Reduction in cerebral blood flow in areas appearing as white matter hyperintensities on magnetic resonance imaging

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Abstract

The purpose of this study was to examine cerebral blood flow (CBF) as measured by arterial spin labeling (ASL) in tissue classified as white matter hyperintensities (WMH), normal appearing white matter, and grey matter. Seventeen healthy older adults received structural and ASL MRI. Cerebral blood flow was derived for three tissue types: WMH, normal appearing white matter, and grey matter. Cerebral blood flow was lower in WMH areas relative to normal appearing white matter, which in turn, was lower than grey matter. Regions with consistently lower CBF across individuals were more likely to appear as WMH. Results are consistent with an emerging literature linking diminished regional perfusion with the risk of developing WMH.

1. Introduction

White matter hyperintensities (WMH) are areas of increased signal intensity detected on T2-weighted magnetic resonance imaging (MRI) scans. These lesions are common among older adults (Brickman et al., 2008a) and are thought to reflect small vessel vascular disease or β-amyloid peptide deposition (Gurol et al., 2006). Increased WMH volume is associated with degree of cognitive impairment among neurologically healthy older adults (Gunning-Dixon and Raz, 2000) and is predictive of rate of cognitive decline among those with Alzheimer’s disease (Brickman et al., 2008b). Although these associative findings have been consistently reported, the functional significance of WMH is unclear.

Increases in total or frontal WMH volume have been associated with decreases in relative glucose metabolism in frontal regions among healthy older adults (DeCarli et al., 1995). In a longitudinal analysis, individuals with increasing severity of WMH over an 8-year period had concomitant increases in relative cerebral blood flow (CBF) in temporal areas and anterior cingulate and decreases in relative CBF in more posterior areas (Kraut et al., 2008). Further, increased severity of WMH was not associated with total parenchymal CBF in a cross-sectional analysis, but decreases in CBF over time increased the risk of developing periventricular WMH (ten Dam et al., 2007). Holland and colleagues recently showed that WMH frequency among healthy older adults and those with AD or cerebral amyloid angiopathy is greater in regions with relatively lower perfusion among younger adults (Holland et al., 2008). These findings raise the possibility that lower regional perfusion increases the risk of developing WMH.

In the current study, we sought to extend these findings by examining arterial spin labeling (ASL)-derived CBF in areas appearing as WMH on T2-weighted fluid attenuated inverse recovery (FLAIR) MRI scans in comparison to CBF in normal appearing white matter (NAWM) and grey matter among neurologically healthy older adults. Two previous studies used perfusion-weighted MRI and found a decline in blood flow in areas appearing as WMH on T2-weighted images compared with NAWM (Marstrand et al., 2002; Sachdev et al., 2004). Thus, we hypothesized that CBF would be decreased in areas appearing as WMH compared with CBF in NAWM and grey matter.

2. Methods

2.1. Subjects

Seventeen older adults ranging in age from 61 to 70 (mean ± S.D. = 64.94 ± 2.95) comprised the study sample. There were 9 women (53%) and the average number of years of formal education was 16.29 (S.D. = 1.72). Men and women did not differ in age (t(15) = 0.235, P = 0.817) or number of years of education (t(15) = 0.371, P = 0.716). All participants were right-handed and spoke English. By self-report and interview, participants did not have past or current histories of medical, neurological, psychiatric disorders or treatment with...
psychotropic medication. Participants were screened for dementia with interview and psychometric assessment (Mattis, 1988); none met diagnostic criteria. Subjects were recruited for participation in an ongoing functional magnetic resonance imaging (fMRI) study of working memory and aging. The study was approved by a local ethics committee, and all participants gave written informed consent to participate in the study.

2.2. Magnetic resonance imaging

2.2.1. Image acquisition

All neuroimaging took place in a single session on a 1.5T Philips Intera scanner. Standard high-resolution T1-weighted anatomical images were acquired in the axial plane using the following parameters: FOV = 256 × 256 mm, matrix = 256 × 256, slice thickness = 1.5 mm, TR/TE = 25 ms/3 ms. T2-weighted FLAIR images were acquired for WMH quantification with the following parameters: FOV = 256 × 256 mm, matrix = 256 × 192, slice thickness = 3 mm, TR/TE = 11,000 ms/144 ms. Detailed description of acquisition of continuous arterial spin labeling (CASL) perfusion images is provided elsewhere (Asllani et al., 2008). Briefly, 30 spin echo, echo planar (EP) CASL control and labeled images were acquired axially (inferior to superior) with a labeling duration of 2000 ms, post-labeling delay of 800 ms, in-plane resolution 3.4 × 3.8 mm, slice thickness of 7.5 mm, FOV = 220 × 220 mm, and TR/TE = 5000 ms/35 ms. Adiabatic inversion labeling of the water spins and correction for off-resonance saturation were implemented following procedures put forth by Alsop and Detre (Alsop and Detre, 1996; see Asllani et al., 2008).

2.2.2. MRI data preprocessing

Preprocessing steps were implemented with the SPM99 software package (Wellcome Department of Cognitive Neurology) with in-house developed software written in Matlab (Mathworks, Natick, MA, USA), described fully in a previous report (Asllani et al., 2007). All CASL and T1-weighted images were coregistered to the first CASL image in the acquisition series. The T1-images were spatially normalized using 7 × 8 × 7 nonlinear basis functions into Talairach space defined by the Montreal Neurologic Institute (MNI) and segmented into grey matter, white matter, and cerebral spinal fluid (CSF) posterior probability images. These tissue segment probability maps were thresholded at 0.80 (80%) to include only voxels with high probability of belonging to a tissue class and reduce the potential influence of partial volume effects. The spatial transformation was applied to the coregistered CASL and tissue segment images; these procedures ensured that all CASL and anatomical images were aligned in the same stereotactic space.

2.2.3. Computation of cerebral blood flow

For computation of cerebral blood flow (CBF), the paired control/label images in Talairach space were used to calculate percent change images following a modified two-compartment formula, as detailed elsewhere (Asllani et al., 2008). In each voxel, CBF values were weighted by the tissue type (i.e., grey matter, white matter, CSF) posterior probability as determined from the T1-weighted image tissue segment. Finally, CBF images were smoothed with a 6-mm spatial kernel.

2.2.4. Quantification of white matter hyperintensities

For quantification of WMH, FLAIR images were imported into the publicly available software package MRICro (Rorden and Brett, 2000), and WMH volumes were determined following a three-step process put forth by Gurol et al. (2006). First, based on a priori knowledge of the distribution of voxel intensities and visual inspection of each individual image, we determined the intensity threshold that labeled voxels appearing as hypointense. Second, on a slice-by-slice basis, regions of interest (ROIs) were manually traced to define gross areas containing WMH while excluding non-WMH areas that were labeled in step 1 (e.g., dural fat). Third, the intersection of the labeled voxels in Step 1 and the manual ROIs yielded the volume of WMH in cm³. These ROIs were treated as a repeated measure (i.e., within-subjects factor). There was a significant overall difference in CBF among these regions (F(2, 32) = 185.61, P < 0.001). Pairwise comparisons indicated that CBF was significantly lower in WMH than in NAWM, which, in turn, was significantly lower than CBF in grey matter (P ≤ 0.001 for all pairwise comparisons). These results are displayed in Fig. 1. Further, the non-parametric Wilcoxon signed ranks test confirmed significant differences among groups (all group comparisons, P < 0.004). Fig. 2 displays an image in which voxels labeled as WMH in at least five subjects are superimposed on the mean CBF image derived by averaging the CASL images from all subjects.

We examined whether the mean CBF value varied as a function of the likelihood that any given voxel was labeled as WMH. To do this analysis, we created a conjunction map of all individual WMH masks from individual subjects in standardized anatomical space. We grouped voxels by frequency and compared mean CBF values across frequency values, which ranged from 1 to 15 (note that no single voxel was labeled as a WMH in all 17 subjects). As displayed in Fig. 3, there was a subtle, but significant (F(14, 21787) = 174.439, P < 0.001) linear decline (P for linear trend = 0.045) across frequency groups. Next we calculated the standard deviation and coefficient of variation (a normalized metric of dispersion, σ/µ) for mean CBF values in each frequency group and correlated those values against frequency. Decreases in both the standard deviation (Spearman’s r = −0.889, P < 0.001) and the coefficient of variation (Spearman’s r = −0.796, P < 0.001) were strongly related to increased frequency. Taken together, these findings suggest that regions with consistently low CBF across individuals are more likely to appear as WMH.

4. Discussion

In the current study, we compared CASL-derived mean CBF across three tissue types — areas appearing as WMH on T2-weighted FLAIR MRI, NAWM, and grey matter — in well-screened, neurologically healthy older adults. Findings indicated that CBF was significantly reduced in areas appearing as WMH compared with both NAWM and grey matter. These results are consistent with previous reports in demonstrating an association between WMH and reduced blood flow, and replicate findings suggesting that this association is specific to the tissue appearing as WMH itself (Marstrand et al., 2002; Sachdev et al., 2004).

White matter hyperintensities are common findings on T2-weighted images among older adults. Although WMH were previously
thought to lack clinical significance, several studies now confirm that they are associated with poorer cognitive function, may put individuals at risk for the development of dementia, and are more prevalent among certain psychiatric diagnoses. Results from the current study suggest that these associations may be due partially to a reduction in perfusion in areas appearing as WMH. However, causal directionality is difficult to infer from cross-sectional studies. It is possible, for example, that small vascular lesions appear first as WMH and then have a secondary effect on underlying perfusion or that the two phenomena occur concomitantly without a causal link. A recent study by Holland et al. (2008) speaks against this possibility. They compared the spatial distribution of WMH among healthy older adults and those with AD or cerebral amyloid angiopathy as it related to a normative atlas of perfusion values derived from neurologically healthy, younger adults and found that WMH frequency was greater in regions with lower normative perfusion values. Similarly, O’Sullivan et al. (2002) showed that CBF was reduced in periventricular NAWM among patients with ischemic leukoaraiosis compared with controls, suggesting that diminished regional white matter perfusion may be a “risk factor” for the development of WMH. Further, a recent report showed an inverse association between cardiac output and subcortical WMH volume, suggesting that systemic hypoperfusion may increase the risk for development of WMH (Jefferson et al., 2007). Future longitudinal or serial MRI studies are required to determine the exact causal relationship between diminished perfusion and WMH.

Visual inspection of Fig. 2 shows reduced CBF in areas labeled as WMH in at least five participants from the current study. The figure also reveals that areas that are most vulnerable to the development of WMH, namely periventricular regions that are confluent with the lateral ventricles, appear to have a more profound reduction in blood flow than white matter that is distributed in deeper cortical areas. Findings displayed in Fig. 3 suggest that areas that are of consistently low CBF are most likely to be labeled as WMH across individuals.

Two potential limitations should be pointed out. First, atrophy-associated ventricular enlargement may have increased the likelihood of contamination by the CSF of the WMH voxels. To limit this possibility, we restricted our analyses to areas with high probability of belonging to a tissue class. Second, although not visualized on T1-weighted images, WMH regions may create subtle signal alterations, which might have affected tissue segmentation used for conversion to CBF of ASL percent change data. Third, it is also conceivable that tissue transit times are longer in WMH, possibly due to associated vasculature changes. This transit time confound would cause an underestimation of CBF. Future work is needed to separate the effect of transit time from absolute CBF reduction in WMH as this would aid in our understanding of the underlying mechanism of the observed CBF reduction in WMH.

Magnetic resonance imaging has emerged as having the utility for identification of the structural and functional correlates of cognitive aging. Converging evidence from animal and human research literature points to age-associated changes in white matter as a particularly salient feature of the aging process. These observations are consistent with a “disconnection” hypothesis of cognitive aging, which posits that integration of coordinated neural systems declines with advancing age (Andrews-Hanna et al., 2007). The current study highlights the possibility that WMH and/or their associated diminution in blood flow may contribute to this disconnection by disrupting neural communication across white matter fibers. Future studies should integrate these MRI modalities with comprehensive cognitive data and evaluate other aspects of white matter integrity.

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