing of the feeding vessels leads to decreased blood flow and ischemia.

Most imaging techniques used for the acquisition of CVR data require the administration of agents that can change resting blood flow.

Acetazolamide challenge is commonly used to assess cerebrovascular autoregulation, and is particularly useful in the assessment of chronic ischemic disease. This drug is an inhibitor of carbonic anhydrase and evokes a shift in the cerebral acid–base balance toward carbonic acidosis, thus increasing CBF.

Another technique that can be used for the same purpose is ETCO₂ manipulation. This technique has been assessed in combination with Xe-CT and, more recently, with BOLD MR in the study of patients with moyamoya disease (Fig. 2E and F).

With most techniques (for example, BOLD MR and Xe-CT), cerebrovascular reserve capacity maps are calculated as the absolute change of CBF from the baseline to the stimulation scan.

The Xe-CT Technique

The Xe-CT technique is used to assess hemodynamic stress by measuring CVR with a tolerance test (acetazolamide or CO₂). Stable xenon is a radiopaque diffusible gas that progressively enters tissues based on blood flow. This examination relies on dynamic CT scanning during inhalation of a gas mixture containing stable xenon and O₂. The Xe-CT technique requires patient cooperation and expensive add-on equipment. There can be associated side effects, including respiratory failure, headaches, nausea, vomiting, and convulsions. A xenon–inhalation assessment can be performed either in combination with acetazolamide or ETCO₂ manipulation. Cerebral blood flow maps can thus be obtained, processed with correction factors for ETCO₂, CO₂ variance, and hematocrit. A CBF value of < 30 ml/min/100 cm³ for adults or < 40 ml/min/100 cm³ for children is defined as an ischemic region, because normal CBF values in children are 20–60% greater than in adults. An increase of blood flow after acetazolamide of < 10 ml/min/100 cm³ is considered to be the threshold. However, for treatment decision-making purposes, these authors used the threshold value of 0, because areas with a value < 0 were found to represent areas with high stroke rates in a prospective study and to improve after indirect EC-IC bypass surgery.

The PET Modality

Positron emission tomography studies for hemodynamic assessment are usually performed with the following tracers: C¹⁵O PET for CBV assessment, H₂¹⁵O for CBF, and O₂ to measure OEF and CMRO₂. Nariai et al. identified differences in hemodynamic characteristics among the various clinical presentations of moyamoya disease. In this study, patients were classified as nonsymptomatic, those with hemorrhage, and those with ischemic symptoms. The latter group was subdivided into patients who presented with TIA only, patients with TIA and infarct TIA, and patients who suffered a permanent deficit following an infarction. Using PET and comparing results to healthy controls, the authors found that CBF in the patient groups classified as nonsymptomatic, TIA, and hemorrhage was not significantly lower than in healthy controls in any region. The CBV in the TIA and infarct TIA groups was significantly higher than in controls. The OEF in the frontal, parietal, and temporal cortex was significantly higher in the infarct TIA group than in the controls. Patients in the hemorrhage and permanent deficit groups had decreased metabolism with normal OEF. A paradoxical steal phenomenon is observed when the vascular bed is maximally dilated and autoregulation is completely lost. It is associated with increased CBV and increased OEF. Interestingly, increased CBV and a normal or mildly increased OEF is rarely observed in atherosclerotic patients, possibly due to the ability to invoke circle of Willis collateral vessels, which are lost in patients with moyamoya secondary to involvement of these collateral vessels by the disease process.
with moyamoya are therefore more dependent on a less effective pial collateral circulation for maintenance of flow. Patients in the infarct TIA group, who showed a marked increase in the OEF, were considered to be at risk for further deterioration. In these patients the MTT (which usually has an inverse relationship with cerebral perfusion pressure), was larger. Evidence of the effectiveness of revascularization was nicely shown in a series of 23 patients who were treated either with synangiosis or with bypass. Postoperative follow-up PET studies showed improved CO2 reactivity in this group.45

The PET modality is one of the most reliable assessment tools because it provides quantitative CBF, CBV, OEF, and CMRO2 information; however, there are still several limits. First, a cyclotron is required near the PET scan to deliver rapidly decaying tracers. Second, PET is associated with delivery of ionizing radiation: a complete 2D PET hemodynamic assessment (with CBF, CBV, CMRO2, and OEF) is associated with an effective dose of 8.9 mSv to the patient. Third, the number of available PET scanners is far less than MR, CT, or SPECT systems, although 3D PET is becoming available, and it offers the advantage of comparable accuracy with shorter scan duration and lower tracer doses. The radiation dose is reduced to approximately one-third: for example, for 15O2 the effective dose becomes 1.2 instead of 4.7 mSv, and a complete hemodynamic study is associated with an effective dose estimate of approximately 2.2 mSv.34

The SPECT Method

This methodology has also been used to assess regional CBF and CVR in moyamoya studies. It is performed using a radioactive blood tracer ([123I–IMP-ARG60 or 99mTc-HMPAO27,82) that is able to cross the blood-brain barrier and to remain fixed for enough time to allow the extracranial tomographic imaging of the gamma emission. Two separate imaging sessions (baseline and acetazolamide challenge) are required, usually separated by 24 hours to enable tracer washout between sessions. The tracer is deposited in proportion to blood flow, predominantly during the first pass through the microcirculation, and it is therefore a marker of flow without providing accurate perfusion in terms of ml/100 g/min. Also, the use of [123I–iomazenil has been described in patients with moyamoya disease, to evaluate neuronal damage.57

The MR Imaging–Based Techniques

For evaluation of stenosis in larger vessels in patients with moyamoya disease, blood flow volume studies in which dynamic 2D cine-phase contrast MR imaging is used have demonstrated a significant decrease in ICA blood flow volume associated with an impressive increase (nearly 2.5 times) in basilar artery blood flow.56 Flow in the ICA was measured at 118 ± 64 ml/minute in patients with moyamoya versus 254 ± 25 ml/minute in age-matched controls. Basilar artery flow measured 434 ± 166 ml/minute in patients with moyamoya versus 173 ± 13 ml/minute in controls.

Two methods are available for assessment of brain perfusion by MR imaging: 1) the "dynamic” susceptibility contrast technique uses principles such as indicator dilution methods in which a decrease in T2 or T2* MR signal intensity is related to the passage of a paramagnetic contrast agent through the capillary bed; and 2) the "arterial spin labeling” technique uses inversion or saturation pulses that label intravascular blood water protons before delivery into the brain slice of interest.33,25

An emerging technique for the assessment of CVR is BOLD MR imaging during manipulation of end-tidal PCO2.51,67 This method does not require the use of a pharmacological agent to change resting blood flow; instead, stimulus application is obtained through a rebreathing circuit in which end-tidal PCO2 levels are manipulated between normocapnea and hypercapnea for measuring vasodilatory reserve, and between normocapnea and hypocapnea to measure vasoconstrictive reserve. This is facilitated using a rebreathing device that permits precision control of ETCO2 in square wave patterns.84 At the same time, BOLD MR sequences identical to those used for functional MR studies are obtained. The CVR maps generated through this method provide spatial, quantitative maps of impaired CVR in terms of percent BOLD signal change per mm Hg change in ETCO2. Thus, the MR imaging method is threshold independent for determining the presence and spatial extent of exhausted autoregulation. Negative CVR responses (decreases in the BOLD MR imaging signal) are observed in patients with uncompensated moyamoya disease, indicating the presence of the steal phenomenon. Under these conditions, hypercapnia diverts blood away from maximally dilated vascular beds to unimpaired beds that retain the capacity for vasodilation.54,69,84 The advantages of this technique are the following: 1) measurements occur at the tissue level, providing spatial maps of impaired CVR; 2) the method does not require measurement of a vascular input function, which can be unreliable in patients with bilateral stenoses; 3) this technique can be added to conventional MR studies already used in the routine workup for these patients; and 4) the technique is safe, with no exposure to ionizing radiation, no injection is needed, and the range of ETCO2 required is within that achieved by patients in daily life situations (30–50 mm Hg).57

Proton-beam MR spectroscopy also deserves a brief note. This MR imaging technique provides insight into different brain metabolites. Interestingly, some authors have noted that patients with moyamoya disease have reduced levels of choline, creatinine, and N-acetylaspartate in the affected brain. A relative increase in these metabolites is then noted 6 months after revascularization surgery.76 Many authors consider N-acetylaspartate to be an important marker for healthy neurons.57

The Ultrasonography Modality

The ultrasonography modality combines both static and dynamic information, and provides supplementary data regarding quantitative flow characteristics (flow velocity), including the ratio between systolic and end-diastolic velocity, flow resistance, and vessel pulsatility. There are several studies confirming the potential of transcranial power Doppler and color Doppler ultrasonography in the detection of arteries that appear occluded on
TABLE 1: Effective radiation dose, acquisition time, and spatial resolution of different brain imaging modalities

<table>
<thead>
<tr>
<th>Imaging Modality</th>
<th>CT</th>
<th>2D DSA</th>
<th>CTA</th>
<th>Xe-CT</th>
<th>Perfusion CT</th>
<th>PET</th>
<th>SPECT</th>
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<tr>
<td>effective dose exposure</td>
<td>1.1–2.5 mSv&lt;sup&gt;16&lt;/sup&gt;</td>
<td>10.5 mSv&lt;sup&gt;3,87&lt;/sup&gt;</td>
<td>3.57–5.73 mSv&lt;sup&gt;16&lt;/sup&gt;</td>
<td>3.5–10 mSv&lt;sup&gt;44&lt;/sup&gt;</td>
<td>2–3 mSv&lt;sup&gt;44&lt;/sup&gt;</td>
<td>2.2–8.9 mSv&lt;sup&gt;44&lt;/sup&gt;</td>
<td>3.5–12 mSv&lt;sup&gt;44&lt;/sup&gt;</td>
</tr>
<tr>
<td>data acquisition time</td>
<td>0.5 min</td>
<td>15 min</td>
<td>20–70 sec&lt;sup&gt;24&lt;/sup&gt;</td>
<td>4 min&lt;sup&gt;44&lt;/sup&gt;</td>
<td>1–2 min&lt;sup&gt;44&lt;/sup&gt;</td>
<td>10 min&lt;sup&gt;44&lt;/sup&gt;</td>
<td>10 min&lt;sup&gt;44&lt;/sup&gt;</td>
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<tr>
<td>spatial resolution</td>
<td>1 mm</td>
<td>0.2 mm&lt;sup&gt;44&lt;/sup&gt;</td>
<td>0.6–1 mm&lt;sup&gt;44&lt;/sup&gt;</td>
<td>4 mm&lt;sup&gt;44&lt;/sup&gt;</td>
<td>1–2 mm&lt;sup&gt;44&lt;/sup&gt;</td>
<td>4–6 mm&lt;sup&gt;44&lt;/sup&gt;</td>
<td>4–6 mm&lt;sup&gt;44&lt;/sup&gt;</td>
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* Data were obtained either from recent literature or from our neuroradiological service. Variation in the radiation doses will occur, based on different manufactures and user-selectable parameters (such as kV and mA for the x-ray techniques). Abbreviations: CTA = CT angiography; DSA = DS angiography.

Conclusions

We have summarized the most recent imaging applications for the study of patients with moyamoya disease. Full characterization of the disease process requires a combined approach that assesses the anatomical and morphological features of the vascular network, the degree to which flow compensation is invoked through development of collateral vessels and lowering of vascular resistance via relaxation of tone in the peripheral resistance vessels (autoregulation), and the extent of metabolic impairment through measurement of O<sub>2</sub> consumption. No single imaging tool is capable of providing all of this information.

For assessing the vascular anatomy, conventional DS angiography continues to offer the highest spatial and temporal resolution; however, MR and CT angiography are adequate and are now commonly used for diagnosis and management.

Several methods are available for assessing autoregulation, each with certain limitations. Ultrasonography cannot provide maps of the extent and degree of CVR impairment at the tissue level. Acetazolamide SPECT and Xe-CT scans have been considered gold standards for the clinical assessment of CVR deficits. However, they suffer from reproducibility differences due to variations in patient response to acetazolamide. The PET method can be used to measure both CVR and tissue respiration, but requires a cyclotron on site. The BOLD MR modality with controlled ETCO<sub>2</sub> is very promising for CVR assessment, with the following advantages: 1) it is easily implemented on all modern MR imaging systems; 2) it is threshold independent for quantitative mapping of impaired autoregulation with vascular steal; and 3) it is not associated with ionizing radiation. Table 1 summarizes information worth considering when using methods based on ionizing radiation.

Currently at our institution we use the following neuroimaging tools: because the detection of diminutive circle of Willis vessels associated with innumerable small flow voids in the basal cisterns and basal ganglia on T2-weighted MR imaging sequences often establishes the diagnosis of moyamoya, we mostly use CT angiography to detail vascular morphology, and we use BOLD MR imaging with control of the ETCO<sub>2</sub> for perioperative and follow-up assessment of CVR.

Finally, in our clinical practice we always treat symptomatic hemispheres showing impaired CVR. Interestingly, we have frequently observed that treatment of one side is often also associated with contralateral CVR im-
provement. Our first-choice treatment modality is always the STA-MCA bypass. Only in the absence of suitable recipient vessels do we perform myosynangiosis. For MCA revascularization, we usually select the branch with the largest diameter. When stenosis includes the M, bifurcation, we analyze the spatial information in the CVR map to target the branch supplying the area with the greatest steal. Future outcome studies will allow us to determine the impact of using these CVR maps for surgical guidance.

Disclaimer

The authors report no conflict of interest concerning the materials or methods used in this study or the findings specified in this paper.

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