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A Mathematical Framework for the Registration and Analysis of Multi-Fascicle Models for Population Studies of the Brain Microstructure

Maxime Taquet*, Benoit Scherrer, Olivier Commowick, Jurriaan M. Peters, Mustafa Sahin, Benoit Macq, Simon K. Warfield

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Abstract

Diffusion tensor imaging (DTI) is unable to represent the diffusion signal arising from multiple crossing fascicles and freely diffusing water molecules. Generative models of the diffusion signal, such as multi-fascicle models, overcome this limitation by providing a parametric representation for the signal contribution of each population of water molecules. These models are of great interest in population studies to characterize and compare the brain microstructural properties. Central to population studies is the construction of an atlas and the registration of all subjects to it. However, the appropriate definition of registration and atlasing methods for multi-fascicle models have proven challenging. This paper proposes a mathematical framework to register and analyze multi-fascicle models. Specifically, we define novel operators to achieve interpolation, smoothing and averaging of multi-fascicle models. We also define a novel similarity metric to spatially align multi-fascicle models. Our framework enables simultaneous comparisons of different microstructural properties that are confounded in conventional DTI. The framework is validated on multi-fascicle models from 24 healthy subjects and 38 patients with tuberous sclerosis complex, 10 of whom have autism. We demonstrate the use of the multi-fascicle models registration and analysis framework in a population study of autism spectrum disorder.

1 Introduction

Diffusion magnetic resonance imaging enables in vivo investigation of the brain microstructure. Diffusion tensor imaging (DTI) has long been used in this context. However, DTI confounds the signal arising from different fascicles and from diffusion of free water, challenging the interpretation of scalar measures such as the fractional anisotropy (FA) and mean diffusivity (MD) [1]. This limitation makes DTI inadequate in the vast majority of the white matter since 60-90% of voxels contain more than one fascicle, according to recent estimates [2].

Various models have been proposed to overcome the limitations of DTI. Among them, generative models such as multi-tensor models [3, 4], CHARMED [5], NODDI [6] and DIAMOND [7] seek to represent the signal contribution from different populations of water molecules. These models are based on underlying biological assumptions and are of great interest to characterize and compare white-matter properties. For example, assessment of the free water diffusion arising from the extracellular space may be useful for the characterization of edema or inflammation [8]. A neuroinflammatory response may indeed lead to an increase in the amount of free diffusion [9]. Modeling of each individual fascicle may be useful to characterize properties such as the fascicle density, the axonal diameter distribution or the myelin integrity [10].

In this context, multi-tensor models are particularly interesting for three reasons. First, they en-
able the direct generalization of conventional measures computed from DTI (FA, MD, etc.) by enabling their computation for each fascicle independently. Second, they provide a model for the unrestricted water diffusion. Third, they can be estimated from short acquisition sequences that are clinically available [3, 11]. At each voxel, multi-tensor models represent the diffusion signal $M$ for a gradient direction $g$ and a b-value $b$ by:

$$M = S_0 \sum_{i=1}^{N} f_i e^{-bgT_i D_i g},$$

(1)

where $N$ is the maximum number of fascicles crossing in one voxel and $f_i$ is the volumetric fraction of fascicle $i$ (with $\sum_i f_i = 1$). Unrestricted water diffusion is represented as one of the compartments with an isotropic tensor ($D = D_{iso} I_3$).

Conducting population studies based on multi-fascicle models (MFM) requires the alignment of all models to a common coordinate system (the atlas). Registering and atlasing multi-tensor images are known to be challenging and many studies attempt to perform population analyses without resorting to them [12, 13, 14, 15]. In [12], a T2-weighted image of the subject is registered to a DWI at $b = 0$ and correspondence between subjects is achieved by segmenting the anatomy based on a T1-weighted atlas. In tract-based spatial statistics (TBSS) [13], single-tensor images are estimated and FA images are used to spatially align subjects. To interpret anisotropies in crossing fiber areas, heuristics based on the mode and FA of the tensor are used. In crossing-fiber TBSS [15], a ball-and-sticks model is estimated but spatial alignment is still based on single-tensor FA images. None of these approaches attempt to register multi-tensor models directly.

Direct registration of multi-fascicle models is important since the latter provide increased contrast in areas where T2-weighted images and FA images are almost constant (as will be shown in Section 4.3). Furthermore, multi-tensor image registration can be made invariant with respect to differences in FA and MD, which is important when those properties need to be compared after alignment. The challenges of registering and analyzing multi-fascicle models stems from difficulties in processing multi-tensors. In particular, interpolating, averaging, smoothing and defining robust similarity metrics for multi-fascicle models cannot be directly extended from the single-tensor case. This is because the $j$-th tensor in one voxel does not necessarily correspond to the same fascicle as the $j$-th tensor in another voxel. Furthermore neighboring voxels may contain different number of fascicles.

Interpolation and smoothing are required in registration to apply transforms and to prevent aliasing in multi-scale approaches. Building an atlas further requires averaging MFM. From a mathematical perspective, interpolating, smoothing and averaging all amount to computing weighted combinations of MFM. In this paper, we propose a mathematical framework to compute weighted combinations of MFM and a similarity metric to register them. These developments enable registration and analysis of multi-fascicle models which open new opportunities for population studies of microstructural properties. These contributions extend our previous work [16] by providing detailed derivations, experiments and discussions.

Section 2 introduces the mathematical framework to process multi-fascicle models. Section 3 details how the mathematical framework is applied to statistically analyze properties of the brain microstructure across populations. Section 4 presents results on in vivo data. Finally, Section 5 concludes.

2 Material

This section introduces the components of the proposed mathematical framework for multi-fascicle models.

2.1 Weighted combinations of multi-fascicle models for interpolation, averaging and smoothing

Computing weighted combinations of multi-fascicle models is at the basis of interpolation, smoothing and averaging. The linear combination of $K$ mixture
models each with \( N \) components results in a mixture with \( KN \) components that we call the complete model:

\[
\mathcal{M}^c = \sum_{k=1}^{K} w_k \mathcal{M}^k = \sum_{k=1}^{K} w_k \sum_{j=1}^{N} f_j^k S_j^k(g) \quad (2)
\]

\[
= \sum_{i=1}^{KN} f_i^c S_i^c(g) = S_0^c \sum_{i=1}^{KN} f_i^c e^{-bg^T D_i^c g} \quad (3)
\]

In most practical applications, increasing from \( N \) to \( KN \) the number of components is not desirable. We therefore estimate a simplified model, \( \mathcal{M}^s \), with \( N \) components which best approximates the complete model:

\[
\mathcal{M}^s = \sum_{j=1}^{N} f_j^s S_j^s(g) = S_0^s \sum_{j=1}^{N} f_j^s e^{-bg^T D_j^s g} \quad (4)
\]

\[
= \arg \min_{\mathcal{M}^s} D(\mathcal{M}^c, \mathcal{M}^s), \quad (5)
\]

where \( D(.,.) \) is some discrepancy measure between the complete and simplified models. This problem is known as mixture model simplification for which efficient approaches have recently been proposed [17, 18, 19]. In [17], the simplified mixture model is defined as that which minimizes the cumulative differential relative entropy between the complete and simplified models:

\[
D(\mathcal{M}^c, \mathcal{M}^s) = \sum_{j=1}^{N} \sum_{i: \pi_i = j} f_i^c D(S_i^c(g) || S_j^s(g)) \quad (6)
\]

\[
= \sum_{j=1}^{N} \sum_{i: \pi_i = j} f_i^c \int_\mathbb{R}^3 S_i^c(g) \log \frac{S_i^c(g)}{S_j^s(g)} dg. \quad (7)
\]

The variables \( \pi_i \) cluster the components \( S_i^c \) of the complete mixture into \( N \) clusters each represented by a single component of the simplified mixture, \( S_j^s \); \( \pi_i = j \) means that \( S_i^c \) is best represented by \( S_j^s \). Banerjee et al. [17] showed that, as long as \( S_i^c(g) \) and \( S_j^s(g) \) belong to the exponential family, Equation (6) can be optimized in an expectation-maximization scheme for which both the E-step and the M-step can be solved in closed form (Fig. 1). This makes the computation of weighted combinations of multifascicle models tractable. Mixtures of distributions from the exponential family is a wide class of mixtures which includes Gaussian mixtures [4], ball-and-sticks models [20], composite hindered and restricted models [21], diffusion directions models [22], Watson and Bingham distributions [6].

In the case of multi-tensor models, the E-step consists in optimizing for the clustering variables \( \pi_i \) assuming \( S_j^s \) are known, based on the Burg divergence \( B(.,.) \) between covariances \( \Sigma = D^{-1} \):

\[
\pi_i = \arg \min_j \left[ \text{Tr} \left( \Sigma_i^c \Sigma_j^{-1} \right) - \log \left| \Sigma_i^c \Sigma_j^{-1} \right| \right] (7)
\]

The M-step then consists in optimizing the parameters of the simplified mixture (that is \( D_j^s \) and \( f_j^s \)) to minimize (6) providing that we know \( \pi_i \). Davis and Dhillon [18] proved that this step amounts to computing the weighted average of covariance matrices and fractions in each cluster:

\[
\Sigma_j^s = \frac{\sum_{i: \pi_i = j} f_i^c \Sigma_i^c}{\sum_{i: \pi_i = j} f_i^c} \quad \text{and} \quad f_j^s = \sum_{i: \pi_i = j} f_i^c. \quad (8)
\]

Alternating the E-step (7) and M-step (8) until convergence provides the parameters \( (f_j^s \text{ and } D_j^s) \) of the weighted combination of mixtures. Initialization is required to control the local minimum to which EM will converge. We initialize the clustering variables \( \pi_i \) by spectral clustering using the cosine similarity matrix between the primary eigenvector \( e_i \) of each tensor \( D_i^c \) [23]. We found this initialization to be efficient in our experiments. However, since the algorithm typically converges in a few steps, one may consider running it multiple times with various initializations and selecting the result that yields the lowest cumulative differential relative entropy.

One may be concerned about the swelling effect due to averaging covariance matrices in (8). This motivates the definition of a log-Euclidean version of the mixture model simplification described above, as it has been defined for single-tensor interpolation [24]. This is achieved by replacing the covariance matrices by their matrix logarithm before performing the EM.
The update of the covariance matrices now reads:

$$\log \Sigma'_j = \frac{\sum_{i: \pi_i = j} f_i^c \log \Sigma_i^c}{\sum_{i: \pi_i = j} f_i^c}.$$ 

Since $\log \Sigma = \log D^{-1} = -\log D$, the logarithmic version of the weighted combination of multi-fascicle models is equivalent to its single-tensor counterpart in voxels with only one tensor. This is not the case in the Euclidean version since covariance matrices, rather than tensors, are averaged. A pseudocode of the method is presented in Algorithm 1.

Importantly, due to the construction of a complete model, the framework described above does not depend on the label $i$ assigned to tensors in the multi-tensor model and accounts for cases where the number of tensors differs between voxels.

### 2.2 A generalized correlation coefficient as a similarity metric for multi-fascicle models

To register multi-fascicle models, a similarity metric between multi-tensor images needs to be defined. Since registration is used for population studies, the similarity metric must be invariant to inter-subject variability. In particular, since mean diffusivity and fractional anisotropy are typically used as potential biomarkers for diseases, the similarity metric must be invariant to changes in FA and MD. This observation has lead Zhang et al. to define a single-tensor similarity metric based on deviatoric tensors, making it invariant to changes in MD [25], though not robust to other differences in diffusivity profiles. In this section, we generalize the correlation coefficient widely used in scalar images when intensities differ between subjects and we show that this similarity metric is invariant under changes in FA and MD.

The correlation coefficient as a similarity metric for block matching is defined as the scalar product between the normalized blocks. For voxels with values in $\mathbb{R}$, the blocks $R$ and $S$ defined over a domain $\Omega$ with $|\Omega|$ voxels are elements of $\mathbb{R} \times \ldots \times \mathbb{R} = \mathbb{R}^{|\Omega|}$ and the correlation coefficient reads:

$$\rho(R, S) = \left\langle \frac{R - \mu_R}{\|R - \mu_R\|}, \frac{S - \mu_S}{\|S - \mu_S\|} \right\rangle,$$ 

\[ \text{(9)} \]

\[ \text{Algorithm 1: Weighted Combinations in one voxel} \]

1. **Input:** $K$ multi-fascicle models $M^k$ with weights $w_k$ and the number $N$ of fascicles in the output.
2. **Output:** A multi-fascicle model: $\sum_{j=1}^N f_j^c e^{-b g^T D_j^c} g$
3. **for** $k$ in 1 to $K$ **do**
   \[\triangleright\] Construct the complete model $M^k$
4. \[\quad\text{for } j \text{ in 1 to } N \text{ do} \]
5. \[\quad i \leftarrow (k - 1)N + j \]
6. \[\quad f_i^c \leftarrow w_k f_j^c \]
7. \[\quad D_i^c \leftarrow D_j^c \]
8. \[\quad \text{end for} \]
9. \[\quad \text{end for} \]
10. \[\pi \leftarrow \text{Initialization}(\{D_i^c, f_i^c\}_{j \leq NK}) \quad \triangleright\text{Initialize clustering} \]
11. **while** $\pi$ has not converged **do**
12. \[\quad \text{for } j \text{ in 1 to } N \text{ do} \quad \triangleright\text{M-Step} \]
13. \[\quad \log D_j^c \]
14. \[\quad \left(\sum_{i: \pi_i = j} f_i^c \log D_i^c\right) \left/ \left(\sum_{i: \pi_i = j} f_i^c\right)\right. \]
15. \[\quad f_j^c \leftarrow \sum_{i: \pi_i = j} f_i^c \]
16. \[\quad \text{end for} \]
17. \[\quad \text{for } i \text{ in 1 to } KN \text{ do} \quad \triangleright\text{E-Step} \]
18. \[\quad \text{for } j \text{ in 1 to } N \text{ do} \]
19. \[\quad B_i(j) \leftarrow \text{Tr} \left(D_i^{-1} D_j^c\right)^- \]
20. \[\quad \pi_i \leftarrow \arg \min_l B_i(l) \]
21. \[\quad \text{end for} \]
22. \[\quad \text{end while} \]
where $\mu$ is the mean of the image values in the block and $\langle .., \rangle$ is the canonical scalar product in $\mathbb{R}^{[M]}$. It is invariant if $R$ is replaced by $aR + b$.

For vector images with values in $\mathbb{R}^n$, blocks are elements of $\mathbb{R}^n \times \ldots \times \mathbb{R}^n = (\mathbb{R}^n)^{[M]}$. The correlation coefficient can be generalized to vector images by redefining the means $\mu_R$ and $\mu_S$ as the projection of the block onto a block $T \in (\mathbb{R}^n)^{[M]}$ that has a constant value at each voxel, i.e. $T(x) = t_0 \forall x \ [26]$: \[
R - \mu_R = R - \frac{\langle R, T \rangle}{\|T\|^2} T. \tag{10}
\]
The factor $\langle R, T \rangle / \|T\|^2$ is a scalar that we call the scalar mean and is equal to $\mu_R$ for scalar images. Equation (9) is therefore a particular case of (10) for $n = 1$ and $t_0 = 1$. This generalized correlation coefficient can be used in any vector space endowed with an inner product. It is invariant if $R$ is replaced by $aR + bT$ where $a$ and $b$ are scalars and $T$ is the chosen constant block.

Let us first generalize the correlation coefficient to single-tensor diffusion images which will prove useful for the generalization to multi-tensor images. Single-tensor blocks are elements of $(S^+_3)^{[M]}$, where $S^+_3$ is the space of $3 \times 3$ symmetric positive definite matrices. It is typically more convenient to work in the log-tensor space in which blocks are elements of $(S_3)^{[M]}$. This space is endowed with the Frobenius inner product and the correlation coefficient of (10) can be readily applied. Choosing $t_0 = I_3$, the correlation coefficient is invariant under linear transformation of the log-tensor eigenvalues: $\log \lambda_1 \rightarrow \log \lambda'_1 = a \log \lambda_1 + \log b$ due to the invariance $\log D \rightarrow \log D' = a \log D + \log bI_3$. It is instructive to observe what the definition of the scalar mean becomes in this space with the Frobenius inner product. We have:

$$\mu_{S} = \frac{\langle D, T \rangle}{\|T\|^2} \tag{11}$$

\[= \frac{\sum_{x \in \Omega} \log \langle D(x), I_3 \rangle_F}{\sum_{x \in \Omega} \langle I_3, I_3 \rangle_F} \tag{12}\]

$$= \log \left( \prod_{x \in \Omega} \lambda_1(x)\lambda_2(x)\lambda_3(x) \right)^{\frac{1}{\|T\|^2}} \tag{13}$$

$$= \log \left( \lambda^T_3 \right). \tag{14}$$

The generalized scalar mean for blocks of single-tensors is therefore the logarithm of the geometric mean $\lambda^T_3$ of diffusivities over the domain $\Omega$.

Defining a scalar product in the space $(S_M)^{[M]}$ of blocks of multi-tensors seems impractical if not impossible. We further generalize the correlation coefficient (10) by substituting the inner product $\langle .., \rangle$, by a more general scalar mapping: $m(\ldots) : S^{[M]} \times S^{[M]} \rightarrow \mathbb{R}$ for any space $S^{[M]}$. The generalized correlation coefficient becomes:

$$\rho(R, S) = m\left( \frac{R - m(R, T)T}{n_m(R - m(R, T)T) - n_m(S - m(S, T)T)} \right), \tag{15}$$

where $n_m(X)^2 = m(X, X)$ is a generalization of the norm, and $T$ is assumed normalized ($n_m(T) = 1$).

This expression does not guarantee the invariance of the generalized correlation coefficient (GCC) with respect to linear changes of the blocks: $R \rightarrow R' = aR + bT$. Furthermore, in order to remain interpretable, the GCC must be symmetric, equal to one in case of perfect match and lower than one in any other case:

$$\rho(aR + bT, S) = \rho(R, S) \tag{16}$$

$$\rho(R, S) = \rho(S, R) \tag{17}$$

$$|\rho(R, S)| \leq 1. \tag{18}$$

These constraints on $\rho$ impose constraints on the scalar mapping $m$. One can show that constraints (15-18) are satisfied if the following constraints are respected by $m$:

$$m(aR + bT, T) = a m(R, T) + b m(T, T) \tag{19}$$

$$m(R, S) = m(S, R) \tag{20}$$

$$n_m(aR) = a n_m(R) \tag{21}$$

$$|m(R, S)| \leq n_m(R)n_m(S). \tag{22}$$

The latter generalizes the Cauchy-Schwartz inequality. Being a scalar product is a sufficient but unnecessary condition to respect these constraints. Therefore, constraints (19-22) as well as the choice of a constant block $T$ and suitable basic operations (to define the multiplication by a scalar and the addition of the constant block $T$), stand together as a model...
to generate correlation coefficients in potentially any space, even when an inner product cannot be defined.

In the case of multi-fascicle models, we further want the similarity metric between two multi-tensor blocks \( M^1(x) = \sum_{j=1}^{N} f^1_j(x) e^{-b g^T D^1_j(x) g} \) and \( M^2(x) = \sum_{i=1}^{N} f^2_i(x) e^{-b g^T D^2_i(x) g} \) to be equal to the single-tensor similarity metric if the blocks contain only one tensor in each voxel. This can be achieved if the scalar mapping is equal to the Frobenius inner product when all but one fractions are equal to zero. We therefore add a fifth constraint on the scalar mapping:

\[
\text{If } f^1_j = 1, f^1_{i\neq j} = 0, f^2_k = 1, f^2_{k\neq j} = 0, \\
\Rightarrow m(M^1, M^2) = \sum_{x \in \Omega} \langle \log D^1_j(x), \log D^2_k(x) \rangle_F . \tag{23}
\]

We define the multiplication of multi-tensors by a scalar \( a \) as the multiplication of all log-tensors by \( a \) and the addition of the constant block \( T \) as the addition of \( t_0 \) to all log-tensors. These definitions naturally generalize the single-tensor case. A generalized scalar mapping \( m \) comes by computing pairwise scalar products between corresponding tensors. This requires to pair tensors between the two blocks at each voxel. We introduce the following notation:

\[
d(p, x) = \sum_{i=1}^{N} f^1_i(x) f^2_{p(i)}(x) \langle \log D^1_i(x), \log D^2_{p(i)}(x) \rangle ,
\]

where \( p \) is the pairing function which associates one and only one tensor of \( M^1 \) to one and only one tensor of \( M^2 \). For \( N \)-fascicle models, there are \( N! \) such pairings. We define the scalar mapping for multi-tensor images as:

\[
m(M^1, M^2) = \sum_{x \in \Omega} d \left( \arg \max_p |d(p, x)|, x \right) . \tag{24}
\]

In practice, the values of \( d(p, x) \) for all \( N! \) pairings \( p \) are computed and we select the one with the highest absolute value. This scalar mapping satisfies (19-23). The absolute value is required by Condition (23) for cases where the Frobenius inner product between the tensors is negative. To better interpret this generalized scalar mapping, it is instructive to assess how it generalizes the concept of scalar means and norm to multi-fascicle models. The generalized scalar mean is given by:

\[
\mu^{SM}_m = \frac{m(M, T)}{m(T, T)} \\
= \frac{\sum_{x \in \Omega} \sum_{i=1}^{N} f_i(x) \frac{1}{N} \langle \log D_i(x), I_3 \rangle_F}{\sum_{x \in \Omega} \sum_{i=1}^{N} \frac{1}{N^2} \langle I_3, I_3 \rangle_F} \\
= \log \left( \prod_{x \in \Omega} \prod_{i=1}^{N} \langle \lambda^3_i(x) \lambda^3_i(x) \rangle^\frac{f_i(x)}{2} \right) . \tag{25}
\]

Remarkably, the generalized scalar mean for multi-fascicle model is the geometric mean of the diffusivities within the block for which all fascicles contribute in a ratio that is equal to their volumetric fraction \( f_i \) in their voxel. As for the generalized norm of multi-fascicle models, it is given by:

\[
n_m^2(M) = m(M, M) = \sum_{x \in \Omega} \sum_{i=1}^{N} f_i(x) \| \log D_i(x) \|_F^2 , \tag{26}
\]

that is the sum of the Frobenius norms of each log-tensor, weighted by the squared fractions. To demonstrate the latter expression, we need to show that the absolute value of \( d \) in (24) is maximized if the pairing \( p \) pairs a fascicle (in \( M_1 = M \)) with itself (in \( M_2 = M \)). The proof is straightforward using the Cauchy-Schwartz inequality on elements \( X = (f_1 \log D_1, \ldots, f_N \log D_N) \) and \( Y_p = (f_{p(1)} \log D_{p(1)}, \ldots, f_{p(N)} \log D_{p(N)}) \). Both the generalized scalar mean and the generalized norm therefore have direct interpretations in terms of multi-fascicle models.

More importantly, the proposed generalized scalar mapping leads to a GCC that is invariant under linear transformations of the eigenvalues of each log-tensor. In other words, \( \rho \) for multi-fascicle models is invariant under the following transformations \((i = 1, 2, 3)\):

\[
\log \lambda_i \rightarrow a \log \lambda_i + \log b, \forall a \in \mathbb{R} \text{ and } \forall b \in \mathbb{R}^+, \\
or, equivalently
\]

\[
\lambda_i \rightarrow b \lambda_i^a, \forall a \in \mathbb{R} \text{ and } \forall b \in \mathbb{R}^+ . \tag{27}
\]
for all fascicles. In particular, this invariance property encompasses differences in mean diffusivity (MD) for unchanged FA if \( a \neq 1 \) and \( b = 1 \). Similarly, changes in FA with unchanged MD can be obtained by varying \( a \) and \( b \) in a specific manner. Indeed, MD is preserved under changes of the eigenvalues following Equation (27), if

\[
b(\lambda_1^a + \lambda_2^a + \lambda_3^a) = \lambda_1 + \lambda_2 + \lambda_3.
\]

For any given set of eigenvalues and any given \( a \), there exists a \( b \) that satisfies this relation. One can therefore fix \( a \) to match the desired FA and subsequently fix \( b \) to respect this MD-preserving relation (since \( b \) does not affect the FA). Finally, by varying both \( a \) and \( b \) in an unconstrained manner, various changes in MD and FA can be accounted for by the invariance property of the GCC.

This similarity metric therefore allows registration of subjects with locally different diffusivity profiles. Importantly, because of the presence of the fractions \( f_1 \) and \( f_2 \) in the scalar mapping, the GCC accounts for cases where the number of tensors is different in different voxels (the corresponding fraction will simply be set to zero).

### 3 Methods

With the framework developed in the previous section, we can now perform population studies by constructing a multi-fascicle atlas and registering all subjects to it. From there, we can employ our novel operators including interpolation and averaging of multi-fascicle models to perform different statistical analyses of brain microstructural properties.

#### 3.1 Registration and Atlasing

Multi-fascicle models are estimated in the coordinate system of a T1-weighted MRI of the same subject (see Section 3.4). The registration between multi-fascicle models is initialized by affine registration of the T1-weighted MRI using the Baladin method [27], yielding a transformation \( T^0 \).

The weighted combinations of multi-fascicle models and the GCC are introduced in a robust multi-scale block matching registration algorithm developed in [28]. A dense deformation field is estimated through the following steps:

- For each pyramid level \( p = 1, \ldots, P \)
- For each iteration \( i = 1, \ldots, N \)
  - Estimate sparse pairings \( C \) between \( R \) and \( F \circ T^{i-1} \) by block-matching
  - Interpolate a dense correction field \( \delta T^i \) from \( C \) using a Gaussian kernel and weighted by the confidence in the matches as in [29].
  - Reject a fixed amount of outliers from \( C \) based on their dissimilarity with the estimated \( \delta T^i \)
  - Estimate an outlier-free correction \( \delta \tilde{T}^i \)
  - Compose the correction \( \delta \tilde{T}^i \) with the current estimate of the transform \( T^i = T^{i-1} \circ \delta \tilde{T}^i \)
  - Apply elastic regularization to the field \( T^i \)

In our implementation, \( P = 4, N = 10 \), block sizes are \( 5 \times 5 \times 5 \), and the outlier removal rate is 20%. The weighted combinations of multi-fascicle models are used to interpolate multi-tensor images when applying the deformation or constructing the multi-scale representation of the image. When warping tensor images (and hence multi-tensor images), tensors need to be reoriented. This reorientation is performed using the finite-strain rationale [30].

Registration is then used iteratively to build an atlas based on the method developed in [31]. This method essentially alternates between aligning and averaging images. To average multi-tensor images, we use the weighted combination of multi-fascicle models described above. Ten iterations are used to build the final atlas. The resulting atlas for single-tensor and multi-tensor images are depicted in Fig. 2.

#### 3.2 Statistical Analysis: Fascicle-Based Spatial Statistics

With all subjects aligned to the multi-fascicle atlas, we can compare properties of the aligned tracts through fascicle-based spatial statistics (FBSS)
(Fig. 3). Tractography is performed once on the atlas using the multi-fascicle tractography method described in [32, 33, 34] and adapted to include the multi-fascicle interpolation. For each registered subject, the tensor most aligned with the tract is selected and its property of interest (FA, MD, etc.) is computed. This provides, for each subject, a vector of length \( n \) (the number of points on the tract), representing the microstructural property along the fascicle.

Point-by-point t-tests are carried out along the tract to compare its properties between the two groups. This yields a vector \( t \) of \( n \) t-scores. Since the smoothness of the tract property depends on the individual, the tract and the resolution of the tractography, we use a non-parametric correction for multiple comparisons based on cluster-based statistics [35]. This method assumes that differences along the tract occur in clusters of adjacent points and proceeds as follows:

1. Define a threshold \( t_0 \) on the t-statistics.
2. Define a binary vector \( b \) of supra-threshold t-statistics:
   \[ b = (t > t_0). \]
3. Detect the connected components \( C = \{c_i\} \) in \( b \).
4. For each connected component \( c_i \), compute its size \( s_i \) (number of points) and its mass \( m_i \) (sum of t-scores).
5. Randomly permute \( N_p \) times the subjects in the groups (i.e. randomly reassign subjects to either groups) and perform Steps 1-4 for each permutation. For each permutation \( k \), record the maximum size \( s^k_p \) and the maximum mass \( m^k_p \) among the detected clusters.
6. The recorded \( s^k_p \) and \( m^k_p \) describe the null distributions of the size and mass of the clusters. Corrected for multiple comparisons, the p-values of each connected component \( c_i \) testing for its likelihood to be due to chance alone are:
   \[
   p^s_i = \frac{1}{N_p + 1} \left( 1 + \prod_{k=1}^{N_p} \delta(s^k_p > s_i) \right)
   \]
   \[
   \text{and } p^m_i = \frac{1}{N_p + 1} \left( 1 + \prod_{k=1}^{N_p} \delta(m^k_p > m_i) \right)
   \]
   for the size statistics and the mass statistics, respectively.

With FbSS, local fascicle segments where the two groups significantly differ can be discovered. Several \( t \)-thresholds \( t_0 \) are typically used to assess the robustness of the findings. Higher \( t_0 \) yield smaller clusters of stronger differences.

3.3 Statistical Analysis: Isotropic Diffusion Analysis

Large isotropic fraction \( f_{iso} \) indicates an excessive extracellular volume [36] which is in turn a surrogate for the presence of edema or neuroinflammation [8]. Isotropic diffusion analysis (IDA), i.e. the statistical analysis of the isotropic fraction, is thus of strong interest for population studies of disease involving these pathologies. The isotropic fraction is non-Gaussian since it ranges between 0 and 1. We apply the logit transform to \( f_{iso} \) prior to computing t-tests. This transform brings the distribution of \( f_{iso} \) closer to normality. Specifically, we transform \( f_{iso} \)-maps into \( l_{iso} \)-maps where:

\[
l_{iso} = \text{logit}(f_{iso}) = \log \left( \frac{f_{iso}}{1 - f_{iso}} \right).
\]

To prevent \( l_{iso} \) to take on infinite values when \( f_{iso} = 0 \) or 1, we bound the latter within \([10^{-6}, 1 - 10^{-6}] \). We then carry cluster-based statistics on the \( l_{iso} \)-maps with the cluster size and cluster masses as quantities of interest, as described in [35].

3.4 In vivo data

In vivo DWI were acquired on a Siemens 3T Trio scanner with a 32 channel head coil using the CUSP-45 gradient sequence [3]. This sequence includes
30 diffusion-encoding gradients on a shell at \( b = 1000s/mm^2 \) and 15 extra gradients in the enclosing cube of constant TE with b-values up to 3000s/mm\(^2\). Eddy current distortion was minimized using a twice-refocused spin echo sequence [37]. Other acquisition parameters were set to FOV= 220mm, matrix= 128 \times 128, number of slices=68, resolution = 1.7 \times 1.7 \times 2mm^3. Data acquisition was conducted using a protocol approved by the Institutional Review Board (IRB). The DW images were aligned to the 1\( \times 1\times1\)mm\(^3\) T1-weighted MRI with rigid registration (using the mean \( b = 0 \) image as a moving image) and the gradients were reoriented appropriately. This compensates for patient head motion and for residual geometric distortions due to magnetic field inhomogeneity and eddy current.

A multi-fascicle model with three tensors including an isotropic compartment were estimated as in [3]. Images were acquired for 24 healthy controls and 38 patients with tuberous sclerosis complex (TSC): 10 diagnosed with autism (TSC+ASD), 17 diagnosed without (TSC-ASD) and 11 too young for diagnosis.

4 Results

In this section, we validate the presented framework for multi-fascicle models. We systematically compare our results with those obtained when multi-tensor images are seen as a stack of single-tensor images on which multi-channel approaches to registration and averaging can be applied.

4.1 Relabeling Invariance Study

Multi-fascicle models assign arbitrary labels \( i \) to tensors in Equation (1). The framework must therefore be invariant under relabeling of tensors. In this experiment, we randomly relabeled tensors 10 times for each of the 24 healthy controls and performed registration between the result and the original image. Using the proposed framework, the deformation fields obtained were exactly the identity. By contrast, using the multi-channel registration, a significantly non-zero deformation field resulted from the registration of relabeled multi-fascicle models (Fig. 4). This result demonstrates the failure of multi-channel registration for multi-fascicle models. In what follows, tensors are labeled based on their FA (\( D_1 \) has the highest FA and \( D_N \) the lowest) to allow a fair comparison between the two approaches.

4.2 \( T^{-1} \circ T \) Study: Assessment of the Interpolation Error

For any transformation \( T \), the composition \( T^{-1} \circ T \) is equal to the identity. Therefore, for any image \( A \), \( T^{-1} \circ T \circ A = A \). However, if we first compute \((T \circ A)\) and then apply \( T^{-1} \) to the result, we do not obtain exactly the original image \( A \) due to interpolation error. Comparing the result \( \tilde{A} = T^{-1} \circ (T \circ A) \) to the original \( A \) thus provides estimates of the interpolation error, independently from the similarity metric.

We investigated the residual error of \( T^{-1} \circ T \circ A \) to compare the interpolation error of the proposed approach for linear combinations and the multi-channel alternative. The experiment was conducted with three different affine transformations \( T \) that were applied to the multi-fascicle models of the 24 healthy controls: (1) a transformation that maps the DWI to the T1-weighted image, (2) a translation by half a voxel in all directions, (3) a rotation of 45 degrees around the vertical axis.

The result \( \tilde{A} \) is compared to the original multi-fascicle model \( A \) in terms of the following four similarity metric computed at each voxel:

\[
\Delta^2_{FA} = \sum_{i=1}^{N} \frac{f_i + \tilde{f}_i}{2} (FA(D_i) - FA(\tilde{D}_i))^2 \tag{28}
\]

\[
\Delta^2_{MD} = \sum_{i=1}^{N} \frac{f_i + \tilde{f}_i}{2} (MD(D_i) - MD(\tilde{D}_i))^2 \tag{29}
\]

\[
\text{Fro}^2 = \sum_{i=1}^{N} \frac{f_i + \tilde{f}_i}{2} \|D_i - \tilde{D}_i\|_F^2 \tag{30}
\]

\[
\Delta_{Dir} = \sum_{i=1}^{N} \frac{f_i + \tilde{f}_i}{2} (1 - |e_{1,i}, e_{1,i}|) \tag{31}
\]

where \( e_{1,i} \) is the principal eigenvector of tensor \( D_i \) with unit norm. The last equation assesses how
aligned the resulting tensors are to the original tensors.

The results, summarized in Fig. 5, demonstrate that the use of the proposed method decreases the interpolation error as compared to the multi-channel alternative, for all similarity metrics and for all three transformations. On average, this decrease varies in magnitude from 38% for $\Delta_{\text{MD}}$ to 73% for $\Delta_{\text{Dir}}$. One-tailed paired t-tests indicate that the decreases are significant in all cases ($p < 10^{-12}$). An example of interpolation obtained with both methods is depicted in Fig. 6 and presents a portion of the corona radiata where fascicles cross. In this region, the multi-channel approach confounds the fascicles and fails to interpolate the multi-fascicle model.

Table 1: Summary statistics of the target registration error in the scan-rescan study

<table>
<thead>
<tr>
<th>Metric</th>
<th>Mean</th>
<th>St. dev.</th>
<th>P(Accuracy ≤ 1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$C_{\text{FA}}$</td>
<td>1.79</td>
<td>2.79</td>
<td>67.3%</td>
</tr>
<tr>
<td>$C_{\text{DTI}}$</td>
<td>2.18</td>
<td>3.05</td>
<td>59.4%</td>
</tr>
<tr>
<td>$C_{\text{MC}}$</td>
<td>3.34</td>
<td>4.18</td>
<td>47.5%</td>
</tr>
<tr>
<td>GCC</td>
<td>0.98</td>
<td>1.66</td>
<td>83.4%</td>
</tr>
</tbody>
</table>

4.3 Scan-Rescan Study: Evaluation of the Similarity Metric

In this section, we independently assess the accuracy of the similarity metric. We exploited two sets of 45 DWI acquired on the same subject during the same scanning session. The subject was required not to move and remained still throughout the acquisition. The two sets of DWI are thus intrinsically aligned. A multi-fascicle model as well as a single-tensor DTI were estimated from each set. The two multi-fascicle models differ due to acquisition noise, artifacts, and estimation errors. This scan-rescan experiment therefore provides a unique opportunity to estimate the accuracy of the proposed similarity metric in a realistic scenario.

A total of 495 landmarks were defined on a regular grid within the first image (Fig. 7(a)). Landmarks were spaced 7 voxels apart in all directions. Blocks of size $5 \times 5 \times 5$ were defined around each landmark and correspondence were sought in a neighborhood of size $21 \times 21 \times 21$ in the second image. Since there is no transformation between the two images, the true correspondence $x_{\text{true}}$ is located at the center of the neighborhood. The accuracy of the best match $x_{\text{max}}$ (that maximizes the similarity $C(x)$) and the saliency of the true match are:

\[
\text{Accuracy} = \| x_{\text{max}} - x_{\text{true}} \|
\]

\[
\text{Saliency} = \frac{C(x_{\text{true}}) - \bar{C}}{\sigma_c},
\]

where $\bar{C}$ and $\sigma_c$ are the mean and standard deviation of the similarity metric within the neighborhood.

Results for these two indices were compared amongst four different metric (1) the correlation coefficient applied to FA images ($C_{\text{FA}}$), (2) the correlation coefficient generalized to single-tensor DTI ($C_{\text{DTI}}$), (3) the multi-channel correlation coefficient applied to multi-tensors ($C_{\text{MC}}$) and the GCC for multi-fascicle models (GCC). Fig. 7(b) depicts the similarity maps for four different neighborhoods with each metric.

Neighborhood 1 in Fig. 7(b) illustrates the case of a specific white matter landmark, for which all four metrics perform equally well. Neighborhood 2 illustrates a case of a specific white matter landmark located at the intersection of crossing fascicles. In this case, both metrics based on multi-fascicle models find the correct match. Matching based on FA has more spurious maxima and matching based on DTI is offset because the single-tensor is a poor model of the diffusion signal in this region. Finally, Neighborhoods 3 and 4 show landmarks located at the boundary between the white and grey matter. In this area the microstructure is more complex. Multi-fascicle models are required in these regions to find a correct match. The multi-channel metric fails to detect the correct correspondence if tensors are not properly paired (see Neighborhood 4).

Fig. 7(c) depicts the volumetric fraction (shown as a color image) in areas where the FA displays no contrast. The fractions show a clear pattern of alternation between isotropic diffusion (blue) and single (green) or multi-fascicle (brown) orientation. These
patterns are repeated in both the scan and the rescan and therefore enable accurate matching.

Quantitatively, the GCC for multi-fascicle models significantly outperforms all other metrics in terms of accuracy (one-tailed paired t-test: $p < 10^{-8}$), as depicted in Fig. 7(d) and summarized in Table 1. The average gain in accuracy $C_{FA}$ is 45%. Importantly, the probability for the accuracy to be lower or equal to 1, that is the fraction of landmarks for which the best match was found in the direct neighborhood of the true match, is $P(\text{Accuracy} \leq 1) > 80\%$ for the GCC while it is lower than 70\% for all other metrics. These results suggest that the remaining registration error would likely be eliminated by regularization and outlier removal in the registration algorithm. No significant difference was observed in terms of saliency except for a significantly larger saliency for $C_{DTI}$ which may partially counterbalance its poorer accuracy in registration.

4.4 Comparison with Single-Tensor DTI Registration

As an alternative to the proposed approach for registration, one may suggest to register single-tensor DTI and use the resulting deformation fields to deform the multi-fascicle models. Such a method would not require any similarity metric between multi-tensors but would still require our framework to apply the deformation field to the multi-fascicle models. In this experiment, however, we illustrate that directly registering multi-fascicle models is more accurate than registering DTI.

We compared DTI registration and multi-fascicle registration under the application of a synthetic deformation field. Using a full dataset of 45 DWI form one subject, resampled to the T1-weighted space at a resolution of 1 mm×0.86 mm×0.86 mm, we estimated both a DTI and a multi-fascicle model. Both models were deformed by the same random synthetic log-Euclidean polyaffine transform (the true field) obtained by drawing parameters from a Gaussian with zero mean and 0.05 standard deviation for 27 regularly spaced affine components [38]. The single-tensor registration was performed using DTI-TK (version 2.3.1) [25] with the default parameters and after properly rescaling tensors. The multi-fascicle registration was performed using our framework. The resulting fields were applied to the original multi-fascicle model and the results were compared, in terms of metrics (28)-(31), to the multi-fascicle model deformed by the true field. We also computed the norm ($\Delta_{\text{field}}$) of the difference between the deformation field obtained by registration and the true field. All comparisons were conducted within a mask that excludes the background.

In terms of all five metrics, registering the multi-fascicle models with our framework significantly improves the alignment accuracy compared to single-tensor registration with DTI-TK (one-sided paired t-test: $p < 10^{-6}$ for all six metrics) as depicted on Fig. 8(a). The differences in accuracy between the two approaches are mostly visible in areas with crossing fascicles, such as the corona radiate as shown in the zoomed-in areas of Fig. 8(b). In these regions, single-tensor DTI have low contrast and DTI-TK is therefore less reliable than our approach which takes advantage of the full multi-fascicle model in the registration.

4.5 Synthetic Fields Study

In this experiment, we compare the registration accuracies when synthetic deformation fields are applied to multi-fascicle models. Ten random log-Euclidean polyaffine deformation fields are generated by drawing parameters from a Gaussian with zero mean and 0.05 standard deviation for 27 regularly spaced affine components [38]. Each of the ten deformations are applied to the 24 multi-fascicle models of the control subjects. Symmetric matrices of Gaussian noise with zero mean and standard deviation at six different levels (0.1, 0.2, 0.3; and 0.5, 1.0, 1.5) were then added to the log of all tensors in both the original and the transformed image, corresponding to SNR of (30dB, 24dB, 21dB, 17dB, 11dB, 7dB). The original and the transformed images were then registered and the resulting deformation field was compared to the initial synthetic field in term of its root mean squared (RMS) error. All 1,440 registrations (24 sub-
jects × 10 deformation fields × 6 noise levels) were performed with both the proposed framework and the multi-channel alternative.

On average, the root mean squared (RMS) error of the deformation field is 17% higher when the multi-channel registration is used instead of the proposed framework. A one-tailed paired t-test on the RMS for the transformation at each SNR shows that the difference in RMS between the approaches is significant: \( p < 10^{-5} \) for all SNR between 11dB and 30dB and \( p = 0.02 \) for SNR=7dB (Fig. 9-Top). The variance of the RMS is also decreased by 45% on average. A one-tailed F-test reveals that this decrease is significant for all SNR (\( p < 0.001 \)) except for SNR=24dB and 7dB (Fig. 9-Bottom). This experiment indicates that even when tensors are labeled based on their FA, the proposed framework outperforms the multi-channel alternative.

### 4.6 Morphometric Contrast Study

The deformation field obtained by registering a subject to an atlas provides a measurement of the local morphometric difference between the subject and a standard anatomy. The determinant of the Jacobian \( |J| \) of the deformation fields at every voxel provides information about the amount of local volume differences (\( |J| < 1 \) indicates a decreased volume and \( |J| > 1 \) indicates an increased volume). Widespread volume deficits in the white and grey matter of patients with TSC have been previously reported [39]. However, the amount of differences detected depends on the accuracy of the registration because the statistical power of the test depends on the registration accuracy [40]. Because of the increased level of microstructure they represent, we expect multi-fascicle models to reveal more morphometric differences than single-tensor models and scalar T1-weighted MRI. To assess the statistical power of all modalities (T1, single-tensor DTI and multi-fascicle models), we used the common voxel-based morphometry method [41]: register all subjects to the atlas, compute the log-jacobian, smooth it by a kernel of 8mm FWHM and correct for family-wise error rate at \( p = 0.05 \).

Results in Fig. 10(a) show that multi-fascicle models reveal more differences than single-tensors and T1-weighted MRI, as expected. This is likely due to an increased statistical power resulting from a higher registration accuracy when the structure of the white matter is better represented. The number of significant voxels does not differ between the proposed approach and the multi-channel alternative. However, the spatial distribution of the volume deficit findings (Fig. 10(b)) better follows the anatomy than the multi-channel alternative as seen, for example, in the left and right internal capsules.

### 4.7 Application: Fascicle-Based Spatial Statistics of the Dorsal Language Circuit

FBSS can detect local abnormalities in white matter pathways, which helps defining foci of neurological disorders. In this section, we investigate whether local decreases in FA along the dorsal language circuit (Fig. 11) can be discovered by FBSS.

Tractography of the dorsal language circuit was performed using the automatic seeding method of [42, 33]. A representative tract that captures the geometry of the bundle was manually selected. One-tailed fascicle-based spatial statistics was first performed between the 38 patients with TSC and the 24 healthy controls to test whether TSC patients have lower FA along the tract than healthy controls (Fig. 12-top row). Results found with both the multi-fascicle approach and the multi-channel approach consistently show that differences between TSC patients and healthy controls are widespread over the tract. This is consistent with recent models of tuberous sclerosis complex presented as a widespread decreased white matter microstructural integrity [32] and a global loss of connectivity [43]. Analysis based on single-tensor images did not reveal significant differences between the groups (except for a small cluster near the dorsal end of the tract). This is probably due to DTI being unable to distinguish the signal arising from each fascicle (one of them generating the group difference) and from free diffusion.

One-tailed fascicle-based spatial statistics was also performed between TSC+ASD and TSC-ASD pa-
tients to further understand the impact of autism on the properties of fascicles in the language system (Fig. 12-bottom row). A cluster of significantly lower FA was found in the middle of the tract, i.e. in the white matter close to the Geschwind’s territory, a region that has previously been associated with the interpretation of facial emotions [44]. Furthermore, using the proposed framework for multi-fascicle registration and analysis, a second cluster of significantly lower FA was found in the white matter close to Broca’s area, a cortical region associated with speech production whose activity was shown to be impaired in patients with autism spectrum disorder [45]. Again, no local difference was observed based on single-tensor images.

Findings of lower FA in TSC+ASD compared to TSC-ASD were previously reported in the literature [32, 33]. However, for the first time, our framework enables the detection of local differences, improving our knowledge of alterations in the brain microstructure related to autism spectrum disorder.

4.8 Application: Isotropic Diffusion Analysis in autism

Isotropic diffusion analysis allows whole-brain inspection of differences in isotropic fraction $f_{iso}$ whose excess relates to the presence of neuroinflammation and edema among others. To investigate in vivo whether autism spectrum disorder may result from a neuroinflammatory response (as suggested by post-mortem studies [46]), we performed isotropic diffusion analysis to compare the TSC+ASD and TSC-ASD groups. Cluster-based statistics was performed at four different thresholds: $t_0 = 2, 2.5, 3,$ and $3.5$ to assess the robustness of the findings with respect to the threshold used.

Consistently for all thresholds, clusters of significantly higher $f_{iso}$ were detected in patients with autism (Fig. 13). Both the size and mass of these clusters show significant departure from the null distribution ($p < 0.05$, Table 2). The multi-channel approach also found significant clusters but these were smaller in size and more sensitive to the choice of threshold (no significant cluster was found for $t_0 = 3.5$). The location of the significant cluster detected with both methods coincide and corresponds to part of the visual system (Fig. 13).

These findings are consistent with recent studies of autism in children which have demonstrated that appropriate maturation of visual system is crucial for social cognition development [47]. Furthermore, while autism is believed to potentially result from a neuroinflammatory process, in vivo evidence of such neurological mechanism are missing. These results illustrate how the proposed techniques for the analysis of multi-fascicle models can provide new insights into the brain microstructure. The validation of the neuroinflammatory process in autism would however require further studies including more subjects and other imaging modalities (such as PET imaging and T2 mapping).

5 Conclusion

Diffusion tensor imaging confounds the diffusion signal arising from different compartments and may therefore not be reliable for population studies of the brain microstructure. In particular, studies based on DTI cannot separate differences in properties of the fascicles due to demyelination or axonal injury, from differences in extracellular volume fraction due to neuroinflammation, edema or partial voluming with CSF. By representing the signal arising from different compartments with distinct parameterizations, multi-fascicle models are able to explain the origin of the observed differences. This property makes multi-fascicle models of great interest for population studies of the brain microstructure.

The cornerstone of image-based population studies is the construction of an atlas and the registration of all subjects to it. In this paper, we introduced a framework for registration and atlasing of multi-fascicle models. A mixture model simplification method was introduced to compute weighted combinations of multi-fascicle models, as used for interpolation, smoothing and averaging. As a similarity metric, a generalized correlation coefficient was developed to be invariant under linear transformations of the eigenvalues of each fascicle in the log-domain, making it robust to inter-subject variability.
Table 2: Size and mass statistics of the significant clusters from isotropic diffusion analysis.

<table>
<thead>
<tr>
<th>Threshold</th>
<th>Cluster</th>
<th>Multi-Channel Value</th>
<th>Multi-Channel p-value</th>
<th>Multi-Fascicle Value</th>
<th>Multi-Fascicle p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Cluster 1</td>
<td>22408</td>
<td>0.046</td>
<td>30927</td>
<td>0.05</td>
</tr>
<tr>
<td></td>
<td>Cluster 2</td>
<td>5126</td>
<td>0.039</td>
<td>14034</td>
<td>0.028</td>
</tr>
<tr>
<td></td>
<td>Total</td>
<td>12574</td>
<td>0.025</td>
<td>14034</td>
<td>0.028</td>
</tr>
<tr>
<td>2.5</td>
<td>Cluster 1</td>
<td>7448</td>
<td>-</td>
<td>22875</td>
<td>-</td>
</tr>
<tr>
<td>3</td>
<td>Cluster 1</td>
<td>1837</td>
<td>0.035</td>
<td>6270</td>
<td>0.017</td>
</tr>
<tr>
<td>3.5</td>
<td>Cluster 1</td>
<td>-</td>
<td>-</td>
<td>1864</td>
<td>0.02</td>
</tr>
</tbody>
</table>

Once all subjects are aligned to the atlas, population studies can be carried out to investigate microstructural properties in brain diseases. We introduce a system of two statistical analyses of the brain microstructure: fascicle-based spatial statistics (FBSS) and isotropic diffusion analysis (IDA). The former allows discoveries of local differences in the microstructural properties of the fascicle in a specific pathway. The latter allows detection of differences in extracellular volume fraction which may relate to neuroinflammation and edema. Together, these analyses allow for comprehensive investigation of the brain microstructure. We illustrated its use in a population study of autism spectrum disorder related to tuberous sclerosis complex and showed that the use of multi-fascicle models in this context increases the sensitivity of the statistical tests.

References


Figure 1: Computing weighted combinations of multi-fascicle models amounts to computing the complete mixture $\mathcal{M}^c$ and simplifying it in an EM scheme to obtain $\mathcal{M}^s$. The E-step is a clustering problem and the M-step consists in averaging log-tensors in each cluster.

Figure 2: Single-tensor and multi-fascicle atlases overlaid on the T1-weighted MRI atlas. The multi-fascicle atlas presents tensors with higher fractional anisotropies than the single-tensor atlas. This is due to the account of both the free water diffusion in the isotropic compartment and the multiple fascicle present in the voxel. The highlighted regions represent the corona radiata where projections of the corpus callosum cross cortico-spinal tracts, and a region where the pyramidal tracts (vertical lines) and the medial cerebellar peduncle (horizontal lines) cross.
Figure 3: Fascicle-based spatial statistics (FBSS) proceeds in three steps. (a) Fascicles (grey line) are drawn on the atlas with a sub-voxel resolution. The point in the middle is at a non-grid location. (b) Multi-fascicle models are interpolated at non-grid locations. (c) At each location along the fascicle, the tensor most aligned to the fascicle is selected to compute the property of interest (FA, MD, etc.).

Figure 4: Average RMS error of the deformation fields obtained by registering a multi-fascicle model with itself after randomly relabeling tensors. Our framework is invariant under relabeling leading to an error that is exactly zero. By contrast, multi-channel registration yields non-zero deformation fields.
Our approach

\[ \Delta_{\text{Dir}} \]

\[ \Delta_{\text{MD}} \]

\[ \text{Fr o} \]

\[ \Delta_{\text{FA}} \]

Figure 5: Estimation of the interpolation errors demonstrates the superiority of the proposed approach compared to the multi-channel alternative to compute linear combinations of multi-fascicle models. The bar plots show the interpolation error for four metrics (\( \Delta_{\text{FA}}, \Delta_{\text{MD}}, \text{Fro} \) and \( \Delta_{\text{Dir}} \)) under three different transformations.

Figure 6: (Left) Performing interpolation for each tensor independently (considered as channels of a multi-channel image) confounds fascicles resulting in an inflated result. (Right) Weighted combination of multi-fascicle models introduced in our mathematical framework clusters similar fascicles to avoid the inflation effect. (a-b) Results obtained on synthetic data by interpolating the multi-fascicle models at the extremities. (c-d) Results obtained on \textit{in vivo} data by applying a linear transform to a multi-fascicle model.
Figure 7: The GCC for multi-fascicle models outperforms other metrics in terms of registration accuracy, as assessed by a scan-rescan experiment. (a) 495 regularly spaced landmarks are used for the experiment. (b) Similarity maps in four neighborhoods (circles indicate true matches) showing that GCC is the most specific metric. (c) In regions with no contrast in FA, the GCC is able to find correct matches due to robust patterns observed in multi-fascicle models. (d) The accuracy of the GCC is significantly better than all other metrics as seen by the cumulative distribution function (CDF). (e) No significant difference in saliency between the metrics are observed, except for a significantly higher saliency with $C_{DTI}$.
Figure 8: Registering multi-fascicle models with our framework leads to higher alignment accuracies than registering single-tensor DTI with DTI-TK. (a) Quantitative assessment shows that registration errors using our framework are significantly lower than those obtained with DTI-TK [25]. (b) The difference in registration error is mostly visible in areas with crossing fascicles, where single-tensor DTI models have low contrast compared to multi-fascicle models. The zoomed-in areas are located in the corona radiata.
Figure 9: Our mathematical framework leads to higher registration accuracies than the multi-channel alternative: the RMS errors (top) and its variance (bottom) are significantly lower. Results are shown for 1,440 registrations performed at various SNR for synthetic deformation fields (* $p < 0.05$, ** $p < 0.005$, *** $p < 0.001$).

Figure 10: Morphometry results show areas with a significant volume deficit within the grey and white matter of TSC patients. (a) Multi-fascicle registration and multi-channel registration reveal more differences than single-tensor and the T1-weighted registrations. The differences observed with multi-fascicle registration are more consistent with the known anatomy than those observed with multi-channel registration, as seen for example in the left and right internal capsules.
Figure 11: The dorsal language circuit is composed of white matter fascicles thought to connect Broca’s area in the frontal lobe (Region 1), Geschwind’s territory in the parietal lobe (Region 2), and Wernicke’s area in the temporal lobe (Region 3). The median tract was manually selected from those tracts to perform fascicle-based spatial statistics (FBSS).

Figure 12: Fascicle-based spatial statistics of multi-fascicle models reveal local differences in the white fascicle properties that single tensor DTI cannot. Curves show the mean FA along the median tract of the dorsal language circuit in each group. Shaded area along the curves represent two standard errors. Grey rectangles indicate that the FA in that cluster is significantly different between the two groups. The top row studies differences between patients with tuberous sclerosis complex (TSC) and healthy controls. The bottom row further investigates differences between TSC patients with (TSC+ASD) and without autism (TSC-ASD). Landmarks 1, 2 and 3 correspond to those in Fig. 11.
Figure 13: Multi-fascicle models reveal clusters of increased isotropic fraction in autism, potentially indicating the presence of neuroinflammation. Clusters found with our framework (top) are larger and more coherent than those obtained with the multi-channel alternative (results shown for $t_0=3$).