Abstract

A new functional magnetic resonance imaging (fMRI) analysis approach based on invariant spatial descriptors was developed for quantitative characterization of brain activation patterns within a region of interest (ROI). In particular, the feasibility of using three dimensional moment invariants (3DMIs) to perform such spatial characterization was examined. The use of spatial descriptors is particularly novel in the field of ROI-based fMRI analysis, since up till now, only magnitude-based features were traditionally employed, which neglect the information encoded by voxel locations within an ROI. The invariance properties of the proposed descriptors to similarity transformations account for inter-subject variability in brain size and subject’s orientation within the MR scanner, thus allowing for spatial distributions of activation statistics to be meaningfully compared across subjects. Enhanced sensitivity in detecting task-related activation differences as compared to traditional magnitude-based methods was demonstrated with real fMRI data.

To handle the issue of feature selection, a modified linear discriminant analysis (LDA) procedure that incorporates leave-one-out cross-validation was developed. Also, methods to deal with the two main issues in ROI-based fMRI group analysis, namely errors in ROI delineation and inclusions of voxels falsely deemed active, were proposed. One method involves remapping the coordinate space with a Gaussian
function, which in effect de-emphasizes voxels near the ROI boundary, thus also accounts for inter-subject variability in brain shapes. The other method detects outlier voxels that exhibit disproportional influence on the proposed invariant spatial descriptors, and deweights or removes those voxels accordingly. Testing these processing methods on real fMRI data showed further increase in discriminability of task-related activation differences compared to the original 3DMIs alone.

To fully exploit the spatio-temporal structure inherent in fMRI data, we extended our spatial characterization approach into the spatio-temporal domain. We showed, for the very first time, that the modulation of the spatial distribution of BOLD signals does in fact correlate with the stimulus, and provides greater sensitivity in detecting activated ROIs and in discriminating task-related differences as compared to traditional mean intensity-based methods. The neuroscience implications of our findings are substantial, and might hence provide brain researchers and clinicians a new promising direction to explore.
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Statement of Co-authorship

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Chapter 1

Introduction

Traditional neuroimaging mainly focuses on detecting structural abnormalities such as brain atrophy in multiple sclerosis (MS) patients, to detect, track, and evaluate neurological disease. However, examining only the structural properties of the brain is inadequate in explaining many neurological disorders such as Parkinson's Disease (PD), that are associated with the brain's functional mechanisms. To facilitate a broader understanding of human brain functions, considerable research efforts have been invested in advancing functional neuroimaging during the past decade with promising results gradually translated to clinical settings [1]. In this thesis, we present our contributions to one of the most prevalent functional neuroimaging modalities, namely functional magnetic resonance imaging (fMRI). In particular, we introduce a novel region of interest (ROI) based fMRI analysis approach for detecting activated ROIs and discriminating activation pattern changes.
1.1 Problem Statement and Motivation

Based on the Global Burden of Disease (GBD) study undertaken by the World Health Organization, World Bank, and Harvard School of Public Health, neurological disorders are ranked as the number one disease burden among other diseases examined including the human immunodeficiency virus (HIV), malignant neoplasm, and digestive diseases [2]. Neurological disorders constitutes 6.3% of the global burden of diseases [2], and is predicted to increase as the global population continues to age. Among the neurodegenerative disorders, Parkinson's disease (PD) is found to be the second most common worldwide, affecting 3,765,000 people [3]. In fact, Canada alone has approximately 100,000 people suffering from PD. Thus, in this thesis, an emphasis is placed on characterizing the brain activity of PD patients [4], [5] as compared to healthy subjects [6], [7], [8], [9]. Nonetheless, the generality of our approach permits other neurological diseases to be examined using the same methods presented.

PD is a chronic, progressive neurodegenerative disease with primary symptoms such as tremor, rigidity, bradykinesia, and postural instability. These symptoms greatly disrupt the daily activities of PD patients. For instance, even something as simple as eating is affected due to tremoring hands. Also, PD can result in depression, slowed reaction time, difficulty in swallowing, and sleep disturbance among many other complications. Most motor-related symptoms are believed to arise from premature death of dopamine-secreting cells in the substantia nigra pars compacta (SNpc) – a brain region involved in motor control. However, the exact cause of premature cell death is unknown, though some studies suggest abnormal accumulation of the protein alpha-synuclein [10].

Currently, the primary clinical tool for diagnosing PD is the Unified Parkin-
son's Disease Rating Scale (UPDRS) [11], which is based on examining patients' symptoms severity. This measure, however, is insufficient in fully detecting and diagnosing PD, partly due to the brain's compensatory mechanisms, which obscure the effects of disease. Also, the symptoms of PD are often confused with the effects of normal aging. To enable better assessment and tracking of PD, alternative markers that directly target the source (i.e. the brain) is required. One possible biomarker is the functional property of the brain, which was examined in this thesis. In particular, the brain activity in motor related regions was studied (see Fig. 1.1).
Prior electrophysiological studies have shown that both cortico-cortical [12] and cortico-subcortical coupling [13] are impaired in PD patients, and are modulated by medication [12], [13]. Also, studies on monkeys with PD showed overactivity of the globus pallidus even before symptoms onset [14]. Furthermore, reduced brain activity was found in the supplementary motor area (SMA), dorsolateral prefrontal cortex (DLPFC), and anterior cingulate during voluntary movements for patients with more advanced disease symptoms [15]. All these preliminary findings confirm the importance of developing sensitive tools to facilitate characterization of brain functions at various stages of disease.

One non-invasive imaging modality that is particularly suitable for studying brain function and permits both spatial and temporal characterization is fMRI. The spatial aspect of fMRI is particularly important, as suggested by McKeown et al. in [16], where brain activation of PD patients was found to focus (i.e. become more localized within a particular brain region) in certain cortical areas upon medication. Thus, extending traditional magnitude-based analysis to the spatial and spatio-temporal domain provides a more complete characterization of brain functions and effects of disease. Such characterization can be transferred to a wide range of clinical applications ranging from disease diagnosis to drug therapy evaluation. For example, by examining at which dosage levels does brain activity change significantly, optimal drug dosage can be inferred. Also, earlier PD detection might be feasible by examining how brain function changes with disease progression. This enables clinicians to develop more effective therapies to better treat PD patients at an earlier stage.
1.2 Thesis Objective

Traditionally, only magnitude-based features such as mean voxel statistics and percentage of activated voxels [17], are used in ROI-based fMRI analysis. This limitation on features is mainly due to inter-subject variability, where nonfunctional related differences in brain shapes, brain sizes, and subject's orientation in the scanner will be amplified if spatial information is naively incorporated. However, prior neuroscience studies suggest that different tasks might be spatially encoded by voxel locations within an ROI [16], [18], [19]. Thus, developing methods to quantitatively characterize the spatial distribution of activation statistics can be beneficial.

In this thesis, the feasibility of using invariant spatial descriptors to account for inter-subject variability was explored. In particular, three dimensional moment invariants (3DMIs) were examined. Also, processing methods based on the proposed invariant spatial descriptors were developed to account for the two main issues in ROI-based fMRI analysis, namely errors in ROI delineation and inclusions of voxels falsely deemed active. Furthermore, to fully exploit the spatio-temporal structure inherent in fMRI data, we extended our spatial characterization into the temporal domain and explored whether the spatial distribution of the blood oxygenation level-dependent (BOLD) signal itself is modulated in time by task performance.

1.3 Magnetic Resonance Imaging

MRI exploits the nuclear magnetic resonance (NMR) phenomenon inherent to hydrogen atoms to non-invasively and safely image soft tissues in living subjects. Since upon excitation of the tissues, all hydrogen atoms will simultaneously respond, a spatial mapping formed by applying different magnetic gradients at different loca-
tions is used to localize each signal source. A brief review of the NMR phenomenon and the MRI principle are provided next.

1.3.1 Nuclear Magnetic Resonance Phenomenon

The human brain is composed of 78% water [20], and each water molecule contains two hydrogen atoms. Each hydrogen atom has a magnetic moment associated with its intrinsic spin. In the absence of an external magnetic field, the orientation of the spin axis of the hydrogen atoms will be random. Thus, no net magnetic moment will be present, as shown in Fig. 1.2(a). When a magnetic field, $B_0$, is applied, the spin axis of the hydrogen atoms will rotate (or precess) around the direction parallel or anti-parallel to $B_0$, as shown in Fig. 1.2(b). More hydrogen atoms will tend to precess around the parallel direction, since the parallel direction corresponds to a lower energy state, thus resulting in a net magnetic moment, $M$ (Fig. 1.2(c)). The

Figure 1.2: Nuclear Magnetic Resonance. (a) Orientation of hydrogen atoms when no magnetic field is present. (b) Orientation of hydrogen atoms when a magnetic field, $B_0$, is applied. (c) Resulting net magnetic moment, $M$, in the direction of $B_0$. (d) Response of $M$ upon excitation by a radio frequency (RF) pulse.
angular frequency of precession, $\nu_0$, is governed by the Larmor relationship:

$$\nu_0 = \gamma \cdot B_0,$$

(1.1)

where $\gamma$ is the gyromagnetic ratio, whose value depends on the nature of the nuclei. For hydrogen, $\gamma = 42.58$ MHz/T. When a radio frequency (RF) pulse matching the precession frequency is applied perpendicular to $B_0$, $M$ will be flipped towards a direction that is perpendicular to $B_0$ as shown in Fig. 1.2(d). The resulting magnetic moment along the $z$-direction, $M_z$, will be zero while the magnetic moment in the $xy$-plane, $M_{xy}$, will be equal to $M$. In this orientation, the hydrogen atoms will be in an excited state. When the RF pulse is switched off, $M_z$ will begin to increase exponentially with a time constant, $T1$, also known as spin-lattice relaxation time, as $M$ returns to its equilibrium state (i.e. aligned with $B_0$). $T1$ differs across tissue types. Hence, different tissues will exhibit different $M_z$, thereby providing the signal contrast required for generating brain images.

Another phenomenon that provides signal contrast is the spin-spin interactions of the hydrogen atoms. At equilibrium, hydrogen atoms precess incoherently, but when excited with an RF pulse, the precessions will become in phase, leading to stronger $M_{xy}$. When the RF pulse is switched off, the hydrogen atoms will begin to de-phase exponentially with a time constant, $T2$, known as spin-spin relaxation time, which also depends on tissue type. Also, inhomogeneities of the magnetic field can lead to faster de-phasing, thus $M_{xy}$ in fact decays with a time constant, $T2^*$ that is shorter than $T2$. This magnetic inhomogeneity-induced contrast is the means by which fMRI images are generated as discussed in Section 1.4.1.
1.3.2 Magnetic Resonance Imaging Principle

If only an external magnetic field, $B_0$, is applied, all hydrogen atoms inside the brain will precess at the same Larmor frequency. Therefore, exciting the brain with an RF pulse matched to this Larmor frequency will excite every hydrogen atom. The measured signal thus will be a mixture of signals from multiple regions of the brain. To localize the signal sources, a spatial mapping is created through controlled manipulations of the external magnetic field, as shown in Fig. 1.3. When $B_0$ is applied, the spin axis of all hydrogen atoms in the brain will be aligned in the direction of $B_0$. If we then apply a magnetic gradient, $G_z$, along the $z$-axis, each hydrogen atom will precess at a new frequency governed by:

$$\nu = \gamma (B_0 + G_z z), \quad (1.2)$$

where $z$ is the $z$-coordinate of a given group of hydrogen atoms in a small volume of tissue. If we then apply an RF pulse matched to $\nu$, the spin axis of the hydrogen atoms precessing at $\nu$ will be flipped onto the $xy$-plane. Thus, a slice of the object will be isolated.

After selecting a slice of an object, we still need to isolate each small volume of tissue in the $xy$-plane. If we apply a magnetic gradient, $G_y$, along the $y$-axis, each hydrogen atom will again precess at a new frequency governed by:

$$\nu = \gamma (B_0 + G_y y), \quad (1.3)$$

where $y$ is the $y$-coordinate of a given group of hydrogen atoms in the selected slice. Based on (1.3), hydrogen atoms residing at higher gradient will precess faster than those residing at lower gradient. If we then remove the magnetic gradient, all hydrogen atoms will resume to their original precession frequencies with a phase
A group of neighboring hydrogen atoms in a small volume of tissue

Figure 1.3: Magnetic Resonance Imaging Principle. (a) Orientation of magnetic moments (black arrows) when a magnetic field, $B_0$, is applied in the z-direction. (b) Slice selection with an RF pulse. All moments precess at the same frequency. (c) Differences in the rate at which the moments precess when a magnetic gradient is applied in the y-direction. More arrows on the circle indicate faster precession. (d) Resulting phase differences between the moments in the y-direction when magnetic gradient is removed. (e) Different precession frequencies when a magnetic gradient is applied in the x-direction.
difference introduced. Thereby, a mapping between phase to location along the y-axis is generated. If we then apply a gradient along the x-axis, the hydrogen atoms along the higher gradient will again precess at a faster rate. Therefore, by examining the phase and frequency of the signal using the Fourier transform, we can determine the spatial location from which the signal originated. This procedure is repeated at a time interval, $TR$, for different slices to image a 3D object such as the human brain.

1.4 Functional Magnetic Resonance Imaging

fMRI is based on the assumption that regional changes in blood flow, blood volume or blood oxygenation level during task performance are related to brain activation. Regional hemodynamic responses can alter the local magnetic fields inside the brain, which creates image contrast. Among the different contrast techniques, BOLD contrast is the most frequently used [21]. To infer brain activation, multiple $T2^*-\text{weighted}$ brain volumes are first acquired over time as a subject perform a certain task (Fig. 1.4). Changes in intensity of a voxel are then tracked over time to generate an intensity time course. Such time course is then compared to an expected response to infer activation. However, as shown in Fig. 1.4, the raw intensity time course of a voxel can be quite different from that of the expected response. Thus, substantial preprocessing is required to unveil the underlying signal of interest. However, the inherently low signal to noise ratio (SNR) and the complex noise structure of BOLD signal makes brain activation detection rather difficult. In particular, noise sources such as random motion artefacts and location-dependent low frequency drifts are especially hard to handle. The spatio-temporal correlations resulting from these non-functional related confounds can greatly complicate fMRI
analysis, and are yet to be resolved.

Figure 1.4: fMRI Analysis Overview. (a) A series of brain images acquired over time. (b) Intensity changes of a voxel over time. Typically, the intensity time course is contaminated by various noise sources such as low frequency drift, high frequency noise, and motion artefacts. (c) Pre-processed time course compared to an expected response.

1.4.1 Hemodynamic Response

BOLD contrast techniques exploit changes in local magnetic field inhomogenities arising from task-induced modulations of blood oxygenation level, to infer brain activation. The underlying assumption is that the measured BOLD signal is approximately proportional to the amount of local neuronal firing [22]. The temporal profile of BOLD signal (i.e. its hemodynamic response) of a small volume of tissue in response to stimulus is shown in Fig. 1.5. During the onset of stimulus, neurons in a given small volume of tissue will begin to fire, leading to oxygen uptake as
required for energy metabolism. De-oxygenated blood is paramagnetic thus introduces inhomogeneity to its surrounding magnetic field, resulting in a decrease (an initial dip shown in Fig. 1.5) in the BOLD signal as was discussed in Section 1.3.1. In response to this demand for oxygen, an oversupply of oxygenated blood rushes into the given small volume of tissue. Oxygenated blood is diamagnetic which does not directly affect the local magnetic field; but since the amount of de-oxygenated blood is reduced, the local magnetic field will become relatively more homogeneous, leading to an increase in BOLD signal. After stimulus ceases, the supply of oxygenated blood diminishes as the BOLD signal gradually returns back to baseline after an undershoot.
1.4.2 Noise Issues in fMRI

The noise structure of BOLD signals is very complex, which greatly complicates brain activation detection. The signal of interest in the raw intensity time courses are often hidden by many confounds. As shown in the sample raw intensity time course in Fig. 1.4, the signal of interest is contaminated by a mean intensity offset, slow frequency drifts, and high frequency noise in addition to motion artefacts which are harder to detect by inspection. The magnitude of the mean intensity offset, however, does not correspond to the level of neural activity. Instead, this offset arises from slight $T_2$ differences of each voxel, and is rather difficult to estimate due to the unpredictable trends of the low frequency drifts.

Low frequency drifts are suspected to be induced by scanner instability, though the physiological state of the subject may also be a cause. Among the different scanner-related noise sources such as magnetic field inhomogeneities and thermal noise, low frequency drifts is probably the most problematic to brain activation detection. The problem arises from the way brain activation is inferred, where the higher the correlation between a voxel time course and an expected response signal (see Fig. 1.4), the more likely the voxel is activated. However, low frequency drifts introduce extra autocorrelations to the voxel time courses, which in turn increases correlation, leading to false activation detections. Methods dealing with autocorrelation in BOLD signal are discussed in Section 1.5.

In addition to scanner noise, fMRI is prone to many physiological noise sources such as motion artefacts, cardiac-linked brain pulsation, respiration, and uncontrolled spontaneous neuronal events. Among these noise sources, motion artefacts is the most challenging to deal with [23]. In general, motion can introduce artefacts into the BOLD signals by two means. First, inhomogeneities of the mag-
netic field induce differences in BOLD signal across positions inside the scanner. Thus, if a subject moves during data acquisition, the BOLD signal of a given voxel will change even if no brain activation is present. Secondly, the strength of the measured BOLD signal depends on spin excitation history [24]. Thus, if a previously excited slice moves into the current acquisition plane, the measured BOLD signal will show reduced amplitude since the previously excited spins would have decayed towards their equilibrium states. This effect, however, is only present for through-plane head motions. Further complicating fMRI analysis is that subjects tend to move in response to stimulus [25], thus introducing stimulus-correlated changes in the BOLD signals that are not related to brain activation. A discussion of methods for handling motion-induced artefacts is provided in Section 1.5.

1.5 fMRI Preprocessing

Extracting relevant neurological information from fMRI data is very challenging due to its complex noise structure. Numerous preprocessing procedures are required to reduce the effects of noise in subsequent analyses. Typical procedures include motion correction, slice timing correction, and temporal autocorrelation correction. Also, due to inter-subject variability in brain shape, brain size, and subject’s orientation in the scanner, to make group statistical inferences, spatial normalization and spatial smoothing are often performed.

1.5.1 Spatial Realignment and Motion Correction

Most existing motion correction methods involve spatially realigning the motion-corrupted fMRI image volumes to a reference volume, often taken as the first volume or the average of the image volumes. The volumes to be registered to the reference
volume are referred to as floating volumes. One approach in realigning the volumes is to define fiducial points and align the fiducial points in the floating volumes to that of the reference volume. However, the low spatial resolution of fMRI images and the lack of distinct fiducial points in the brain makes the fiducial point-based approach inferior to intensity-based approaches [26].

The overall intensity-based realignment process involves three major steps. First, six rigid transformation parameters are estimated to rotate and translate a floating volume so that it becomes aligned with the reference volume. Resampling is then performed to interpolate the intensity values of the floating volume at each voxel location of the reference volume. The optimal resampling method is Fourier interpolation [27], which is equivalent to sinc interpolation in the spatial domain. However, sinc interpolation involves using every voxel in a volume to interpolate each voxel's new intensity value, thus computationally impractical. Alternative interpolation methods include trilinear and B-spline interpolations [28].

Following rigid transformation and resampling, a criterion to determine the spatial correspondence between the floating volume and the reference volume is evaluated. Many criteria have been proposed including image difference [23], image ratio uniformity [29], correlation [30], and mutual information (MI) [31], but no census has been reached as to which criteria is optimal. After criterion evaluation, the described procedures including rigid transformation, resampling, and criterion evaluation are iterated until a desired matching criterion is achieved.

Spatial realignment corrects for the differences in orientation of the brain volumes, but does not account for motion-induced intensity changes as discussed in Section 1.4.2. To account for signal changes due to magnetic field inhomogeneities and spin excitation history effects, Friston et al. in [24] proposed using an autore-
gressive moving average (ARMA) model, which incorporates the position of a given voxel in current and previous scans in correcting the signal intensity of a given voxel. Alternatively, Liao et al. in [32] proposed a method called, "Motion-corrected independent component analysis" (MCICA), which corrects for motion artefacts by determining a linear combination of spatial transformations applied to certain basis motion-corrupted volumes that maximizes a joint entropy-based criterion. Their simulation results demonstrated that MCICA is robust to activation level, additive noise, and random motions in the reference volume.

1.5.2 Slice Timing Correction

During analysis, all slices in a brain volume are assumed to be collected simultaneously. However in practice, each slice is collected at a slightly different time. This artefact can lead to phase differences between voxel intensity time courses. For instance, assuming brain activation onsets at time $t$, if an activated voxel resides in a future slice, the first signal measurement will be made at $t + dt$. This intensity value will be interpreted as measured at time $t$, thus the voxel will appear to have begun responding prior to stimulus onset. To account for this artefact, slice timing correction is often performed through Fourier transform, where the magnitude spectrum of the signals is preserved, but the phase is adjusted accordingly. This procedure is equivalent to sinc interpolation in time, which is used in standard fMRI analysis packages such as statistical parametric mapping (SPM) [33].

1.5.3 Temporal Autocorrelation Correction

To deal with temporal autocorrelations in the BOLD signals that are not related to brain activation, two approaches are conventionally used – coloring and whiten-
Coloring involves high-pass filtering to remove the majority of the low frequency autocorrelations followed by temporally smoothening the BOLD signals with a Gaussian kernel matched to the hemodynamic response function to increase the signal to noise ratio (SNR) [35]. This procedure imposes an autocorrelation structure onto the BOLD signals and presumes that the imposed autocorrelation overwhelms any other noise-induced autocorrelations.

The alternative approach is whitening, which involves regressing out the task-related response from the BOLD signals and estimating the autocorrelations in the residual signals for generating a whitening matrix. This matrix is multiplied by the BOLD signal to minimize residual autocorrelations. Existing methods for estimating residual autocorrelation includes AR(1) [36], AR(1) plus white noise [37], AR(\(\rho\)) [38], ARMA [39], and 1/f noise model [40].

In addition to the two described approaches, methods based on component analyses have also been recently explored [41]. Such approach involves decomposing the BOLD signals into signals of interest and confounds, as discussed in Section 1.6.1. Also, methods exploiting the whitening or decorrelating property of wavelets have been proposed [42], [43].

1.5.4 Spatial Normalization

To make group inference in fMRI studies, inter-subject variability in brain shape, brain size, and subject's orientation in the scanner must first be accounted for. One widely used approach is to co-register all subjects' brain images to a common brain template such as the Talairach space [44]. This co-registration is performed by spatially warping or normalizing the structural brain image of each subject to a template, and then applying the same spatial warping to all functional images.
of the subject. The advantage of warping the structural image instead of directly warping the functional images is that the structural image has significantly higher spatial resolution.

The overall spatial normalization procedure involves two steps. First, an optimal 12-parameters affine transform is performed to coarsely register the structural image to a template. To ensure that implausible transformations will not be produced, prior information of the parameter values are often incorporated through a Bayesian framework. The additional 6 parameters (zoom and shear along the 3 coordinate directions) compared to rigid transform as discussed in Section 1.5.1, account for brain shape and size differences between a subject’s brain and a template. Local refinements can then be performed through non-linear deformations as modeled by a linear combination of smooth spatial basis functions [45].

Alternatively, instead of normalizing the whole brain, approaches for aligning a subject’s brain at the ROI level have recently been proposed. Miller et al. [46] used large deformation diffeomorphic metric mappings (LDDMM) for ROI registration and demonstrated increased statistical power over conventional whole brain normalization to the Talairach space. An alternative approach based on continuous medial representations (cm-rep) of the ROIs has been proposed [47].

Several drawbacks to spatial normalization are worth noting. First, the brain shape and size of a subject could be quite different from that of a template, especially for diseased patients. Therefore, improbable deformations of a subject’s brain may result. Also, any interpolation involved in the normalization procedure might introduce false information into subsequent analysis. Secondly, estimating a huge number of parameters is typically required during non-linear deformations (ranging from hundreds to thousands), and poor parameter estimates can easily lead to
imperfect registration results [48]. Furthermore, normalization artefacts can cause
signals from functionally distinct areas, especially small subcortical structures, to
be misaligned [49]. Thus, the functional overlap across subjects will be reduced
[50]. The most conventional method to deal with this issue is to spatially smooth
the data to increase functional overlap. This method, however, may inappropriately
pool responses from functionally dissimilar regions, thus reducing the spatial reso-

1.5.5 Spatial Smoothing

Spatially smoothing the data is intended to serve several purposes. First, the spatial
scale of hemodynamic response is approximately 2 to 5 mm, based on high resolu-
tion optical imaging experiments [52]. Therefore, by the matched filter theorem,
smoothing the data with a kernel of similar size as the hemodynamic response can
increase the SNR. Secondly, the Gaussian random field theory used for multiple
comparison corrections as to be discussed in Section 1.7.1, requires estimating the
spatial smoothness of the data, which is often assumed to be similar to that of the
smoothing kernel [52]. Lastly, spatially smoothing the data helps increase the func-
tional overlaps across subjects for making group inferences as discussed in Section
1.5.5. However, this procedure may pool functionally dissimilar regions together,
thus rendering subsequent analysis erroneous as pointed out in Section 1.5.4.

An alternative approach for increasing SNR without sacrificing as much spa-
tial resolution, is to apply spatial wavelet transforms [53], [54]. This approach involves representing clusters of voxels with a few wavelet coefficients and de-weighting or removing those coefficients that fall under a certain threshold (i.e., soft or hard thresholding) chosen based on the level of Type I errors (voxels falsely deemed activated) the experimenter is willing to accept. The inverse wavelet transform is then applied to the surviving coefficients to estimate a denoised version of the data. Also, a multifiltering approach based on combining spatially smoothened data at multiple scales was proposed with enhanced brain activation detection demonstrated [55].

1.6 fMRI Analysis Approaches

In general, fMRI analysis approaches can be classified into two categories, namely exploratory and hypothesis-driven. The exploratory approach attempts to identify and group interesting patterns in the data, whereas hypothesis-based approach examines the correspondence between the data and a hypothesized response to experimental stimulus.

1.6.1 Exploratory Approach

The explorative approach seeks to uncover the underling patterns in the data without the need to hypothesize an expected stimulus response, which is often quite difficult. Using this analysis approach, unforeseen signals of interest can be discovered, in addition to task-related signals and confounds. Frequently used methods include clustering [56], [57], [58], [59], [60], [61], [62], [63], principal component analysis (PCA) [64], independent component analysis [41], and variants of these methods.

Clustering involves identifying and grouping voxels with similar temporal
patterns. Common techniques including K-means [56], fuzzy K-means [57], and
dynamical cluster analysis [59] require specifying the number of clusters and a sim­
ilarity metric a priori. However, the correct choice of the number of clusters and
a similarity metric is not obvious, and the quality of the extracted clusters is diffi­
cult to assess. Thus, even though these clustering techniques are computationally
efficient, the generated results might be hard to interpret and justify.

An alternative approach is component analysis based method, such as PCA
and ICA, which involves estimating a matrix, $W$, to project the voxel intensity time
courses onto a space where task-related components and confounds can be separated:

$$S = W \cdot X, \quad (1.4)$$

where $X$ is a $N \times T$ matrix with each row containing a voxel intensity time course,
and $S$ is a $N \times T$ matrix with each row containing the intensity time course of a
decomposed component. $N$ and $T$ are the number of voxels within an ROI (e.g.
the whole brain) and number of time points, respectively. To determine which voxel
might be activated, components that resemble confounds are first removed from
$S$ with only task-related components projected back into the voxel intensity time
course domain, i.e. $X' = W^{-1}S$. Non-zero rows of $X'$ thus corresponding to the
activated voxels.

PCA projects the voxel intensity time courses onto a space where the vari­
ance of the data is maximized along the projection directions. This criterion lacks
neurological motivation, but can nonetheless generate task-related signals and com­
ponents that resemble scanner and physiological effects as was demonstrated in a
study on the effects of nicotine on sustained attention [65].

A more neurologically motivated approach is ICA, which takes advantage of
the sparse nature of neural coding [66], i.e. only a portion of the brain activates
at a given instant in time. Conventional ICA methods either generate components that are independent in time (mutually independent rows of $S$) [67] or independent in space (mutually independent columns of $W$) [41]. However, seeking independent components in time without any spatial constraints can lead to physically improbable forms of $W$ and vice versa [68]. Instead, Stone et al. in [68] proposed using skewed-spatiotemporal ICA to analyze fMRI data, which maximizes statistical independence in both time and space. Task-related time courses generated with skewed-spatiotemporal ICA showed superior correspondence with the expected response as compared to PCA, temporal ICA, and spatial ICA [68].

### 1.6.2 Hypothesis-based Approach

The hypothesis-based approach examines the correspondence between a voxel intensity time course and a hypothesized response to experimental stimulus to infer whether a voxel is activated. The most widely used hypothesis-based method by far is the general linear model (GLM) as popularized by the software, SPM [33]: Each column of $Y$ contains a voxel's intensity time course, and each column of the design matrix $X$ contains a regressor, which can be the time courses of the signal of interest or the time course of a confound. Only a boxcar time-locked to stimulus and convolved with the hemodynamic response is illustrated in the example in Fig. 1.6, assuming that motion artefacts and temporal autocorrelations were removed during the preprocessing stage. If confounds were not removed prior to analysis, other regressors can be horizontally concatenated to $X$ to account for those confounds.

Each element in $\beta$ represents the correspondence between a voxel's intensity time course (i.e. a column of $Y$) and a regressor (i.e. a column of $X$). Each column of $\beta$ represents the residual of the corresponding voxel. To generate an activation
Figure 1.6: General Linear Model. The matrix $Y$ contains a voxel intensity time course along each column, and the matrix $X$ contains the regressor(s) such as the expected response. Each element in $\beta$ represents the correspondence between a voxel time course and the expected response, and $\varepsilon$ is the residual matrix.

statistics map (or statistical parametric map), each element in $\beta$ is scaled by the variance of the corresponding column of $\varepsilon$. A threshold is then set to determine which voxels are activated as to be discussed in Section 1.7.1. We note that hypothesizing $X$ is not trivial. Addressing this concern, McKeown in \cite{69} proposed a hybrid approach, where regressors are predicted using spatial ICA.

### 1.7 Activation Detection using Inferential Statistics

After an activation statistics map is generated, one can either separately examine whether each voxel is activated (i.e. has statistical value above a certain threshold) or can examine the statistical properties of a group of voxels (i.e. an ROI).
1.7.1 Voxel-based Approach

The voxel-based approach examines each voxel's statistical value separately, thus provides better localization of brain activation as compared to the ROI-based approach. However, to make group inference under such approach, the activation statistics map must first be spatially warped into a common space to generate exact correspondence between voxels across subjects. This warping procedure can introduce artefacts as discussed in Section 1.5.4. Also, the choice of threshold above which a voxel is considered activated is not obvious, though oftentimes a threshold corresponding to an uncorrected p-value of 0.05 is used across all voxels, which is inappropriate due to the multiple comparisons problem.

With thousands of voxels within an activation statistics map, tens or even hundreds of the voxels could easily be falsely deemed active by chance (e.g. 5% of 10,000 voxels = 500 voxels). To address this problem, most fMRI studies now incorporate some form of multiple comparisons correction such as Bonferroni correction or methods based on random field theory [70] or false discovery rate (FDR) [71]. Bonferroni correction is based on simple probability rules as follows:

Let $\alpha_{FW}$ be the family-wise Type I error rate, which is defined as the probability of making at least one Type I error in a family of tests (i.e. 1 voxel out of $N$ falsely declared activated) when all the null hypothesis are true. If we let $\alpha$ be the probability threshold for each test, then the probability of not making a Type I error is $1 - \alpha$. For $N$ tests, the probability would be $(1 - \alpha)^N$. Thus, $\alpha_{FW} = 1 - (1 - \alpha)^N \approx N \cdot \alpha$ since $\alpha$ is small. Thus, for 10,000 voxels and $\alpha_{FW} = 0.05$ for example, an $\alpha$ of $0.05/10,000 = 0.000005$ is required. However, this threshold is too conservative for most cases, since the voxels are spatially correlated. To account for this spatial correlation, a less stringent threshold based on Gaussian random field
(GRF) theory have been proposed [70].

In brief, determining a threshold based on GRF theory involves first estimating the smoothness of an activation statistics map, which is often assumed to be similar to that of the spatial smoothing kernel. The estimated smoothness value is then used to define the size of a voxel cluster and calculate the expected Euler Characteristic (EC), where EC is the number of voxel clusters that remain after a certain activation statistics threshold is applied. As we increase the threshold, the number of clusters that remain will be either one or zero. Recall that the family-wise Type I error is the probability of making one Type I error in a family of tests. Since the expected EC corresponds to the probability of finding only one cluster above a particular threshold by chance, the expected EC is thus approximately equal to the family-wise error. If we now reverse the argument by first choosing a certain \( \alpha_{FW} \), we can determine the corresponding activation statistics threshold at which the family-wise error will be less than \( \alpha_{FW} \).

The described GRF based procedure requires spatially smoothing the data, which might inappropriately pool functionally dissimilar regions together, as mention in Section 1.5.5. An alternative method that is also less stringent than Bonferroni correction uses FDR to determine which voxels are likely to be activated. FDR is defined as the proportion of false positives (voxels falsely deemed active) among those tests for which the null hypothesis is rejected. Bonferroni correction, on the other hand, controls the amount of false positives among all tests whether or not the null hypothesis is rejected. This FDR-based procedure was introduced in [72] and works as follows:

Let \( q \) be the maximum FDR between 0 and 1 that a researcher is willing to ac-
cept on average. Order the p-values of the $N$ voxels from smallest to largest: $p(1) \leq \ldots \leq p(N)$. For $i$ from 1 to $N$, voxels with $p(i) \leq i \cdot q/V$ are declared activated.

The advantage of the FDR approach is that the threshold is adaptive to each subject’s data, thus relieves the need to find an arbitrary threshold that works for all subjects. However, spatial correlation is not accounted for, which implies the calculated threshold will still be too conservative depending on the level of spatial correlation present.

1.7.2 ROI-based Approach

Voxel level inferences have the advantage of providing precise localization of activated voxels within a subject. However, to make group inferences under such scheme, both spatial normalization and spatial smoothing are required which in fact reduces spatial resolution [51]. The alternative approach is to draw inference at the ROI level, which involves segmenting out ROIs for each subject individually, and examining the statistical properties of regional activations across subjects. This approach has been shown to offer finer localization and increased sensitivity to task-related effects [51]. However, any chosen regional statistics must account for inter-subject variability in brain shape, brain size, and subject’s orientation in scanner for inference to be meaningful. Towards this end, past studies have used magnitude-based features, such as mean voxel activation statistics and percentage of activated voxels [17], which neglect any spatial information to avoid complications arising from the differences in subject’s orientation in the scanner. The spatial information encoded by the location of voxels within an ROI, however, might be an important attribute of brain activations as suggested by several studies. For example, Thickbroom et
al. [18] showed that the spatial extent of activation, as opposed to response magnitude, was modulated by different levels of force during a sustained finger flexion task. Also, McKeown et al. [16] showed by applying ICA to electroencephalogram (EEG) data that L-Dopa medication appears to have a 'focussing' effect on the cortical activity of PD patients. Furthermore, Haxby et al. demonstrated that brain response can be inferred from the pattern of activity even at the macroscopic scale [19], analogous to the idea of population coding where brain response is encoded by the overall pattern of activity in a set of (spatially-disparate) neurons [73]. Thus, developing tools to quantitatively characterize the spatial distribution of activation statistics will definitely be fruitful. Also, a major issue in ROI-based fMRI analysis is errors in ROI delineation. Conventionally, ROIs are manually segmented with the aid of anatomical atlases. However, any errors introduced by the segmentation process will increase inter-subject variability. In this thesis, we propose new methods to incorporate spatial information into ROI-based fMRI analysis that simultaneously accounts for ROI segmentation errors, as to be discussed in the next section.

1.8 Contributions to ROI-based fMRI Analysis

The contributions of this thesis are three-folds:

1. We demonstrated for the very first time that spatial information encoded by the location of voxels within an ROI can be meaningfully incorporated into ROI-based fMRI analysis.

2. Processing methods to deal with ROI delineation errors, brain shape inter-subject variability, and inclusion of voxels falsely deemed activated were developed.
3. We extended the spatial analysis into the temporal domain, and showed that the spatial distribution of BOLD signal itself is in fact modulated in time by task performance.

The use of spatial descriptors is particularly novel in the field of ROI-based fMRI analysis, since up till now, only magnitude-based features are considered. The invariant nature of the proposed spatial descriptors to similarity transformations (i.e. translations, rotations, and scaling) accounts for inter-subject variability in brain size and subject’s orientation in the scanner, thus enables spatial characterization without resorting to spatial normalization.

In [6], we demonstrated that incorporating spatial information using 3DMIs (as described in detail in Chapter 2 and 3) enhances sensitivity in discriminating task-related activation changes. Our proposed analysis framework also allows one to systematically compare two sets of brain activations. The comparison can be across subject groups (e.g. diseased group versus control group) or within group where the same set of subjects perform two different tasks or perform the same task before and after medication. Hence, our method can prospectively be translated to a wide range of clinical applications such as disease diagnosis, prognosis, and drug evaluation. In fact, the method proposed in [6] and its extension [7] were used to examine the effects of L-Dopa medication on PD patients [4], [5].

With infinitely many 3DMIs, deciding which ones to use is not trivial. In [7], we developed a modified LDA procedure that incorporates leave-one-out cross-validation to deal with this feature selection problem. Furthermore, we modified the 3DMIs by remapping the coordinate space with a Gaussian function. This remapping stresses voxels near the activation centroid over voxels near the ROI boundary, which in effect accounts for both ROI delineation errors and inter-subject
variability in brain shapes. However, all voxels near the ROI boundary are affected regardless of whether they belong to the ROI. To more directly address the problem associated with ROI delineation errors, a processing method based on 3DMIs was proposed [8] that simultaneously refines ROI delineation and spatially denoises the fMRI activation statistics. The underlying idea is that no voxel in isolation should exhibit exceedingly disproportional influence on the 3DMIs. Thus, voxels deemed overly influential are either deweighted or removed. Applying this procedure to real fMRI data demonstrated further increase in sensitivity for discriminating task-related activation differences over using 3DMIs alone.

The results in [6], [7], and [8] all suggest that incorporating spatial information into ROI-based fMRI analysis is beneficial. These analyses were performed on T-maps where the spatial information is collapsed over time, thus only the time-averaged spatial patterns of brain activity were examined. To fully exploit the spatio-temporal structure of fMRI data, we also extended our spatial analysis to the spatio-temporal domain to explore whether the spatial distribution of the BOLD signal itself is modulated in time by task performance [9]. The spatial feature time courses generated by calculating the 3DMIs at each time point, were used to infer ROI activation, in addition to discriminating activation differences as in [6], [7], and [8]. Our results demonstrated that the spatial distribution of BOLD signal is in fact modulated by task performance. The neuroscience implication of this finding is substantial, as this is the first time task-related spatial modulation of brain activation is demonstrated. In all past studies, the traditional belief is that only the magnitude of BOLD signal is modulated by task. This finding can potentially be extended to functional connectivity analysis to more meaningfully summarize the temporal response of an ROI as opposed to averaging the intensity time courses of
all voxels within an ROI.

All our results suggest the importance of incorporating spatial information in ROI-based fMRI analysis. Exploiting the information encoded by voxel locations might hence provide brain researchers a whole new area of research to explore.
Bibliography


2.1 Introduction

Most conventional functional magnetic resonance imaging (fMRI) analysis methods generate statistics to determine the probability that a given voxel is being activated during the performance of the underlying behavioral task(s). To combine fMRI statistics across subjects, there are generally two broad approaches. One involves warping the brains of each individual subject to a common exemplar shape [1]. The other involves drawing the regions of interest (ROIs) of each individual subject, and then examining the statistical properties of activations in the ROIs across subjects. Determining how to summarize the different activation statistics within an ROI is not obvious. Mean value, percentage of activated voxels, and variations of these...

\footnote{A version of this chapter has been published. B. Ng, R. Abugharbieh, X. Huang, and M. J. McKeown, “Characterizing fMRI Activations within Regions of Interest (ROIs) using 3D Moment Invariants,” IEEE Computer Society Workshop on Mathematical Models in Biomedical Image Analysis, New York, NY, June 17-18, 2006, pp. 63 (8 pages).}
two metrics are typically used to represent the level or extent of activation inside a ROI. For instance, Allen and Courchesne [2] used the percentage of activated voxels and mean percent signal changes to study the activation differences between subjects with and without autism. Aizenstien et al. [3] used the mean of the intensity time courses of significantly activated voxels and percentage of voxels with negative T-values to compare the activation distributions of young and elderly subjects. Similarly, Bogorodzki et al. [4] used features extracted from mean time-intensity curves to discriminate between different groups of marijuana smokers. All these discrimination techniques are essentially based on comparing activation statistics amplitudes, and will be referred to as intensity-based methods for the remainder of this chapter. While this allows the results from different subjects to be combined in an uncomplicated manner, the spatial information related to the location of the activated voxels is ignored. This spatial information, however, could provide more discriminative power over traditional approaches. Thus, we propose a method for characterizing the distribution of fMRI activation statistics of each voxel inside an ROI using three-dimensional (3D) shape descriptors based on 3D moment invariants. The invariant nature of any shape descriptor is critical, as it would not be desirable to have any shape information that was dependent upon the particular coordinate system used. A recent study by Mangin et al. [5] demonstrated the discriminative power of 3D moment invariants on structural data. In this thesis, we proposed extending the application of these invariants to functional data. In this chapter, we show that applying the proposed method to fMRI data recorded from healthy subjects performing externally and internally guided tasks reveals significant handedness-related and task-related activation differences in the supplementary motor areas, cerebellum, primary motor cortex, prefrontal cortex, and caudate. In
contrast, many of these activation differences were not detected using conventional intensity-based methods.

2.2 Imaging and Data Pre-processing

2.2.1 Study Subjects and Experimental Conditions

In this study, fMRI data was collected from 8 healthy subjects. SMA, cerebellum, primary motor cortex, prefrontal cortex, and caudate were chosen as the regions of interest. Each patient was asked to perform two motor tasks in 20 s blocks first with their right hand and then with their left hand:

A. In the first task, patients were externally guided, where they had to follow a finger tapping sequence shown on a screen.

B. In the second task, patients were internally guided, where they had to perform a finger tapping sequence that they had seen on a screen earlier.

2.2.2 fMRI Data Acquisition

The fMRI data were acquired on a 3.0 Tesla Siemens scanner (Siemens, Erlangen, Germany) with a birdcage type standard quadrature head coil and an advanced nuclear magnetic resonance echoplanar system. The participant’s head was positioned along the canthomeatal line. Foam padding was used to limit head motion within the coil. High-resolution T1 weighted anatomical images (3D SPGR, TR=14ms, TE=7700 ms, flip angle=25°, voxel dimensions 1.0 mm×1.0 mm×1.0 mm, 176×256 voxels, 160 slices were acquired for co-registration and normalization of functional images. A total of 49 co-planar functional images were acquired using a gradient
echoplanar sequence (TR=3000 ms, TE=30 ms, flip angle=80°, NEX=1, voxel dimensions 3.0 mm×3.0 mm×3.0 mm, imaging matrix 64×64 voxels). Each functional run consisted of 128 time points, and two radio frequency excitations were performed prior to image acquisition to achieve steady-state transverse relaxation.

2.2.3 fMRI Pre-Processing

The fMRI data of each individual was pre-processed independently for motion correction, smoothing, and time realignment using Statistical Parametric Mapping (SPM-99) [6]. The time series of functional images were aligned for each slice in order to minimize the signal changes related to small motion of the subject during the acquisition. Spatial filtering of functional time series was performed by convolving each EPI image with a two-dimensional (2D) Gaussian smoothing kernel with full width at half maximum (FWHM) of 2.8 mm×2.8 mm. Temporal filtering of functional time series included removal of the linear drifts of the signal with respect to time from each voxel’s time-course and low-pass filtering of each voxel’s time-course with a one-dimensional (1D) Gaussian filter with FWHM of 6 s. After preprocessing, a voxel-based T-statistical map was generated for each individual by correlating the time course of each voxel in the epochs of interest with an empirically determined hemodynamic response (HDR) obtained from [7].

2.3 Methods

2.3.1 Framework of Proposed Method

The overarching goal of the method is to provide sensitive linear discriminant analysis for fMRI activation statistics within a given ROI. This may either be across
subjects, where the same ROI is compared across subjects belonging to different
groups (e.g. for the ROI, “right cerebellar hemisphere”, “disease group” is com-
pared to “control group”), or may represent an ROI in the same group of subjects
across different tasks (e.g. for the ROI, “right cerebellar hemisphere”, the activ­
ations statistics for “internally guided task” are compared to “externally guided
task”). For simplicity, in the remainder of this chapter we will simply refer to both
these situations as comparing “Group A” to “Group B”.

We first assume that T-statistics associated with activation have already been
calculated using, e.g. the General Linear Model as utilized in SPM, and that “acti­
vated” voxels have been determined by finding those voxels exceeding an arbitrary
threshold (e.g. 1.96).

Fig. 2.1 summarizes the framework of the proposed method. The 3D moment
invariants of the activated-voxel statistics for each subject/condition are then col­
clected into two groups. The invariants across all groups are then dimension-reduced
and projected onto a space with maximum discrimination using linear discriminant
analysis (LDA). A permutation test is then applied to the resulting feature vectors
to estimate the level of significance (p-value) of the null hypothesis that the activa­
tion distributions of the two groups are the same. The implementation details of
each module are described in Sections 2.3.2 to 2.3.4.

2.3.2 3D Moment Invariants

The 3D moments of a density function $\rho(x,y,z)$, (representing functional activation
levels in the present context), are defined as:

$$m_{pqr} = \int_{-\infty}^{+\infty} \int_{-\infty}^{+\infty} \int_{-\infty}^{+\infty} x^p y^q z^r \rho(x,y,z) \, dx \, dy \, dz$$

(2.1)
where $n = p+q+r$, is the order of the moment, $(x,y,z)$ are the coordinates of each voxel, and $\rho(x,y,z)$ are the T-statistics representing the level of activation of each voxel inside the ROI. Like the one dimensional (1D) case where the moments describe the shape of the density function, 3D moments, similarly, provide the general 3D shape information of $\rho(x,y,z)$. However, potential spatial misalignments in the T-maps would be problematic to the analysis if the 3D moments were naively applied directly, as the misalignments would be inappropriately interpreted as shape changes. Therefore, metrics invariant to rotational, translational, and scaling artifacts are required. In this study, 3D moment invariants were used to ensure that only true shape differences in the activation are captured, independent to the particular co-ordinate system used.

Translational invariance can be obtained by using central moments defined as:

$$
\mu_{pqr} = \int_{-\infty}^{+\infty} \int_{-\infty}^{+\infty} \int_{-\infty}^{+\infty} (x - \bar{x})^p(y - \bar{y})^q(z - \bar{z})^r \cdot \rho(x,y,z) \, dx \, dy \, dz
$$

where $\bar{x}$, $\bar{y}$, and $\bar{z}$ are the centroid co-ordinates of the density function, calculated
as:

\[
\bar{x} = \frac{m_{100}}{m_{000}}, \quad \bar{y} = \frac{m_{010}}{m_{000}}, \quad \bar{z} = \frac{m_{001}}{m_{000}}.
\]  

(2.3)

Scale invariance can be obtained by normalizing the moments as follows:

\[
\eta_{pqr} = \frac{m_{pqr}}{m_{000}^{\frac{1}{2}} + 1}
\]  

(2.4)

To obtain rotational invariance, the 3D moments need to be summed in a certain fashion as given in (2.5) to (2.7) and (2.13) to (2.16).

Seven invariants were used in this study, three of which are based on 2\textsuperscript{nd} order moments derived by Sadjadi and Hall [8]:

\[
J_1 = m_{200} + m_{020} + m_{002}
\]  

(2.5)

\[
J_2 = m_{200}m_{020} + m_{200}m_{002} + m_{020}m_{002} - m_{101}^2 - m_{110}^2 - m_{011}^2
\]  

(2.6)

\[
J_3 = m_{200}m_{020}m_{002} - m_{002}m_{110}^2 + 2m_{110}m_{101}m_{011} - m_{020}m_{101}^2 - m_{200}m_{011}^2
\]  

(2.7)

The remaining four higher order invariants used are based on moment tensor contraction [9].

Considering only affine transforms, general tensors become Cartesian tensors, thus covariant and contravariant tensors are equal [10], meaning if \( a_{ij} \) is a Cartesian tensor, then:

\[
a_{ij} = a_{ij}^T = a_{ji}.
\]  

(2.8)

By contracting \( a_{lmn}, a_{mnop}, a_{ijklm}, \) and \( a_{opqr}^{ijklmn} \), 3\textsuperscript{rd}, 4\textsuperscript{th}, 5\textsuperscript{th}, and 6\textsuperscript{th} order 3D moment invariants were derived. Below we detail the algorithm for generating a 3\textsuperscript{rd} order 3D moment invariant based on moment tensor contraction:

Let \( M_{lmm} \) be a 3\textsuperscript{rd} order moment tensor as defined in [11], where \( l, m, n = 1, 2, 3 \), since 3D moment invariants were pursued. We can obtain a 3\textsuperscript{rd} order 3D
moment invariant, $MI_{3D}$, by contracting $a_{lmn}^{ijk}$:

$$a_{lmn}^{ijk} = a^{ijklmn} = M_{ij}^{lm} M_{mn}^{ijkl} = M_{jkn}^{lmn} = \Sigma_{i=1}^{N} \alpha_i \cdot m_{piqr}^{2} = MI_{3D}, \quad (2.9)$$

where $N = (order + 1) \cdot \frac{2+order}{2} = (3 + 1) \cdot \frac{2+3}{2} = 10$ and $p_i, q_i, r_i$ can be calculated as follows:

$$p_i(l, m, n) = \delta(l - 1) + \delta(m - 1) + \delta(n - 1), \quad (2.10)$$

$$q_i(l, m, n) = \delta(l - 2) + \delta(m - 2) + \delta(n - 2), \quad (2.11)$$

$$r_i(l, m, n) = \delta(l - 3) + \delta(m - 3) + \delta(n - 3), \quad (2.12)$$

$\alpha_i$ is calculated by counting the number of times the same pattern of $p_i, q_i, r_i$ is generated by the above algorithm. For example, $\alpha_1$ corresponding to $m_{300}$ equals to 1 since in only the case when $l = 1, m = 1,$ and $n = 1$ can the pattern $\{3, 0, 0\}$ be generated. The derived higher order moment invariants used in this study are as follows:

$$B_3 = m_{300}^2 + m_{030}^2 + m_{003}^2 + 3m_{210}^2 + 3m_{201}^2 + 6m_{120}^2 + 6m_{111}^2 \quad (2.13)$$

$$+ 3m_{102}^2 + 3m_{021}^2 + 3m_{012}^2$$

$$B_4 = m_{400}^2 + m_{040}^2 + m_{004}^2 + 4m_{310}^2 + 4m_{301}^2 + 6m_{220}^2$$

$$+ 12m_{211}^2 + 6m_{202}^2 + 4m_{210}^2 + 12m_{121}^2 + 12m_{112}^2 + 4m_{103}^2 \quad (2.14)$$

$$+ 4m_{031}^2 + 6m_{022}^2 + 4m_{013}^2$$

$$B_5 = m_{500}^2 + m_{050}^2 + m_{005}^2 + 5m_{410}^2 + 5m_{401}^2 + 10m_{320}^2$$

$$+ 20m_{311}^2 + 10m_{302}^2 + 10m_{230}^2 + 30m_{221}^2 + 30m_{212}^2 + 10m_{203}^2$$

$$+ 5m_{140}^2 + 20m_{131}^2 + 30m_{122}^2 + 20m_{113}^2 + 5m_{104}^2 + 5m_{041}^2$$

$$+ 10m_{032}^2 + 10m_{023}^2 + 5m_{014}^2 \quad (2.15)$$
\[ B_6 = m_{600}^2 + m_{660}^2 + m_{006}^2 + 6m_{510}^2 + 6m_{501}^2 + 15m_{420}^2 \]

\[ + 30m_{411}^2 + 15m_{302}^2 + 20m_{330}^2 + 60m_{321}^2 + 60m_{312}^2 + 20m_{303}^2 \]

\[ + 15m_{240}^2 + 60m_{231}^2 + 90m_{222}^2 + 60m_{213}^2 + 15m_{204}^2 + 6m_{150}^2 \]

\[ + 30m_{141}^2 + 60m_{132}^2 + 60m_{123}^2 + 30m_{114}^2 + 6m_{105}^2 + 6m_{051}^2 \]

\[ + 15m_{042}^2 + 20m_{033}^2 + 15m_{024}^2 + 6m_{015}^2 \] (2.16)

Other higher order invariants can be derived using similar procedures to increase the number of features which might enhance discriminative power of the feature vector. However, the chosen set of 3D moment invariants provided adequate shape discriminative capability for the purposes of this study, thus higher order invariants were not used.

### 2.3.3 Linear Discriminant Analysis (LDA)

Once the 3D moment invariants of the activation distributions of each subject are calculated, the next step is to test whether those feature vectors (corresponding to the spatial distribution of the activation regions) differ between the two experimental groups. To maximize discriminability, the 3D moment invariants are projected onto a 1D space, where the between group variance is maximized, and the within group variance is minimized. This projection is achieved using LDA [12]. The cost function used in LDA is the generalized Rayleigh quotient defined as:

\[ J(w) = \frac{w^T S_B w}{w^T S_W w}, \]  

(2.17)

where \( w \) is the vector pointing in the direction of maximum discrimination of the data, and \( S_B \) and \( S_W \) are the between-group and within-group scatter matrices defined as:

\[ S_B = (m_1 - m_2)(m_1 - m_2)^T \]  

(2.18)
\[ S_W = \sum_{i=1}^{2} \sum_{x \in \Omega_i} (x - m_i)(x - m_i)^t, \quad (2.19) \]

where \( m_i \) is a vector containing the mean of each 3D moment invariant of group \( i \), \( x \) is a matrix containing the 3D moment invariants of all subjects, and \( \Omega_i \) denotes group \( i \). The \( w \) that maximizes \( J(w) \) must satisfy:

\[ S_B w = \lambda S_W w \quad (2.20) \]

for some constant \( \lambda \). By transforming \( J(w) \) into a generalized eigenvalue problem, the potential singularity problem associated with \( S_B \) and \( S_W \) (due to limited number of samples) is mitigated. The vector \( w \) can be found by determining the eigenvector corresponding to the maximum eigenvalue of (2.20).

### 2.3.4 Permutation Test

To determine the probability of the activation distributions of the two experimental groups being the same, permutation was used. The permutation test does not require a priori assumptions about the data distribution, and thus is preferred over the T-test and F-test [13]. The procedures of the permutation test algorithm are as follows [14]:

1. Given \( m \) \( n \)-dimensional feature vectors for each group, \( X \) and \( Y \) where \( m > n \), calculate the Mahalanobis distances between each vector in \( X \) and the centroid of \( Y \) as follows:

\[ d_{Mahal}(i) = \sqrt{\left( \frac{x_{ij}^1 - \bar{y}}{s_1} \right)^2 + \cdots + \left( \frac{x_{ij}^m - \bar{y}}{s_m} \right)^2}, \quad (2.21) \]

where \( \bar{y} \) is the centroid of \( Y \), \( x_{ij}^1 \) is the \( j \)th element of the \( i \)th sample feature vector of \( X \), and \( s_k \) is square root of the \( k \)th diagonal element of the pooled covariance matrix of \( X \) and \( Y \).
2. Calculate mean of $d_{\text{Mahal}}(i)$ and denote this mean as $d_{\text{orig}}$.

3. Pool all sample feature vectors, stripped of their group labels.

4. Randomly draw $m$ samples from a random distribution and label these samples $X'$ and label the remaining samples $Y'$.

5. If the mean $d_{\text{Mahal}}(i)$ between $X'$ and $Y'$ is $> d_{\text{orig}}$ increment counter $q$ by 1.

6. Repeat steps 3 to 6 $(N-1)$ times (ex. $N = 5000$).

7. The $p$-value representing the significance of the differences between $X$ and $Y$ is estimated by $q/N$.

2.3.5 Visualization of Activation Patterns

Since the voxel distributions are 3D, the T-statistics of each voxel can only be represented by color or intensity. However, plotting the T-statistics in this form does not provide much insight into the spatial distribution of the activation, since shapes in the 4th dimension are hard to visualize. Therefore, a different approach is taken.

Projecting the 3D structural data onto a 2D plane through PCA and using the T-statistics as the 3rd dimension, surface plots of the activation distribution were generated through cubic interpolation. Note that the first two principle components were used in generating these surface plots to ensure that most of the structural information is retained Fig. 2.2.

2.3.6 Comparison of ROI Analysis Methods

Given the T-values of each voxel inside an ROI, metrics for representing the characteristics of the activation within the ROI are required so that comparisons can be
Figure 2.2: Methods used to compare activation statistics across ROIs. The proposed method captures the rich shape information of the activated regions using 3D moment invariants. Note that irrespective of the orientation of the activation (as demonstrated by the 2 different perspectives shown) the same 3D Moment Invariants will be generated. Conventional methods (right) collapse the spatial information into a single metric such as the mean thresholded T-statistic (upper right) or percentage of activated voxels (lower right), ignoring the texture of the activation, and potentially reducing discriminability across subjects.
made between subjects and across groups under different experimental conditions. Conventional methods involve using mean T-values and percentage of activated voxels as metric, whereas we propose incorporating spatial information of the activated voxels. Methods based on these three metrics were compared for detecting the activation differences during externally and internally guided tasks:

- **Thresholded Mean**: Taking T-values greater than 1.96, compute the mean T-values during externally and internally guided tasks. Calculate the difference between the means of these two conditions, and apply T-test to determine if the differences are significant.

- **% Activated Voxels**: Compute the percentage of voxels with T-values greater than 1.96 during externally and internally guided tasks. Calculate the difference between the percentages of these two conditions, and apply T-test to the differences to determine level of significance.

- **3D Moment Invariants**: Proposed method as described in Section 2.3.1.

The first two methods are intensity-based, whereas the 3D Moment Invariant method incorporates the extensive spatial information of the activated voxels.

In this study, 10 ROIs were examined, consisting of the left and right supplementary motor areas (SMAs), cerebellar hemispheres, primary motor cortices, prefrontal cortices, and caudates. The activation distributions of these ROIs during externally versus internally guided finger sequential movements were compared.

### 2.4 Results and Discussion

The results generated using the three methods described in Section 2.3.5 are summarized in Table 2.1. As suggested by Fig 2.2, conventional methods, by collapsing the
Table 2.1: p-values of Activation Differences During External versus Internal Task

<table>
<thead>
<tr>
<th>ROIs</th>
<th>Experimental Conditions</th>
<th>Right Hand</th>
<th></th>
<th>Left Hand</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Thresholded</td>
<td>% Activated</td>
<td>Thresholded</td>
<td>% Activated</td>
</tr>
<tr>
<td>Left SMA</td>
<td></td>
<td>0.0466</td>
<td>0.2134</td>
<td>0.0352*</td>
<td>0.0328*</td>
</tr>
<tr>
<td>Right SMA</td>
<td></td>
<td>0.2955</td>
<td>0.0880</td>
<td>0.0796</td>
<td>0.0454*</td>
</tr>
<tr>
<td>Left CER</td>
<td></td>
<td>0.5853</td>
<td>0.7093</td>
<td>0.0158*</td>
<td>0.0828</td>
</tr>
<tr>
<td>Right CER</td>
<td></td>
<td>0.7510</td>
<td>0.8793</td>
<td>0.0174*</td>
<td>0.1280</td>
</tr>
<tr>
<td>Left PMC</td>
<td></td>
<td>0.1496</td>
<td>0.5681</td>
<td>0.3188</td>
<td>0.4301</td>
</tr>
<tr>
<td>Right PMC</td>
<td></td>
<td>0.6656</td>
<td>0.9110</td>
<td>0.1076</td>
<td>0.1027</td>
</tr>
<tr>
<td>Left PFC</td>
<td></td>
<td>0.8641</td>
<td>0.7796</td>
<td>0.0334*</td>
<td>0.2537</td>
</tr>
<tr>
<td>Right PFC</td>
<td></td>
<td>0.7140</td>
<td>0.0741</td>
<td>0.0328*</td>
<td>0.5955</td>
</tr>
<tr>
<td>Left CAU</td>
<td></td>
<td>0.1423</td>
<td>0.0469*</td>
<td>0.1014</td>
<td>0.0440*</td>
</tr>
<tr>
<td>Right CAU</td>
<td></td>
<td>0.6369</td>
<td>0.0757</td>
<td>0.0048*</td>
<td>0.2358</td>
</tr>
</tbody>
</table>

CER = cerebellum, PMC = primary motor cortex, PFC = prefrontal cortex, and CAU = caudate. *Statistically significant at $\alpha = 0.05$.

T-statistics within an ROI into a single value has the advantage of being invariant to the co-ordinate system used and enables values across subjects/conditions to be combined in a straightforward manner. Nevertheless, these advantages are mitigated by the loss of potentially useful spatial information. Unless the spatial information – in effect, the 3D texture – of an ROI is utilized, the changes in T-statistics during externally and internally guided tasks might be very difficult to detect using only intensity information (Fig. 2.2, right). On the other hand, examining the shapes of the activation distributions from different angles reveals that the patterns of activation are substantially different during externally and internally guided tasks. Thus, exploiting this spatial information may provide higher sensitivity to activation differences, provided these differences are not subject-specific.

A key result of this study is that the enhanced sensitivity of the 3D moments does not necessarily amplify inter-subject differences to the same extent. We found consistent, significant differences in the spatial distribution of ROI-based activation statistics even when combining across subjects (Table 2.1). This suggests that some
spatial aspects of fMRI activation within ROIs are modulated by the task performed, and do not simply reflect inter-subject variability.

There is increasing interest in the role of BOLD signal decreases reflecting neuronal de-activations (e.g. [15]). Although not examined in the current study, we note that the proposed method can easily incorporate negative T-values, by concatenating 3D moment invariants associated with the negative T-values into the feature vector for classification.

2.4.1 Supplementary Motor Areas

Both the thresholded mean method and the 3D invariant method detected activation differences between externally and internally-guided tasks in the left SMA when the dominant hand was used. Since the SMA is known to be preferentially invoked during internally guided tasks [16] [17] [18], detecting activation differences in the left SMA is expected, based on prior neuroscience knowledge, when the task involves the right hand.

During the left handed task, all three methods detected activation differences in the right SMA. This again matches expectation since the SMA is more involved during internally guided tasks, and the left hand is used. However, the thresholded mean T-statistic also detected significant activation differences in the left SMA. This might have arisen from connections between the left and right SMA, but requires connectivity analysis to verify.

2.4.2 Cerebellum

Only the proposed method was able to detect significant activation differences in the cerebellum during the right handed task. This activation difference might be
associated with handedness as seen in past studies [19].

2.4.3 Primary Motor Cortex

Only the 3D moment invariant method detected differences using the non-dominant hand. As recent evidence has suggested distinct mechanisms are utilized with the dominant and non-dominant hand [20], [21], we suggest that the different activation between tasks with the non-dominant hand is expected.

2.4.4 Prefrontal Cortex

Again, only the proposed method was able to detect significant activation differences in both prefrontal cortices during the right handed task and in the left prefrontal cortex during the left handed task. Fig. 2.2 suggests why only the proposed method detected activation differences (i.e. the mean T-value and percentage of activated voxels differ by a minuscule amount during the two tasks, whereas substantial changes in the shapes of the activation distributions are shown in the Fig. 2.2). Prefrontal cortex has been previously shown to be differentially activated during externally guided and internally guided tasks [22].

2.4.5 Caudate

Both intensity-based methods detected activation differences in the left caudate during the left handed task. In addition, percentage activated voxel method detected activation differences in the left caudate during the right handed task. 3D moment invariant method, on the other hand, detected no activation differences in the left caudate, but detected activation differences in the right caudate in both tasks. Since the right caudate is associated with timing information [23] and the load placed on
the right caudate to process the timing information during the two tasks are quite different, the detected activation differences is expected.

2.5 Conclusion

We have demonstrated that certain spatial aspects of activation statistics appear to be relatively well conserved across subjects yet are significantly modulated by task performance. Incorporating spatial information of activation statistics within an ROI by utilizing 3D moment invariants appears to complement traditional activation based methods. A direct extension of the methods is to look at the spatial characteristics of combinations of ROIs that maximally discriminate between groups, an approach currently being pursued.
Bibliography


Chapter 3

Feature Selection with Modified LDA

3.1 Introduction

One main application of functional magnetic resonance imaging (fMRI) is to make statistical inferences about regions of the brain responsible for different behavioral tasks. Typically, this entails assigning a statistical value to each voxel related to its probability of being activated during the performance of the task with all voxel statistics assembled into a statistical parametric map. The individually-derived fMRI statistical maps are then either compared across subject groups or across different tasks to draw statistical inferences about group activations. Due to inter-subject variability in brain shape, brain size, and orientation of the subjects in the MR scanner, combining fMRI statistics across subjects to make group inferences is

\footnote{A version of this chapter is under revision. B. Ng, R. Abugharbieh, X. Huang, and M. J. McKeown, “Characterization of fMRI Activation Maps Using Invariant 3D Moment Descriptors,” IEEE Transactions on Medical Imaging, invited paper (under revision).}
not trivial. To this end, two broad approaches have been traditionally employed. The most common approach involves warping the brain of each individual subject to a common exemplar template [1]. However, spatial normalization, typically followed by spatial smoothing, may inappropriately pool responses from functionally dissimilar regions [2], especially for small, subcortical structures, thus degrading important spatial information. The alternative approach to whole brain warping involves drawing the regions of interest (ROIs) for each individual subject, and examining the statistical properties of regional activations across subjects within those ROIs. Castanon et al. demonstrated that this latter approach offers finer localization and increased sensitivity to task-related effects [2], thus this subject-specific ROI-based approach is adopted in this thesis.

Defining a metric to rigorously characterize and compare ROI activation maps across groups of subjects without merely amplifying inter-subject variability is also very challenging. Mean activation statistics, percentage of activated voxels, and mean voxel time course averaged over an ROI have all been previously used [3]. For example, Allen and Courchesne used the percentage of activated voxels and mean percent signal changes to study the activation differences between subjects with and without autism [4]. Aizenstin et al. used the mean intensity time courses of the significantly activated voxels and the percentage of voxels with negative T-values to compare activation distributions of young and elderly subjects [5]. Similarly, Bogorodzki et al. used features extracted from mean intensity time curves to discriminate between different groups of marijuana smokers [6]. All these group discrimination techniques are essentially based on comparing the magnitudes of the activation statistics with the underlying assumption that the spatial distribution of voxel activation statistics within an ROI is not meaningful. These features will
thus hereafter be referred to as ‘magnitude-based’ features in the remainder of this chapter.

Characterizing the activation statistics within an ROI using magnitude-based features provides a simple means for comparing activation maps across different subjects, since these methods are insensitive to variability in brain shape, brain size, subjects’ orientation inside the MR scanner, and location of the activated voxels. However, we have previously demonstrated that exploiting the spatial information encoded by the location of the activated voxels within an ROI can significantly increase discriminative power [7]. We note an interesting parallel between magnitude-based features, which ignore the spatial distribution of activation statistics within an ROI, and neural coding methods in the brain. At the neuronal level, the mean firing rate (known as “rate coding”) is only one of the many possible ways in which a particular brain response might be encoded. Other models, such as “population coding”, suggest that the brain response could be inferred from the overall pattern of activity in a set of (spatially-disparate) neurons [8], where this population coding may also be present at the macroscopic scale of neuroimaging [9]. Similarly, instead of only looking at mean changes in the blood-oxygenation level dependent (BOLD) signal averaged over an ROI, it may be beneficial to also explore the spatial distribution changes of BOLD signals during different task-performances. This is because the spatial distribution of activation statistics within an ROI may actually be altered by task performance even when no change in mean activation statistics or percentage of activated voxels is present as illustrated in Fig. 3.1. In this thesis, we propose a novel method for characterizing the spatial distribution of fMRI activation statistics within an ROI using three dimensional moment invariants (3DMIs). The invariance properties of the proposed descriptors are critical in ensuring that
Figure 3.1: Simplified synthetic example of ROI activation maps illustrating the underlying idea of the proposed method. Assuming 100 voxels are within the ROI and only 3 voxels are active in both conditions. Mean activation statistics = 6/100 and percentage of activation = 3% are the same for both conditions A and B. However, these two tasks are spatially encoded quite differently. This difference in the spatial distribution of activation statistics could be characterized using 3D moment invariants as proposed in this thesis to better discriminate changes in activation patterns between the two conditions.
the extracted spatial features are independent of the subject's orientation in the MR scanner. Also, using spatially invariant features accounts for some of the intersubject variability such as differences in brain sizes. A recent study by Mangin et al. demonstrated the discriminative power of 3DMIs on structural MRI data [10]. In this thesis, we extend the use of such invariant descriptors to functional MRI data.

Our proposed method is applied to real fMRI data recorded from twenty one healthy subjects performing externally and internally-guided motor tasks. The actual motor task in both cases is the same (see Section 3.3), but the brain processes involved are different depending on whether the subject is mimicking a visual cue or recalling the movement from memory. This relatively minor but nonetheless crucial distinction makes such a task ideal for testing the sensitivity of different activation discriminant analysis methods. Using 3DMIs, we demonstrate significant task-related activation differences in the left lentiform nucleus, right cerebellar hemisphere, left supplementary motor areas (SMA), left primary motor cortex (M1), and left prefrontal cortex (PFC). In contrast, percentage of activated voxels did not detect any significant task-related activation differences, and thresholded mean activation statistics only detected activation differences in the SMAs. These results suggest that different tasks might be encoded by the 3D 'spatial texture' of the activation statistics within an ROI in addition to the magnitude and extent of activation.
3.2 Materials

3.2.1 Study Subjects and Experimental Conditions

All research was approved by the appropriate Institutional and Ethics Review Boards. After obtaining informed consent, twenty one healthy subjects were asked to perform two motor tasks in 20 s blocks with their right hand while in the fMRI scanner:

- In the first task, subjects were externally guided (EG), following a finger tapping sequence displayed as a movie of a hand on a screen.

- In the second task, subjects were internally guided (IG), performing a finger tapping sequence they had seen on a screen earlier during the first task, but recalling the sequence from memory.

3.2.2 Data Acquisition

Images were acquired on a 3.0 Tesla Siemens scanner (Siemens, Erlangern, Germany) with a birdcage-type standard quadrature head coil and an advanced nuclear magnetic resonance echoplanar system. The participant’s head was positioned along the canthomeatal line. Foam padding was used to limit head motion within the coil. High-resolution T1-weighted anatomical images (3D SPGR, TR=14 ms, TE=7700 ms, flip angle=25°, voxel dimensions 1.0×1.0×1.0 mm, 176×256 voxels, 160 slices) were acquired for co-registration with the functional images. Co-planar T2*-weighted functional images were acquired at each time point using a gradient echoplanar sequence (TR=3000 ms, TE=30 ms, flip angle=80°, NEX=1, voxel dimensions 3.0×3.0×3.0 mm, imaging matrix 64×64 voxels, 49 slices). Each functional run consisted of 128 time points, and two radio frequency excitations were performed prior to image acquisition to achieve steady-state transverse relaxation.
3.2.3 fMRI Pre-Processing and Statistical Map Generation

The fMRI data was preprocessed for each subject independently for slice timing and motion correction using the standard parametric mapping software (SPM 2) [11]. Spatial filtering of functional time series was performed by convolving each EPI image with a two-dimensional Gaussian smoothing kernel with full width at half maximum (FWHM) of 2.8 mm×2.8 mm. Temporal filtering of functional time series included removal of the linear drifts of the signal with respect to time from each voxel’s time course, and low-pass filtering of each voxel’s time course with a one dimensional (1D) Gaussian filter with a FWHM of 6 s. No spatial normalization (brain warping) was performed.

After preprocessing, each time point in all runs was coded as a particular type of task (such as rest, right hand EG finger sequence movements (FSM), etc.). To emphasize the proposed spatial analysis, we chose to use the simplest method in generating the statistical parametric maps: a simple T-test applied to each voxel based on the BOLD signal changes in all runs between two tasks: 1) right hand EG FSM versus rest and 2) right hand IG FSM versus rest. The resulting voxel activation statistics were then mapped onto the anatomical ROIs, which were manually drawn by a trained observer according to standard atlases [12]. Regions examined included bilateral caudate, lentiform nucleus, thalamus, SMA, M1, PFC, and cerebellar hemisphere. More details about the ROI specification procedures can be found in [13].
3.3 Methods

3.3.1 Proposed Framework for Spatial fMRI Analysis

The overarching goal of the proposed method is to provide sensitive discriminant analysis of fMRI activation statistics within an ROI. This may either be across subjects, where the same ROI is compared across subjects belonging to different groups (e.g. for the ‘right cerebellar hemisphere’ ROI, all subjects in a ‘disease group’ are compared to all subjects in a ‘control group’), or may be within the same group of subjects across different tasks (e.g. for the ‘right cerebellar hemisphere’ ROI, the activations statistics for the IG task are compared to that of the EG task for all subjects). For simplicity, in the remainder of this chapter, we will simply refer to both situations as comparing ‘Group A’ to ‘Group B’.

Fig. 3.2 summarizes the framework of the proposed method. Assuming that activation statistics have been calculated using e.g. the General Linear Model
(GLM) as utilized in SPM, features characterizing the activation statistics within an ROI are first extracted for each subject and collected into two groups. To enhance robustness to potential outliers, a winsorization procedure is performed to generate estimates of the feature values. For the case of single ROI feature analysis using traditional magnitude-based features, linear discriminant analysis (LDA) is not necessary since in a 1D feature space, there is no alternative axis orientation for increasing discriminability. Instead, a permutation test is directly applied to the single ROI features to estimate the probability (p-value) of the null hypothesis that the activation maps of the two groups are drawn from the same distribution. For the case of multiple ROI feature analysis using the proposed spatial features, LDA is first applied to all subjects' ROI features to determine the relative weightings of these features in the maximal discriminant 1D space. These weightings are then used to sort the features in decreasing level of discriminability. A modified LDA procedure that incorporates leave-one-out cross-validation is then performed to choose the number of features to include in the analysis without over-fitting. A permutation test is then applied to the resulting feature vectors to estimate the probability that the activation maps are from the same distribution.

3.3.2 ROI Spatial Feature Extraction: 3D Moment Invariants

Two types of ROI features are contrasted in this thesis. One is based on activation statistics magnitude, whereas the other is based on the spatial distribution ('3D spatial texture') of activation statistics within an ROI. More specifically, we have chosen to compare two magnitude-based features, thresholded mean activation statistics and percentage of activated voxels, which are widely-used in ROI-based fMRI analysis [3], against the proposed spatial features, namely 3DMIs. The thresh-
Olded mean activation statistics are calculated by averaging the voxel activation statistics (or T-statistics as used in this thesis) greater than a particular threshold, \( \tau \), over an ROI. The percentage of activated voxels is defined as the number of voxels with T-values above \( \tau \) divided by the total number of voxels within the ROI. We have tested a range of \( \tau \) from 1 to 4 at 0.5 increments. The ROIs detected with significant activation differences were consistent for this range of thresholds, thus only results generated with \( \tau = 1.96 \), which corresponds to an uncorrected p-value of 0.05, are included for comparisons with the proposed 3DMI-based method. The 3D spatial features we employed (3DMIs) are calculated based on 3D moments defined as:

\[
m_{pqr} = \int_{-\infty}^{+\infty} \int_{-\infty}^{+\infty} \int_{-\infty}^{+\infty} x^p y^q z^r \rho(x, y, z) dx dy dz,
\]

where \( n = p + q + r \) is the order of the 3D moment, \( (x, y, z) \) are the coordinates of a voxel, and \( \rho(x, y, z) \) is the T-value of a voxel located at \( (x, y, z) \) within an ROI. We have restricted our analysis to positive T-values, since there is little consensus on the interpretation of negative T-values [14]. As for the choice of T-threshold, we have decided to include all voxels with T-values greater than zero. One might argue that using a T-threshold of zero will include many non-active voxels that are falsely deemed active (false positives) into the analysis. However, false positives will not likely be located at the same position within an ROI across subjects. Hence, inclusion of these voxels will only reduce discriminability of our spatial features. On the other hand, truly active voxels that are corrupted by noise may reside in similar location within an ROI across subjects. If a high T-threshold is set, some of these active voxels will be excluded from the analysis. Therefore, we argue that using a T-threshold of zero provides a suitable, conservative estimate of activation differences, while retaining most of the original information of the ROIs, and is thus
appropriate for our spatial activation statistics analysis.

Naively using 3D moments in discriminating activation statistics is problematic, since potential spatial misalignments of the T-maps would be inappropriately interpreted as spatial changes. Therefore, metrics invariant to similarity transformations (rotation, translation, and scaling) are needed. Furthermore, by using spatially invariant features, inter-subject variability in brain size or slight translational/rotational shifts of the activated region would be accounted for. In this study, 3DMIs are used to ensure that only true spatial differences in activation statistics are captured in the fMRI data analysis, i.e. independent of the subjects' orientation in the MR scanner and brain size.

Translational invariance of 3D moments can be obtained by using central moments defined as:

$$\mu_{pqr} = \int_{-\infty}^{+\infty} \int_{-\infty}^{+\infty} \int_{-\infty}^{+\infty} (x - \bar{x})^p (y - \bar{y})^q (z - \bar{z})^r \cdot \rho(x, y, z) dx dy dz \quad (3.2)$$

where $\bar{x}$, $\bar{y}$, and $\bar{z}$ are the centroid co-ordinates of $\rho(x, y, z)$, calculated as:

$$\bar{x} = \frac{m_{100}}{m_{000}}, \quad \bar{y} = \frac{m_{010}}{m_{000}}, \quad \bar{z} = \frac{m_{001}}{m_{000}}. \quad (3.3)$$

Scale invariance can be obtained by normalizing the moments as follows:

$$v_{pqr} = \frac{m_{pqr}}{m_{000}^{\frac{p+q+r}{2}}}. \quad (3.4)$$

Examining (3.2), if we consider the distance from the centroid (e.g. $(x - \bar{x})$) as a weighting of the T-values $\rho(x, y, z)$, the T-values of voxels closer to the ROI boundary will be more highly weighted. However, some of the outer voxels may not belong to the ROI, but are included due to manual segmentation errors. Also, T-values of voxels near the activation centroid are less prone to noise as higher signal strength is expected at those voxel locations. Thus, to mitigate manual ROI segmentation
errors and to stress voxels near the activation centroid (i.e. to ensure errors in T-values of voxels near the ROI boundary are not given higher weighting), the following remapping is performed:

\[
\begin{align*}
x_g &= \frac{1}{\sigma_x \sqrt{2\pi}} \exp\left(-\frac{(x - \bar{x})^2}{2\sigma_x^2}\right), \\
y_g &= \frac{1}{\sigma_y \sqrt{2\pi}} \exp\left(-\frac{(y - \bar{y})^2}{2\sigma_y^2}\right), \\
z_g &= \frac{1}{\sigma_z \sqrt{2\pi}} \exp\left(-\frac{(z - \bar{z})^2}{2\sigma_z^2}\right),
\end{align*}
\]  

where \((x - \bar{x})\), \((y - \bar{y})\), and \((z - \bar{z})\) in (3.2) are substituted by \(x_g\), \(y_g\), and \(z_g\), and \(\sigma_x\), \(\sigma_y\), and \(\sigma_z\) are set as 2 times the size of the ROI along each direction to account for inter-subject variability in brain sizes. Values lower than 2 can be used to further emphasize the voxels near the centroid and vice versa, but we empirically found that a value of 2 deweights the outer voxels without totally neglecting them. Also, since this remapping bounds \(x_g\), \(y_g\), and \(z_g\) between 0 and 1, which accounts for inter-subject variability in brain sizes, the traditional way for obtaining scale invariance (3.4) is no longer required. We denote the centralized 3D moments with \((x - \bar{x})\), \((y - \bar{y})\), and \((z - \bar{z})\) in (3.2) replaced by \(x_g\), \(y_g\), and \(z_g\) as \(\eta_{pqr}\).

To obtain rotational invariance, the 3D moments need to be combined in specific ways. Sixteen 3DMIs are examined in this study. Three of which are based on 2nd order moments derived by Sadjjadi and Hall [15]:

\[
J_1 = \eta_{200} + \eta_{020} + \eta_{002}
\]  

\[
J_2 = \eta_{200}\eta_{020} + \eta_{200}\eta_{002} + \eta_{020}\eta_{002} - \eta_{101}^2 - \eta_{110}^2 - \eta_{011}^2
\]  

\[
J_3 = \eta_{200}\eta_{020}\eta_{002} - \eta_{002}\eta_{110}^2 + 2\eta_{110}\eta_{101}\eta_{011} - \eta_{020}\eta_{101}^2 - \eta_{200}\eta_{011}^2
\]  

Another eleven 3DMIs (\(J_{22}^2, J_{222}, J_{33}^2, J_{111}, J_{233}, J_{123}, J_{112}, J_{333}, J_{133}, J_{1133}, J_{1113}\)) derived by Lo and Don based on complex moments are also used [16]. In addition, we
derived two other higher order 3DMIs ($B_3$ and $B_4$) using moment tensor contraction similar to the approach described in [17] as described next.

Considering only rigid transformations, general tensors become Cartesian tensors, thus covariant and contravariant tensors are equal [18], meaning if $a_{ij}$ is a Cartesian tensor, then:

$$a_{ij} = a^i_j = a^{ij}. \quad (3.9)$$

Thus, by contracting $a^{ijk}_{lmn}$ and $a^{ijkl}_{mnop}$, 3rd and 4th order 3DMIs can be derived as follows:

For the 3rd order case, let $M_{lmn}$ be a 3rd order moment tensor as defined in [19], where each of the subscripts: $l, m, n_i$ goes from 1 to 3 since 3DMIs are pursued. By contracting , we obtain a 3rd order 3DMI, $B_3$:

$$a^{ijk}_{lmn} = a^{ijlmn} = M^{ijk} M^{lmn} = M^{lmn} M^{lmn} = \sum_{i=1}^{N} \alpha_i \cdot m^{p_i q_i r_i} = B_3, \quad (3.10)$$

where $N = \frac{\text{order} + 1}{2} = (3 + 1) \cdot \frac{2+3}{2} = 10$ and $p_i, q_i, r_j$ can be calculated as follows:

$$p_i(l, m, n) = \delta(l - 1) + \delta(m - 1) + \delta(n - 1) \quad (3.11)$$

$$q_i(l, m, n) = \delta(l - 2) + \delta(m - 2) + \delta(n - 2) \quad (3.12)$$

$$r_i(l, m, n) = \delta(l - 3) + \delta(m - 3) + \delta(n - 3) \quad (3.13)$$

where is the Kronecker’s delta function.

$\alpha_i$ is calculated by counting the number of times the same pattern of $p_i, q_i, r_i$ is generated by the above algorithm. For example, $\alpha_1$ corresponding to $m_{300}$ equals to 1 since in only the case when $l = 1, m = 1, n = 1$ can the pattern $\{3, 0, 0\}$
be generated. Based on the above tensor contraction procedures, the resulting 3rd
order 3DMI is defined as below:

\[ B_3 = \eta_{300}^2 + \eta_{030}^2 + \eta_{003}^2 + 3\eta_{1210}^2 + 3\eta_{1201}^2 + 3\eta_{1202}^2 + 3\eta_{1211}^2 + 3\eta_{1021}^2 + 3\eta_{0212}^2 + 3\eta_{0122}^2 \]  
\[ (3.14) \]

\[ B_4 = \eta_{400}^2 + \eta_{040}^2 + \eta_{004}^2 + 4\eta_{1310}^2 + 4\eta_{1301}^2 + 6\eta_{1220}^2 + 12\eta_{1211}^2 + 6\eta_{2020}^2 + 4\eta_{1301}^2 \]
\[ + 12\eta_{1211}^2 + 12\eta_{1122}^2 + 4\eta_{1031}^2 + 4\eta_{0311}^2 + 6\eta_{0222}^2 + 4\eta_{0132}^2 \]  
\[ (3.15) \]

In fact, other higher order invariants can be derived using similar procedures. However,
to ease interpretation, we have decided to restrict our analysis to 3DMIs that are combinations of 2nd, 3rd, and 4th order 3D moments, which corresponds to combinations of spatial variance, skewness, and kurtosis. For example, \( J_1 \) can be interpreted as the sum of spatial variance along the x, y, and z direction, and \( B_3 \) can be interpreted as a non-linear combination of skewness and other spatial descriptive terms. Note also that higher order moments are less robust to noise [20], thus are not used in this study.

3.3.3 Feature Winsorization

Many linear procedures such as LDA are prone to outliers. To enhance robustness, we first winsorized [21] all the feature values for each ROI separately prior to subsequent processing or analysis. The winsorization procedure is as follows: Let \( F \) be a vector with each element, \( F_i \), corresponding to a single ROI feature value (e.g. \( J_1 \)) of the \( i^{th} \) subject in e.g. group A. \( F_i \) of all subjects from both groups are pooled into \( F \). To winsorize \( F \), we first determine which subject(s) are outliers using (3.16):

\[ I_{out} = \text{find}(\mid F - F_{med} \mid > N_{std} \cdot \sigma), \]  
\[ (3.16) \]

where \( I_{out} \) is the set of outlier subjects, \( F_{med} \) is the median of \( F \), and \( N_{std} \) is the number of standard deviations from \( F_{med} \) beyond which \( F_i \) is considered an out-
lier. Assuming $F_i$ is normally-distributed (since LDA is derived assuming the input features are Gaussian), $N_{std}$ is set to 3 to retain 99.7% of the possible $F_i$ values as non-outliers. $\sigma$ is a robust estimate of the standard deviation of $F$ calculated as:

$$\sigma \approx \lambda \cdot \text{median}(|F - F_{med}|), \quad (3.17)$$

where $\lambda$ converts the median of the absolute differences (MAD) of $F$ to standard deviation and is set to 1.483 assuming $F_i$ is normally-distributed [21]. For only the outlier $F_i$'s, we winsorize each of these values using (3.18):

$$F_i^{\text{win}} = F_{med} + \text{sgn}(F_i - F_{med}) \cdot N_{std} \cdot \sigma, \quad i \in I_{out}. \quad (3.18)$$

This winsorization procedure is applied to each ROI and each feature separately. We note that for the data examined in this study, typically only 1 or 2 subjects were detected as outliers for each feature. Also, we have chosen to winsorize as opposed to throwing away the outliers for two main reasons. First, the number of subjects in most fMRI studies is typically quite low (i.e. like 8 or 10), which renders it impractical to eliminate a given subject’s data just because it is an outlier for a particular (single) ROI or feature. However, one might argue that we can always just remove a subject for a particular ROI or feature (i.e. retaining that subject when analyzing the other ROIs and features). The problem is that changing the sample size across ROIs will introduce variability in statistical power, thus fair comparisons across ROIs cannot be made (i.e. not detecting significant differences in activation statistics with a certain ROI could be attributed to reduced number of subjects). Therefore, to avoid complications arising from changes in sample size, we have chosen to winsorize instead of eliminating outliers from further analysis.
3.3.4 Feature Selection: Modified LDA

A major issue in fMRI study is the low number of subjects available for making statistical inferences. If we naively include all described 3DMIs in the analysis, overfitting is likely to occur. Therefore, we developed a modified LDA procedure that incorporates leave-one-out cross-validation for choosing which 3DMIs to include and in validating our results.

Before feature selection can be performed, the 3DMIs must first be sorted based on their overall relative discriminability across ROIs. For every subject and for a particular ROI, we concatenate all the 3DMIs into a single vector, and apply LDA [22] to the resulting feature vectors of the two groups. To mitigate potential singularity problems associated with the within-group scatter matrix, $S_W$, the projection matrix, $W$, is determined by solving the following generalized eigenvalue problem:

$$S_B W = \lambda S_W W,$$

where $S_B$ is the between-group scatter matrix and $\lambda$ is a diagonal matrix with its diagonal elements corresponding to the eigenvalues of the columns of $W$. The projection vector, $w$, that points in the direction of maximal discriminability is then determined by finding the column of $W$ associated with the maximum eigenvalue. Since $|w|$ can be interpreted as the relative weightings of the 3DMIs for which the two groups of feature values are maximally discriminated, the 3DMIs can be ranked in descending level of discriminability based on the magnitude of $|w|$. In this study, 16 3DMIs are examined, so we first rank the 3DMIs from 1 to 16 for each ROI (16 corresponding to the largest $|w|$). We then sum up the ranks of each 3DMI across ROIs and sort the 3DMIs in descending order of their rank sums. The reason for ordering the 3DMIs based on the overall discriminability across ROIs is to ensure
that the same features are used to analyze each ROIs. Else, fair comparisons across ROIs cannot be made.

Once the 3DMIs are sorted in decreasing level of discriminability, we employ a modified LDA procedure that incorporates leave-one-out cross-validation to determine the number of 3DMIs, $K$, to be used in the analysis. The procedure is as follows:

1. Let $k$ be the number of features (ordered in decreasing level of discriminability) to be included in each subject's feature vector, and initialize $k$ to 2.

2. Apply LDA to the two groups of feature vectors, but with the $i^{th}$ subject from both groups excluded, to determine $w$.

3. Project the $i^{th}$ subject's feature vectors onto $w$, and denote the projected features as $F_i^A$ and $F_i^B$.

4. Repeat steps 2 and 3 for every other subject, vertically concatenate the projected features (i.e. $F_i^A$ and $F_i^B$), and denote the resulting column vectors as $F_i^A$ and $F_i^B$.

5. Calculate the Mahalanobis distance between $F_i^A$ and $F_i^B$ using (3.20):

$$d_{Mahal}(K) = \sqrt{d \cdot \Sigma^{-1} \cdot d},$$  \hspace{1cm} (3.20)

$$d = mean(F^A - F^B),$$  \hspace{1cm} (3.21)

$$\Sigma = Cov(F^A - F^B),$$  \hspace{1cm} (3.22)

6. Repeat 2 to 5 for every ROI, and denote the sum of all ROIs' $d_{Mahal}$ as $d_{Mahal}^{all}$.

7. Repeat steps 2 to 6 for $k = 3$ to 16.
8. The $k$ corresponding to the maximum $d_{\text{Mahal}}^\text{all}$, $K$, is the number of features that best discriminates the two groups of activation maps across different ROIs in general.

We note that since in step 2, the $i$th subject's information is not used in generating $w$, the projected feature values calculated in step 3 are "test data" and not "training data", thus provides validation to our results. A plot of $d_{\text{Mahal}}^\text{all}$ versus the number of features is provided in the Section 3.4. Also note that the $K$ most discriminant 3DMIs are first projected onto the maximal discriminant 1D space using the above modified LDA procedure prior to applying permutation test.

### 3.3.5 Permutation Test

To determine the probability that the features extracted from the ROI activation maps of the two experimental groups are drawn from the same distribution, we decided to use the permutation test. Permutation test does not require a priori assumptions about the data distribution, and is thus preferred over T-test and F-test [23]. The permutation test procedure is as follows [24]:

1. Given $m$ $n$-dimensional feature vectors (assuming row vectors) for each group, vertically concatenate the $m$ feature vectors of the two groups into matrices, $X$ and $Y$. In the present study, $m = 21$ (i.e. 21 subjects) and $n = 1$ (i.e. each subject's 3DMI feature vector has been projected onto a 1D space during LDA, and mean activation statistics magnitude and percentage of activation are 1D features).

2. Calculate the Mahalanobis distances between $X$ and $Y$ using (3.20), and denote it as $d_{\text{orig}}$. 
3. Pool all sample feature vectors, stripped of their group labels.

4. Randomly draw \( m \) samples from the pool of feature vectors, label these samples \( X' \), and label the remaining samples \( Y' \).

5. If \( d_{Mahal} \) between \( X' \) and \( Y' \) is > \( d_{orig} \), increment a counter \( q \) by 1.

6. Repeat steps 3 to 6 \((N-1)\) times, e.g. let \( N = 100,000 \).

7. The p-value representing the probability that \( X \) and \( Y \) are drawn from the same distribution is estimated with the value \( q/N \).

3.3.6 Comparison of ROI Features

As mentioned in Section 3.3.2, two types of features are examined in this thesis: magnitude-based features including thresholded mean activation statistics and percentage of activated voxels, and spatial texture-based features, 3DMIs. We combined the described 3DMIs using the proposed method, and compared the results with that generated using thresholded mean activation statistics and percentage of activated voxels as single features as well as with these two magnitude-based features combined using the proposed analysis framework. We note that one might argue that any enhanced discriminability with the proposed method can easily be due to more features being used. However, in past ROI-based fMRI studies, only a limited number of magnitude-based features were considered due to inter-subject variability in brain shape, brain size, and subject's orientation in the MR scanner. Hence, we are extending the prior set of features with invariant spatial descriptors to make multiple feature analysis feasible.
3.4 Results

In this study, we are interested in comparing magnitude-based features (i.e. thresholded mean activation statistics and percentage of activated voxels) against the proposed spatial features, 3DMIs. To determine whether the differences in activation statistics during EG versus IG are significant, permutation test was applied to the described features with the results summarized in Table 3.1.

Table 3.1: p-values of Activation Differences During External versus Internal Tasks based on Positive T-statistics

<table>
<thead>
<tr>
<th>ROI</th>
<th>Thresholded Mean</th>
<th>Percentage of Activated Voxels</th>
<th>Proposed 3DMIs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Left LEN</td>
<td>0.1565</td>
<td>0.5576</td>
<td>0.0383*</td>
</tr>
<tr>
<td>Left CER</td>
<td>0.3143</td>
<td>0.7178</td>
<td>0.4412</td>
</tr>
<tr>
<td>Left SMA</td>
<td>0.0036*</td>
<td>0.1461</td>
<td>0.0184*</td>
</tr>
<tr>
<td>Left THA</td>
<td>0.2822</td>
<td>0.2902</td>
<td>0.1144</td>
</tr>
<tr>
<td>Left M1</td>
<td>0.1709</td>
<td>0.5256</td>
<td>0.0409*</td>
</tr>
<tr>
<td>Left PFC</td>
<td>0.5254</td>
<td>0.6588</td>
<td>0.0469*</td>
</tr>
<tr>
<td>Left CAU</td>
<td>0.4146</td>
<td>0.3759</td>
<td>0.8351</td>
</tr>
<tr>
<td>Right LEN</td>
<td>0.4559</td>
<td>0.9640</td>
<td>0.3330</td>
</tr>
<tr>
<td>Right CER</td>
<td>0.5079*</td>
<td>0.2656</td>
<td>0.0252*</td>
</tr>
<tr>
<td>Right SMA</td>
<td>0.0416*</td>
<td>0.0709</td>
<td>0.9895</td>
</tr>
<tr>
<td>Right THA</td>
<td>0.2674</td>
<td>0.0942</td>
<td>0.3465</td>
</tr>
<tr>
<td>Right M1</td>
<td>0.8176</td>
<td>0.7541</td>
<td>0.3794</td>
</tr>
<tr>
<td>Right PFC</td>
<td>0.6828</td>
<td>0.3062</td>
<td>0.8906</td>
</tr>
<tr>
<td>Right CAU</td>
<td>0.4587</td>
<td>0.9291</td>
<td>0.2097</td>
</tr>
</tbody>
</table>

LEN = lentiform nucleus, CER = cerebellar hemisphere, SMA = supplementary motor area, THA = Thalamus, M1 = primary motor cortex, PFC = prefrontal cortex, and CAU = caudate. *statistically significant at $\alpha = 0.05$.

The percentage of activated voxels did not detect any significant activation differences in any of the ROIs, and the thresholded mean only detected significant differences in the SMAs. In contrast, by first arranging the 3DMIs in decreasing order of discriminability $I_{113}, I_{133}, I_{333}, I_{123}, I_{112}, I_{33}, J_{3}, I_{11}, B_{3}, I_{3333}, I_{233}, I_{222}, B_{4}, I_{22}, J_{2}, J_{1}$ and using the proposed modified LDA procedure to choose the optimal
number of features ($K = 5$ as shown in Fig. 3.3), the proposed 3DMI method was able to detect significant task-related differences across subjects in the left lentiform nucleus, right cerebellar hemisphere, left SMA, left M1, and left PFC.

![Spatial feature selection](image)

Figure 3.3: Spatial feature selection (see description in Section 3.3.4). With the 3DMIs ordered based on their relative discriminability, 5 appears to be the optimal number of features for the dataset examined in this study. When more than five 3DMIs are used, discriminability is reduced due to overfitting.

### 3.5 Discussion

Our results suggest that the proposed 3DMI-based method is much more sensitive in detecting task-specific group changes in fMRI activation statistics within an ROI than using magnitude-based features. The ROIs detected with task-specific differences are consistent with previously-described neurobiological knowledge, as briefly
3.5.1 Neurobiological Relevance of Task-specific ROI Differences

The lentiform nucleus is known to be involved in diseases such as Parkinson’s disease, where subjects can more easily perform EG tasks [25],[26],[27]. This notion is further supported by a recent study which reported increased activity in the basal ganglia-thalamo-cortical loop during IG tasks that was not present during EG tasks [28]. Thus, detecting significant differences in the lentiform nucleus is consistent with prior imaging findings.

Significant differences in activation statistics were also detected in the right cerebellar hemisphere. This difference might be associated with handedness as seen in past studies [29]. Previous research has also indicated that sequential (as opposed to single) tasks often recruit the contralateral cerebellar hemisphere [29], whereas the typical motor pathway involves the ipsilateral cerebellar hemisphere [30]. It has been postulated that this increased activity in the contralateral cerebellar hemisphere may be related to the encoding of complex tasks [31].

Detecting significant differences in the left SMA can be explained by the fact that this neural region is known to be preferentially invoked during internally guided tasks [32], [33], [34]. This difference attributes to the increased contralateral activity for right-hand tasks.

Significant differences between EG and IG tasks were also detected in the left M1 and left PFC. The failure of magnitude-based methods in detecting significant differences in the left M1 could be explained by the fact that the M1 is the final pathway for motor movements and the actual movements performed during the EG and IG conditions were similar, thus requiring about the same level of activation.
The findings of significant differences in the left M1 with the 3DMI-based method is intriguing, and suggests that the spatial distribution within the M1 is different despite the similarity in level of activation.

The dorsolateral prefrontal cortex (DLPFC) has been shown to be involved in many cognitive-motor tasks, including sequence learning [35], spatial working memory [36], and movement planning and execution based on visual cues [37]. The finding of differential involvement in the DLPFC between EG and IG movements supports the idea of task-related differences in the functional geographic distribution of activation within this relatively large anatomical area.

### 3.5.2 Spatial Analysis of fMRI Data

Our results suggest that although the level of activation might have been modulated by task, the changes in the spatial distribution appeared to be much more discriminant. While it may be possible to envisage a sensitive method that merely amplifies inter-subject variability, the proposed method based on invariant spatial descriptors was able to capture task-modulated spatial aspects of activation statistics that are consistent across subjects. Thus, incorporating spatial information could in fact enhance the sensitivity of the activation discrimination analysis.

An important aspect of ROI-based fMRI analysis is the choice of T-threshold. In this thesis, we have decided to use a T-threshold of zero for two reasons. First, there is little consensus on the interpretation of negative T-values [14]. Secondly, false positives will not likely reside in the same location within an ROI for multiple subjects. Thus, inclusion of these voxels will only decrease discriminability. However, if a high T-threshold is set while some active voxels are buried by noise, these active voxels, which would be consistent across subjects, will be excluded from the
analysis. Therefore, using a T-threshold seems to be the most appropriate, and would provide a conservative estimate of the activation differences, while retaining most of the original information of the ROIs.

A drawback of the proposed spatial features is that we cannot easily state whether a given region is "more active" or "less active", just that the activation pattern is similar or different. This is analogous to a population-coding perspective at the neuronal level, where stating that a given region is more or less active is very difficult when only the spatial distribution of activation changes. In most cases where there is a significant difference in the spatial distribution of activation statistics, however, the changes in the thresholded mean activation statistics may give some indication of which condition results in "more" activation, even if the difference in the thresholded mean activation statistics does not reach statistical significance due to, e.g. inappropriate selection of threshold.

3.6 Conclusions

In this chapter, we presented a novel approach for fMRI data analysis that incorporates spatial information in contrast to traditional magnitude-based methods. The proposed technique uses 3DMIs to characterize the spatial distribution of brain activations within an ROI. Applying the proposed method to real fMRI data demonstrated that certain spatial aspects of activation statistics appear to be significantly modulated by task performance and being well conserved across subjects. Naively incorporating the spatial information would normally worsen inter-subject variability, but the invariance properties of the proposed spatial descriptors account for variability in subject's orientation in the scanner, while conserving the task-related 'textural' changes in brain activations. Thus as suggested by our results, incorporat-
ing spatial information can greatly enhance the sensitivity in detecting changes in activation statistics, as compared to using mean activation statistics and percentage of activated voxels as features. A direct extension of the proposed method is to look at the spatial distribution changes in time to infer functionally connectivity, a path currently being explored.
Bibliography


Chapter 4

Joint Spatial Denoising and
Active ROI Delineation

4.1 Introduction

One main application of functional magnetic resonance imaging (fMRI) is in making statistical inferences about the brain mechanisms responsible for different behavioral tasks. Conventionally, this entails generating activation statistics maps with each voxel assigned a statistical value related to its probability of being activated. To make group inferences under this approach, spatial warping of each subject’s brain to a common exemplar shape is often performed [1]. However, spatial normalization, typically followed by spatial smoothing, may pool responses from functionally dissimilar regions [2]. An alternative approach is to manually draw regions of interest (ROIs) individually for each subject, and examine the statistical properties...
of regional activation across subjects. Though this ROI-based approach deals with inter-subject anatomical variability, it suffers from two main drawbacks. First, delineating the boundaries of the ROIs is typically guided by anatomical structures [3], which ignores functional properties. Secondly, any outlier voxels can skew the statistical analysis. To deal with Type I errors (i.e. voxels falsely detected as activated) and to increase signal to noise ratio (SNR), a fixed Gaussian filter is conventionally used to spatially smooth the activation statistics maps. This procedure assumes that the MR scanner has a Gaussian-like point spread function (PSF) that smears the fMRI signal over a small neighborhood. Thus based on matched filter theorem [4], the SNR will be increased. However, smoothing the activation maps may affect voxels that do not contain Type I errors, and degrade the spatial resolution. To enhance localization, researchers have investigated spatial wavelet transforms as an alternative [5]. This approach involves representing clusters of voxels with a few wavelet coefficients and de-weighting or removing those coefficients that fall under a certain threshold (i.e. soft or hard thresholding). The inverse wavelet transform is then applied to the surviving coefficients to estimate a denoised activation map. Under this framework, better localization is obtained since not all voxels are altered as in the case of Gaussian smoothing. In this chapter, we propose a method for simultaneously denoising activation statistics within an ROI and refining ROI delineation. Assuming that activation statistics within a small neighborhood are spatially correlated, voxels within each neighborhood would thus exhibit similar level of influence on the overall ROI's response. To impose this condition, isolated voxels at region boundaries with disproportional influence on a derived ROI feature are removed, thereby refining the ROI delineation. Other overly influential voxels are de-weighted based on their influence relative to their neighbors to reduce the
biasing effects of these voxels in later analysis. Thus, only voxels detected as exceedingly influential are affected, while most of the original information is retained. The validity of the proposed approach is suggested by the fact that using one feature for denoising (e.g. spatial variance) resulted in greater effect size in another feature (e.g. average activation statistics magnitude).

4.2 Data Acquisition

4.2.1 Study Subjects and Experimental Conditions

After obtaining informed consent, fMRI data were collected from 8 healthy subjects. Each subject was required to perform a right-handed motor task that involved squeezing a bulb with sufficient pressure such that an 'inflatable ring', shown as a black horizontal bar on a screen, was kept within an undulating pathway. The pathway is straight during rest periods and becomes sinusoidal at time of stimulus. Each run lasted 260 s, alternating between rest and stimulus of 20 s duration. At time of stimulus, the subject was required to squeeze the bulb at three different frequencies: 0.25 Hz (slow), 0.5 Hz (medium) or 0.75 Hz (fast). The data were collected as part of a larger experiment exploring the rate of change of force production in older subjects and subjects with Parkinson's disease.

4.2.2 Functional MR Imaging

Functional MRI was performed on a Philips Gyroscan Intera 3.0 T scanner (Philips, Best, Netherlands) equipped with a head-coil. We collected echo-planar (EPI) T2*-weighted images with BOLD contrast. Scanning parameters were: repetition time 1985 ms, echo time 3.7 ms, flip angle 90°, field of view (FOV) 216×143×240 mm, in
plane resolution 128×128 pixels, pixel size 1.9×1.9 mm. Each functional run lasted 4 minutes where 36 axial slices of 3 mm thickness were collected in each volume, with a gap thickness of 1 mm. We selected slices to cover the dorsal surface of the brain and included the cerebellum ventrally. A high resolution 3D T1-weighted image consisting of 170 axial slices was acquired of the whole brain to facilitate anatomical localization of activation for each subject.

4.2.3 Preprocessing and Activation Statistics Generation

The fMRI data was pre-processed for each subject, using Brain Voyager’s (Brain Innovation B.V.) trilinear interpolation for 3D motion correction and sinc interpolation for slice time correction. Further motion correction was performed using motion corrected independent component analysis (MCICA) [6]. No temporal or spatial smoothing was performed on the data.

Using Amira (Mercury Computer Systems, San Diego, USA), custom scripts to co-register the anatomical and functional images were generated. Ten ROIs were manually drawn on each structural scan based upon anatomical landmarks and guided by a neurological atlas [3]. ROIs chosen included the thalamus, cerebellum, primary motor cortex (M1), supplementary motor area (SMA), and prefrontal cortex (PFC). The labels on the segmented anatomical scans were resliced at the fMRI resolution.

A hybrid Independent Component Analysis (ICA) General Linear Model (GLM) scheme [7] was used to contrast each of the three frequency blocks with the periods when subjects were producing static force in generating each subject’s activation statistics map. The underlying idea in this scheme is that the GLM regressors (reference and confounds) are derived using ICA, and leave-one-out validation
is used to select the best model.

4.3 Methods

The goal of our proposed method is twofold; denoising a subject's ROI activation map (hereafter referred to as T-maps) to reduce the biasing effects of voxels deemed falsely active as well as refining the ROI delineation. Assuming that the ROI T-maps have been calculated for two groups (e.g. left SMA for subjects performing two different motor tasks), the proposed method proceeds as follows (Fig. 4.1):

![Figure 4.1: Framework of proposed spatial denoising method](image)

For each subject's ROI T-map, we first calculate the influence of each voxel on the value of a chosen ROI feature. Average magnitude, percentage of activation, and spatial variance are examined in this study. We then use a robust outlier detection method to determine which voxel(s) appear to be overly 'influential.' Each outlier voxel is then de-weighted or removed by examining its relative influence as compared to its neighbors.

4.3.1 ROI Feature Extraction

Three features are examined in this thesis, namely average magnitude, percentage of activation, and spatial variance. We chose these features, because average magnitude and percentage of activation are widely used in ROI-based fMRI analyses.
and spatial variance has recently been shown to provide enhanced sensitivity in characterizing the activation distribution within an ROI [9]. Average magnitude is calculated by averaging voxel T-values greater than or equal to zero over the ROI. Percentage of activation is defined as the number of voxels with T-values above 1.96 divided by the total number of voxels within the ROI. Spatial variance, $J_1$, which is based on 3D moments, is calculated as follows:

$$J_1 = \eta_{200} + \eta_{020} + \eta_{002},$$

$$\eta_{pqr} = \frac{\mu_{pqr}}{\mu_{000}},$$

$$\mu_{pqr} = \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} (x - \bar{x})^p(y - \bar{y})^q(z - \bar{z})^r \rho(x, y, z) dx dy dz,$$

where $n = p+q+r$ is the order of the centralized 3D moment, $\mu_{pqr}$, $(x, y, z)$ are the coordinates of a voxel, $\rho(x, y, z)$ is the T-value of a voxel located at $(x, y, z)$, and $\bar{x}$, $\bar{y}$, and $\bar{z}$ are the centroid coordinates of $\rho(x, y, z)$. We note that $J_1$ is invariant to translation, rotation, and scaling, which accounts for differences in brain size across subjects and ensures that the calculated feature values does not depend on subject’s orientation in the scanner during acquisition [9]. The invariance properties, thus, allow the $J_1$ values to be easily compared across subjects.

### 4.3.2 Voxel Influence Map Generation

To generate the voxel influence map of an ROI for a given subject, we first calculate the feature value, $F_{all}$, using all voxels in that subject’s T-map. We then remove a voxel from the original T-map, and recalculate the ROI feature, $F_i$, with that voxel removed. The ‘influence’ of the removed voxel is then defined as the absolute difference between $F_{all}$ and $F_i$, which we denote by $dF_i$. The voxel influence map
of a subject’s ROI is then generated by performing this procedure for every voxel within the ROI.

4.3.3 Outlier Voxel Detection

To detect outliers, i.e., overly influential voxels within a subject’s ROI, a robust criterion based on the median and the median of absolute deviations (MAD) of $dF_i$ is used:

$$I_{out} = \text{find}(|dF_i - dF_{med}|) > N_{std} \cdot \sigma,$$

(4.4)

where $I_{out}$ is the set of voxels detected as outliers relative to the other voxels in the ROI, $dF_{med}$ is the median of $dF_i$, and $N_{std}$ is the number of standard deviations from $dF_{med}$ beyond which the value is considered an outlier. Assuming $dF_i$ is normally-distributed, we set $N_{std} = 3$ to retain 99.7% of the possible $dF_i$ values as non-outliers. $\sigma$ is a robust estimate of the standard deviation of $dF_i$, defined as:

$$\sigma \approx \lambda \cdot \text{median}(|dF_i - dF_{med}|),$$

(4.5)

where $\lambda$ converts MAD to standard deviation and is set to 1.483 assuming $dF_i$ is normally-distributed [10].

It is worth noting that our defined criterion (4.4) is robust to outliers, since the median is not as sensitive to extreme values as the mean. If $dF_{mean}$ is instead used in (4.5), extreme outliers can cause $\sigma$ to become arbitrarily large, thus intermediate outliers will likely be undetected.

4.3.4 Outliner Voxel De-weighting

To deweight the T-value ($T_k$) of each outlier voxel $k \in I_{out}$ within a subject’s ROI, we first separate its 26-connected neighbors (including itself) into two groups based
on (4): outliers ($V_{\text{out}}$) and non-outliers ($V_{\text{non-out}}$). The number of neighbors may be less than 26, especially if voxel $k$ lies at the ROI boundary. Also, errors in ROI delineation can skew the analysis, thus outliers located at region boundaries are treated more stringently than those residing inside the ROI.

If voxel $k$ lies inside the ROI and has only outlier neighbors, we leave $T_k$ unmodified, since a group of voxels in the same neighborhood exhibiting huge influence on an ROI’s feature are more likely to correspond to ROI response. In contrast, if some of voxel $k$’s neighbors are non-outliers, we deweight $T_k$ as follows:

$$T_k = \%_{\text{non-out}} \sum_{j \in V_{\text{non-out}}} w_j T_j + \%_{\text{out}} \sum_{m \in V_{\text{out}}} w_m T_m,$$  \hspace{1cm} (4.6)

$$w_j = \frac{1/dF_j}{\sum_{j \in V_{\text{non-out}}} 1/dF_j}, w_m = \frac{1/dF_m}{\sum_{m \in V_{\text{out}}} 1/dF_m},$$  \hspace{1cm} (4.7)

where $\%_{\text{non-out}}$ and $\%_{\text{out}}$ are the percentage of non-outlier and outlier neighbors. Our choice of $w$ is based on the assumption that activation statistics within a small neighborhood are spatially correlated. Therefore, no single voxel should exhibit exceedingly greater influence than its neighbors. To impose this condition, we scale down $T_i$ by $1/dF_i$, which proportionally reduces the influence of voxel $i$. Hence, each of the two sums in (4.6) corresponds to a representative T-value estimate of the non-outlier and outlier voxels. $T_k$ is then approximated by summing the two T-value estimates weighted by $\%_{\text{non-out}}$ and $\%_{\text{out}}$, since if $\%_{\text{out}} \gg \%_{\text{non-out}}$, $T_k$ is more likely to correspond to ROI response and vice versa.

If voxel $k$ lies on the ROI boundary, $T_k$ is set to zero, except if all its neighbors are outliers, where we deweight $T_k$ using (4.6). Zeroing the T-values in effect refines the ROI boundary, thus accounts for slight errors in ROI delineation.
4.3.5 Activation Discriminant Analysis

In this study, we are interested in detecting frequency-related activation differences (fast versus slow frequency) within various ROIs. Given either the raw or the denoised T-maps, we first calculate the average magnitude, $T_{ave}$, percentage of activation, $\%_{Act}$, and spatial variance, $J_1$, and test whether the differences in feature values between fast and slow frequencies, which we denote as activation map feature differences (AMFD), are significant using a paired T-test. The significant level is set at $\alpha = 0.05$, which corresponds to $|T_{crit}|$ of 2.36 for 8 subjects.

4.3.6 Comparison of Spatial Denoising Methods

With raw ROI T-maps as baseline, we compared our proposed method to the effects of traditional Gaussian spatial smoothing with a kernel size of 2 voxels (6 mm) full width half max (FWHM). In our approach, the feature used for denoising can be different from that used for activation discriminant analysis. We therefore compared the effects of using $T_{ave}$ versus $J_1$ as denoising feature. Also, using different features for denoising and activation discriminant analysis serves as a validity check. Note that $\%_{Act}$ is not a suitable denoising feature, since a voxel is either greater or less than 1.96, thus the resulting influence map will have only two possible values.

4.4 Results and Discussion

Table 4.1 summarizes the activation discriminant analysis results. In general, $T_{ave}$ and $\%_{Act}$ increased with task-frequency (i.e. positive T-values), and $J_1$ decreased with increasing frequency (i.e. negative T-values), thus demonstrating a 'focusing effect'. With the raw ROI T-maps, significant AMFD were detected only in the left
Table 4.1: T-values of Activation Map Feature Differences (AMFD) Comparing Fast versus Slow Frequency

<table>
<thead>
<tr>
<th>Methods</th>
<th>Raw $T_{ave}$</th>
<th>Raw $%Act$</th>
<th>Raw $J_1$</th>
<th>Gaussian Smooth $T_{ave}$</th>
<th>Gaussian Smooth $%Act$</th>
<th>Gaussian Smooth $J_1$</th>
<th>Proposed Method $T_{ave}$</th>
<th>Proposed Method $%Act$</th>
<th>Proposed Method $J_1$</th>
</tr>
</thead>
<tbody>
<tr>
<td>LTHA</td>
<td>1.27</td>
<td>2.15</td>
<td>-1.64</td>
<td>1.33</td>
<td>2.60</td>
<td>-1.98</td>
<td>1.18</td>
<td>3.14</td>
<td>-1.77</td>
</tr>
<tr>
<td>RTHA</td>
<td>1.33</td>
<td>2.28</td>
<td>-1.33</td>
<td>1.50</td>
<td>3.18</td>
<td>-1.74</td>
<td>1.65</td>
<td>3.73</td>
<td>-2.57</td>
</tr>
<tr>
<td>LCER</td>
<td>1.18</td>
<td>1.47</td>
<td>-0.50</td>
<td>1.55</td>
<td>1.59</td>
<td>-1.02</td>
<td>1.39</td>
<td>1.34</td>
<td>-0.15</td>
</tr>
<tr>
<td>NCER</td>
<td>1.35</td>
<td>2.03</td>
<td>-1.16</td>
<td>1.55</td>
<td>1.98</td>
<td>-1.03</td>
<td>1.56</td>
<td>1.90</td>
<td>-1.19</td>
</tr>
<tr>
<td>LMI</td>
<td>1.37</td>
<td>1.62</td>
<td>-2.17</td>
<td>1.49</td>
<td>1.74</td>
<td>-2.22</td>
<td>1.62</td>
<td>1.61</td>
<td>-1.71</td>
</tr>
<tr>
<td>RMI</td>
<td>1.80</td>
<td>2.20</td>
<td>-1.89</td>
<td>2.00</td>
<td>1.82</td>
<td>-2.09</td>
<td>1.66</td>
<td>2.61</td>
<td>-2.24</td>
</tr>
<tr>
<td>LCMA</td>
<td>1.97</td>
<td>2.61*</td>
<td>-2.41</td>
<td>2.15</td>
<td>2.43</td>
<td>-2.44</td>
<td>2.23</td>
<td>2.94</td>
<td>-2.77</td>
</tr>
<tr>
<td>RMCA</td>
<td>1.64</td>
<td>1.49</td>
<td>-1.17</td>
<td>1.71</td>
<td>1.76</td>
<td>-1.07</td>
<td>1.54</td>
<td>1.52</td>
<td>-1.59</td>
</tr>
<tr>
<td>LLFC</td>
<td>2.43*</td>
<td>1.00</td>
<td>-1.54</td>
<td>2.66*</td>
<td>1.70</td>
<td>-1.78</td>
<td>3.20*</td>
<td>3.46*</td>
<td>-2.74*</td>
</tr>
<tr>
<td>RLFN</td>
<td>2.19</td>
<td>1.99</td>
<td>-1.93</td>
<td>2.82*</td>
<td>2.22</td>
<td>-1.79</td>
<td>2.58*</td>
<td>2.52*</td>
<td>-1.71</td>
</tr>
</tbody>
</table>

THA = thalamus, CER = cerbellum, L = left, R = right, $T_{ave}$ = average magnitude, $%Act$ = percentage of activation, $J_1$ = spatial variance. First row under "Proposed Method" = feature being tested, second row = feature used for denoising. *significant at $|T_{crit}| = 2.36$. PFC (with $T_{ave}$) and left SMA (with $%Act$ and $J_1$). Spatially smoothing the T-maps with a Gaussian kernel only additionally detected the right PFC (with $T_{ave}$). In contrast, the proposed method detected significant AMFD in substantially more ROIs, including the bilateral thalamus, M1, PFC and the left SMA. Activation in the thalamus has been shown to scale with the rate of change of force production [11], and increasing movement speed is known to recruit widespread cortical areas including the right M1 [12] and the SMA. Thus, the results generated using the proposed method, are quite consistent with previous neuroscience studies.

Comparing $T_{ave}$ and $J_1$ as denoising feature, though $T_{ave}$ led to some increase in discriminability (i.e. increase in the T-value magnitude, especially when $%Act$ was used as the test feature) by merely modifying 15% of the voxels within each ROI on average (5% removed, 10% deweighted), substantially greater increase in discriminability was gained by using $J_1$ as the denoising feature (first and third column from the right in Table 4.1), which modified about 20% of the voxels within each ROI (10% removed, 10% deweighted). The reason for the greater discriminability increase is that $J_1$ accounts for voxel location, whereas $T_{ave}$ would modify any voxel.
within an ROI regardless of its location. In fact, using $J_1$ as denoising feature, we observed that on average, 80% of the outlier voxels were at the ROI boundary (e.g. Fig. 4.2), compared to 70% when $T_{ave}$ was used. This illustrates the importance of accurate ROI delineation in ROI-based fMRI analysis. Also, we note that using one feature for denoising resulted in greater discriminability in another feature, thus validating our proposed method.

4.5 Conclusion

We presented a new method for refining ROI delineation and denoising fMRI activation statistics within an ROI. Assuming voxels inside a small neighborhood are spatially correlated, we deweight only those voxels that exhibit exceedingly greater influence on an ROI's feature. Applying our method to real data demonstrated enhanced discriminability in detecting subtle activation differences as compared to
Gaussian spatial smoothing. A direct extension would be to apply our method to raw fMRI signals, which could spatially and temporally denoise the data.
Bibliography


Chapter 5

Temporal Dynamics of Spatial Distributions in fMRI BOLD Signals

5.1 Introduction

The most common application of functional magnetic resonance imaging (fMRI) is in mapping neural region(s) to particular function(s) by examining which brain areas activate when a certain task is performed. Most conventional analysis methods, such as statistical parametric mapping (SPM) [1], analyze each voxel’s timecourse independently and assign a statistics value to that voxel based on its probability of being activated. To make group inferences under this approach, spatial warping of each subject’s brain to a common exemplar shape is often performed to create a corre-

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4A version of this chapter has been accepted. B. Ng, R. Abuharbieh, S. J. Palmer, and M. J. McKeown, "Characterizing Task-Related Temporal Dynamics of Spatial Activation Distributions in fMRI BOLD Signals," 10th International Conference on Medical Image Computing and Computer Assisted Intervention, Brisbane, Australia, Oct. 29-Nov. 2, 2007 (accepted, 8 pages).
spondence between voxels across subjects [2]. However, spatial normalization, which is typically followed by spatial smoothing, may inappropriately pool responses from functionally dissimilar regions [3], thus degrading important spatial information. An alternative approach that involves drawing regions of interest (ROIs) individually for each subject, and examining the statistical properties of regional activation across subjects, has been shown to offer finer localization and increased sensitivity to task-related effects [3]. This subject-specific ROI-based approach is thus followed in this study.

To determine whether an ROI is activated or not, a simple approach is to calculate the average intensity over an ROI at every time point, and determine if the resulting average time course significantly correlates with the stimulus [4]. This approach, however, ignores any spatial information of activity within an ROI and assumes that only signal amplitude is modulated by task. However, spatial information might be an important attribute of brain activity. Preliminary evidence supporting this idea of spatial characterization was first shown by Thickbroom et al. [5], where the spatial extent of activation, as opposed to response magnitude, was found to be modulated by different levels of force during a sustained finger flexion task. Their results were based on visual inspection and counting the number of activated voxels within an ROI. Recently, we presented a more elaborate study of the spatial patterns of activity within an ROI where quantitative measures of invariant spatial properties were used to discriminate task-related differences in brain activity [6]. Results demonstrated that, by examining changes in different spatial aspects of an activation distribution, sensitivity in detecting functional changes is enhanced as compared to using intensity means only.

Previous analyses examining spatial patterns of activation, including that in
[6], were performed on T-maps where the spatial information is collapsed over time, thus only considered the time-averaged spatial patterns of brain activity. In this chapter, we extend our previously proposed spatial characterization approach to the temporal domain to explore whether the spatial distribution of the blood oxygenation level-dependent (BOLD) signal itself is modulated in time by task performance. We note an important difference between our current and previous work [6] is that the generated spatial feature time courses can be used to infer ROI activation, as opposed to only comparing 2 groups of time-averaged activation statistical maps. To characterize the spatial changes, three dimensional (3D) moment descriptors were used as features and were calculated at each time point. The magnitudes of these features, however, are normally not comparable across subjects due to inter-subject variability in brain shapes and sizes, but are comparable for the same subject over time. Thus, any detected modulations of the spatial features over time for a given subject may in fact represent meaningful spatial changes in activation.

In this study, eight healthy subjects were recruited to perform a bulb-squeezing task at various frequencies. The cerebellum, primary motor cortex (M1), supplementary motor area (SMA), prefrontal cortex (PFC), and anterior cingulate cortex (ACC) were chosen as regions of interest. We demonstrate that our method can both detect activation within an ROI, as well as discriminate differences in activation patterns at the various task frequencies. This confirms previous findings of the value in incorporating spatial information into traditional intensity-based fMRI analyses.
Figure 5.1: Experimental task and stimulus timing. (a) Subjects were required to keep the side of the black ring on the gray path (see text). (b) R = rest, Slow, Med, and Fast = stimulus at 0.25, 0.5, and 0.75 Hz, respectively. Each block is 20 s in duration.

5.2 Data Acquisition and Preprocessing

In this study, after informed consent was obtained, fMRI data were collected from 8 healthy subjects. Each subject was required to perform a right-handed motor task that involved squeezing a bulb with sufficient pressure such that an ‘inflatable ring’, shown as a black horizontal bar on a screen, was kept within an undulating pathway (Fig. 5.1(a)). The pathway remains straight during rest periods and becomes sinusoidal at time of stimulus. Each run lasted 260 s, consisting of a 20 s rest period at the beginning and end, 6 stimuli of 20 s duration, and 20 s rest periods between the stimuli, as shown in Fig. 5.1(b). At time of stimulus, the subject was required to squeeze the bulb at 0.25, 0.5 or 0.75 Hz, corresponding to ‘Slow’, ‘Med’, and ‘Fast’ in Fig. 5.1(b). The data were collected as part of a larger experiment exploring the rate of change of force production in older subjects and subjects with Parkinson’s disease.
5.2.1 fMRI Data Acquisition

Functional MRI was performed on a Philips Gyroscan Intera 3.0 T scanner (Philips, Best, Netherlands) equipped with a head-coil. We collected echo-planar (EPI) T2*-weighted images with BOLD contrast. Scanning parameters were: repetition time 1985 ms, echo time 3.7 ms, flip angle 90°, field of view (FOV) 216×143×240 mm, in-plane resolution 128×128 pixels, pixel size 1.9×1.9 mm. Each functional run lasted 4 minutes where 36 axial slices of 3 mm thickness were collected in each volume, with a gap thickness of 1 mm. We selected slices to cover the dorsal surface of the brain and included the cerebellum ventrally. A high resolution 3D T1-weighted image consisting of 170 axial slices was acquired of the whole brain to facilitate anatomical localization of activation for each subject.

5.2.2 fMRI Preprocessing

The fMRI data was preprocessed for each subject, using Brain Voyager’s (Brain Innovation B.V.) trilinear interpolation for 3D motion correction and sinc interpolation for slice time correction. Further motion correction was performed using motion corrected independent component analysis (MCICA) [7]. To handle temporal autocorrelations, a ‘coloring’ scheme was used [8], where the time series were high-pass filtered at 0.02 Hz (task-frequency being 0.025 Hz) to remove the majority of the low frequency noise, and temporally smoothened with a Gaussian of width 2.8s [8]. The first and last 20 s of the time series were truncated to mitigate transient effects. No spatial smoothing was performed.

The Brain Extraction Tool (BET) in MRicro [9] was used to strip the skull off of the anatomical and first functional image from each run to enable a more accurate alignment of the functional and anatomical scans. Custom scripts to co-register the
anatomical and functional images were generated using the Amira software (Mercury Computer Systems, San Diego, USA).

Ten specific ROIs were manually drawn on each unwarped structural scan using Amira. The following ROIs were drawn separately in each hemisphere, based upon anatomical landmarks and guided by a neurological atlas [10]: cerebellum, M1 (Brodman Area 4), SMA (Brodman Area 6), PFC (Brodman Area 9 and 10), and ACC (Brodman Area 28 and 32). The labels on the segmented anatomical scans were resliced at the fMRI resolution. The raw time courses of the voxels within each ROI were then extracted for analysis as described in the next section.

5.3 Methods

The main goal of the proposed method is to demonstrate that temporal dynamics of the spatial distribution in BOLD signals can be used to infer whether an ROI is activated, as well as to discriminate differences in activation patterns at various task frequencies. Details of feature time course extraction, activation detection, and activation pattern discrimination are discussed below.

5.3.1 Feature Time Course Extraction

The spatial feature descriptors used in this thesis are based on centralized 3D moments, defined as:

\[
\mu_{pqr}(t) = \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} (x - \bar{x})^p(y - \bar{y})^q(z - \bar{z})^r \rho(x, y, z, t) \, dx \, dy \, dz, \tag{5.1}
\]

where \( n = p + q + r \) is the order of the moment, \((x, y, z)\) are the coordinates of a voxel, \( \rho(x, y, z, t) \) is the intensity of a voxel located at \((x, y, z)\) inside a given ROI at time \( t \), and \( \bar{x}, \bar{y}, \) and \( \bar{z} \) are the centroid coordinates of \( \rho(x, y, z, t) \). To untangle the effect
of amplitude changes, $\rho(x, y, z, t)$ is normalized such that the intensity values of the voxels within the ROI sums up to one at every time point $t$. This step ensures that the mean ROI intensity does not change with time. Thus, any detected modulations in the spatial feature will be purely due to spatial changes in the BOLD signal. To ease interpretation of the results and since higher order moments are less robust to noise [11], only $2^{\text{nd}}$ and $3^{\text{rd}}$ order 3D moment descriptors characterizing spatial variance [12] and skewness, respectively, were used:

$$J_1(t) = \mu_{200}(t) + \mu_{020}(t) + \mu_{002}(t),$$

$$S(t) = \mu_{300}(t) + \mu_{030}(t) + \mu_{003}(t),$$

To compare with the results obtained using the proposed spatial feature time courses, the traditionally used mean intensity time course, $\bar{I}(t)$, for each ROI of a given subject is calculated by averaging the intensity over the ROI at every time point.

### 5.3.2 Activation Detection

To make group inference as to whether a given ROI is activated, each subject’s ROI feature time courses (spatial or mean intensity) are first correlated with a box-car that is time-locked to stimulus with a delay of 4 s [13]. We did not convolve the box-car with a haemodynamic response function since spatial changes, as governed by the different onsets of the active voxels, may exhibit a different temporal profile than that of the haemodynamic response. For each subject, this results in thirty correlation values, one per combination of feature and ROI (e.g. $J_1(t)$, left M1). Each correlation value is then converted into a T-value (5.4):

$$T = |r| \sqrt{\frac{N - 2}{1 - r^2}},$$
where \( r \) is the correlation value and \( N \) is the number of samples used in generating \( r \). The set of T-values of a particular combination of feature and ROI from all subjects is then tested against 1.96 using a T-test to determine the probability (p-value) that the T-values are lower than 1.96 (i.e. the probability that ROI is not activated). The critical p-value was chosen at 0.05.

### 5.3.3 Activation Pattern Discrimination

To discriminate the differences in activation pattern at the various task frequencies, each subject’s ROI feature time courses (spatial or mean intensity) are first segmented according to Fig. 5.2. Except for the first and last segments, each segment consists of a 10 s rest before and after the 20 s stimulus. Segments of the same task frequency are concatenated and correlated with the corresponding segments of the shifted reference signal (see Fig. 5.2). This results in ninety correlation values per subject, one for each combination of frequency, feature, and ROI (e.g. slow, \( J_1(t) \), left M1). Each correlation value is then converted into a T-value using (5.4).

For each combination of feature and ROI, the set of T-values of a particular frequency from all subjects are tested pair-wise against the other two frequencies (i.e. fast versus slow, fast versus medium, medium versus slow). This is performed using a T-test to determine the probability (p-value) that the sets of T-values from the two task frequencies are the same (i.e. the probability that activation patterns at the two frequencies are the same). The critical p-value was chosen at 0.05.

### 5.4 Results and Discussion

Table 5.1 summarizes the activation detection results generated by extracting the spatial and intensity features from real fMRI data as described in Section 5.3.1, and
Figure 5.2: Feature time course segmentation. The box-car curve corresponds to timing of the stimulus delayed by 4 s. The solid line is a sample feature time course (spatial variance, $J_1(t)$, of the left M1 averaged over subjects with its temporal mean removed and divided by its standard deviation). The dotted lines show how the feature time courses are parsed into 6 segments. Slow, Med, and Fast correspond to the task frequencies of 0.25, 0.5, and 0.75 Hz, respectively.

correlating the resulting feature time courses with the reference signal.

Using spatial variance, $J_1(t)$, the left cerebellum, left M1, both SMAs, left PFC, and left ACC were detected as active. We expected the left M1 to be activated, as typically observed for right-handed motor tasks. It is worth noting that the left M1's BOLD signal distribution shown reduced spatial variance (i.e. focuses) during the time of stimulus (see Fig. 5.2). Skewness, $S(t)$, additionally detected activation in the right M1. These results demonstrate that the spatial distribution of BOLD signals is, in fact, modulated by stimulus, which supports our hypothesis that spatial changes in BOLD signals are task-related and can be used to infer activation within an ROI.

Using the traditional mean intensity measure, the right cerebellum, both SMAs, left PFC, and both ACCs were detected as active. Comparing to the results generated with the proposed spatial features, some consistencies are shown. In fact, based on the results in Table 5.1, activation within an ROI appears to modulate
Table 5.1: p-values of ROI Activation

<table>
<thead>
<tr>
<th>Feature</th>
<th>$J_1(t)$</th>
<th>$S(t)$</th>
<th>$I(t)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>LCER</td>
<td>0.002*</td>
<td>0.077</td>
<td>0.469</td>
</tr>
<tr>
<td>RCER</td>
<td>0.220</td>
<td>0.066</td>
<td>0.042*</td>
</tr>
<tr>
<td>LM1</td>
<td>0.002*</td>
<td>0.051</td>
<td>0.162</td>
</tr>
<tr>
<td>RM1</td>
<td>0.157</td>
<td>0.047*</td>
<td>0.164</td>
</tr>
<tr>
<td>LSMA</td>
<td>0.027*</td>
<td>0.034*</td>
<td>0.021*</td>
</tr>
<tr>
<td>RSMA</td>
<td>0.044*</td>
<td>0.035*</td>
<td>0.034*</td>
</tr>
<tr>
<td>LPFC</td>
<td>0.036*</td>
<td>0.020*</td>
<td>0.034*</td>
</tr>
<tr>
<td>RPFC</td>
<td>0.341</td>
<td>0.185</td>
<td>0.058</td>
</tr>
<tr>
<td>LACC</td>
<td>0.001*</td>
<td>0.301</td>
<td>0.023*</td>
</tr>
<tr>
<td>RACC</td>
<td>0.056</td>
<td>0.112</td>
<td>0.045*</td>
</tr>
</tbody>
</table>

CER = cerebellum, $J_1(t) =$ spatial variance, $S(t) =$ skewness, $I(t) =$ mean intensity, L = left, R = right, * statically significant at $\alpha = 0.05$.

both in amplitude and in space.

Segmenting the feature time courses according to task frequencies and using the proposed spatial features, significant frequency-related activation differences were detected in the right cerebellum and right M1 when comparing fast versus slow frequencies (Table 5.2).

These results matched our expectations since the modulation of movement speed is known to involve a complex network of brain areas, including the right cerebellum and right M1 [14]. Also, significant activation differences were found in the right PFC and right ACC using the proposed spatial features. In contrast, no significant activation differences were found using mean intensity. Also, no significant activation differences were detected when comparing fast versus medium frequencies and medium versus slow frequencies for any of the features, thus these results were excluded in Table 5.2.

Examining the results in Table 5.2, spatial changes appear to provide greater sensitivity in detecting subtle activation differences as compared to intensity.
Table 5.2: p-values of Activation Differences Comparing Fast versus Slow Frequencies

<table>
<thead>
<tr>
<th>Feature</th>
<th>$J_i(t)$</th>
<th>$S(t)$</th>
<th>$I(t)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>LCER</td>
<td>0.1834</td>
<td>0.1931</td>
<td>0.1528</td>
</tr>
<tr>
<td>RCER</td>
<td>0.9993</td>
<td>0.0096*</td>
<td>0.2757</td>
</tr>
<tr>
<td>LM1</td>
<td>0.2932</td>
<td>0.4017</td>
<td>0.4190</td>
</tr>
<tr>
<td>RM1</td>
<td>0.0442*</td>
<td>0.3223</td>
<td>0.1524*</td>
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<tr>
<td>LSMA</td>
<td>0.6872</td>
<td>0.2499</td>
<td>0.1836</td>
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<td>RSMA</td>
<td>0.7887</td>
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<td>0.0579</td>
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<tr>
<td>LPFC</td>
<td>0.5981</td>
<td>0.3059</td>
<td>0.2760</td>
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<tr>
<td>RPFC</td>
<td>0.0398*</td>
<td>0.3668</td>
<td>0.7416</td>
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<tr>
<td>LACC</td>
<td>0.2551</td>
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<td>0.2126</td>
</tr>
<tr>
<td>RACC</td>
<td>0.0311*</td>
<td>0.1028</td>
<td>0.3248</td>
</tr>
</tbody>
</table>

CER = cerebellum, $J_i(t) = \text{spatial variance}, S(t) = \text{skewness}, I(t) = \text{mean intensity}, L = \text{left}, R = \text{right}, *\text{statically significant at } \alpha = 0.05.

5.5 Conclusion

In this chapter, we proposed using 3D moment-based spatial descriptors to characterize the temporal dynamics of spatial activation distribution within an ROI for fMRI analysis. We demonstrated with real fMRI data that certain spatial aspects of activation, as opposed to just amplitude, are modulated by stimulus—a result that appeared consistent across subjects. Furthermore, we showed that our method was able to better discriminate frequency-related differences in activation patterns during motor task performance when compared to using mean intensity only. These results suggest that spatial characterization of BOLD signal can complement traditional intensity-based fMRI analysis. A direct extension of the proposed method would be to examine functional connectivity and phase relations between ROIs using spatial feature time courses, an approach currently being pursued.
Bibliography


A novel ROI-based fMRI analysis paradigm is proposed for quantitatively characterizing the spatial distribution of brain activation within an ROI. We demonstrated for the very first time that the information encoded by voxel locations can be meaningfully incorporated in ROI-based fMRI analysis by using invariant spatial descriptors. The essence of our method is based on the invariant nature of the proposed descriptors, which accounts for inter-subject variability in brain shape, brain size, and subject’s orientation in the scanner. Our proposed analysis paradigm facilitates both spatial and spatio-temporal analysis of fMRI data.

6.1 Spatial Analysis of Activation Statistics Maps

Incorporating spatial information into ROI-based fMRI analysis by using invariant spatial descriptors (3DMIs) demonstrated enhanced discriminability of task-related activation differences over using traditional magnitude-based features [1], [2], [3]. However, several important issues remained to be addressed, namely the choice of activation statistics threshold for isolating activated voxels, the multiple comparisons...
problem, and further validation of the results.

In the current method, all voxels with positive T-statistics are used, and the justification is that false positives (voxels falsely deemed active) will not reside at the same location within an ROI across subjects. Hence, inclusion of these voxels will only reduce discriminability. Yet, truly active voxels corrupted by noise may reside in similar location within an ROI across subjects. Thus, if a "high" T-threshold is set, some of these active voxels will be excluded from the analysis. Therefore, using a T-threshold of zero provides a conservative estimate of activation differences, while retaining most of the original information of the ROIs. However, developing methods to first isolate the activated voxels before spatial characterization might further enhance discriminability. In fact, the processing method in [3] partially deals with this issue, where overly influential voxels that reside in isolation within an ROI (thus likely to be non-active) are deweighted. By further exploiting the idea in [3], other non-active voxels can be identified.

Regarding the multiple comparisons correction issue, all current results are thresholded at an uncorrected p-value of 0.05. The justification is that only tens of ROIs are examined as opposed to thousands of voxels in voxel-based analysis. Also, analogous to the assumption of spatial correlation between voxels in the GRF multiple comparison correction approach, the selected ROIs are all motor related, thus their response might also be correlated. Thereby, using Bonferroni or FDR correction might be too conservative. Instead, perhaps through examining the functional connectivity between ROIs, correction can be applied to networks of ROIs, which accounts for ROI-level spatial correlation, leading to less conservative thresholds.

Concerning extended validation of results, all our results were compared to prior neuroscience findings. However, most prior studies only suggest that the
magnitude of the ROI responses are modulated by task, whereas we are suggesting that different tasks are encoded by the spatial distribution of activation. Since the ground truth is not known, we are currently relying on indirect supporting evidence from prior studies. To more elaborately validate our method, realistic synthetic data is needed. One way to generate such data is to segment an ROI that is expected to incur no activation, and synthetically inject activation into that ROI. The drawback is that the noise structure across ROIs could be quite different, thus such approach does not guarantee that our results are valid for all ROIs. Alternatively, we can spatially permute the voxel locations and check if significant activation differences are still detected. This procedure is only appropriate for spatial feature analysis, but nonetheless provides a new validation approach to explore.

6.2 Spatio-temporal Analysis of BOLD fMRI Signal

In [4], we demonstrated that the spatial distribution of BOLD signal is modulated in time by task performance, which has significant neuroscience implications since all prior studies assume only the magnitude of BOLD signal is modulated by task. In functional connectivity analysis, an ROI response is often summarized by averaging voxel intensity time courses over an ROI. A more meaningful way to summarize an ROI might be to use spatial feature time course. In fact, examining the phase relations between spatial feature and mean intensity time courses might elucidate other functional properties of the brain. Also, examining the phase relations between ROIs might help infer causal relations.

We note that in [4], the issue of removing temporal autocorrelations was not fully addressed, since only the coloring scheme was tested. Comparing results generated by other preprocessing approach like whitening, will be beneficial. Also,
we suspect that the temporal autocorrelation problem might be less pronounced in the spatial domain. Thus, comparing the amount of residual autocorrelations in the spatial feature and mean intensity would be interesting.

The range of applications of our proposed analysis paradigm is very significant. In fact, with adequate validation, we foresee our approach being prospectively applied in clinical settings for diagnosing neurological diseases and evaluating drug therapies. Also, our paradigm can be easily transferred to examine other neurological diseases due to its generality.
Bibliography


