Severe asymptomatic carotid stenosis is associated with robust reductions in homotopic functional connectivity

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ABSTRACT

Severe (>70% narrowing) asymptomatic carotid stenosis (SACS) is associated with cognitive impairment and future strokes, and connectivity basis for the remote brain consequences is poorly understood. Here we explored homotopic connectivity and parenchymal lesions measured by multimodal magnetic resonance imaging (MRI) parameters in patients with SACS. Twenty-four patients with SACS (19 males/5 females; 64.25 ± 7.18 years), 24 comorbidity-matched controls (19 males/5 females; 67.16 ± 6.10 years), and an independent sample of elderly healthy controls (39 females/45 males; 57.92 ± 4.94 years) were included. Homotopic functional connectivity (FC) of resting-state functional MRI and structural connectivity (SC) of deterministic tractography were assessed. Arterial spin labeling based cerebral perfusion, susceptibility weighted imaging based micro-hemorrhagic lesions, and T2-weighted white matter hyperintensities were also quantified. Significant and robust homotopic reductions (validated by the independent dataset and support vector machine-based machine learning) were identified in the Perisylvian fissure in patients with SACS (false discovery rate corrected, voxel p < 0.05). These involved regions span across several large-scale brain systems, which include the somatomotor, salience, dorsal attention, and orbitofrontal-limbic networks. This significantly reduced homotopic FC can be partially explained by the corrected white matter hyperintensity size. Further association analyses suggest that the decreased homotopic FC in these brain regions is most closely associated with delayed memory recall, sensorimotor processing, and other simple cognitive functions. Together, these results suggest that SACS predominantly affects the lower-order brain systems, while higher-order systems, especially the topographies of default mode network, are least impacted initially, but may serve as a hallmark precursor to vascular dementia. Thus, assessment of homotopic FC may provide a means of noninvasively tracking the progression of downstream brain damage following asymptomatic carotid stenosis.

1. Introduction

Patients with severe (>70% narrowing) asymptomatic carotid stenosis (SACS) are at moderate to high risk of progressive cognitive deterioration and cerebrovascular events (de Weerd et al., 2014). Therefore, the clinical focus for these patients is the management of future strokes. However, best management practices, and which populations might benefit from interventions, are still debated (Rosenfield et al., 2016). One reason for this lack of consensus stems from the poorly understood neural basis of SACS.

Compromised hemodynamics and connectivity may promote the etiology of cognitive decline and vulnerability to strokes. Pathologically, severe stenosis is associated with potential hemodynamic changes, silent cerebrovascular disease, and varying degrees of structural damage (van Veluw et al., 2017). Functionally, patients with SACS have reported altered connectivity that involves intra- and inter-hemispheric large-scale systems (Avelar et al., 2016; Baradaran et al., 2016; Cheng et al., 2012; Lin et al., 2014; Panczykowski et al., 2014). Despite existing documentation, there is still a lack of objective imaging metric that may be noninvasively utilized to understand and

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track the progress of the disease. It is known that the most critical endpoint of SACS is the occurrence of ischemic strokes and vascular dementia. Both SACS and ischemic strokes preferentially affect low-level brain functions that are closely related to sensorimotor processing, and patients with SACS show similar declines in the same cognitive domains to those with strokes (Grefkes and Fink, 2014). Moreover, as a type of silent cerebrovascular disease, SACS can be considered a predecessor to strokes. For example, patients with SACS show altered functional as well as structural connectivity that mimics ischemic stroke (Cheng et al., 2012; Lin et al., 2014; Panczykowski et al., 2014). Therefore, imaging measures that best capture stroke pathophysiology may also be beneficial for investigating SACS.

Considering the complexity and heterogeneous nature of carotid stenosis, metrics used for tracking the brain state should reflect multiple dimensions, such as cognition and disease sensitivity. Since the pioneering work of Biswal and colleagues (Biswal et al., 1995), resting-state functional magnetic resonance imaging (R-fMRI) has promoted the use of noninvasive assessments of the brain's functional architecture into clinical experimentation. Functional connectivity (FC) of R-fMRI measures hemodynamic correlations between regional low frequency (<0.1 Hz) blood oxygen level dependent (BOLD) signals. One architectural feature, functional homotopy, or voxel-mirrored homotopic interhemispheric connectivity (VMHC) (Zuo et al., 2010), quantifies voxel-wise functional connectivity of the mirror-symmetrical pairs of voxels between hemispheres. VMHC represents a robust metric (Zuo et al., 2013; Zuo and Xing, 2014) that reflects interhemispheric integration, which has been linked to stroke recovery (Grefkes and Fink, 2014; Siegel et al., 2016; van Meer et al., 2010) and other neurological and psychiatric disorders (Anderson et al., 2011; Hopman et al., 2012).

In this study, we aimed to explore the homotopic FC changes of R-fMRI in patients with SACS, while considering: homotopic structural connectivity (SC) with deterministic tractography, white matter hyperintensity (WMH) loads with T2-weighted Fluid-attenuated inversion recovery (FLAIR), and cerebral blood flow (CBF) measured by pulsed arterial spin labeled (PASL). We hypothesized that patients with SACS would demonstrate significant and classifiable reduced functional homotopy, predominately in the somatomotor, attention, and executive systems, and the cerebral perfusion and silent lesions can partially explain these changes.

2. Methods

2.1. Participants

Twenty-four SACS patients were recruited between January 2015 and June 2016. The inclusion criteria were: (a) 55 – 80 years; (b) unilateral internal carotid artery (ICA) stenosis ≥70%; (c) right-handed; (d) free of stroke, transient ischemic attack, or dementia; (e) Modified Rankin Scale (SALTER et al., 1999) score of 0 or 1. Exclusion criteria were: (a) contralateral ICA stenosis ≥ 50%; (b) posterior circulation diseases; (c) Mini-Mental State Examination (MMSE) (FOLSTEIN et al., 1975) score < 26; (d) functional disability (Modified Rankin Scale ≥ 2); (e) severe systemic diseases and neuropsychiatric diseases; (f) contraindications for MRI; and (g) poorly educated (< 6 years). In this same timeframe, we recruited 24 comorbidities- and demographically- matched healthy controls (HCs). Additionally, we included an independent dataset of elderly HCs (age, 57.92 ± 4.94 years; 39 females/ 45 males) to validate our results. The present study was approved through the Medical Ethics Committee of the Affiliated Hospital of Zhongnan Hospital of Wuhan University, and all participants gave written informed consent. Detailed demographics and clinical measures are listed in Table 1.

| Table 1 | Demographics and clinical characteristics. |
|---------|------------------------------------------|
|          | Healthy (n = 24) | Patients (n = 19) | p |
| Gender (Male/Female) | 19/5 | 15/4 | 0.99* |
| Age (yrs.) | 67.16 ± 6.10 | 64.25 ± 7.18 | 0.17 |
| Education (yrs.) | 11.13 ± 3.46 | 9.58 ± 2.83 | 0.12 |
| Hypertension | 18/6 | 17/2 | 0.23 |
| Diabetes | 4/20 | 4/15 | 0.71 |
| Hyperlipidemia | 11/13 | 9/10 | 0.92 |
| Smoke | 6/18 | 9/10 | 0.13 |
| Affected side | – | 7L/12R | |
| MMSE | 27.38 ± 0.65 | 26.84 ± 0.69 | 0.015* |
| MoCA | 24.21 ± 1.56 | 23.26 ± 1.15 | 0.017* |
| Word fluency | 37.13 ± 4.01 | 33.42 ± 6.20 | 0.258 |
| Digit symbol substitution | 31.50 ± 5.51 | 28.00 ± 4.74 | 0.029* |
| Backwards digit-span | 6.46 ± 0.93 | 5.79 ± 0.98 | 0.042* |
| Forwards digit-span | 4.46 ± 0.78 | 3.84 ± 0.83 | 0.021* |
| Immediate recall | 35.79 ± 5.63 | 31.00 ± 4.45 | 0.004** |
| Delayed recall | 6.50 ± 1.14 | 4.63 ± 1.57 | <0.001*** |
| WMHCcorrected | -0.59 ± 1.01 | 0.64 ± 0.71 | <0.001*** |
| WMHsize (ml) | 1.56 ± 3.05 | 10.76 ± 13.48 | 0.005** |
| WMHnumber | 5.87 ± 6.31 | 13.42 ± 5.84 | <0.001*** |

Data are presented as mean ± SD. Abbreviations: MMSE, Mini-Mental State Examination; MoCA, Montreal Cognitive Assessment; a, p-value was obtained using the two-tailed Chi-squared test; WMH = white matter hyperintensity; WMHCcorrected, white matter hyperintensity (corrected by total cranial volume, and log10 transformed); WMHsize, white matter hyperintensity volumes; WMHnumber, number of WMH.

2.2. Neurobehavioral assessments

Neurobehavioral assessments, including the MMSE, the Montreal Cognitive Assessment (MoCA) (NASREDDINE et al., 2005) Beijing Version, the Digit Symbol, and the Rey Auditory Verbal Learning Tests, were performed within seven days of MRI scan. The MMSE and MoCA Beijing Version were utilized to assess global cognition. The Digit Symbol Test required subjects to translate numbers to paired symbols as quickly as possible, and number of correct translations within 90 s was recorded. The Rey Auditory Verbal Learning Test (RAVLT) was used to evaluate memory and verbal learning ability. Participants were given a list of 15 unrelated words and asked to recall as many words as he/she could remember. This procedure was then repeated four more times. After a thirty-minute delay, the participants were once more asked to recall as many word as possible from the original list. The total number of the words immediately recalled during the first five repeats and the sum of the delayed recall were recorded. In the Verbal Memory Test, participants were required to repeat orally presented lists of numbers, beginning with a 2-number sequence, and increasing by one additional number for each correct performance. In the forward span, participants were asked to recite the span in forward order. In the backward span, participants were asked to recite the span in reverse order.

2.3. MRI data acquisition

MRI data were collected using a MAGNETOM Trio 3.0 T MR scanner (Siemens, Germany). The scanning sessions included (1) T1 high-resolution anatomical images (176 sagittal slices, 1-mm in-plane resolution); (2) resting-state BOLD volumes (33 axial slices, 3.8-mm slice thickness with a 0.3-mm gap, repetition time (TR) = 2 s, 240 volumes); (3) diffusion tensor imaging (DTI) (TR / echo time (TE) = 6000/93 ms, matrix = 128 × 128, 44 axial slices, 2.0 mm slice thickness, b values = 0 and 1000s/mm²) in 64 directions; (4) susceptibility-weighted imaging (SWI) (TR/TE = 29 / 20 ms, voxel size 0.5 × 0.5 × 0.2 mm, 128 slices, 2 mm thickness); (5) fluid attenuated inversion recovery (FLAIR) imaging (TR/TE / inversion time (TI) = 6000/396 /2200 ms, voxel size 0.5 × 0.5 × 1 mm, 160 slices, 1 mm thickness); and (6) perfusion imaging using Q2TIPS pulsed arterial spin labeling (PASL) sequence (TR / TE = 2500 / 11 ms, matrix = 64 × 64, flip angle = 90°).

L. Gao, et al. NeuroImage: Clinical 24 (2019) 102101
slice thickness = 6 mm). During the R-fMRI scan, subjects were asked to remain as still as possible, keep their eyes closed, not fall asleep, and avoid thinking systematically.

2.4. MRI data processing

2.4.1. Preprocessing of R-fMRI data

R-fMRI data were preprocessed using the Data Processing & Analysis for Brain Image (DPABI, rfmri.org/dpabi) (Yan et al., 2016), Statistical Parametric Mapping (SPM12, http://www.fil.ion.ucl.ac.uk), and MATALB 2013a (https://www.mathworks.com), following conventional methods as previously described (Li et al., 2017). Briefly, these included (1) removal of the first 10 functional volumes, (2) slice timing correction, (3) realignment and correction for small head movements, (4) coregistration between individual T1 anatomical and the realigned functional volumes, (5) normalization into Montreal Neurological Institute (MNI152) space through Diffeomorphic Anatomical Registration Through Exponentiated Lie Algebra ( DARTEL) (Ashburner, 2007), using an elderly sample representative template (60–70 years), (6) spatial smoothing with 6 mm full width half maximum (FWHM) Gaussian blur, (7) reduction of confounding factors via linear regression, including the signals from the white matter and cerebrospinal fluid, and linear and quadratic trends, (8) temporal filtering (0.01–0.1 Hz) of the time series, and finally (9) motion scrubbing (Power et al., 2012; Yan et al., 2013, 2016a) with a frame-wise displacement (FD) threshold of 0.5.

2.4.2. Quality assurance

To correct for head motion, we calculated the FD and took the following measures to reduce motion effects: (i) ensure that no subject was found with maximum displacement in one or more of the orthogonal directions > 0.5 mm or a maximum rotation > 0.5°; (ii) the voxel-specific FD (FDvox) described as Yan et al. (2013) was calculated for each subject, and considered as nuisance covariate in the statistical analyses. For better whole-brain coverage, we also made a 90% group-specific mask. Before calculating functional connectivity, we first generated an individual-specific functional brain mask that included only voxels (in MNI standard space) present in each participant. We then generated a group-specific mask that included all voxels present in at least 90 percent (90%) of participants to ensure common coverage. This was then further constrained by a 99% gray-matter probability mask.

2.4.3. Homotopic FC

Homotopic FC was computed in two manners: (1) voxel-wise mirrored homotopic connectivity (i.e., VMHC) as developed and provided by Zuo et al. (2010), by registering the resting-state functional volumes onto a symmetric anatomical template and then calculating the Pearson’s correlation coefficients (functional connectivity) of BOLD signals between the mirror-symmetric interhemispheric voxels, i.e., the individual VMHC map; (2) a less fine-grained ROI-wise functional homotopy, by registering the resting-state functional volumes to a recently validated homotopic brain parcellation based atlas (AICHA) (Joliot et al., 2015), which parcellates the brain into 384 parcels (192 homotopic pairs) based on clustering algorithms. The correlation coefficients were then z-score transformed.

2.4.4. Seed-based FC

Seed-based FC was computed based on the 16 significant clusters of between-group VMHC comparison (Table 2 & Fig 2). The seeds were 6-mm radius spheres centered at peak coordinates from the significant clusters: bilateral putamen (MNI coordinates: X = ± 24, Y = 9, Z = −9), bilateral supramarginal gyr (MNI coordinates: X = ± 63, Y = −33, Z = 27), bilateral planum temporale (MNI coordinates: X = ± 57, Y = −12, Z = 6), bilateral opercular (MNI coordinates: X = ± 39, Y = 6, Z = 9), bilateral lateral occipital (MNI coordinates: X = ± 42, Y = −60, Z = 51), bilateral postcentral gyr/PFC (MNI coordinates: X = ± 48, Y = −12, Z = 36), bilateral postcentral gyr (MNI coordinates: X = ± 6, Y = −36, Z = 57), and bilateral superior parietal lobules (MNI coordinates: X = ± 42, Y = −42, Z = 10). These additional seed-based functional connectivity analyses were performed using the DPABI toolbox with conventional correlation analysis.

2.4.5. Homotopic SC

Homotopic SC was defined as white matter microstructural connectivity between each homotopic pair, including mean fractional anisotropy (FA), fiber number, and fiber length, using the deterministic tractography algorithm (Descoteaux et al., 2009). Processing was performed by using the FMRIB Software Library (FSL, https://www.fmrib.ox.ac.uk/fsl), the Pipeline for Analyzing brainDifussion imAges (PANDA, https://www.nitrc.org/projects/panda) (Cui et al., 2013), and MATALB 2013a under a Linux-Ubuntu environment. The main steps included: (1) motion and eddy current artifacts correction; (2) brain extraction; (3) diffusion tensors calculation, in which the resulting FA volume of each subject was then registered to the FMRIB58_FA template using the FSL nonlinear registration tool FNIRT to obtain the warp field from native to standard space; (4) the nonlinear warp was initialized with an affine registration generated with the FSL linear registration tool FLIRT; (5) deterministic whole-brain streamline counts were generated with the fiber assignment by continuous tracking algorithm using the Diffusion Toolkit (trackvis.org/dtk) and TrackVis (www.trackvis.org); parameters included a 35° angle threshold, 20 seeds were placed in every voxel and a minimum streamline length of 20 mm was used. Network nodes were based on the 192 homotopic pairs (a total of 384 cortical and subcortical regions) comprising the AICHA atlas. The AICHA was mapped to an individual’s native space using the inverse of the warp field previously computed. Each individual’s network was represented with a symmetric 384 × 384 connectivity matrix. Each element of the connectivity matrix was populated with the number of streamlines between the corresponding pair of regions (streamline count), which served as a measure of inter-regional connection strength. The resultant average fiber FA, fiber number, and fiber length between each homotopic pair were imported into the SPSS 16.0 (SPSS Inc., Chicago, IL, USA) for further statistical analysis.

2.4.6. Brain perfusion and quantification of brain lesions

Brain perfusion was assessed with the PASL data using SPM12-based batch scripts ASLtbx (https://cfn.upenn.edu/~zewang/ASLtbx.php) including motion correction, temporal filtering, coregistration and spatial normalization, CBF image generation, and 6-mm spatial smoothing for subsequent group statistics.

We also calculated cerebral microhemorrhages and WMH by using

| Regions | MNI coordinates | Voxels | BA | T values |
|---------|-----------------|--------|----|---------|
| Putamen | x ± 24, Y = 9, Z = −9 | 27 | - | -6.892 |
| Supramarginal gyr | X = ± 63, Y = −33, Z = 27 | 40 | - | -8.203 |
| Parietal temp | X = ± 57, Y = −12, Z = 6 | 22 | - | -8.092 |
| Opercular cort | X = ± 39, Y = 6, Z = 9 | 13 | - | -7.345 |
| Posterior parietal cort | X = ± 42, Y = −60, Z = 51 | 16 | - | -6.505 |
| Postcentral gyr/PMC | X = ± 48, Y = −12, Z = 36 | 35 | 6/43 | -6.993 |
| Postcentral gyr | X = ± 6, Y = −36, Z = 57 | 12 | 5 | -6.868 |
| Superior parietal lobules | X = ± 42, Y = −42, Z = 10 | 40 | - | -6.489 |

Table 2 shows all local maxima. x, y, and z = MNI coordinates. Abbreviations: BA: Brodmann’s area; T-value: the voxel of maximal intensity in this cluster; MNI: Montreal Neurological Institute; x, y, z: coordinates of primary peak locations in the MNI space. The threshold was set at voxel p < 0.05, FDR corrected. These results were reported with BSPMVIEW (http://www.bobspunt.com/software/bspmview/).
the SWI, FLAIR, and T1-weighted images. After visual inspection, brain lesions were automatically identified to create lesion maps for each individual’s SWI and FLAIR images. The LST (www.applied-statistics.de/lst) for SPM12 were used to spatially normalize each individual’s SWI and FLAIR lesion maps to the MNI152 template.

2.4.7. Brain behavior relationship

To examine whether there were any associations between the changes of VMHC and behavioral assessments in patients with SACS, we conducted Pearson’s linear correlation analyses between behavioral assessments and mean VMHC values within each significant cluster. To explore the contribution of parenchymal lesion load to the VMHC, we also performed a correlation analysis between the VMHC and the WMH in the patients with SACS. All correlations were Bonferroni corrected at a significance level of \( p < 0.05 \).

2.4.8. Results verification and replication

We repeated the VMHC analysis using a range of Gaussian kernel (FWHM) values \((4, 6, 8, 10, \text{and} 12 \text{ mm})\) and found that our results were robust. To examine the extent to which the patients could be discriminated from the controls based on the inter-group differences of VMHC, we applied a linear support vector machine (SVM) classification to the VMHC maps using the Pattern Recognition for Neuroimaging Toolbox (PRoNTo; www.mlnl.cs.ucl.ac.uk/pronto). To assess model performance, we applied a leave one-out cross-validation scheme. Under this cross-validation strategy, the classifier’s accuracy was computed by leaving one subject out at a time and predicting this subject’s group label based on a training set that includes all remaining subjects. Permutation testing was used to estimate the null distribution and examine the statistical significance of the classification accuracy \((N \text{ permutations} = 10,000; \ p < 0.05)\).

2.5. Statistical analysis

For all clinical variables, inter-group statistics were carried out using SPSS 16.0 (SPSS Inc., Chicago, IL) with a significance level set at \( p < 0.05 \). For the imaging measures, independent two-sample t-tests were used for functional connectivity and CBF, and nonparametric analyses were used for voxel-wise WMH and SWI analyses. These imaging measures were corrected using either voxel-wise false discovery rates (FDR) correction \((p < 0.05)\) or cluster-level family-wise error (FWE) correction \((p < 0.001 \text{ and cluster} \ p < 0.05)\) and controlled for age, gender, and head motion parameters (mean frame displacement).

2.6. Distribution of functions through the resulting t statistic VMHC

Finally, to examine brain functions subserving the significant clusters that showed between-group differences in the VMHC, we searched for research terms in the NeuroSynth database (www.neurosynth.org) (Yarkoni et al., 2011). A total of 32 terms were selected, and the corresponding brain activation regions were binarized into masks. The terms included “somatosensory, pain, motor, movements, noxious, motor imaginary, execution, action, auditory, speech production, speech, dorsal premotor, hands, listening, emotional, visual, working memory, stroke, language, default mode, semantic, reading, memory, autobiographic memory, dementia, encoding, episode memory, eye, face, navigation, tapping, and visual word”, etc. The statistical T map of VMHC was then binned into 5 percentile increments with its T values organized from most negative to most positive. These 20 gradient divisions form the heat map matrix of the topographies and the Z values of the topic terms. These terms were then ranked based on their statistical association with gradient T values for visualization. Heatmap colors encode the strength of correlation between each equally divided part of the VMHC T map and the Z statistic associated with the topic terms. The overall data processing flow is shown in Fig. 1.
Due to our relatively small sample size, it was important to assess the reliability of the results. To this end, we used cross-validation and SVM-based machine learning for classification. Balanced accuracy: $95.29\%$ ($p_{BA} = 0.02$); total accuracy $95.53\%$, class accuracy (CA) $94.74\%$, $95.83\%$ ($p_{CA} = 0.02$), class predictive values were $94.74\%$ and $95.83\%$ (Fig. 7).

3.7. Qualification of brain lesions and perfusion

For the PASL-derived CBF, patients with SACS showed significant hypoperfusion in the left pars opercularis, left orbitofrontal, bilateral thalami, right dorsal anterior cingulate, and left middle occipital cortices (Table 4). For the voxel-wise comparison of SWI, we identified several brain clusters with hypointensity in patients, including frontal, parietal, temporal, and periventricular white matter (Table 5 & Fig. 8).
For WMH measured by T2-FLAIR, we found that the distribution and number of WMH in SACS patients were significantly higher than the controls, especially in the periventricular areas. Areas of overlap included the left inferior frontal gyrus (Brodmann area 22), the left inferior parietal lobule (Brodmann areas BA40/2), the left precentral gyrus/postcentral gyrus, the left putamen, the left medial frontal gyrus (Brodmann area 6) and the left middle temporal gyrus.

Fig. 3. ROI-based homotopic connectivity. (A). Seed ROI selection from the AICHA atlas. Each hemisphere includes 192 homotopic regions. (B). Functional connectome and homotopic connectivity. First, functional connectivity matrix seeding from the AICHA atlas were thresholded into individual homotopic matrices. Individual homotopic matrices were then input into the second statistic model. (C). Between-group comparison of the homotopic connectivity. Matrix and surface-view of homotopic connectivity between each homotopic ROI. These results were corrected at a connection level, $p < 0.05$ with FDR correction.

For WMH measured by T2-FLAIR, we found that the distribution and number of WMH in SACS patients were significantly higher than the controls, especially in the periventricular areas. Areas of overlap included the left inferior frontal gyrus (Brodmann area 22), the left inferior parietal lobule (Brodmann areas BA40/2), the left precentral gyrus/postcentral gyrus, the left putamen, the left medial frontal gyrus (Brodmann area 6) and the left middle temporal gyrus. To aid in visualization, the three suprathreshold statistical parameter maps are

Fig. 4. Seed-based functional connectivity. (A). Schematic display of the selection of seed regions. Each sphere represents a seed region, and the lines connecting them represent homotopic functional connectivity with significant inter-group differences. (B). The topography of the seven average large-scale resting brain networks provided by Yeo et al. (2011). (C). Between-group comparison of seed-based functional connectivity with seeds selection from A. Purple circle represents a seed region. These results were FDR corrected at a voxel-level, $p < 0.05$. Here, we only list the seed-based FC with significant inter-group differences and do not include the seeds with only homotopic FC differences. Theoretically, each seed should yield at least one significant cluster with its homotopic region.
3.8. Association of VMHC to behavioral and lesion parameters

We first quantified the relationship between the VMHC and behavioral measurements. VMHC between the bilateral supramarginal gyri (SACS patients, \( n = 19 \), \( r = 0.591, p = 0.0078 \); all subjects, \( r = 0.538, p = 0.0002 \)) and the bilateral superior temporal gyri (SACS patients, \( n = 19, r = 0.664, p = 0.0019; \) all subjects, \( r = 0.651, p < 0.001 \) were both positively correlated with delayed recall (Fig. 9). We next assessed the relationships between WMH loads and the cognitive variables, as well as WMH loads and VMHC. We did not find any significant correlation between the WMH loads (including WMH number, absolute WMH size, and relative WMH volume with TIV) and the behavioral variables. However, the corrected WMH loads (i.e., the ratio of WMH size and total intracranial volume) was significantly negatively correlated with the VMHC between the bilateral angular (SACS patients, \( n = 19, r = -0.675, p = 0.002; \) all subjects, \( r = -0.644, p < 0.001 \)), insula (SACS patients, \( n = 19, r = -0.715, p = 0.001; \) all subjects, \( r = -0.610, p < 0.001 \)), supramarginal gyrus (SACS patients, \( n = 19, r = -0.776, p < 0.001; \) all subjects, \( r = -0.678, p < 0.001 \)), and bilateral putamen (SACS patients, \( n = 19, r = -0.742, p < 0.001; \) all subjects, \( r = -0.632, p < 0.001 \)) (Fig. 9).

3.9. Distribution of functions through the resulting T statistic VMHC

Finally, we examined whether the functional processes ascribed to the resulting T statistic VMHC specifically involved sensory inputs, perception, and concrete cognitive functions. We performed meta-analytic decoding using the Neurosynth database to correlate the research terms and the T statistical maps. Top terms exhibiting the strongest associations included basic functions, such as "somatosensory, pain, motor, movements, noxious, motor imaginary, execution, action, auditory, speech production." (Fig. 10).

4. Discussion

4.1. Summary of findings

Our investigation demonstrates robust homotopic FC reductions in patients with SACS in brain regions dominated by the middle cerebral arteries, especially in the Perisylvian fissure. These involved regions span across the large-scale intrinsic networks including somatomotor, salience, dorsal attention, and orbitofrontal-limbic. Results showed that the significantly reduced homotopic FC can be partially explained by white matter hyperintensity size. Further association analyses suggested that decreased homotopic FC in these brain regions is most closely associated with delayed memory recall, sensorimotor processing, and other simple cognitive functions. These results suggest that SACS may preferentially affect lower-order cognitive processes, while...
the higher-order systems, including regions in the DMN that subserve abstract thought, appear to be least impacted. However, decreased homotopic FC within the DMN may be a precursor to vascular dementia. Thus, measurements of homotopic FC may provide clinical utility by noninvasively tracking the progression of downstream brain damage following SACS.

Prominent homotopic FC is a significant phenomenology in human (Stark et al., 2008; Zuo et al., 2010), non-human primates (O’Reilly et al., 2013), and some other known mammals’ brains (Layden et al., 2019). Reports on agenesis (Tyszka et al., 2011), absence (Split-brain) (Johnston et al., 2008), and resection (callosotomy) (Pizoli et al., 2011; Roland et al., 2017) of the corpus callosum suggest that this prominent homotopy in the association (higher-order) cortices is heavily dependent on the integrity of the corpus callosum, while homotopy in the primary cortices may be supported via multiple extracallosal pathways.

Homotopic FC exhibits increasing primary cortical integration (i.e., increasing homotopic connectivity) and higher-order segregation (i.e., decreasing homotopic connectivity) across lifespan development (Zuo et al., 2010), and has been implicated in several neuropsychiatric and neurological conditions (Carter et al., 2010; Hoptman et al., 2012; Kelly et al., 2011; Li et al., 2017). Our current findings show significant homotopic reductions in patients with SACS, specifically, in the topographies of the Perisylvian fissure, temporal-occipital-parietal conjunction, lateral parietal, insular, and basal ganglia. These regions comprise parts of the somatomotor, dorsal attention, frontoparietal, and salience networks, whose functions putatively subserve somatosensory, memory, attention, executive, fine motor, and visceral processes (also see the metanalytic decoding). These findings are generally consistent with earlier reports on both human and rodents (Cheng et al., 2012; Lin et al., 2014; Panczykowski et al., 2014) and our recent studies (Wang et al., 2017).

Our results of the VMHC were validated through three additional analyses (preprocessing scheme, independent controls, and SVM-based cross validation, Figs. 2, 3, 6, and 7), which demonstrated the robustness of our results, and together suggested that VMHC reduction is an

### Table 4

| Regions                        | MNI coordinates | Voxels | BA | T values |
|--------------------------------|-----------------|--------|----|----------|
| Left middle occipital gyrus    | −26 −74         | 24     | 196| −3.75    |
| Left pars opercularis          | −52 12 −2       | 1066   | 48 | −3.18    |
| Left orbitofrontal cortex      | −44 30 −8       | 1066   | 47 | −2.98    |
| Thalamus                       | 34 −48 4        | 207    | −3.00|        |
| Right dACC                     | 14 44 −4        | 140    | 32 | −2.99    |

shows all peak coordinates. Abbreviations: BA: Brodmann’s area; T-value: the voxel of maximal intensity in this cluster; MNI: Montreal Neurological Institute; x, y, z: coordinates of primary peak locations in the MNI space; dACC, dorsal anterior cingulate cortex. The threshold was set at a corrected voxel p < 0.001, cluster level p < 0.05, FWE cluster-level corrected. These results were reported with BSPMVIEW (http://www.bobspunt.com/software/bspmview/).
Table 5

| Regions                  | MNI coordinates | Voxels | BA | T values |
|--------------------------|-----------------|--------|----|----------|
| Left Caudate             | −15 −4 24       | 21,952 | −5.10 |
| Left Hippocampus         | −31 −13 −10     | 21,952 | 20  | −4.426   |
| Left SMA                 | −4 −9 58 330    | 6      | −4.37 |
| Left Thalamus            | −4 −24 11 21,952| −4.23  |
| Right Superior Temporal  | 44 −8 −8        | 604    | 48  | −4.11    |
| Left Inferior Parietal   | −51 −39 58 1042 | 2/40   | −3.961|
| Right Supramarginal gyrus| 64 −24 31 284   | 2      | 3.72  |
| Right Parietal Operculum | 47 −25 21 268   | 48     | −3.67 |
| Left Supramarginal gyrus | −45 −33 25 1141 | 48     | 3.63  |
| Right Supramarginal gyrus| 16 −25 35 560   | 48     | −3.65 |
| Right Fusiform           | 38 −48 −21 307  | 37     | 3.20  |

Between-group comparison on SWI (SACS v.s. Controls). Table 5 shows all peak coordinates. Abbreviations: BA: Brodmann's area; T-value: the voxel of maximal intensity in this cluster; MNI: Montreal Neurological Institute; x, y, z: coordinates of primary peak locations in the MNI space; dACC, dorsal anterior cingulate cortex. The threshold was set at a corrected voxel p < 0.001, cluster-level p < 0.05, FWE cluster-level corrected. These results were reported with BSPMVIEW (http://www.bobspunt.com/software/bspmview/).

Fig. 8. Qualification of brain lesions and hypoperfusion. (A), Between-group comparison on CBF. The blue tone indicates significant brain regions with hypoperfusion in the patients with SACS. No significant hyperperfusion was found. (B), Between-group comparison in SWI signal intensities. The blue tone indicates significant low-signal foci in the patients with SACS. (C), Between-group comparison on T2-FLAIR based WMH. The hot tone indicates significant brain regions with increased WMH in the patients with SACS. These results (A-C) were corrected by FWE cluster-level p < 0.05 and voxel-level p < 0.001. (D), Cumulative WMH distribution for the patients with SACS. From left to right (cold to hot hues), the color indicates number of subjects (up to 9) with WMH in the same brain region. (E), Cumulative WMH distribution for the healthy controls (HC). From left to right (cold to hot hues), the color indicates number of HC subjects (up to 4) with WMH in the same brain region. (F), Superimposed map of brain lesions and hypoperfusion. Suprathreshold between-group differences of PASL-based hyperperfusion (Red label), significant high load of WMH measured by FLAIR (Green label), and hypointensities measured by SWI (Blue label) in the patients with SACS. Their main overlapping areas include the left inferior frontal gyrus (BA 22), the left inferior parietal lobule (BA40/2), the left precuneus/postcentral gyrus, the left putamen, the left medial frontal gyrus (BA6) and the left middle temporal gyrus. Abbreviations: L, left; R, right; BA, Brodmann area.

important feature of SACS patients. As noted in the primary analysis (Fig. 2), the main results are largely consistent across smoothing kernels (Fig. 6). However, as the smoothing kernel increased, or possibly due to a threshold effect, new brain regions with significant between-group difference emerged, particularly those located in the medial portions of the DMN (i.e., the posterior cingulate and ventromedial prefrontal cortices) (Raichle, 2015). These results suggest the possibility that an early change in SACS initiates a marginal change in DMN, which might subsequently involve the core regions within the DMN, speculation that coincides with vascular dementia and patterns of disease propagation (Dichgans and Leys, 2017). In addition, we found that in the classification of VMHC differential brain regions, one elderly control fell into the “patient” class, which may indicate undetected hypoperfusion in subpopulations of healthy older adults.

Our results on homotopic connectivity can be considered in the context of ischemia-hypoxia and hemodynamic changes. Reduced CBF and impaired cerebrovascular reactivity have been consistently reported in asymptomatic higher-grade ICA stenosis patients (Chen et al., 2017; Hartkamp et al., 2017; Hu et al., 2018), particularly in the anterior circulation of the affected hemisphere (Silvestrini et al., 2000). Evidence suggests that similar homotopic reductions, as shown in the current study, can be seen in acute cerebral hypoxia (experimental hypopcapnia) (Marshall et al., 2015). Surprisingly, our examination of PASL-based CBF did not show prominent reductions in the brain territories dominated by the MCA in the patients with SACS. We speculate that this might be due to blood transit delays in severe ICA stenosis with a single labeling/inversion time. This consideration is both supported by previous reports (Bokkers et al., 2009; Hendriks et al., 2004), and confirmed by our unpublished analysis of lag structure on resting-state time series, with average time delays of ~ 2 s. This asymmetrical change results in neural propagation and phase delay (mismatch), which corresponds to the dramatically decreased functional homotopy measured by resting-state BOLD time series.
significations in VMHC in the Perisylvian
figurations, motor actions, and physical
low-level of the functional hierarchy and anchored to sensory percep-

Fig. 9. Association with behavior. VMHC between the bilateral superior parietal lobules (SACS patients, n = 19, r = 0.591, p = 0.0078; all subjects, r = 0.538, p = 0.0002) and the bilateral superior temporal gyri (SACS patients, n = 19, r = 0.664, p = 0.0019; all subjects, r = 0.651, p < 0.001) were both positively correlated with delayed recall. The VMHC between the bilateral angular gyri (SACS patients, n = 19, r = -0.675, p = 0.002; all subjects, r = -0.644, p < 0.001), insula (SACS patients, n = 19, r = -0.715, p = 0.001; all subjects, r = -0.610, p < 0.001), supramarginal gyri (SACS patients, n = 19, r = -0.742, p < 0.001; all subjects, r = -0.672, p < 0.001) were significantly negative correlated with the corrected WMH loads (i.e., the ratio of WMH size and total intracranial volume).

Organizationally, we note that SACS patients show prominent reduc-
tions in VMHC in the Perisylvian fissure, typically considered at the
low-level of the functional hierarchy and anchored to sensory percep-
tions, motor actions, and physical fitness (Huntenburg et al., 2018; Margulies et al., 2016). Further, the angular, supramarginal, and superior temporal gyri—considered at a higher-level in the functional hierarchy—also showed significant reductions of VMHC, and were significantly correlated with memory, processing speed, and attention deteriorations. The mammalian cerebral cortex is a functional hierarchy with spatiotemporal organization. At the low-level of this hierarchy sit the primary sensorimotor cortices, with transmodal association cortices considered higher up and the DMN located near the apex (Huntenburg et al., 2018; Margulies et al., 2016). Movement from lower to higher in this functional hierarchy generally corresponds with concrete to abstract cognitive processing, respectively. From this perspective, patients with SACS primarily show decreased FC in low-level areas of this functional hierarchy, with corresponding limited impairment in the abstract cognitive functioning domain. Our observations and speculations are also consistent with previous reports (van Meer et al., 2010). For example, similar homotopic reductions can be seen in acute cerebral hypoxia (experimental hypercapnia) (Marshall et al., 2015), or stroke (van Meer et al., 2010), following the hierarchy from basic somatomotor or concrete cognitive functions (Fig. 10), including the research topics of somatosensory, pain, motor, movements, noxious, motor imaginary, execution, action, etc. These results reinforce that SACS preferentially affects the lower hierarchy of the brain’s functional organization, i.e., regions subserving somatomotor and simple cognitive functions, while areas further up on the hierarchy such as the DMN are least impacted.

Pathologically, SACS represents a type of silent cerebrovascular disease, which includes silent lacunar infarct, cerebral microbleeds, and WMH (van Veluw et al., 2017). To explain the underlying causes of the VMHC reductions, we further explored the relationship between the VMHC and white matter microstructural integrity, WMH load, and cerebral microbleeds. Using deterministic tractography, we found that only five of 192 homotopic regions showed significantly lower SC in the SACS patients, in the brain regions of superior medial frontal, hippocampus, paracentral, and superior parietal cortex. These regions are consistent with regions showing significant VMHC reductions. Moreover, among the three DTI indices, the difference in mean FA was the most significant, followed by fiber length, with fiber number being the least significant. These results suggest that mean FA may be a sensitive indicator that is more likely to reflect white matter damage in patients with SACS (Avelar et al., 2016; Baradaran et al., 2016). Despite the significant reduction in these significant homotopic structural connections, we believe that this does not account for the full scope of homotopic FC reductions observed, since the number of the homotopic pairs is disproportionately matched. We also identified hypointensities of SWI in the frontal, parietal, and temporal gray matter, and periventricular white matter regions in patients with SACS (Table 5 & Fig. 8). The distribution and number of WMH in SACS patients were
significantly higher than the matched healthy aging controls, especially in the periventricular areas. Furthermore, the VMHC between the bilateral angular, insula, and supramarginal gyri, and bilateral putamen were significantly negatively correlated with the corrected WMH loads (i.e., the ratio of WMH size and total intracranial volume) (Fig. 9). This suggests that the brain lesion load of SACS patients is higher than that of healthy controls, and may at least partially explain the reduction of functional connectivity. However, these relationships should be replicated and further investigated by future studies.

Regarding the application of VMHC in brain diseases, the reliability and validity of VMHC, and particularly the use of SACS as a potential biomarker, warrants additional investigations. A recent study has indicated that in the application of VMHC to gender differences, the intraclass correlation coefficient is roughly 0.44 (Chen et al., 2018), suggesting that the test-retest reliability and generalizability still needs to be explored in depth. However, in this study and previous studies, in the context of multiple white matter lesions, VMHC is a better indicator of heterogeneous brain disease. Recently, an important focus in the neuroimaging community is reliability and validity (Xing and Zuo, 2018; Zuo et al., 2019b; Zuo and Xing, 2014, 2019). In particular, in the last decade, the field has witnessed the escalation of multi-center, large-sample inter-individual data for exploring individual differences and obtaining general conclusions. Yet, still, the sample length of the intra-individual data is still relatively small, only a few minutes, showing poor reliability and validity. Therefore, future research should extend the scan times of intra-individual data and perform repeated-measurements of resting scans (Zuo et al., 2019b).

We must address the limitations of this study. Since this study was not primarily designed for the assessment of sensorimotor functions, we did not assess any brain-behavior relationships in the motor ability domain. Second, because a single-inflow-time pASL sequence was used, we may not be able to calculate individual CBF maps and mean transit time accurately. Third, the threshold effect of silent brain lesions and their relationship with functional connectivity are rather complicated, which is one of the active research topics in this field and one of the future research directions. Finally, due to the number of statistical items in this study, we used two of the most popular correction methods, voxel-wise FDR correction with \( p < 0.05 \), and voxel-wise uncorrected \( p < 0.001 \) with cluster-level FWE correction with \( p < 0.05 \). Although these are the most current recommended methods, the disadvantage is diminished localizing ability and sensitivity.

**Conclusion**

Homotopic FC alterations, specifically in the somatomotor, salience, dorsal attention, and executive systems are characteristic changes in patients with SACS. These results suggest that the SACS preferentially affects the lower tier of the functional hierarchy, while cognitive systems further up on the hierarchy (i.e., the default mode network) are least impacted by SACS, but may be a hallmark precursor to vascular dementia. Thus, measurements of homotopic FC may help to non-invasively track the progression of downstream brain damage following asymptomatic carotid stenosis.

**Declaration of Competing Interest**

All authors disclosed no relevant relationships.
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