The Use of Quantitative Magnetic Resonance Perfusion for Assessment of CBF in The Perioperative Management of Carotid Stenosis: Case Illustration


Department of Neurological Surgery, Northwestern University Feinberg School of Medicine, Feinberg School of Medicine, 676 N. St. Clair, Suite 2210, Chicago, IL 60611, USA

Abstract: Objective and Importance: Internal Carotid Artery (ICA) stenosis is a common condition with a high prevalence in the normal population and carries a high risk of stroke. Hemodynamic impairment has been proven to be a powerful and independent risk factor for stroke. Understanding the hemodynamics beyond a stenosis may play a role in selecting patients who would benefit from treatment, determining the success of the treatment, and monitoring for disease recurrence. Our group has developed a unique quantitative approach to MR perfusion. We report on the use of Quantitative Magnetic Resonance Perfusion (MRP) for assessment of Cerebral Blood Flow (CBF) pre- and post- Carotid Angioplasty and Stenting for symptomatic ICA stenosis.

Clinical Presentation: A 71-year-old female with significant comorbidities presented with two episodes of left transient ischemic attacks (TIAs) over the course of one month related to high grade left ICA stenosis. Preoperative quantitative MRP showed asymmetry in CBF between the two hemispheres. Carotid angioplasty and stenting was recommended.

Intervention: Left ICA angioplasty and stenting was performed and reduced the stenosis to <10%. Quantitative MRP done after the procedure revealed improved CBF to the affected side and normalization in the asymmetry of perfusion between the two hemispheres. Quantitative MRP values showed that affected regions were hypoperfused, but flow was sufficiently maintained so as to avoid infarct.

Conclusion: Relative MRP images allowed us to elucidate improvement in perfusion resulting from carotid angioplasty and stenting. This novel quantitative MRP allows quantification of CBF and the potential to provide additional information on the degree of hemodynamic compromise.

Keywords: Carotid artery angioplasty and stenting, cerebral blood flow, intracranial atherosclerosis, magnetic resonance perfusion, quantitative magnetic resonance perfusion.

INTRODUCTION

Internal Carotid Artery (ICA) stenosis, caused by atherothrombotic deposition, is a common condition with a high prevalence in the normal population (~0.5% in patients over 65-year-old and ≥ 10% between ages 75-85). The annual average risk of stroke for an asymptomatic stenosis varies between 1% and 3%, and may reach up to 15% if the stenosis is symptomatic [1]. Stenosis of the extracranial ICA accounts for less than 20% of all strokes and that of the intracranial arteries alone or in combination with extracranial stenosis accounts for another 20% [1]. In the presence of stenosis, the reduction in the mean arterial pressure of the cortical arteries varies with the degree of stenosis and depends on the status of the collateral circulation [2]. Knowing that the Circle of Willis and/or the external to internal carotid artery collaterals may maintain the perfusion pressure beyond a stenosis makes it impossible to correlate a percent of asymptomatic stenosis (Example > 70%) with hemodynamic impairment. Although most ischemic strokes secondary to ICA stenosis are attributed to embolic phenomena, the presence of hemodynamic impairment has been proven to be a powerful and independent risk factor for stroke [2-4]. For this reason, understanding the hemodynamics beyond a stenosis may play a role in selecting patients who would benefit from treatment, determining the success of the treatment, and monitoring for recurrence. Available techniques for assessing CBF include positron-emission tomography (PET), single-photon emission computed tomography (SPECT), and various Magnetic Resonance techniques that use relative quantification. We report on the use of Magnetic Resonance Perfusion (MRP) for assessment of CBF pre- and post-Carotid Angioplasty and Stenting (CAS) in a 71-year-old woman who presented with symptomatic left ICA stenosis. Quantitative MRP values were calculated using a recently developed technique [5,6].

MATERIALS AND METHODOLOGY

Patient

The patient is a 71-year-old female with a history of severe coronary artery disease and Chronic Obstructive Pul-
Pulmonary Disease (COPD) who presented with two episodes of left hemisphere TIAs (transient right hand weakness, but no aphasic trends) over the course of one month related to high grade left carotid artery bifurcation stenosis (Fig. 1A). Carotid angiography revealed a 90% stenosis of the left ICA at the carotid artery bifurcation as well as 50% stenosis more proximally in the left common carotid artery. CAS was recommended and performed due to her co-morbidities. ICA angioplasty and stenting reduced the stenosis to <10% (Fig. 1B). The more proximal stenosis was not treated. No complications occurred due to the procedure. At one month follow-up the patient denied any TIAs and was neurologically intact. Follow-up carotid duplex showed no re-stenosis.

MRI Studies

Quantitative CBF (qCBF) images were acquired preoperatively using the standard stroke imaging in our institute. This includes T1, T2, FLAIR, Time Of Flight (TOF) and perfusion/diffusion weighted images. Preoperatively the perfusion scans were acquired using a recently reported MRI technique [7] which quantifies cerebral perfusion based on a simple modification of the standard deconvolution analysis [5, 6, 8]. Preoperative assessment of qCBF acquired a standard gradient recalled echo planar imaging (GE-EPI), MRI perfusion (TR/TE=1440 ms/47 ms, 12, 5.0 mm slices) with a 20 ml injection of contrast agent. Post-operatively an additional contrast enhanced MR angiogram was acquired (3D FLASH, FOV = 310 mm x 251 mm, matrix= 512 x 416, BW=490 Hz/pixel, 80 1.0 mm slices, TR/TE/F.A.=3.36 ms/1.18 ms/25°, contrast = 20 ml GD-DTPA). The additional contrast injection served as a preloading contrast dose that allowed the acquisition of a slightly modified MR qCBF scan. In this case, a single shot spin-echo EPI (TR/TE=1500 ms/72 ms, 13, 5.0 mm slices) was acquired in combination with an 8-channel head coil and parallel imaging technique (iPAT, 2X acceleration). Spin-echo perfusion images have been shown to be less prone to inhomogeneity artifact than
GRE-EPI sequences [9]. In both cases, 1.78 x 1.78 mm voxels provide spatial resolution sufficient to distinguish white and gray matter.

RESULTS

The observed asymmetry in cerebral perfusion was quantified using a region of interest analysis. Regions of interest were placed bilaterally to cover deep white matter in the mid-ventricular region. Pre-operatively, the relative cerebral perfusion showed the right side to be 25% higher than the affected side. Post operatively, perfusion in the right and left sides were normalized to be within 3% of each other. Quantitative perfusion for the right/left hemispheres were 24.42 +/- 9.41 ml/100g-min and 19.53 +/- 5.76 ml/100g-min respectively (pre-operatively) and 21.96 +/- 8.05 and 21.52 +/- 6.98 ml/100g-min, respectively (post-operatively).

DISCUSSION

Stroke is the leading cause of disability and the third most common cause of death in the United States. Despite the various sources of thromboembolism, atherosclerosis of the carotid artery is the most common cause of ischemic stroke [10]. At the carotid bulb wall, low shear stress and flow separation with stasis and nonlaminar flow creates an environment favorable for blood-borne lipid particles to interact with the vessel wall and leads to the formation of fatty streaks containing mononuclear cells and lipid containing macrophages creating an atherosclerotic plaque [10]. On the other hand, hyperplastic fibrosclerotic changes in the sub-intima leads to a fibrotic thickening in the intima-media complex which hosts atherothrombotic deposition at those intimal sites leading to the development of stenosis [1]. When the deposition exceeds 1.5 mm it forms a plaque consisting of a fibrous cap encapsulating various cellular (inflammatory cells e.g. leukocytes and macrophages), extracellular (collagen, fibrin, and smooth muscle cells), and lipid components [1]. As the plaque develops and enlarges it increases the degree of stenosis until it becomes hemodynamically significant when exceeding 70% of the lumen thus jeopardizing CBF. However, most patients, due to the chronic nature of the disease, develop compensatory collateral supply and remain asymptomatic [10, 11]. It is recognized that the presence of carotid stenosis does not predict the presence or degree of hemodynamic compromise in the distal cerebral circulation. Rather, it is the impaired cerebrovascular reactivity that is predictive of cerebral ischemia [12]. One of the goals of an operation or an intervention in patients with atherosclerotic carotid disease is to prevent ischemic stroke by improving CBF.

In patients with severe carotid stenosis, whether the mechanism of ischemia is hemodynamic or embolic may not be important [2]. Especially that, those patients with the improvement of cerebral hemodynamics after treatment will have a decreased chance of ischemic injury. “Hemodynamic impairment” is the term used to describe the presence of reduced cerebral perfusion pressure (CPP) [2]. In any region of the brain the CPP equals the mean regional arterial pressure minus either the venous pressure or the intracranial pressure, depending on which is higher [2]. Under normal circumstances CPP would be normal and the CBF would be closely matched to the resting metabolic rate of the brain tissue (Stage 0 hemodynamic impairment), consequently the Oxygen Extraction Fraction (OEF) shows little regional variation. When CPP is moderately reduced, cerebrovascular autoregulation maintains a constant CBF by arteriolar vasodilation (Stage 1 hemodynamic impairment). This may cause increase in the intravascular (Cerebral Blood Volume) CBV. With severe CPP reduction, the autoregulation fails to compensate and the CBF begins to decline, but the cerebral brain metabolism is maintained by the increase in OEF (Stage II hemodynamic impairment = ‘Misery Perfusion’) [2, 13]. The average baseline OEF is approximately 30% but may increase up to 80% [2]. Stage I hemodynamic compromise was found by some investigators to correlate with an increased stroke risk while others found no correlation [13]. A positive association of Stage II cerebral hemodynamic compromise and stroke risk has been documented [13, 14].

We do not routinely measure the mean arterial pressure or the CPP in brain arteries to identify hemodynamic impairment, rather we rely on identifying compensatory mechanisms by physiological imaging [2]. The main reason is that imaging modalities (such as PET and SPECT) capable of quantifying these hemodynamic variables are expensive,
not commonly available, use ionizing radiation, and may have low spatial and temporal resolution (as is the case with SPECT) [15]. The physiologic imaging modalities in use are usually divided into three major categories. The first are those that compare the baseline measurement of blood flow or velocity to the measurement after a vasodilatory stimulus [2]. The second are those that identify autoregulatory vasodilation by either measuring the mean transit time (MTT) or calculating it from the ratio of CBF and CBV [2]. The last category of imaging modalities measure the OEF [2].

Among the commonly used physiologic imaging modalities are the Transcranial Doppler sonography (TCD) and the MR. The TCD is a noninvasive technique that has been proven reliable for measuring CBF velocity in large cerebral vessels [15]. This technique is not without limitations such as the variation in the measurement in the setting of cardiac arrhythmia and the inability to perform the exam on the middle cerebral artery (MCA) in 5-20% of the patients due to the insufficient ultrasound transmission through the skull [15]. As for the perfusion-weighted MR, it is a semiquantitative technique that has a good spatial resolution and is capable of assessing the microcirculation like PET or SPECT without the need for ionizing radiation [15]. Another advantage of perfusion-weighted MRI is that it does not consume a lot more time than routine MRI. There have been some reports on the use of MRI to quantitate CBF. One example is the work by Lythgoe and colleagues [3]. This group quantitated CBF using MR, based on empirically derived scale factor. The use of empirically derived scale factors has been shown to be less reliable than patient based qCBF measurement [16]. This is particularly relevant in this case where compensatory mechanisms increase perfusion to the contralateral hemisphere.

On the other hand, another MRI technique that has the potential to determine quantitative CBF is arterial spin labeling (ASL). In ASL imaging, the magnetization of flowing blood is inverted upstream of the region in which the perfusion measurement is desired. As the blood flows into the imaging regions, the inverted magnetization is transferred to the tissue, altering local magnetization in proportion to CBF. By measuring the change in local magnetization, CBF can be calculated. A plethora of tagging and readout schemes have been developed [17-20] and have shown the potential for measuring quantitative CBF [21-23]. The signal changes observed in ASL are small, typically 1-2%, so that acquiring useful images requires multiple signal averages and scan times of several minutes. Furthermore, with ASL perfusion imaging, calculation of quantitative CBF depends on a variety of physiologic parameters (T1 of blood, T1 of tissue, water diffusion rate, etc) [24]. In particular, the accuracy of CBF depends on the time it takes for the blood to flow from the labeling region to the readout slice. Some ASL techniques can minimize this dependence when transit times are within “normal ranges”. In patients with severely altered hemodynamics, this transit time can confound the accurate determination of CBF [25]. Therefore, although ASL perfusion has been shown to measure quantitative CBF, its utility in the setting of hemodynamic compromise, where label-to-tissue transit times cannot be predicted, has not been established.

Conversely, the limited availability and low spatial resolution of PET perfusion scans has prompted the development of perfusion image techniques that have the potential to be disseminated in the broader medical community. MRI based qCBF images used in this case report are relatively inexpensive, safe to use and can be readily added to a comprehensive neurological MRI examination which includes: T1- and T2-weighted anatomic images, angiography and diffusion images. This report uses a unique approach to quantify CBF using readily available MRI technology known as the “bookend technique”. The bookend technique combines a single-image two-point quantitative blood volume image with whole brain dynamic CBF images to yield whole brain quantitative CBF images. As with more traditional approaches that quantify perfusion, the bookend technique ultimately depends on tracer kinetics and mathematical deconvolution to calculate CBF, CBV and MTT. However unlike the commonly used MRI-based relative measures of CBF, the bookend perfusion scans are able to quantify cerebral perfusion. Furthermore, the bookend technique does not rely on empirical correction factors derived from population averaged CBF values. Instead, the bookend technique can determine a patient specific quantification based on the T1 changes of the parenchyma resulting form the injection of contrast agent. A large prospective study comparing the quantitative results from this technique with the relative measurements obtained from MRP and the clinical status of the patient will be needed to better estimate the value of this new technique.

CONCLUSION

MR images of relative cerebral perfusion reflect changes in local hemodynamic impairment resulting from surgical intervention. It is hoped that our ability to quantify changes in cerebral perfusion using an innovative quantitative MR will allow us to determine which patients have hemodynamically significant stenosis and would benefit from intervention. Future work involve further validation of this technique in selected patients allowing the inclusion of qCBF into the diagnostic algorithm and selection of patient who would benefit from treatment.

REFERENCES

[8] Ostergaard L, Sorensen AG, Kwong KK, Weisskoff RM, Gyldensted C, Rosen BR. High resolution measurement of cerebral blood flow using intravascular tracer bolus passages. Part II: Experimen-