Improved gray matter surface based spatial statistics in neuroimaging studies

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ABSTRACT

Neuroimaging often involves acquiring high-resolution anatomical images along with other low-resolution image modalities, like diffusion and functional magnetic resonance imaging. Performing gray matter statistics with low-resolution image modalities is a challenge due to registration artifacts and partial volume effects. Gray matter surface based spatial statistics (GS-BSS) has been shown to provide higher sensitivity using gray matter surfaces compared to that of skeletonization approach of gray matter based spatial statistics which is adapted from tract based spatial statistics in diffusion studies. In this study, we improve upon GS-BSS incorporating neurite orientation dispersion and density imaging (NODDI) based search (denoted N-GSBSS) by 1) enhancing metrics mapping from native space, 2) incorporating maximum orientation dispersion index (ODI) search along surface normal, and 3) proposing applicability to other modalities, such as functional MRI (fMRI). We evaluated the performance of N-GSBSS against three baseline pipelines: volume-based registration, FreeSurfer’s surface registration and ciftify pipeline for fMRI and simulation studies. First, qualitative mean ODI results are shown for N-GSBSS with and without NODDI based search in comparison with ciftify pipeline. Second, we conducted one-sample t-tests on working memory activations in fMRI to show that the proposed method can aid in the analysis of low resolution fMRI data. Finally we performed a sensitivity test in a simulation study by varying percentage change of intensity values within a region of interest in gray matter probability maps. N-GSBSS showed higher sensitivity in the simulation test compared to the other methods capturing difference between the groups starting at 10% change in the intensity values. The computational time of N-GSBSS is 68 times faster than that of traditional surface-based or 86 times faster than that of ciftify pipeline analysis.

1. Introduction

Gray matter (GM) in the cerebral cortex is key to many sensory, cognitive, and motor functions of the brain. Detecting cortical alterations with neuropathologic conditions could provide potential biomarkers to facilitate early diagnosis and assessment of disease severity. In recent years, the development of neuroimaging techniques, such as high-resolution magnetic resonance imaging (MRI), functional magnetic resonance imaging (fMRI), diffusion weighted magnetic resonance imaging (dMRI), positron emission tomography (PET) or single photon emission computed tomography (SPECT), have promoted the identification of structural and functional characteristics of the developing brain and underlying mental disorders [1–7]. An increasing number of studies have shown structural and functional gray matter changes in clinical applications - e.g., amyotrophic lateral sclerosis [8], schizophrenia and bipolar disorder [9,10], age related effects [11], attention deficit hyperactivity disorder [12], and Alzheimer’s disease [13]. While T1 images can be acquired at high resolution (e.g., 1 mm isotropic), clinical imaging in other modalities (such as dMRI and fMRI) are constrained by imaging and physiological factors leading to lower resolution (2–3 mm isotropic). As the cortex is about 1.6–4.5 mm thick [14–16] within the gray matter tissue region between white and...
Spatial coherence in the volumetric images (see Fig. 1-a). In particular, rigid volumetric registration since it is quite challenging to incorporate individual cortical anatomy may not be sufficient. A majority of studies focus on volume- or surface-based analysis on a particular modality [11,37,38]. Few studies [32,38-40] have incorporated multi-modalities into a single integrated pipeline of surface-based analyses. The desire to better understand structural-functional relationships drives the need for robust analysis frameworks. The Human Connectome Project (HCP) minimal preprocessing pipeline [38] is one such approach that integrates multimodal data for cross subject analysis. It is built upon the FreeSurfer software tool (https://surfer.nmr.mgh.harvard.edu) for surface generation and alignment to standard space in addition to defining Connectivity Informatics Technology Initiative (CIFTI) format and grayordinate system that incorporates cortical and subcortical information. In a recent study, multimodal surface matching (MSM) [41] registration is incorporated into a pipeline that uses multimodal registration features containing myelin maps [59], resting-state networks (RSNs) and visuotopic features to drive alignment to a group template. In the HCP approach [38], the data acquisition protocol is customized and often requires newly developed preprocessing methods unlike conventional data acquisition schemes.

There is huge amount of clinical data that is already acquired from healthy individuals and also in different clinical populations that is not acquired as per the HCP proposed standards. Having tools that could provide HCP-style analyses to leverage the existing data to the extent possible will be beneficial for clinical research. The ciftify pipeline [42] bridges the gap for making HCP-style analysis applicable to non-HCP (i.e., legacy) datasets by adapting the key modules from HCP pipeline into existing structural workflows. For functional/diffusion MRI data, the alignment with anatomical T1 plays an important role to map volume data onto the surface. Thus, preprocessing choices need to be made to maximize the data quality given its limitations in legacy datasets. The ciftify pipeline takes the preprocessed data from other modalities and converts it into needed grayordinate format for further analysis.
In this paper, we propose N-GSBSS for carrying out localized statistical testing of neuroimaging data across multiple modalities in GM. Unlike the skeletonization approach in GBSS, cortical surfaces reconstructed from high resolution T1 images are employed to facilitate cross-subject analysis. This method provides a bridge between volume and surface registration approaches to achieve cross-subject correspondence of low resolution image data. This method is an extension of our previous work, GS-BSS [24]. While conceptually similar, improvements are made in registration methodology that allow mapping of the metrics of interest in subject space. The key idea in this method is to incorporate normal search from the cortical surface to get metrics from highly probable GM voxels using the orientation dispersion index (ODI) from the NODDI model. ODI is higher in GM compared to that of WM [43], thus searching for higher ODI could help to locate underlying highly probable GM. Toward this end, we show an application to statistical analysis of fMRI data. To test the sensitivity of the approach, a simulation study is performed by varying region of interest (ROI) size and percentage change of intensity values within the ROI. It is presented as a full end-to-end pipeline to perform such spatial statistics in group analysis. We evaluated the performance of N-GSBSS against three baseline pipelines: volume-based registration (VBR), FreeSurfer’s surface registration (SBR) and ciftify pipeline for fMRI and simulation studies. The source code for N-GSBSS is made available at https://github.com/MASILab/N-GSBSS/. The computational time of N-GSBSS is 68 times faster than that of traditional SBR or 86 times faster than the ciftify pipeline method [42].

2. Methods

2.1. Background

GS-BSS method was proposed to perform voxel-based statistical analysis of diffusion microstructure features acquired at low resolution on GM surfaces using high-resolution T1 images. Structural images are segmented and normalized to MNI template space using diffeomorphic anatomical registration using exponentiated lie algebra (DARTEL) method [44]. Diffusion metrics of interest are co-registered to structural T1 and transformed to MNI template space using forward deformation. GM surfaces are deformed to MNI template space using inverse deformation obtained from the registration step. Correspondence between cortical surfaces is obtained with diffeomorphic spectral matching DSM [45] and the mapping is applied to the deformed diffusion microstructure data in MNI template space to project onto the target surface for group analysis. GS-BSS is shown to yield better performance compared to that of VBM or the skeletonization approach of GBSS, which is based on alignment invariant skeleton projection. However, there are some methodological limitations that could impact the sensitivity of such analysis. First, the possibility of having any misalignment between diffusion microstructure and structural images after co-registration, could impact the sensitivity of the analysis to be performed on highly probable GM region. Second, the diffusion metrics of interest are projected onto the GM cortical surface in MNI template space that could allow the prospect of including distortions caused in the data from the volume registration step. Finally while the GM surfaces are used for achieving cortical correspondence, all the data is mapped back into voxel-space before performing statistical analysis.

In this paper, the goal is to improve spatial statistics in GM by projecting all the metrics of interest from each modality onto a single target cortical surface and carry out vertex based statistical analysis. Current work addressed the limitations of GS-BSS and provided improvement in the following areas,

- To overcome possible alignment issues from co-registration step and improve intra-subject correspondence, cortical search is proposed that can further improve the sensitivity of the method.
- To minimize distortions and keep the data as close to the raw images that are acquired as possible, metrics of interest are mapped onto the cortical surface in subject space unlike the GS-BSS method where the metrics of interest are mapped from the volume image in MNI space onto the deformed cortical surface in MNI template space.

- To perform spatial statistics on vertices, unlike the voxel based spatial statistics that is performed in GS-BSS.
- To show applicability of the method in additional modalities like fMRI.

2.2. Subjects and neuroimaging data acquisition

Neuroimaging data were collected on 30 healthy subjects (average age = 31.94 (male, n = 18)/35.83 (female, n = 12)) who participated in an on-going study of brain connectivity in neuropsychiatric disorders. The Vanderbilt University Institutional Review Board approved the study and all participants provided written informed consent prior to enrolling in the study. Neuroimaging data were acquired on a 3 T scanner (Achieva, Philips Medical Systems, Best, The Netherlands) equipped with a 32-channel head coil located at the Vanderbilt University Institute of Imaging Sciences. The following neuroimaging data were acquired on each subject: 1) a T1-weighted 3D-TFE anatomical scan (1 mm isotropic resolution, TE = 2 ms, TR = 8.95 ms and TI = 643 ms), 2) up to 6 functional EPI scans (3 mm resolution during which subjects completed an event related spatial working memory task (described below), and 3) a diffusion-weighted imaging scan protocol (2.5 mm isotropic resolution, with a matrix of 96 × 96, 50 slices, TR = 2.65 s, TE = 101 ms, Gmax = 37.5 mT/m) that included two diffusion shells with b-values of 1000 s/mm² (24 directions) and 2000 s/mm² (60 directions). Two subjects are excluded from the diffusion processing due to motion-related quality issues in diffusion MRI acquisition. HARDI from one subject is marked unusable due to zipper artifact in B0. Second subject is excluded based on quality checking measures due to subject movement (15 mm movement). Cardiac and respiratory gating were not used.

2.3. Preprocessing

2.3.1. T1 anatomical data processing

Each structural scan was segmented into GM, WM, and cerebrospinal fluid (CSF) tissue classes using the VBM8 toolbox (http://dbm.neuro.uni-jena.de/vbm/) from SPM12 (http://www.fil.ion.ucl.ac.uk/spm/). Additionally, each voxel of the images was automatically labeled using multi-atlas segmentation [46] according to the BrainCOLOR protocol [47] into 132 brain regions and 1 background that was used as a preprocessing step for MacROUIE. The white, central and pial cortical surfaces were reconstructed by MacROUIE [48] using the topology-preserving geometric deformable surface model. The central surfaces were used in further surface-based processing including registration and mapping volume data onto the surfaces.

2.3.2. Diffusion data processing

Diffusion-weighted images (DWI) were preprocessed using EDDY [49] tool from FMRIB Software Library FSL [50] for eddy current correction and subject motion correction. The registration matrix of each DWI was used to measure patient movement, and the gradient table was rotated accordingly. For diffusion data processing, the data from 2 shells were combined into a single DWI file and corresponding b-values and b-vectors were concatenated accordingly. A scheme file was generated using the fsls2scheme command from Camino (http://camino.cs.ucl.ac.uk). A brain mask was created using the FSL brain extraction tool [51].

For NODDI processing, the DWI file, scheme file and mask (generated as described above) were passed to the AMICO package (https://github.com/daducci/AMICO/), which is a fast implementation of NODDI [43] with linear approximation. Single transformation was
derived using b0 image to co-register to structural T1-weighted scan using spatial normalization from SPM12 with 12-parameter affine registration. Corresponding transformation is applied to NOODI-derived maps of intracellular volume fraction, isotropic volume fraction ($V_{iso}$), and orientation dispersion index (ODI). These ODI and $V_{iso}$ maps from multiple subjects were used in further analysis and validation of N-GSBSS.

### 2.3.3. Working memory fMRI processing

During the functional EPI scans, subjects completed a slow event-related spatial working memory task. Briefly, on each trial, three filled circles were presented sequentially, one at a time, during a 3-s encoding phase. The encoding phase was followed by a 16 s delay period during which a fixation dot was shown. Following the delay period, a probe (open circle) was presented for 1 s and subjects had to indicate with a button press whether or not the probe matched one of the previously encoded locations. Each trial was followed by a 14 s inter-trial interval. Subjects complete 30 working memory trials and 18 control trials. The working memory and control trials were identical, except for the fact that subjects were asked not to memorize the locations during the cue period of the control trials and pressed both the yes and no button during the probe period. Different colored circles, red and gray, were used to alert subjects to working memory and control trials, respectively. Preprocessing and generation of first-level, subject-specific statistical parametric maps were performed using spatial normalization in SPM12 [52]. Preprocessing included slice timing and motion correction, and co-registration of each subject’s functional EPI scans to their anatomical T1-weighted scans. Subject-specific, voxel-wise maps showing relative difference in the BOLD response between working memory and un-modelled baseline for cue, maintenance, and probe conditions were generated by modeling each subject’s time series data. Note, the contrast maps for cue, maintenance, and probe conditions were kept in the individual subject-specific space co-registered to T1 prior to being entered into the N-GSBSS pipeline described below.

### 2.4. N-GSBSS pipeline

The steps involved in carrying out the spatial statistics starting from the preprocessed multi-modal data to transferring all the metrics of interest onto a single target surface are illustrated in this section. The data from the co-registered volume images is projected onto the GM cortical surface using enclosing voxel approach. Alignment issues after co-registration would introduce partial volume effects or outliers by fetching data from the voxels that may not belong to highly probable GM. In order to overcome this limitation, cortical search is implemented using ODI measure as it has been shown to be higher in GM compared to that of WM [43].

#### 2.4.1. Cortical search using NOODI maps

Diffusion microstructure indices from NOODI including ODI and $V_{iso}$ are used in the cortical search. First ODI is masked with $V_{iso}$ to exclude any voxels with isotropic volume fraction of $> 0.5$ indicating CSF regions. The surface normal is calculated at each vertex on the central surface. As the T1 was acquired at 1 mm resolution and the cortical thickness is $< 5$ mm thick, we search the maximum ODI at each vertex along positive and negative normal directions (2 mm at maximum range with an interval of 1 mm). We create a search map by collecting these enclosing voxels that the normal directions point out. The metrics of interest in other modalities are finally transferred onto the central surface via the search map. Fig. 2(a) illustrates this approach and corresponding histogram of masked ODI is shown in Fig. 2(b) before and after search.

#### 2.4.2. Cortical correspondence on the target surface

Cortical surfaces are highly variable, so roughly similar surfaces would be useful for surface registration. As preprocessing volume registration can provide reasonably well-aligned surfaces, structural T1 is non-linearly registered with MNI template using ANTs’s SyN registration method (52). Corresponding inverse deformation is applied to the surface as the first step. The vertex coordinates of the surface are converted to RAS format before applying “antsApplyTransformsToPoints” from ANTs toolbox. The deformed coordinates are converted back into original format thus transforming the surface from subject space to MNI space (#2 from Fig. 3). However, as shown in Fig. 1(a), the cortical anatomy is not yet well aligned across the subjects after volume deformation. Then, we refine/update the correspondence using surface registration step [45] in the same way as (24), which is expected to establish better correspondence. It provides mapping information of the cortical surface from each subject onto the target surface (#3 from Fig. 3) on which spatial statistics can be performed.

#### 2.4.3. Project metrics of interest on target surface

As cortical anatomical properties such as cortical thickness were derived from the surface, they were already assigned to each vertex. These properties were then projected onto the target surface via the established shape correspondence from step 3. Images from different modalities are co-registered to T1 anatomical images before proceeding with further analysis as shown in step 4. Cortical ODI search is performed by taking in ODI and $V_{iso}$ measures from the NOODI model to get the corresponding map of highly probable GM vertices for co-registered images (step 5 in Fig. 3). Step 6 illustrates the first level analysis carried out on each modality to derive metrics of interest. In the volume images, the metrics of interest were mapped onto the individual GM surface (step 7 in Fig. 3) from the voxel that encloses the corresponding vertex coordinate obtained from the cortical ODI search (step 5 in Fig. 3). Both dMRI based NOODI metrics and fMRI based working memory contrast maps were projected via the vertex coordinates and the mapped properties were then transferred onto a common target surface (Step 8 in Fig. 3). Spatial statistics across the subjects are performed on the target surface by applying 2 mm smoothing kernel for cross subject analysis. We adapted the Gaussian kernel smoothing proposed by [38], where each vertex was weighted based on data from the neighboring vertices and scaled by the vertex area.

#### 2.4.4. Summary highlighting enhancements

A novel ODI search along surface normal for maximum ODI value is used to probe for highly probable GM regions in the co-registered image. Additionally, enhancements that are made to the earlier method [24] are the transfer of metrics of interest on to the GM cortical surface in the individual subject space instead of MNI space, to reduce the error that could occur with volume and surface deformation to the MNI template. While [24] showed the application to diffusion micro-architecture features, this work extends the applications to fMRI data, thus enabling multimodal analysis across structural and functional changes. Group analysis is performed at vertex level on the target surface.

The evaluation of the approach is carried out in the following ways.

1. We compare qualitative mean ODI, a diffusion microstructure feature, for N-GSBSS with and without cortical ODI based search in comparison with ciftify pipeline.
2. We perform non-parametric permutation testing on contrast maps obtained from first level analysis of working memory tasks in fMRI.
3. We perform a simulation study in structural MRI to evaluate sensitivity and specificity of the approach.

### 2.5. Spatial statistics

Once all the properties from different modalities were projected on the target surface, GM based vertex-wise spatial statistics were calculated using the Permutation Analysis of Linear Models (PALM) [53] package from the FSL software library (FMRIB; [http://www.fmrib.ox.ac.uk/fsl/](http://www.fmrib.ox.ac.uk/fsl/)) which performs inference through permutation. Significant
results are reported after controlling for family-wise error (FWE) with $p < 0.05$ through threshold free cluster enhancement (TFCE).

2.6. Baseline methods

2.6.1. Volume based registration (VBR) processing

Volume images of metrics of interest from other modalities were registered to MNI template by applying the non-rigid transformation obtained from anatomical T1-weighted images. Gaussian kernel smoothing of 2 mm was applied before performing spatial statistics. Nonparametric permutation based testing was performed on smoothed volume data within a brain mask using FSL PALM [53] (https://fsl.fmrib.ox.ac.uk/fsl/fslwiki/PALM). Statistical results were projected

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Fig. 2. (a) ODI overlaid with cortical surface mapping based on enclosing voxels, 1 mm above, 2 mm above, 1 mm below and 2 mm below of central surface obtained using normal search. At each vertex, maximum ODI value is selected from these 5 values along the vertex normal (white arrow in zoomed in box) and corresponding map is used for projecting the diffusion metrics on to the cortical surface. (b) Histogram of ODI projected on to the cortical surface on single subject before and after ODI search.

Fig. 3. Flowchart of the N-GSBSS data processing for each subject. (1) The central surface is reconstructed via MaCRUISE (red) (2) and transformed to the MNI space (yellow) using ANTs volume registration. (3) These volumes are diffeomorphically registered to a single target surface. (4) Metrics of interest in other modalities are co-registered to corresponding anatomical T1-weighted image. (5) Cortical ODI search is performed using ODI and $V_{iso}$ from NODDI metrics to search for higher ODI excluding $V_{iso}$ within a given range (6) Data are processed for each modality (NODDI for diffusion microstructure and first level analysis for working memory tasks) to derive metrics of interest for cross-sectional analysis. (7) Metrics of interest are mapped onto the individual surface. (8) The mappings from shape correspondence are used to project intensity values of metrics of interest to the target surface (blue). (9) Vertex-wise spatial statistics on all projected data are performed on the target surface. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
onto the target surface based on the enclosing voxel approach for visualization and comparison with surface based results.

2.6.2. Surface based registration (SBR) processing

In order to compare the proposed approach, we used the FreeSurfer software registration method [30] for cortical shape correspondence. Metrics of interest from volume data in subject space were projected onto the central surface using the enclosing voxel approach. These metrics were transferred to the target surface via the shape correspondence and smoothed on the target surface for cross-sectional analysis. In order to make a fair comparison with N-GSBSS results with optimal multiple comparison correction, metrics of interest from two hemispheres were considered as a single dataset before carrying out the permutation based statistical tests.

2.6.3. Ciftify pipeline processing

The ciftify pipeline [42] has been developed to facilitate grayordinate-based analysis in CIFTI format for legacy datasets. In preprocessing, surface reconstruction is carried out using ciftify_recon_all command that takes recon_all FreeSurfer 6.0 (https://surfer.nmr.mgh.harvard.edu) outputs and generate CIFTI file for structural measures (e.g., cortical thickness) from the surface. The distortion corrected dMRI images are registered to their own structural T1 images by FMRIB Software Library’s (FSL 5.0) FLIRT [54]. First rigid alignment is performed followed by the boundary-based registration by supplying WM segmentation obtained from FreeSurfer as an input argument. For fMRI processing, preprocessed first level images are co-registered to their own structural T1 image using SPM12. Conversion tools provided in ciftify toolbox are used to put preprocessed dMRI data and fMRI data into grayordinates in CIFTI format for further analysis. To project diffusion measures from volume onto the cortical surfaces, a ribbon mapping method is used, in which the volumetric measures are collected along the GM ribbon defined by white and pial surfaces, as described in [16]. Unfortunately, there are no T2 weighted images available in our custom dataset. Thus, myelin-style volume to surface mapping is infeasible for our diffusion analysis since myelin maps are unavailable. The grayordinates are based on the low-resolution standard mesh (with ~32 k vertices in each hemisphere) at 2 mm resolution with a total of ~64 k cortical vertices for both hemispheres obtained with the default settings. The low-resolution mesh standard template is the suggested template that is appropriate for cross-subject analysis of low-resolution data like dMRI or fMRI.

Processing time comparison between N-GSBSS and SBR using FreeSurfer are reported in Table 1. We used a single thread (Intel Xeon CPU E5-2630 v4 @ 2.40GHz and 32 GB of RAM) on an Ubuntu 16.04 LTS Linux Workstation.

2.7. Simulation study setup

The spherical masks with a radius of 3, 4, and 5 mm were created in template space and transferred back to subject space via the inverse transformation from ANTs SyN [55] registration for each subject. This range was chosen since the cortex is around < 5 mm thick and because capturing the ROIs with different radii could reflect the differences in accounting for partial volume effects in the GM and WM border regions. The location was chosen to contain cortical folding that is variable across multiple subjects to account for partial volume effects when performing cross subject studies.

The GM probability maps for the 30 subjects were randomly divided into two groups, G1 and G2, with 15 subjects in each group. The GM probability data in G2 were then modified in the subject space to simulate percentage change of intensity values in intervals of 10% in the corresponding mask regions. A total of 27 combinations (3 masks and 9 different scalings) were considered for evaluation.

With 0% change, the images in G2 were the same as original images. Thus, we considered the difference between the groups as a baseline. We excluded 100% change of the region of interest in G2, which is completely reduced to zero. With 50% change, the intensity values were half of the original values in ROIs from G2 images.

GM probability data from each of the 27 combinations in G2 were then processed through N-GSBSS to place all the data on the target surface for cross-sectional analysis. GM probability data were also evaluated for VBR, SBR and ciftify for comparison with the same parameter/experimental settings, including 2 mm Gaussian kernel smoothing. Non-parametric permutation tests were then performed between G1 and G2 for all combinations using FSL’s PALM [53] package with 5000 iterations.

To assess the sensitivity of the approaches, we examined the ratio of maximum t-statistic (“t-stat ratio”), which was defined as the amount of scaling with respect to the baseline. To have a single metric with comparable result across all the methods, we reported the ratio with respect to baseline. Baseline is where we performed second level analysis for group differences across the 2 groups where no changes are applied to original GM probability maps.

3. Results

In this section, we present the results of all the N-GSBSS analysis as follows: 1) Qualitative results of mean ODIs with and without search in comparison with the ciftify pipeline 2) Application in fMRI to identify active regions in task based working memory. 3) GM simulation results in structural MRI based on varying ROI size and intensity differences.

Mean ODI values across 30 subjects are shown on the target surface (Fig. 4) for N-GSBSS without search, with cortical ODI search and the ciftify pipeline. With cortical ODI search, partial volume effects are addressed reflecting higher ODI across the cortex compared to that of other two approaches.

3.1. Working memory fMRI results

As an application of N-GSBSS in fMRI, working memory data was processed for 30 healthy subjects in cue, probe and delay tasks. We compared significant regions revealed by VBR, SBR, the ciftify pipeline and N-GSBSS methods as shown in Fig. 5. For all these tasks, the overall activation pattern is comparable across different methods. As expected, the significant vertices in VBR are fewer and more scattered than the cortical surface-based approaches of SBR, ciftify and N-GSBSS.

Quantitative representation of the number of significant vertices with p < 0.05 for all the three tasks are shown in Fig. 6. Note that N-GSBSS has a higher number of significant vertices in all the tasks than VBR, SBR and ciftify pipeline results. The ciftify pipeline results are comparable to that of N-GSBSS more than VBR or SBR approaches. Applying cortical ODI search further improved the activation percentage in N-GSBSS.

<table>
<thead>
<tr>
<th>Pipeline</th>
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<th>Total time</th>
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<td>SBR</td>
<td>Per hemisphere:</td>
<td>~273.6 min</td>
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<td></td>
<td>FSRUNTIME@mrhspike.1.48 h, 1 thread</td>
<td></td>
</tr>
<tr>
<td></td>
<td>FSRUNTIME@mrhspike.0.80 h, 1 thread</td>
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<tr>
<td>Ciftify</td>
<td>ReconAll (mrhspike and mrhspike): 4.71 h, 1 thread</td>
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<td></td>
<td>Ciftify: 1 h 5 min, 1 thread</td>
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<tr>
<td>N-GSBSS</td>
<td>ANTs volume registration: ~2.12 min, 1 thread</td>
<td>~4 min</td>
</tr>
<tr>
<td></td>
<td>DSM surface registration: ~1.49 min, 1 thread</td>
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</table>
3.2. Simulation study in structural MRI with changes in regions of interest

Here, we evaluate N-GSBSS with respect to VBR, SBR and ciftify pipeline techniques in identifying sensitivity and specificity of changes in GM voxels located in spherical ROIs of 3, 4, and 5 mm radius located in a region of the frontal cortex. Fig. 7 illustrates spheres with a radius of 5 and 3 mm.

Quantitative results in Fig. 8 show the t-statistics ratio for varying ROI sizes of 3 mm, 4 mm, and 5 mm, and percentage change in the GM probability values from 10% to 90% in the intervals of 10%. T-stat ratio is the maximum t-statistic for each scenario with respect to the baseline to reflect how much it was scaled with induced changes in the region of interest. The baseline is chosen to be the differences between the 2 groups in the current experiment. For VBR, to capture the intensity difference between groups, the probability change must be at least 40% with 5 mm spherical ROI, 50% for 4 mm, and 60% for 3 mm ROI. SBR results showed sensitivity for 20% change with 5 mm ROI. However little difference is observed between baseline and 4 mm ROI from 40% and no difference was captured with 3 mm ROI. N-GSBSS results are much more sensitive starting at 10% with 5 mm ROI, 20% with 4 mm and 30% for 3 mm spherical ROI. N-GSBSS also showed higher maximum t-statistics than SBR. With higher intensity differences starting at 70%, VBR results have higher t-statistic ratio than that of N-GSBSS. In all other cases N-GSBSS has higher maximum t-statistic ratio and better sensitivity.

4. Discussion

Herein, we describe an approach for carrying out multi-modal spatial statistics in low resolution images by taking advantage of high resolution T1 weighted images that are acquired as part of the scan protocol. This approach favorably compares with traditional volume based analyses and with respect to the FreeSurfer surface registration approach along with the ciftify pipeline. Our approach offers an advantage over VBM by achieving improved cortical alignment in agreement with other surface-based registration techniques [25,28–33]. Moreover, in comparison with FreeSurfer, SBR, and ciftify pipelines, the N-GSBSS approach showed an improvement in sensitivity. It suggests that the initial alignment obtained by non-rigid deformation from the T1 image provides a deformed cortical shape that makes surface registration much easier. Consequently, this improves the statistical power compared to existing approaches.

The key aspect of this work is the addition of NODDI based search, which ensures that metrics from low-resolution images are retrieved from highly probable GM. It is achieved by making use of the ODI measure from NODDI which is known to be higher in GM compared to that of WM [43]. Thus by searching for maximum ODI, alignment issues after co-registration or PVE effects from underlying voxels is addressed. The patterns of mean ODI are comparable between these methods with higher values along the gyral regions. The overall mean ODI values in ciftify approach appear to be less than that of the GSBSS approach with or without search (Fig. 4). Lower values could be due to the partial
effects arising from thinner cortex regions as acknowledged in Fukutomi et al.’s paper [56] indicating the possibility of residual PVE effects in the regions of thinner cortex. When compared to mean ODI values reported in Fukutomi et al.’s paper, the results indicated in our study have higher ODI values across all the methods. Possible reason for this deviation could be due to the number of differences between the two datasets like demographics, data acquisition, and processing. Also we followed the original NODDI model which has empirical settings as mentioned below where $d_|| = 1.7 \times 10^{-3} \text{mm}^2/\text{s}$, to be representative of both white and gray matter on two-shell data ($b = 1000/2000 \text{ s/mm}^2$), while in Fukutomi et al., paper [56] $d_||$ is calculated to be $1.1 (0.1) \times 10^{-3} \text{mm}^2/\text{s}$ for gray matter from an empirically chosen range and the results reported are based on three-shell data ($b = 1000/2000/3000 \text{ s/mm}^2$). While the preliminary normal search proposed based on higher ODI seems to improve sensitivity for the changes occurring in pure gray matter, these results may have to be carefully reviewed if a regional variation is essential for the study of interest.

As we are interested in low resolution with dMRI acquired at 2.5 mm resolution and fMRI at 3 mm resolution, we are assuming that after co-registration to T1, the underlying data is roughly aligned at voxel level. Thus we utilize the search map obtained from diffusion modality to apply to fMRI for getting the data based on enclosing voxel approach. The reported fMRI t-statistics suggest an improvement in sensitivity with N-GSBSS. While there is no ground truth for validating the implication of the higher activation, since the contrast maps are relative to that of the un-modeled baseline across 30 subjects, the activation could indicate that the proposed method could be highly sensitive to capture underlying variations that are indirectly contributing to the activations instead of capturing the false positives.

The simulation study is set up to perform sensitivity or specificity check for N-GSBSS to the underlying changes in tissue microstructure. As we are interested in performing analysis in psychiatric applications including schizophrenia [57,58] that are known to have changes in prefrontal region, the ROI is chosen from this region. The GM

![Fig. 6. Percentage activation of working memory fMRI data were processed for 28 healthy controls and results are reported for (a) correct cue, (b) correct delay (c) correct probe tasks with 2 mm smoothing for VBR, SBR, ciftify, N-GSBSS-S0 with no search and N-GSBSS-S2 with 2 mm search methods. The number of significant vertices, with p-values < 0.05 after FWE correction based on nonparametric randomize one sample t-test with 10,000 iterations, is divided by total number of vertices and the percentage is reported.](image1)

![Fig. 7. The gray matter probability map shows the simulated effect as an overlay mask of 5 mm (red) and 3 mm (dark blue) spheres. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)](image2)
probability map is chosen as the parameter of interest and the intensity changes are simulated within an ROI region. Compared to the baseline methods, N-GSBSS showed superior sensitivity to the underlying changes in both intensity and the size of the ROI as shown in Fig. 8. While volume-based analysis was not able to detect any significant differences between groups for at least up to 50% change in the GM probability values, N-GSBSS was able to capture differences starting from 10% change with ROI size of 5 mm, 20% for 4 mm and 30% for 3 mm. The low performance of VBM could be potentially due to partial volume effects prevalent in the volume-based approach even after applying the GM mask to limit the analysis to highly probable GM regions.

In the simulation study, SBR analysis showed a similar pattern as N-GSBSS. However, the sensitivity of this approach is not as high as N-GSBSS. Differences between the methods are likely due to different registration approaches since both of them used the same surface to obtain corresponding GM probability values from the volume image. The ciftify pipeline results are similar to those of SBR, which is expected since the ciftify pipeline uses FreeSurfer registration. The subtle difference between ciftify and SBR are observed likely due to the different surface reconstruction in each of these pipelines. For a fair comparison, we used the ciftify pipeline with default parameters to the extent possible. For example, the analysis results in the ciftify pipeline are based on the “gray ordinates” with 64 k vertices (the suggested tessellation for cross subject analysis of low resolution data) on both left and right hemispheres. This surface tessellation differs from that of the target central surface used in SBR and N-GSBSS analysis (about 261 k vertices for both hemispheres). This could have contributed to the lower sensitivity of ciftify pipeline in this simulation study due to the limited ability to capture smaller ROI regions with less number of vertices. The higher sensitivity of N-GSBSS to capture GM probability percentage changes as low as 10% for 5 mm ROI and 40% for smaller ROI of 3 mm ROI could indicate that it is able to capture more number of highly probable vertices accurately. In future, additional validations could be performed to evaluate the performance for different resolutions and also at different ROI locations.

5. Conclusion

Overall significant regions captured by N-GSBSS agree with those of VBR, SBR, and ciftify pipelines across different modalities while achieving high spatial specificity. It is highly likely that the volumetric transformation already deformed cortical surfaces into similar shapes (geometry) before the surface registration, which results in better shape correspondence by reducing the local anatomical ambiguity in the surface registration. N-GSBSS possesses high flexibility that allows any registration method as well as multiple modalities. We expect that such a feature can be generally extended to various modalities in general neuroimaging studies.

An operational virtual machine and source code for N-GSBSS are posted in a Docker image: (https://github.com/MASILab/N-GSBSS/).
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Appendix A. Supplementary data

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References


