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Generalised coherent point drift for group-wise multi-dimensional analysis of diffusion brain MRI data

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Abstract

A probabilistic framework for registering generalised point sets comprising multiple voxel-wise data features such as positions, orientations and scalarvalued quantities, is proposed. It is employed for the analysis of magnetic resonance diffusion tensor image (DTI)-derived quantities, such as fractional anisotropy (FA) and fibre orientation, across multiple subjects. A hybrid Student's t-Watson-Gaussian mixture model-based non-rigid registration framework is formulated for the joint registration and clustering of voxel-wise DTI-derived data, acquired from multiple subjects. The proposed approach jointly estimates the non-rigid transformations necessary to register an unbiased mean template (represented as a 7-dimensional hybrid point set comprising spatial positions, fibre orientations and FA values) to white matter regions of interest (ROIs), and approximates the joint distribution of voxel spatial positions, their associated principal diffusion axes, and FA. Specific white matter ROIs, namely, the corpus callosum and cingulum, are analysed across healthy control (HC) subjects (K=20 samples) and patients diagnosed with mild cognitive impairment (MCI) (K=20 samples) or Alzheimer's disease (AD) (K=20 samples) using the proposed framework, facilitating intergroup comparisons of FA and fibre orientations. Group-wise analyses of the

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latter is not afforded by conventional approaches such as tract-based spatial statistics (TBSS) and voxel-based morphometry (VBM).

1 1. Introduction

Group-wise registration of multi-dimensional unstructured point sets com-2 prising different types of data such as directional/axial and scalar-valued 3 quantities is useful for a variety of medical imaging and computer vision 4 applications. This study proposes a probabilistic approach for group-wise 5 registration of generalised point sets comprising positions, associated axial 6 orientations and scalar-valued measures. This is achieved through formulation of a hybrid mixture model (HdMM), combining suitable probability 8 distributions to model disparate data features within a cohesive framework. 9 As an exemplar application, the proposed framework is employed for the 10 joint registration and clustering of magnetic resonance (MR) diffusion tensor 11 image (DTI)-derived data, acquired from multiple subjects. The generality 12 of the proposed framework however, makes it suitable for registering other 13 types of hybrid point sets comprised of feature vectors containing principal 14 curvatures, surface normals, integral descriptors, etc. High-dimensional fea-15 ture vectors are in general more descriptive (than spatial positions alone, for 16 example) and discriminative when establishing correspondences, due to the 17 low probability of matching all features for non-corresponding points. 18

MR-DTI has found widespread use for studying structural changes within 19 brain white matter (WM), and the potential of such changes as biomarkers for 20 dementia and other neurodegenerative diseases. DT fields are estimated from 21 diffusion weighted images (DWIs), which encode diffusion of water molecules 22 along different gradient directions. MR-DTIs use a diffusion tensor model 23 (Basser et al., 1994) that, under some assumptions, can be related to lo-24 cal tissue microstructure. They aid in voxel-wise quantification of diffusion 25 characteristics, which may be expressed in terms of principal eigenvectors and 26 eigenvalues of the estimated diffusion tensors. Tissue microstructure affects 27 local diffusion properties. For example, water diffuses preferentially parallel 28 to the major axis of a fibre bundle, as opposed to perpendicular to it and, 20 consequently, gives rise to the sense of tissue anisotropy commonly observed 30 in major WM tracts. Fractional anisotropy (FA), a measure frequently em-31 ployed to describe tissue anisotropy (Pierpaoli and Basser, 1996), represents 32 the degree of directional dependence in diffusion at a specific voxel. The pri-33 mary eigenvector of a diffusion tensor represents the preferred direction for 34

the diffusion of water at any given voxel, and is often interpreted as reflecting
 the local fibre orientation within tissue.

Region of interest (ROI)-based analyses have been used to assess changes 37 in local (Salat et al., 2005) and global (Cercignani et al., 2001) tissue diffusion 38 properties. A limitation of such approaches is the need to accurately delin-39 eate ROIs across multiple patients'/subjects' images. Consequently, they 40 are affected by low reproducibility, leading to discrepancies across studies. 41 Tract-based spatial statistics (TBSS) (Smith et al., 2006) and voxel-based 42 morphometric (VBM) approaches (Ashburner and Friston, 2000) are suit-43 able alternatives that are fully automatic and enable analysis of localised 44 changes to FA and other diffusion measures, across the entire WM volume. 45 The quality of non-rigid registration used in VBM significantly influences 46 the subsequent voxel-wise analysis. To overcome this issue, (Smith et al., 47 2006) proposed the widely used TBSS approach, which ensures that registra-48 tion quality has less influence on subsequent statistical analysis of FA (and 49 other diffusion-derived quantities). TBSS constructs an alignment invariant 50 mean FA skeleton following registration of subjects' FA images to a template. 51 Neighbouring voxels located perpendicular to the skeleton are identified for 52 each subject, and the highest FA values are assigned to each skeleton voxel. 53 The resulting projections to the skeleton enable statistical analysis across 54 multiple subjects. 55

Alternative probabilistic techniques that jointly register and cluster WM 56 fibre trajectories (obtained from diffusion tractography), and which enable 57 quantitative analysis of diffusion measures over fibre pathways (rather than 58 voxel-wise quantification), have also been proposed. For example, registra-59 tion of curves and fibre bundles using diffeomorphisms and currents, and 60 a statistical framework to assess variability in geometry and fibre density 61 across a population, was proposed in (Durrleman et al., 2009), (Durrleman 62 et al., 2011). Maddah et al. (2008) employ a Gamma mixture modelling 63 framework to register fibre trajectories by establishing probabilistic corre-64 spondences, and jointly cluster them into representative fibre bundles. The 65 authors also note therein, through use of a suitable fibre tract atlas as a prior 66 during the clustering procedure, correspondences may be estimated across fi-67 bre trajectories obtained from multiple subjects, thereby enabling statistical 68 analysis of FA and other diffusion quantities across populations. Similarly, 60 (Mayer et al., 2011) proposed a supervised approach for joint registration 70 and segmentation WM tracts, wherein, the iterative closest fiber algorithm 71 (Mayer and Greenspan, 2008) was used to register fibre sets between a manu-72

ally annotated tractography atlas and a subject's reconstructed set of fibres. 73 The resulting segmentation was subsequently refined using a probabilistic 74 boosting tree-based classifier. In (Zvitia et al., 2010), the authors propose a 75 combined adaptive mean shift and Gaussian mixture model (GMM) formu-76 lation to jointly cluster fibre trajectories into compact fibre sets, and subse-77 quently register fibre sets obtained from multiple subjects. The registration 78 of two clustered fibre sets is formulated as a problem of aligning two distinct 79 GMMs, analogous to point set registration using GMMs (Jian and Vemuri, 80 2005). Similar approaches to clustering fibre trajectories across a population, 81 using spectral embedding, have also been proposed (O'Donnell and Westin, 82 2007), facilitating the estimation of WM atlases and enabling automatic seg-83 mentation of major WM tracts. An unbiased, group-wise, whole-brain trac-84 tography registration approach was proposed by (O'Donnell et al., 2012). 85 Kernel density estimation was used to approximate the probability distribu-86 tion of fibre trajectories within each brain and the overall distribution of the 87 atlas, was modelled as a mixture of the former. Alignment of WM tracts was 88 achieved by minimizing an entropic measure defined on the atlas distribution. 89 In a follow up study ODonnell et al. (2017), this group-wise registration ap-90 proach was combined with their previous work on spectral clustering of fibre 91 trajectories, to formulate an end-to-end automated framework for automated 92 WM tract identification, thereby enabling statistical analyses of DTI-derived 93 quantities. Garyfallidis et al. (2015) proposed a linear registration framework 94 to align WM bundles directly in the space of streamlines. They also demon-95 strated the viability of their approach to construct bundle specific atlases. 96 In a recent study (Benou et al., 2018), novel descriptors called Fiber-Flux 97 Diffusion Density (FFDD), which jointly describe fibre bundle geometry and 98 diffusivity measures were proposed, to facilitate localized quantification of 99 WM fibre bundles. Additionally, a FFDD dissimilarity measure was formu-100 lated and a novel registration framework (based on the fast marching method) 101 for WM tract-profiles was proposed, enabling inter-subject comparisons and 102 group-wise statistical analysis. Such techniques are however, dependent on 103 the tractography algorithm employed to estimate fibre trajectories, introduc-104 ing an additional potential source of error, and typically require some degree 105 of user intervention (to define seeds for streamline generation for example). 106

¹⁰⁷ Applications of the various methods described above have included, for ¹⁰⁸ example, identification of relationships between mild cognitive impairment ¹⁰⁹ (MCI) and Alzheimer's disease (AD), and localised changes to WM diffusion ¹¹⁰ characteristics. For example, in (Zhang et al., 2007), ROI-based analysis was used to identify significant reduction in FA in the cingulum for patients
diagnosed with MCI and AD, relative to healthy controls (HC). In (Medina
et al., 2006), VBM was used to identify significant reduction in FA in posterior regions of the brain, for MCI and AD patient groups, using VBM. While
(Liu et al., 2011) used the TBSS-approach and found reduced FA in the
cingulum, corpus callosal and inferior/superior longitudinal fasiculus tracts,
among others.

This study proposes a probabilistic approach to enable statistical anal-118 vsis of diffusion-derived measures, as an alternative to existing VBM- and 119 TBSS-based approaches. The latter are based on non-rigid registration of 120 subjects' FA images to a standard space to perform such analysis. Instead, 121 our approach uses group-wise non-rigid point set registration based on a 122 novel mixture modelling framework, which approximates the joint probabil-123 ity density of: (1) spatial positions (of voxel centroids within a region/tract 124 of interest), (2) primary diffusion axes (henceforth referred to as fibre orien-125 tations for brevity), and (3) fractional anisotropy, estimated at the voxels of 126 interest. The proposed framework is flexible and can be used to model other 127 diffusion-derived data such as mean/radial diffusivity, relative anisotropy, 128 tensor-eigenvalues, etc. — a functionality also afforded by TBSS. However, 129 the proposed approach also enables analysis of the variation in fibre orienta-130 tions, across multiple subjects, which is not possible with conventional TBSS 131 and VBM approaches. 132

Statistical analysis of fibre orientations across multiple subjects and com-133 parisons between patient groups was pursued in a previous study (Schwartz-134 man et al., 2005). Here, the authors followed a VBM-style approach where 135 DTIs from multiple subjects were spatially normalized to a reference template 136 using a spline-based tensor interpolation approach together with a tensor re-137 orientation mechanism designed to preserve the principal diffusion direction. 138 Subsequently, Watson distributions were fitted by maximum likelihood es-139 timation to the fibre orientations observed across a group, at each voxel, 140 independently. This provides a measure of the mean orientation and disper-141 sion, observed across the group of subjects. A drawback of such an approach 142 however, is the need to choose a single, appropriate template, for spatial 143 normalization, which is particularly difficult for images exhibiting varying 144 degrees of pathology-induced morphological changes. All subsequent reg-145 istrations performed and correspondences estimated are biased towards the 146 chosen template. VBM-based approaches in general, are dependent on the ac-147 curacy of non-rigid registration and the exact estimation of correspondences, 148

to ensure validity in the subsequent voxel-wise statistical analyses. TBSS 149 and our proposed approach are less restrictive in this regard. Registration of 150 WM regions defined by hybrid point sets (comprising voxel positions, asso-151 ciated fibre orientations and FA values) across subjects, is achieved using a 152 group-wise rigid, and subsequent non-rigid point set registration procedure, 153 based on a HdMM. In the proposed approach, correspondence probabilities 154 are estimated by approximating the joint probability density of position, 155 fibre orientation and FA, which are iteratively revised as the registration 156 progresses. Consequently, three distinct sources of information are leveraged 157 to guide the registration of an unbiased, study-specific atlas (iteratively re-158 vised as the registration progresses), onto each subject's WM tract/ROI. The 159 evolving soft correspondences provide model-based estimates for the mean fi-160 bre orientation and FA value (for a given population) at each component in 161 the mixture model and help mitigate any misalignment incurred during reg-162 istration. 163

164 1.1. Motivation and Contributions

The primary motivation for this study is to enable quantitative compar-165 isons of both voxel-wise scalar-valued (such as FA) and vector-valued (such as 166 position and orientation) DTI data, across multiple subjects. Although the 167 proposed framework is used to analyse voxel-wise diffusion-derived quantities 168 in this study, the method itself is not intrinsically dependent on voxel-wise 169 (or structured grid-wise) data, i.e. the framework could be used to register 170 and analyse unstructured data as well. The proposed hybrid mixture model 171 approximates the joint probability density function (PDF) of spatial posi-172 tions, associated fibre orientations and FA values, using Student's t, Watson 173 and Gaussian distributions, respectively. The proposed approach models the 174 *PDF* of fibre orientations, rather than the directions of the observed primary 175 diffusion eigenvectors, which tend to be random (as diffusion tensors are an-176 tipodally symmetric). To the best of our knowledge, this is the first study to 177 formulate such a hybrid mixture model-based registration framework, which 178 employs Watson distributions to model fibre orientations. 179

180 2. Methods

181 2.1. Pre-processing

MR-DWIs were acquired for 60 subjects (20 HC, 20 MCI, 20 AD), as part of prospective cohort of the VPH-DARE@IT project (vph-dare.eu).

All images used in this study were acquired using identical protocols: 2 184 diffusion-weighted b-values (0, 800), with diffusivity gradients applied along 185 32 directions; image size of $(240 \times 240 \times 120)$ slices, 2.5mm thick in the right-186 left, anterior-posterior and inferior-superior directions, respectively. DTIs 187 were estimated from these for each subject using TORTOISE v 2.5.0 (Pier-188 paoli et al., 2010), which employs state-of-the-art algorithms for motion and 189 eddy current correction, correcting B0 susceptibility induced EPI distortions 190 and B-matrix re-orientation artefacts. Tensor-fitting was then achieved us-191 ing iRESTORE (Chang et al., 2012), based on non-linear iterative least-192 squares. TORTOISE registers each subject's DWIs to their corresponding 193 T2-weighted structural MRI during the aforementioned pre-processing steps. 194 As the latter were acquired at resolutions of $(1.5 \times 1.5 \times 1.5 mm)$, all estimated 195 DTIs (and correspondingly, DTI-derived images) were up-sampled relative to 196 their raw DWIs. Finally, tensor-derived measures such as the eigenvector and 197 fractional anisotropy images were also estimated using TORTOISE. 198

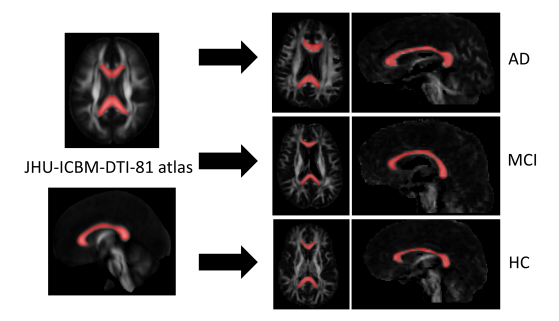


Figure 1: Nifty-Reg used to propagate labels for WM regions of interest from JHU-ICBM-DTI-81 atlas to each subject in AD, MCI and HC groups. Images depict propagation of the corpus callosum label from the atlas to subjects in AD, MCI and control groups.

¹⁹⁹ The proposed framework is flexible and can consider the entire WM vol-

ume as the region of interest, eliminating the need for pre-processing steps 200 in the form of a priori definition of the ROIs (using atlas-based label prop-201 agation for example). However, such an automated approach to analysing 202 the entire WM volume across multiple subjects carries significant computa-203 tional burden. Consequently, for the purpose of this study, we restrict our 204 attention to two WM regions, namely, the cingulum and corpus callosum. 205 An atlas-based label propagation approach is used to segment the WM ROIs 206 from all subjects' FA images. The fractional anisotropy image of the JHU-207 ICBM-DTI-81 atlas¹ (Mori et al., 2008) - (Hua et al., 2008) is non-rigidly 208 registered to each subject's FA image (following an initial affine alignment), 209 using Nifty-Reg v 1.3.9 (Ourselin et al., 2001), (Modat et al., 2010), a de-210 formable image registration algorithm based on cubic B-splines. Following 211 FA image registration, the segmented labels for the cingulum and corpus cal-212 losum defined on the atlas (available along with the FA atlas), are resampled 213 to the space of each subject's FA image. In this way, labels delineating the 214 cingulum and corpus callosum in the atlas image, are propagated to each 215 subject's image, segmenting the ROIs (as illustrated in Fig. 1). 216

¹Available at: http://www.loni.usc.edu/ICBM/Downloads/Downloads^{*}DTI-81.shtml

217 2.2. Algorithm Overview

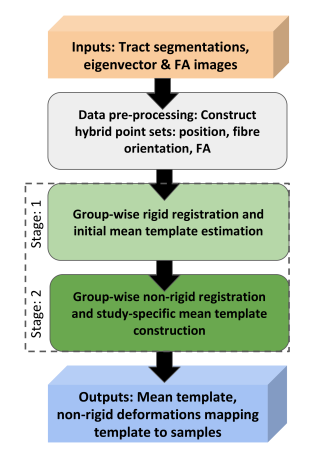


Figure 2: Summary of steps involved in the proposed framework to jointly register and cluster hybrid point sets comprising spatial positions, fibre orientations and FA values, for a WM tract/ROI. Dashed box outlines the two stages of the proposed algorithm.

The steps involved in the proposed approach are summarised by Fig. 2. 218 For a group of k = 1...K subjects to be analysed (e.g. comprising control, 219 MCI and AD sub-groups), their tract segmentations, eigenvector and FA im-220 ages were used to construct hybrid point sets \mathbf{D}_k , where each data point is a 221 7-dimensional vector denoted as $\mathbf{d}_{ki} = [\mathbf{x}_{ki}, \mathbf{n}_{ki}, f_{ki}]$. Here \mathbf{x}_{ki} represents the 222 spatial co-ordinate, \mathbf{n}_{ki} represents the primary diffusion eigenvector and f_{ki} 223 denotes the FA value for the i^{th} voxel, in the k^{th} subject's image. \mathbf{x}_{ki} are 224 consequently, densely distributed points within the volumes/ROIs. The re-225 sulting hybrid point sets were, subsequently, jointly registered and clustered 226

by fitting an *M*-component hybrid mixture model (comprising Student's t, 227 Watson and Gaussian distributions) to the data. This was achieved over 228 two stages (as depicted in Fig. 2): (1) Group-wise rigid registration of the 229 hybrid point sets \mathbf{D}_k and mean template **M** construction; and (2) Group-230 wise non-rigid registration, wherein the mean template estimated in stage 231 1 was non-rigidly registered to each sample from all patient groups simul-232 taneously. The similarity transformation and the non-rigid transformation, 233 corresponding to stage 1 and 2 of the algorithm respectively, are both repre-234 sented by \mathbf{T}_k throughout this study. For the former, $\mathbf{T}_k = [s_k, \mathbf{R}_k, \mathbf{t}_k]$. Here, 235 $s_k, \mathbf{R}_k, \mathbf{t}_k$ represent the scaling, rotation and translation (for the k^{th} sam-236 ple), respectively, estimated in stage 1. These are used to align the hybrid 237 point sets to the estimated mean template and initialise the subsequent non-238 rigid registration step (stage 2) by correcting global pose differences across 239 the data set. Stage 2 of the algorithm estimates non-rigid transformations 240 \mathbf{T}_k , defined by a linear combination of radial basis functions (with a Gaus-241 sian kernel). Together with a Gaussian kernel, the basis function weights 242 \mathbf{W}_k estimated define point-wise displacements that map the mean template 243 to each sample within a subject group. In both stages of the algorithm, 244 estimation of the desired registration parameters was accompanied by the 245 joint clustering of positions, orientations and FA values. The parameters to 246 be estimated for each of the j = 1...M components of the hybrid mixture 247 model include: $\{\mathbf{m}_{i}^{p}, \sigma_{p}^{2}, \nu_{j}\} = \Theta_{p}$, which represent mean spatial positions, 248 their variance and the degrees of freedom, respectively, for the Student's t-249 distributions; $\{\mathbf{m}_{i}^{d}, \kappa_{i}\} = \Theta_{n}$, which represent the mean fibre orientations 250 and concentration around the means, respectively, for the Watson distribu-251 tions; $\{m_i^f, \sigma_f^2\} = \Theta_f$, which denote the mean FA values and FA variance, 252 respectively, for the Gaussian distributions; and π_j which denote the mix-253 ture coefficients. Following non-rigid registration, the study-specific mean 254 template estimated (for each WM ROI) M thus comprises positions, \mathbf{m}_{i}^{p} 255 orientations \mathbf{m}_{i}^{d} and FA values m_{i}^{f} . 256

257 2.3. Joint Probabilistic Model of Position, Orientation and Anisotropy

The problem of joint registration and clustering of hybrid point sets is formulated as one of maximum likelihood parameter estimation, using a hybrid mixture model that approximates the joint *PDF* of spatial positions (of voxel centroids), fibre orientations, and fractional anisotropy. By assuming voxel positions, fibre orientations, and FA values to be independent and identically distributed (i.i.d), for each subject and across multiple subjects, the

joint PDF can be approximated as a product of the individual conditional 264 densities (Bishop, 2006) for position, orientation and FA. Consequently, by 265 considering all data points $\mathbf{d}_{ki} \in \mathbf{D}_k$, from all K subjects, to be i.i.d. the con-266 ditional probability of an observation being sampled from an *M*-component 267 HdMM is given by equation 1a. The set of all transformations (similarity 268 or non-rigid) is represented by $\mathbf{T}_k \in \mathbb{T}$; Θ_p represents the set of model pa-269 rameters associated with the Student's t-distributions \mathcal{S} , used to model the 270 distribution of voxel spatial positions; Θ_n represents the parameters of the 271 Watson distributions \mathcal{W} (modelling fibre orientations); Θ_f denotes the set 272 of parameters of the Gaussian distributions \mathcal{N} (modelling FA); and $\pi_i \in \Pi$ 273 represents the set of mixture coefficients, of the HdMM. Here and through-274 out, subscript i = 1...M denotes mixture components and the choice of 275 distributions indicated earlier will be justified later in this Section. Using 276 equation (1a) the log-likelihood function is formulated as shown in equation 277 (1b), which defines the cost function to be optimised with respect to the mix-278 ture model and transformation parameters $\{\Theta_p, \Theta_n, \Theta_f \Pi, \mathbb{T}\} \in \Psi$, to jointly 279 register and cluster the hybrid point set data $\mathbf{D}_k \in \mathbb{D}$. 280

$$p(\mathbf{d}_{ki}|\Theta_p,\Theta_n,\Theta_f,\mathbf{T}_k) = \sum_{j=1}^M \pi_j \mathcal{S}(\mathbf{x}_{ki}|\Theta_p,\mathbf{T}_k) \mathcal{W}(\mathbf{n}_{ki}|\Theta_n,\mathbf{T}_k) \mathcal{N}(f_{ki}|\Theta_f,\mathbf{T}_k)$$
(1a)

$$\ln p(\mathbb{D}|\Psi) = \sum_{k=1}^{K} \sum_{i=1}^{N_k} \ln p(\mathbf{d}_{ki}|\Theta_p, \Theta_n, \Theta_f, \mathbf{T}_k)$$
(1b)

$$P_{kij}^{t} = \frac{\pi_{j} p(\mathbf{d}_{ki} | \Theta_{p}^{t}, \Theta_{n}^{t}, \Theta_{f}^{t}, \mathbf{T}_{k})}{\sum_{l=1}^{M} \pi_{l} p(\mathbf{d}_{ki} | \Theta_{p}^{t}, \Theta_{n}^{t}, \Theta_{f}^{t}, \mathbf{T}_{k}^{t})}$$
(1c)

$$Q(\Psi^{t+1}|\Psi^{t}) = \sum_{k=1}^{K} \sum_{i=1}^{N_{k}} \sum_{j=1}^{M} P_{kij}^{t} \Big[\ln \pi_{j} + Q(\Theta_{p_{j}}^{t+1}, \mathbf{T}_{k}^{t+1}|\Theta_{p_{j}}^{t}, \mathbf{T}_{k}^{t}) + Q(\Theta_{n_{j}}^{t+1}, \mathbf{T}_{k}^{t+1}|\Theta_{n_{j}}^{t}, \mathbf{T}_{k}^{t}) + Q(\Theta_{f_{j}}^{t+1}, \mathbf{T}_{k}^{t+1}|\Theta_{f_{j}}^{t}, \mathbf{T}_{k}^{t}) \Big]$$
(1d)

Algorithm 1 Hybrid Mixture Model: HdMM

Inputs: Group of hybrid point sets $\mathcal{D}_{k=1K}$, number of mixture
components M, max.iterations
Outputs: Set of HdMM parameters $\{\Theta_p,\Theta_n,\Theta_f\}$ \in $\Psi,$ soft
correspondences
1: INITIALIZATION
2: Initialize $\mathbf{M}, \sigma_p^2, \sigma_f^2$ using K-means clustering.
3: All $\pi_j = 1/M$ and $\nu_j = 3.0, \kappa_j = 1.0$
4: procedure Stage 1 EM:
5: GROUP-WISE RIGID REGISTRATION $(\mathbf{D}_k, \Theta_p, \Theta_n, \Theta_f, \Pi, \mathbf{T}_k)$ \triangleright EM
initialized
6: while Iteration < max.iterations do
7: Compute P_{kij} \triangleright E-step
8: Update $\mathbf{R}_k, s_k, \mathbf{t}_k$ \triangleright M-step
9: Update $\Theta_p, \Theta_n, \Theta_f$ \triangleright M-step
10: end while
11: return $\Theta_p, \Theta_n, \Theta_f, \Pi, \mathbf{T}_k$
12: end procedure
13: Estimated mean template \mathbf{M} , mixture coefficients Π and similarity trans-
formations $\{\mathbf{T}_k\}_{k=1K}$ initialise Stage 2.
14: procedure Stage 2 EM:
15: GROUP-WISE NON-RIGID REGISTRATION $(\mathbf{D}_k, \Theta_p, \Theta_n, \Theta_f, \Pi, \mathbf{W}_k) \triangleright \mathrm{EM}$
non-rigid initialized
16: while Iteration $< \max$.iterations do
17: Compute P_{kij} \triangleright E-step
18: Update \mathbf{W}_k \triangleright M-step
19: Update $\mathbf{M}, \sigma_p^2, \nu_j, \Theta_n, \Theta_f$ \triangleright M-step
20: Update spatial positions of each \mathbf{D}_k
21: end while
22: return $\Theta_p, \Theta_n, \Theta_f, \Pi, \mathbf{W}_k$
23: end procedure

```
A tractable approach to maximising equation 1b is achieved using the
expectation-maximisation (EM) framework (Dempster et al., 1977), which
iteratively alternates between: the expectation (E)-step, which evaluates the
mixture component membership probabilities as shown in equation 1c (i.e.
posterior probabilities P_{kij}^t, that define soft correspondences and are expec-
```

tations of the latent variables in the model) for the observed data, given 286 an estimate of the model parameters Ψ^t , at the tth EM-iteration; and the 287 maximisation (M)-step, which uses the computed posterior probabilities P_{kij}^t 288 to maximise the conditional expectation of the complete-data-log-likelihood 289 function Q (refer to equation 1d), with respect to each model parameter, 290 resulting in revised estimates Ψ^{t+1} . As shown in equation 1d, Q for the hy-291 brid mixture model can be expressed as a sum of contributions from each 292 distribution and corresponding data feature (i.e. position, orientation and 293 FA), denoted, $Q(\Theta_p^{t+1}|\Theta_p^t), Q(\Theta_n^{t+1}|\Theta_n^t), Q(\Theta_f^{t+1}|\Theta_f^t)$, respectively. The com-294 plete algorithm for the proposed hybrid mixture model, to jointly register 295 and cluster a group \mathbb{D} of hybrid point sets, is summarized in Algorithm 1. 296 Subsequent sections discuss each probability distribution and estimation of 297 their associated parameters, within the proposed framework, in more detail. 298

299 2.4. Mixture Model for Primary Diffusion Axes

In addition to modelling the spatial distribution of voxels defining ROIs, 300 the proposed approach also deals with axial data distributed over the S^2 301 sphere, i.e. fibre orientations defined by primary diffusion eigenvectors. 302 GMMs and TMMs, comprising Gaussian and Student's t-distributions, re-303 spectively, are inappropriate for clustering such data and consequently, a 304 mixture of Watson distributions, also defined over the spherical domain, is 305 employed in this study. While Von-Mises-Fisher distributions are frequently 306 used for clustering directional data, they are unsuitable for axial data, as 307 they lack of antipodal symmetry. Watson distributions on the other hand, 308 are naturally suited to model diffusion data as they are antipodally symmet-309 ric (i.e. the probability density is the same along an axis in either direction) 310 and as the aim here is to model the PDF of diffusion axes at correspond-311 ing spatial locations, rather than any specific direction along the axes (Jupp 312 and Mardia, 1989). They are fully defined by two parameters, namely, the 313 mean/principal axis ($\pm \mathbf{m}^d$, about which the distribution is rotationally sym-314 metric) and a scalar concentration parameter κ . The latter describes the 315 degree of concentration about the mean axis of the distribution, with high 316 values indicating high concentration. The PDF of a Watson distribution 317 with mean direction \mathbf{m}^d and concentration κ is expressed as equation 2a, 318 for antipodally symmetric 3D unit vectors $\pm \mathbf{n}$. Here, $M(\cdot)$ represents the 319 Kummer function. Watsons are in general more flexible than Fisher distri-320 butions as there is no positivity constraint on κ and they can be used to 321 model both directional and axial data. (Bijral et al., 2007) proposed an ef-322

ficient EM-based clustering framework for axially-distributed data, using a
 WMM, employed in this study to cluster fibre orientations.

$$p(\pm \mathbf{n} | \mathbf{m}^d, \kappa) = M(\frac{1}{2}, \frac{D}{2}, \kappa)^{-1} \exp^{\kappa (\mathbf{m}^{d^T} \mathbf{n})^2}$$
(2a)

$$p(\mathbb{N}|\Theta_n) = \sum_{k=1}^{K} \sum_{i=1}^{N_k} \ln \sum_{j=1}^{M} \pi_j p(\pm \mathbf{n}_{ki} | \mathbf{m}_j^d, \kappa_j)$$
(2b)

The joint likelihood of the diffusion eigenvectors $\pm \mathbf{n}_{ki} \in \mathbf{N}_k$ observed 325 across all N_k points in all K hybrid point sets, given Watson distributions 326 with mean directions and concentrations $\{\mathbf{m}_{j}^{d}, \kappa_{j}\}_{j=1...M} \in \Theta_{n}$, is evaluated as 327 shown in equation 2b. Here, $\mathbf{N}_k \in \mathbb{N}$ denotes the set of all observed diffusion 328 vectors across the entire population. It is important to note at this point 329 that, as the clustering of fibre orientations is initially performed jointly with 330 rigid registration of the hybrid point sets \mathbf{D}_k , the estimated rotations $\mathbf{R}_k^{(t)}$ at the t^{th} EM-iteration, are applied to the current estimate of the mean fibre 331 332 orientations $\mathbf{m}_{j}^{d^{(t)}}$, prior to the evaluation of the posterior probabilities P_{kij} , 333 and concentrations κ_i , in the E- and M-steps, respectively. Additionally, for 334 the estimation of \mathbf{m}_{i}^{d} the inverse of the estimated rotations \mathbf{R}_{k}^{T} were applied 335 to their corresponding sample's diffusion eigenvectors \mathbf{n}_{ki} , to align the k^{th} 336 sample to the current estimate of the mean template (refer to equation 3c). 337

$$Q(\Theta_n^{t+1}|\Theta_n^t) = \sum_{k=1}^K \sum_{i=1}^{N_k} \sum_{j=1}^M P_{kij}^{(t)} \ln p(\pm \mathbf{n}_{ki} | \mathbf{R}_k^{(t)} \mathbf{m}_j^{d^{(t)}}, \kappa_j^{(t)})$$
(3a)

$$Q(\Theta_{n}^{t+1}|\Theta_{n}^{t}) = \sum_{k=1}^{K} \sum_{i=1}^{N_{k}} \sum_{j=1}^{M} [P_{kij}^{(t)} \ln p(\pm \mathbf{n}_{ki} | \mathbf{R}_{k}^{(t)} \mathbf{m}_{j}^{d}, \kappa_{j}) +$$
(3b)
$$\lambda_{j}(1 - (\mathbf{R}_{k}^{(t)} \mathbf{m}_{j}^{d})^{T} \mathbf{R}_{k}^{(t)} \mathbf{m}_{j}^{d})]$$
$$\mathbf{m}_{j}^{d^{(t)}} - \frac{\sum_{k=1}^{K} \sum_{i=1}^{N_{k}} P_{kij}^{(t)}((\mathbf{R}_{k}^{T^{(t+1)}} \mathbf{n}_{ki})^{T} \mathbf{m}_{j}^{d^{(t)}}) \mathbf{R}_{k}^{T^{(t+1)}} \mathbf{n}_{ki}}{||\sum_{k=1}^{K} \sum_{i=1}^{N_{k}} P_{kij}^{(t)}((\mathbf{R}_{k}^{T^{(t+1)}} \mathbf{n}_{ki})^{T} \mathbf{m}_{j}^{d^{(t)}}) \mathbf{R}_{k}^{T^{(t+1)}} \mathbf{n}_{ki}||} = 0$$
(3c)

$$\left[\frac{M'(\kappa_j)}{M(\kappa_j)}\right]^{(t+1)} = \frac{\sum_{k=1}^{K} \sum_{i=1}^{N_k} P_{kij}^{(t)} (\mathbf{n}_{ki}^T \mathbf{m}_j^{d^{(t+1)}})^2}{\sum_{k=1}^{K} \sum_{i=1}^{N_k} P_{kij}^{(t)}}$$
(3d)

$$\kappa_{j}^{(t+1)} \approx \frac{1}{2} \left[\frac{1 - \left[\frac{M'(\kappa_{j})}{M(\kappa_{j})}\right]^{(t+1)} D}{\left[\left(\frac{M'(\kappa_{j})}{M(\kappa_{j})}\right)^{2}\right]^{(t+1)} - \frac{[M'(\kappa_{j})}{M(\kappa_{j})}\right]^{(t+1)}} \right]$$
(3e)

Maximum likelihood estimates for the associated parameters are evalu-338 ated at each M-step of the algorithm by maximising the expectation of the 339 complete data likelihood (equation 3a), with respect to \mathbf{m}_{i}^{d} and κ_{j} , subject 340 to the constraint $\mathbf{m}_j^{d^T} \mathbf{m}_j^d = 1$ (Bijral et al., 2007). This is achieved by max-341 imising the Lagrangian form of Q shown in equation 3b. Mean directions \mathbf{m}_{i}^{d} 342 are estimated numerically, using fixed-point iteration, to solve the non-linear 343 equation (shown in equation 3c) obtained from differentiating Q (3b) with 344 respect to \mathbf{m}_{i}^{d} . κ_{i} on the other hand is approximated (refer to equation 3e) 345 using the continued fraction representation for the ratio of, the derivative 346 $\frac{M'(\kappa_j)}{M(\kappa_j)}$ (equation 3d). of the Kummer function and the function itself, i.e. 347 In a recent study (Sra and Karp, 2013) derived two-sided bounds for ap-348 proximating κ , particularly useful when dealing with high dimensional data. 349 However, for 3D data (as in this study) the approximation presented in equa-350 tion 3e is sufficient (as noted by (Bijral et al., 2007), (Sra and Karp, 2013)). 351 Better approximations for κ_i may be obtained using numerical techniques 352 such as Newton's method, however, at the expense of significant increase in 353 computational burden. 354

³⁵⁵ 2.5. Mixture Model for Fractional Anisotropy

The distribution of voxel-wise FA in WM ROIs across a population, is 356 modelled using a univariate GMM. GMM was chosen as the resulting model-357 predicted FA values at the estimated spatial correspondences, across subjects, 358 is guaranteed to be normally distributed — a useful property for subsequent 359 statistical analyses, as noted in (Smith et al., 2006), where the authors also 360 show that FA values at corresponding spatial positions across populations are 361 indeed approximately normally-distributed. Additionally, GMMs are com-362 putationally efficient, as analytical solutions exist for revising estimates of 363 the associated model parameters (mean m_j^f and variance σ_f^2 of FA), at each 364 EM-iteration. Assuming the observed FA values f_{ki} at voxels in ROIs, across 365

a group of subjects $\mathbf{F}_k \in \mathbb{F}$ are i.i.d, the joint log-likelihood log $p(\mathbb{F}|\Theta_f)$, is 366 expressed as equations 4a, 4b. Consequently, the conditional expectation of 367 the complete data log likelihood Q, maximised with respect to the model pa-368 rameters associated with the Gaussian distributions in the mixture, is given 369 by equation 4c (only terms dependent on m_i^f and σ_f^2 are retained in Q). As 370 GMM-based clustering of FA values is performed jointly with the registra-371 tion of WM ROIs, and clustering of voxel positions and the associated fibre 372 orientations, the influence of a Gaussian component in the mixture model 373 is automatically limited to its local neighbourhood. This helps ensure that 374 only voxels in close proximity to each other contribute significantly to the 375 estimation of mean FA values at each mixture component. Estimates for 376 the GMM parameters m_j^f and σ_f^2 in the M-step of the algorithm are derived 377 analytically, as shown in (Bishop, 2006). 378

$$p(\mathbf{F}_{k}|m_{j}^{f},\sigma_{f}^{2}) = \prod_{i=1}^{N_{k}} \sum_{j=1}^{M} \pi_{j} \mathcal{N}(f_{ki}|m_{j}^{f},\sigma_{f}^{2})$$
(4a)

$$\ln p(\mathbb{F}|\Theta_f) = \sum_{k=1}^{K} \ln p(\mathbf{F}_k|\Theta_f)$$
(4b)

$$Q(\Theta_f^{t+1}|\Theta_f^t) = -\frac{1}{2} \sum_{k=1}^K \sum_{i=1}^{N_k} \sum_{j=1}^M P_{kij}^t \Big[\frac{(f_{ki} - m_j^f)^2}{\sigma_f^2} \Big]$$
(4c)

379 2.6. Rigid Alignment and Template Construction

Previously, we proposed a group-wise rigid point set registration frame-380 work based on Student's t-mixture model (Ravikumar et al., 2016), (Raviku-381 mar et al., 2018), which exploits the inherent robustness of Student's t-382 distribution for robust registration of shapes in the presence of missing data 383 and significant proportions of outliers. Additionally, in a more recent study 384 (Ravikumar et al., 2017) we proposed a variant of the hybrid mixture model-385 based registration framework formulated in this study. In (Ravikumar et al., 386 2017) Von-Mises-Fisher distributions were used in place of the Watson distri-387 butions used in this study, to model directional data such as surface normal 388 vectors, for rigid and non-rigid shape registration. A Watson distribution-380 based variant of (Ravikumar et al., 2017) is employed in the present study 390 as an initial step, to rigidly align WM ROIs (hybrid point sets representing 391 voxel centroid positions, fibre orientations and FA values), segmented from 392

all subjects' images, whilst simultaneously estimating a mean model. The 393 latter subsequently serves as an unbiased, study-specific template for non-394 rigid registration. Rigid group-wise registration is preferred to a pair-wise 395 approach as it enables estimation of a mean template and the desired sim-396 ilarity transformations in an unbiased manner. Rigid alignment also helps 397 initialise the subsequent non-rigid registration by recovering global differ-398 ences in pose between sample shapes, and establishes soft correspondences 399 across subjects. 400

Group-wise point set registration using mixture models assumes that the 401 point sets to be aligned are transformed observations of a central mixture 402 model (which we refer to as the mean template) (Gooya et al., 2015). Con-403 sequently, the optimal transformations that align the template to the group 404 of shapes are those that maximise the likelihood of the data (or equivalently, 405 minimise the negative log-likelihood function). The desired similarity trans-406 formations are thus iteratively refined along with the template itself at each 407 M-step of the algorithm. The main differences between EM-based estima-408 tion of parameters for TMMs and GMMs are: (1) TMMs have two associated 409 latent variables (as opposed to just one with GMMs, which represent the mix-410 ture component membership of the data), whose expectations are evaluated 411 in the E-step and used to compute a set of corrected posterior probabilities 412 P_{kij}^{\star} , estimated identically to (Ravikumar et al., 2016), (Ravikumar et al., 413 2018) (refer to the Appendix); and (2) Student's t-distributions are defined 414 by three parameters (as opposed to two for Gaussians). The additional pa-415 rameter is referred to as the degrees of freedom/shape parameter ν , which is 416 responsible for controlling the heaviness of the tails of the distribution (and 417 consequently, the degree of robustness to outliers). The behaviour of the 418 t-distribution tends towards that of a Gaussian as $\nu \to \infty$. 419

$$\log p(\mathbb{X}|\Theta_p, \mathbb{T}) = \sum_{k=1}^{K} \sum_{i=1}^{N_k} \log \sum_{j=1}^{M} \pi_j \mathcal{S}(\mathbf{x}_{ki} | \mathbf{T}_k(\mathbf{m}_j^p), \sigma_p^2, \nu_j)$$
(5a)

$$Q(\Theta_p^{t+1}, \mathbb{T}^{t+1} | \Theta_p^t, \mathbb{T}^t) \propto -\frac{1}{2\sigma_p^2} \sum_{k=1}^K \sum_{i=1}^{N_k} \sum_{j=1}^M P_{kij}^{\star t} \| \mathbf{x}_{ki} - s_k \mathbf{R}_k \mathbf{m}_j^p - \mathbf{b}_k \|^2$$
(5b)

⁴²⁰ The joint *PDF* of voxel positions $\mathbf{x}_{ki} \in \mathbf{X}_k$, across all *K* subjects in ⁴²¹ a group (denoted, $\mathbf{X}_k \in \mathbb{X}$), is given by equation 5a (assuming they are ⁴²² i.i.d transformed observations of a TMM). In equation 5a, \mathbf{T}_k represents the

similarity transformation (comprising rotation \mathbf{R}_k , scaling s_k and translation 423 \mathbf{b}_k), to align the positions \mathbf{m}_i^p defining the mean template, to the k^{th} sample 424 in the group. In our recent work (Ravikumar et al., 2016), (Ravikumar et al., 425 2018), we showed that the form of Q to be maximised, to estimate the desired 426 similarity transformations $\mathbf{T}_k \in \mathbb{T}$ and mixture component parameters Θ_p , 427 is given by equation 5b. Closed form expressions are derived for the M-428 step update equations of all TMM and transformation parameters, which 429 are presented in the Appendix. Fibre orientations and FA are invariant to 430 translation \mathbf{b}_k and scaling s_k , consequently, these transformation parameters 431 are estimated identically as in (Ravikumar et al., 2016), (Ravikumar et al., 432 2018). Although the former are rotationally dependent, the contribution of 433 fibre orientations to the estimation of \mathbf{R}_k is ignored as the direction of the 434 observed diffusion eigenvectors tend to be random. Consequently, rotations 435 \mathbf{R}_k are derived based on the spatial positions of hybrid point sets alone, 436 by optimising the form of Q shown in equation 5b, similar to (Ravikumar 437 et al., 2016), (Ravikumar et al., 2018). However, following estimation of 438 the desired rotations \mathbf{R}_k at each EM-iteration, the current estimate of the 439 mean template is transformed by rotating both spatial positions \mathbf{m}_{i}^{p} and 440 their associated fibre orientations \mathbf{m}_{i}^{d} , to align it with the k^{th} sample in the 441 group. Additionally, it is important to note that, while the fibre orientations 442 and FA values are ignored in the derivation of the desired transformation 443 parameters, they are intrinsic to the estimation of the posterior probabilities 444 P_{kij} at each E-step of the algorithm. Consequently, they drive the estimation 445 of soft correspondences, which in turn affect the transformations evaluated 446 at each M-step of the algorithm. 447

448 2.7. Non-rigid Point Set Registration

Coherent point drift (CPD) (Myronenko and Song, 2010) is a well known 449 pair-wise, non-rigid point set registration technique based on motion coher-450 ence theory. The spatial transformation between two point sets is considered 451 to be an initial position (of the moving point set) plus some unknown dis-452 placement (or velocity) function mapping it to the target point set. This un-453 known transformation is regularized using Tikhonov regularization, to ensure 454 estimation of a smooth displacement function, and is expressed in the Repro-455 ducing Kernel Hilbert Space (RKHS). Using variational calculus, Myronenko 456 and Song (2010) showed that the optimal displacement function under such 457 smoothness constraints, can be expressed as a linear combination of kernel 458 functions (i.e. Gaussian radial basis functions). Similarly, our approach also 459

employs Gaussian radial basis functions to parametrize the non-linear trans-460 formation, and the associated basis function weights are estimated by max-461 imising the likelihood function using EM (similar to estimation of rotation, 462 translation and scaling, in the rigid registration approach discussed in the 463 previous section). CPD models the target point set as a transformed obser-464 vation of the source point set (i.e. the point set to be registered). The latter 465 is consequently considered to represent the centroids of a Gaussian mixture 466 model, which is fit to the former using EM, and the transformation necessary 467 to register the source to the target set is estimated as parameters of the mix-468 ture model. In addition to the Gaussian components in the mixture model, 469 CPD incorporates a uniform distribution component to model noise/outliers 470 present in the data. This confers added robustness to the registration pro-471 cess. However, a user-defined parameter is used to balance the weight of the 472 uniform distribution component relative to its Gaussian counterparts, which 473 needs to be tuned for different applications and data sets, for optimal regis-474 tration. To ameliorate the need for parameter tuning, we employ Student's 475 t-distributions in place of the Gaussian and uniform distributions used in 476 CPD and re-formulate the approach in a group-wise non-rigid registration 477 framework. As stated previously, the robust nature of t-distributions makes 478 them well suited to registration applications requiring automatic robustness 479 to outliers. A similar approach for pair-wise registration of 2D/3D point sets 480 was proposed previously, by (Zhou et al., 2014). 481

The mean tract template estimated during the initial group-wise rigid 482 registration step (discussed in section 2.6), is non-rigidly registered to each 483 patient group (AD, MCI and HC) independently. The desired non-rigid 484 transformations are defined with respect to the template \mathcal{M} as: $\mathbf{M} + v^k(\mathbf{M})$ 485 (considering spatial positions \mathbf{m}_{i}^{p} alone), where v is a displacement func-486 tion mapping the template to the k^{th} sample in the group. In (Myronenko 487 and Song, 2010) the authors show that the desired displacement field is con-488 strained to be smooth by employing Tikhonov regularization (or regularizing 489 the norm of v, expressed in RKHS). This forces points in close proximity, to 490 move together. Regularization of this nature is akin to employing a prior on 491 the displacement field of the form $p(v) = \exp^{-\frac{\lambda}{2}\phi(v)}$, where $\phi(v)$ represents the 492 regularization term and λ controls the trade-off between registration accuracy 493 and smoothness of the deformation field. The prior on the displacement field 494 is incorporated into the TMM, resulting in a log-likelihood function expressed 495 as equation 6a. As stated previously, (Myronenko and Song, 2010) show that 496

the function v, which maximises the data likelihood, can be expressed as a 497 linear combination of radial basis functions (refer to equation 6b). Conse-498 quently, to register the study-specific mean template to each sample from 499 all patient groups simultaneously, the objective function to be maximised 500 with respect to the basis function weights $w_{kj} \in \mathbf{W}_k$, is expressed as shown 501 in equation 6c, where \mathbf{G} represents the Gaussian kernel/Gram matrix. The 502 basis function weights required to register the study-specific mean template 503 to each sample are estimated as shown in 6d, by computing the derivative of 504 Q with respect to the weights, similarly to (Myronenko and Song, 2010). In 505 equation 6d $\mathbf{P}_k^s = \sum_{i=1}^{N_k} P_{kij}^{\star t}$, \mathbf{P}_k^T is the transpose of the posterior probability matrix for the k^{th} sample, diag is a diagonal matrix, and **I** is the identity 506 507 matrix. Subsequently, the mean template is deformed to match each k^{th} 508 sample (in the entire population) as described by equation 6e. 509

$$\log p(\mathbb{X}|\Theta_p) = \sum_{k=1}^{K} \sum_{i=1}^{N_k} \log \sum_{j=1}^{M} \pi_j \mathcal{S}(\mathbf{x}_{ki}|v^k(\mathbf{m}_j^p), \sigma^2, \nu_j) + \frac{\lambda}{2} \phi(v^k) \quad (6a)$$

$$v^{k}(\mathbf{q}) = \sum_{j=1}^{M} w_{kj} G(\mathbf{q} - \mathbf{m}_{j}^{p})$$
(6b)

$$Q(\Theta_p^{t+1}, \mathbf{W}_k^{t+1} | \Theta_p^t, \mathbf{W}_k^t) = -\frac{1}{2\sigma_p^2} \sum_{i=1}^{N_k} \sum_{j=1}^M P_{kij}^{\star t} \| \mathbf{x}_{ki} - (\mathbf{m}_j^p + v^k(\mathbf{m}_j^p)) \|^2 + \frac{\lambda}{2} \mathbf{W}_k^T \mathbf{G} \mathbf{W}_k$$
(6c)

$$\mathbf{W}_{k}^{(t+1)} = [\operatorname{diag}(\mathbf{P}_{k}^{s^{t}})\mathbf{G} + \lambda\sigma_{p}^{2}\mathbf{I}]^{-1}[\mathbf{P}_{k}^{T^{t}}\mathbf{X}_{k} - \operatorname{diag}(\mathbf{P}_{k}^{s^{t}})\mathbf{M}^{t}]$$
(6d)

$$\mathbf{M}_{k}^{(t+1)} = \mathbf{T}_{k}^{t}(\mathbf{M}_{k}^{t}, \mathbf{W}_{k}^{t}) = \mathbf{M}_{k}^{t} + \mathbf{G}\mathbf{W}_{k}^{t}$$
(6e)

Following convergence of the non-rigid registration step, a study-specific mean template comprising, mean spatial positions, mean fibre orientations and mean FA values (representative of the entire population of AD, MCI and HC subjects), is estimated. Additionally, point-wise displacements mapping this mean template to each sample in the entire population (as described

by equation 6e), is also obtained, thereby establishing the spatial correspon-515 dences used for any subsequent inter-group statistical comparisons. These 516 correspondences play a similar role to the mean FA skeleton estimated in 517 TBSS. In addition to these spatial correspondences, we also compute "model-518 predicted" values for FA and fibre orientation, at each correspondence, for all 519 subjects. These model-predicted values are probabilistic weighted averages 520 of the FA values and fibre orientations associated with the voxels in the orig-521 inal DTI-derived FA and eigenvector images (i.e. the original hybrid point 522 sets). The weighted averages are assigned to each spatial correspondence 523 point and are analogous to the 'soft/probabilistic spatial correspondences' 524 estimated in previous studies, such as (Hufnagel et al., 2008), (Gooya et al., 525 2015) for example. Here, the weights are defined by the posterior probabil-526 ities estimated for each voxel, of each subject's original FA and eigenvector 527 images (P_{kij}) , following non-rigid registration. Equations describing the es-528 timation of model-predicted FA values and fibre orientations are included in 529 the Appendix (refer to equations 19a - 19b). Although point set registration 530 techniques are typically employed to register 3D point sets (comprising only 531 spatial positions) representing the surface/boundary of an object, this study 532 incorporates additional image-based features (such as fibre orientations and 533 FA values), that enable registration of dense point sets, defined by voxel 534 centroids located at the boundary of, and within a region of interest. 535

⁵³⁶ 3. Results and Discussion

537 3.1. Rigid Registration Accuracy

Rigid registration accuracy of the proposed framework and the robustness of Student's t-distributions to outliers is assessed using synthetic data comprising point sets containing positions, associated fibre orientations and FA values. The synthetic data set was generated by rigidly transforming a corpus callosum hybrid point set by varying amounts. Four distinct synthetic samples (Samples 1-4) were generated in this manner from the original ground truth point set (referred to as Sample 0), as illustrated by Fig. 3.

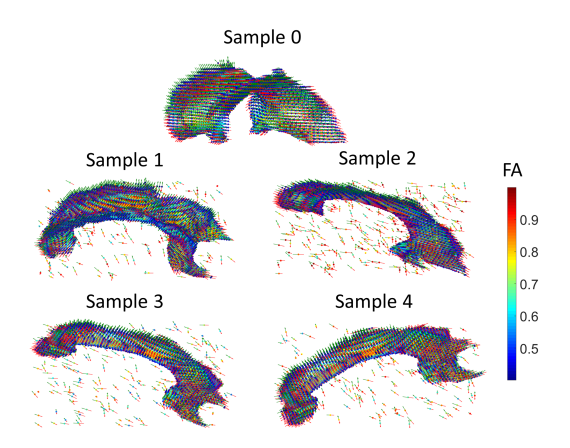


Figure 3: Synthetic corpus callosum data set comprising: Sample 0, the ground truth hybrid point set; and Samples 1-4, which are rotated and modified versions of Sample 0.

The rigidly transformed point sets were also modified by the addition of 545 varying proportions of random outliers (comprising positions, orientations 546 and FA values). Fibre orientations associated with the outliers were gen-547 erated from normalized 3D points. While their FA values were uniformly 548 sampled within the range [0.2, 0.8]. The FA values associated with the voxels 549 of each modified hybrid point set were also varied by ± 0.1 , relative to the 550 ground truth point set. This was necessary in order to emulate real data 551 as FA values typically vary at corresponding anatomical locations, between 552 subjects. This process was repeated 10 times, to generate 10 unique syn-553 thetic data sets (each comprising one ground truth and 4 modified, unique 554 samples), which were subsequently rigidly aligned using the proposed Wat-555 son distribution-based HdMM algorithm (i.e. 10 distinct registration exper-556

iments). Random rotations and proportions of outliers were generated for 557 each experiment, within the range of $[-30^\circ, 30^\circ]$ and [2%, 5%], respectively 558 (as illustrated in Fig. 3). Table 1 summarises the mean ground truth eu-559 clidean distances between Samples 1-4 and Sample 0 across all 10 experiments 560 (prior to registration), and the axes about which rotations were applied to 561 generate each sample in each experiment. The average rigid registration er-562 rors following alignment of the synthetic data sets (with M = 2000 mixture 563 components) using the proposed framework are also reported in Table 1. 564

Rigid registration accuracy was evaluated by: (a) computing the in-565 trinsic distance between the estimated and ground truth rotations (Huynh, 566 2009), for easy interpretation of the rotation errors (θ_{err}), in degrees (refer to 567 equation 7); and (b) computing the mean Euclidean distance (ED) between 568 (transformed) Samples 1-4 and Sample 0 (averaged across all points). Ta-569 ble 1 summarises average rotation and Euclidean distance errors (computed 570 across all 10 experiments). Point-wise Euclidean distances are first evaluated 571 between each modified sample (Samples 1-4) and Sample 0, following rigid 572 registration, and subsequently averaged across all points. The resulting mean 573 Euclidean distance is then averaged once again across all 10 experiments and 574 is reported in Table 1. 575

$$\theta_{err} = \arccos\left[\frac{\operatorname{tr}((\mathbf{R}_{k}^{g}(\mathbf{R}_{k}\mathbf{R}_{1}^{T})^{T}) - 1}{2}\right]$$
(7)

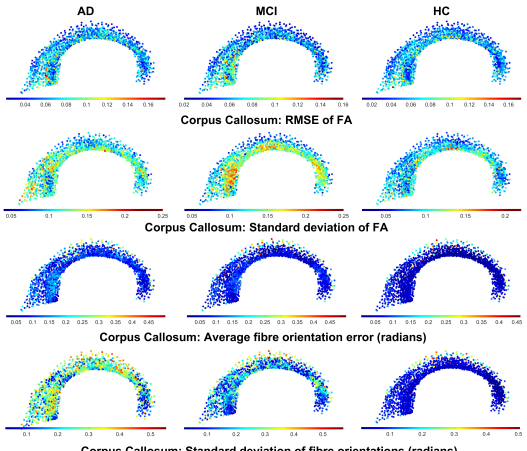
Sample #	Rotated Around	Ground Truth Euc. Dist. (mm.)	Rot. Err. (degrees)	Euc. Dist. (mm.)
1	x,y	43.57 ± 19.85	0.06 ± 0.03	0.34 ± 0.15
2	y,z	42.85 ± 13.12	0.05 ± 0.03	0.30 ± 0.16
3	z,x	42.77 ± 8.74	0.04 ± 0.03	0.23 ± 0.13
4	x,y,z	35.52 ± 17.19	0.04 ± 0.03	0.25 ± 0.17

 Table 1: Summary of rigid registration errors across 10 experiments using synthetic corpus

 callosum data sets.

The average Euclidean distance errors reported in Table 1 indicate that 576 the proposed Watson-based HdMM framework achieved very low errors (de-577 spite the presence of random outliers) as all values are substantially lower 578 than the voxel size of the original eigenvector and FA image (refer to section 579 2.1), from which the ground truth corpus callosum hybrid point set (sample 580 0) was generated. Robustness to outliers may be attributed to the con-581 stituent t-distributions in the HdMM, modelling spatial positions. Similarly 582 the proposed approach was also able to accurately recover the applied ground 583 truth rotations, resulting in very low rotation errors for all samples (as shown 584 in Table 1), relative to the magnitude of the rotations applied to generate 585 the synthetic data set. The proposed approach therefore, is considered to 586 successfully approximate the joint density of position, fibre orientation and 587 FA, for the synthetic corpus callosum data set, and accurately recover the 588 applied rigid transformations. 580

590 3.2. Model Quality



Corpus Callosum: Standard deviation of fibre orientations (radians)

Figure 4: Model quality evaluated for the corpus callosum, independently for AD, MCI and HC groups, using M = 2000 mixture components. Rows one and two: RMSE of FA and standard deviations of the same computed across subjects; Rows three and four: Angular errors for fibre orientations (in radians) and standard deviations of the same computed across subjects.

The ability of the HdMM to model DTI-derived quantities was assessed using clinical data, acquired from the VPH-DARE@IT prospective cohort, described in section 2.1. Specifically, model quality was quantified by evaluating the similarity between the estimated correspondences (resulting from non-rigidly registering the the unbiased study-specific mean template to each

sample from all patient groups) and the nearest neighbour voxels in the corre-596 sponding subject's original FA and eigenvector images. FA accuracy is quan-597 tified as the root-mean-squared error (RMSE), evaluated between the model-598 predicted and original voxel-wise FA values, across all correspondences, for 599 each subject. The group-wise average error (for each subject group) of FA 600 was subsequently computed. The minimum arc length (measured in radians) 601 between two unit vectors is used to measure the accuracy of local fibre orien-602 tation in a similar manner. As discussed in section 2.4, the proposed frame-603 work models axial data rather than directional data. When computing fibre 604 orientation errors, corresponding unit vectors between the model-predicted 605 and original voxel-wise eigenvectors are first identified. This is achieved by 606 evaluating their scalar product and ensuring it is positive — i.e. if the dot 607 product is negative, the antipodal counterpart of the model-predicted vector 608 is used instead. The resulting measure thus quantifies the angular error in 600 fibre orientation between the model-predicted and original voxel-wise data 610 (in the eigenvector image), for each subject. These measures represent reg-611 istration residuals which describe the quality of correspondences established 612 by the proposed HdMM (i.e. how well the HdMM can model the observed 613 DTI-derived data), and only indirectly reflect registration 'accuracy'. To pro-614 vide a more general view of registration accuracy, the mean-squared distance 615 (MSD, formulated as shown in the Appendix), quantifying spatial position 616 errors was also evaluated between the registered study-specific mean template 617 and the original hybrid point sets from all patient groups (Note: MSD values 618 were evaluated between dense volumetric point sets). It is important to note 619 that the model-predicted values for FA and fibre orientation assigned to the 620 spatial correspondences established using the proposed approach, are proba-621 bilistic in nature (as discussed in section 2.7). Consequently, they reflect the 622 DTI-derived quantities of voxels located in the local spatial neighbourhood 623 of the correspondences. 624

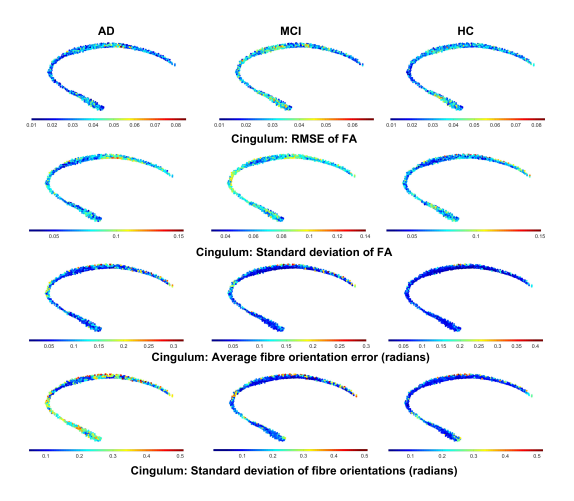


Figure 5: Model quality evaluated for the cingulum, independently for AD, MCI and HC groups, using M = 1500 mixture components. Rows one and two: RMSE of FA and standard deviations of the same computed across subjects; Rows three and four: Angular errors for fibre orientations (in radians) and standard deviations of the same computed across subjects.

Results summarizing the ability of the proposed framework to model DTIderived quantities across all 60 subjects are presented in Fig. 4 - Fig. 7 and Tables 2 - 7. Fig. 4 and Fig. 5 help visualise the spatial distribution of mean registration errors and the standard deviations of FA values and fibre orientations observed across subjects within each patient group, for the corpus callosum and cingulum, respectively. We would like to highlight that while samples from all patient groups were registered simultaneously, the registra-

tion errors presented in Fig. 4 - Fig. 7 and Tables 2 - 7 alone were evaluated 632 for each patient group separately. This was done in order to identify any 633 group-specific trends that exist in the registration accuracy afforded by the 634 proposed approach. In Fig. 4 and Fig. 5 the RMSE values of FA were 635 computed by averaging across subjects in each group, at each corresponding 636 position. Similarly, the standard deviations were also evaluated point-wise 637 across subjects for each group. The depicted mean angular errors were av-638 eraged across subjects, quantifying the fibre orientation accuracy at each 639 corresponding position, and point-wise estimates for the standard deviations 640 in fibre orientation were also evaluated. The presented standard deviations 641 in Fig. 4 and Fig. 5 aid in interpretation of the error measures evaluated, 642 and provide a frame of reference, for both WM regions. The spatial dis-643 tribution of the variation in FA across subjects within each patient group, 644 was evaluated as follows: (a) the nearest neighbour voxel in the original hy-645 brid point sets were first identified based on the spatial positions estimated 646 by non-rigid registration of the study-specific mean template, to each cor-647 responding sample ; (b) the FA values associated with the voxels identified 648 for each subject were in turn used to compute the standard deviation across 649 subjects, within each patient group; and (c) these values were subsequently 650 mapped on to the study-specific mean template estimated for the corpus cal-651 losum and cingulum, for easy comparison with the registration errors plotted 652 in a similar manner, as shown in Fig. 4. Similarly, the standard deviations 653 in fibre orientations about the mean, were also evaluated across subjects. 654 within each patient group, for both WM regions. Here, the difference be-655 tween the mean fibre orientation estimated at each correspondence point in 656 the study-specific mean template, and the nearest neighbour voxels identified 657 (refer to (a) above) in the original hybrid point sets, was evaluated as the 658 minimum arc length (in radians) between each other. This in turn was em-659 ployed to compute the standard deviation in fibre orientations and visualize 660 their spatial distribution across both WM regions. 661

Based on these results, the proposed HdMM is considered to establish 662 valid correspondences across patients, as the estimated fibre orientation and 663 FA errors are low across the majority of correspondences. Fibre orientation 664 errors were consistently < 0.2 radians across most correspondences for both 665 WM ROIs (refer to first and third row in Fig. 4). FA errors meanwhile, were 666 < 0.1 for the corpus callosum and cingulum (refer to second and fourth row in 667 Fig. 4), across all patient groups. For the former WM region, FA errors below 668 0.1 were produced for > 92% of all established correspondences. While for 669

the latter, all correspondences, had errors below 0.1. Fibre orientation errors 670 were < 0.2 across > 94% of correspondences estimated for both WM ROIs, 671 in all patient groups. Errors of this magnitude are considered reasonable as 672 the model-predicted FA values and fibre orientations evaluated at correspon-673 dences are based on the soft-assignment approach (refer to section 2.7), using 674 the estimated posterior probabilities. Consequently, they reflect weighted av-675 erages of FA and fibre orientations of neighbouring voxels. FA variations of 676 ≈ 0.1 may occur due to partial volume effects at WM-GM and WM-CSF 677 interfaces (Smith et al., 2006), particularly when WM tracts/ROIs are very 678 thin compared to the voxel size (often the case following dementia-related 679 atrophy of brain tissue), potentially further contributing to the observed er-680 rors. Additionally, significant variations in DTI-data in a select few cases 681 within individual patient groups may be another source of the high average 682 errors evaluated, in a small proportion of correspondences. These results are 683 further supported by the standard deviations of FA and fibre orientations 684 depicted in Fig. Fig. 4 and Fig. 5, which highlight the high degree of vari-685 ation in FA and fibre orientations (across subjects), respectively, relative to 686 the corresponding errors evaluated across both WM regions. 687

These results are further verified by the histograms of errors in fibre 688 orientation and FA presented in Fig. 6 and 7, respectively, summarising the 689 correspondence-wise errors evaluated for each subject in the population. In 690 this case, fibre orientation errors were computed as in preceding experiments, 691 while FA errors were evaluated as the root-squared-error (RSE) between 692 the model-predicted values and the closest voxels in the corresponding FA 693 images. In general, high errors occur at only a few correspondences, across 694 both the cingulum and corpus callosum. Registration errors for the AD and 695 MCI groups were higher than for the HC group for both ROIs. This is 696 attributed to the presence of varying degrees of pathology-induced changes 697 in a few subjects in these groups, verified by Figs. 6 and 7, and by computing 698 region-wise mean and standard deviations of FA and fibre orientation errors, 699 presented in Tables 3 - 7. 700

Tables 2 - 7 report the average spatial position, fibre orientation and FA errors evaluated across correspondences and subjects. Statistically significant reduction in mean spatial position errors across experiments conducted using differing model complexities (i.e. different number of mixture components) are highlighted in bold in Tables 2 and 5, considering a significance level of 5%. In Tables 4 and 7 the reported mean FA errors were estimated by first computing the RMSE, this time averaging across correspondences,

and subsequently computing the mean RMSE across subjects. Tables 3 and 708 6 summarise the mean angular error values, first averaged across correspon-709 dences and subsequently across subjects. These alternate error measures are 710 presented to assess model quality of the HdMM across regions, and comple-711 ment the correspondence-wise errors presented in Fig. 4 - 5. From Tables 2 712 - 7, the number of mixture components required to adequately characterise 713 the entire population was identified as M = 1500 and M = 2000 for the cin-714 gulum and corpus callosum, respectively. The fibre orientation and FA errors 715 depicted in Fig. 4 - 5 were evaluated using these values. All subsequent inter-716 group statistical analyses conducted employed these model complexities for 717 the respective WM regions. 718

Table 2: Model quality of HdMM for the cingulum, assessed in terms of the mean spatial position error evaluated across correspondences and subjects, using the MSD metric, for each patient group, and for varying model complexities. Bold values indicate statistically significant reduction in errors.

# Mixture Components	Spatial Position Error: MSD (mm.)			
# Mixture Components	AD	MCI	HC	
500	0.86 ± 0.11	0.84 ± 0.09	0.82 ± 0.09	
1000	0.73 ± 0.10	0.72 ± 0.08	0.71 ± 0.08	
1500	0.67 ± 0.09	0.66 ± 0.07	$\boldsymbol{0.64\pm0.07}$	
2000	0.65 ± 0.09	0.63 ± 0.07	$\boldsymbol{0.62}\pm\boldsymbol{0.07}$	

Table 3: Model quality of HdMM for the cingulum, assessed as the mean fibre orientation error evaluated across correspondences and subjects, for each patient group, and for varying model complexities.

#	Mean Fibre Orientation Error (radians)				Mean Fibre Orientation Error (radians		
Mixture Components	AD	MCI	HC				
300	0.11 ± 0.10	0.08 ± 0.02	0.07 ± 0.02				
600	0.09 ± 0.08	0.07 ± 0.02	0.06 ± 0.01				
1200	0.09 ± 0.08	0.06 ± 0.01	0.06 ± 0.01				
1500	0.08 ± 0.08	0.06 ± 0.01	0.05 ± 0.01				

Table 4: Model quality of HdMM for the cingulum, assessed as the average RMSE of FA evaluated over correspondences and averaged across subjects, for each patient group, and for varying model complexities.

#	Mean RMSE of FA			
Mixture Components	AD	MCI	HC	
300	0.06 ± 0.01	0.06 ± 0.01	0.06 ± 0.01	
600	0.06 ± 0.01	0.06 ± 0.01	0.06 ± 0.01	
1200	0.05 ± 0.01	0.05 ± 0.01	0.05 ± 0.01	
1500	0.05 ± 0.01	0.05 ± 0.01	0.05 ± 0.01	

Table 5: Model quality of HdMM for the corpus callosum, assessed in terms of the mean spatial position error evaluated across correspondences and subjects, using the MSD metric, for each patient group, and for varying model complexities. Bold values indicate statistically significant reduction in errors.

# Mixture Components	Spatial Position Error: MSD (1		ASD (mm.)
# Mixture Components	AD	MCI	HC
500	1.15 ± 0.17	1.14 ± 0.10	1.09 ± 0.12
1000	0.99 ± 0.15	0.98 ± 0.09	0.94 ± 0.10
1500	0.91 ± 0.13	0.90 ± 0.08	0.85 ± 0.09
2000	0.86 ± 0.12	0.85 ± 0.07	$\boldsymbol{0.81} \pm \boldsymbol{0.08}$

Table 6: Model quality of HdMM for the corpus callosum, assessed as the mean fibre orientation error evaluated across correspondences and subjects, for each patient group, and for varying model complexities.

#	Mean Fibre Orientation Error (radians)			
Mixture Components	AD	MCI	HC	
500	0.13 ± 0.19	0.10 ± 0.14	0.06 ± 0.01	
1000	0.13 ± 0.19	0.13 ± 0.16	0.05 ± 0.01	
1500	0.12 ± 0.19	0.09 ± 0.13	0.05 ± 0.01	
2000	0.12 ± 0.18	0.09 ± 0.13	0.05 ± 0.01	

Table 7: Model quality of HdMM for the corpus callosum, assessed as the average RMSE of FA evaluated over correspondences and averaged across subjects, for each patient group, and for varying model complexities.

#	Me	Mean RMSE of FA			
Mixture Componen	ts AD	MCI	HC		
500	0.11 ± 0.03	0.11 ± 0.02	0.10 ± 0.01		
1000	0.10 ± 0.03	0.10 ± 0.02	0.09 ± 0.01		
1500	0.09 ± 0.03	0.09 ± 0.03	0.08 ± 0.004		
2000	0.09 ± 0.03	0.08 ± 0.03	0.07 ± 0.01		

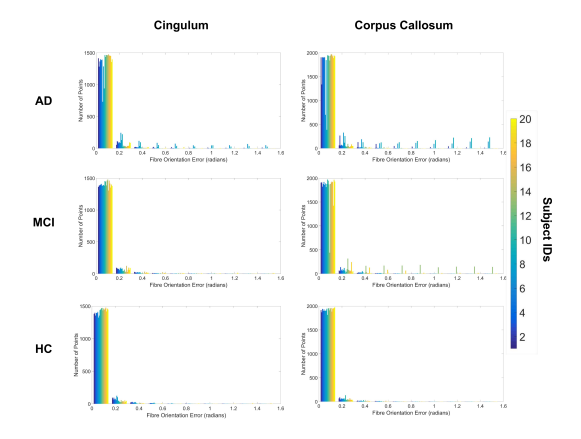


Figure 6: Histograms of fibre orientation errors for each subject in AD, MCI and HC groups, evaluated between established correspondences and ground truth voxels.

Results in Fig. 6 and 7 indicate that the proposed framework achieves low fibre orientation and FA errors at each estimated correspondence, for all

subjects in the HC group (for both WM ROIs). The estimated correspon-721 dences were less accurate for two cases in the AD group (for both cingulum 722 and corpus callosum) and for one case in the MCI group (only corpus cal-723 losum), which is attributed to significant variation in fibre orientations and 724 FA values in these cases and ROIs, relative to the remaining samples in their 725 corresponding patient groups. As discussed previously, this may be a re-726 sult of varying degrees of pathology-induced changes in these cases relative 727 to the rest of their group. Consequently, the accuracy of the HdMM when 728 fitting to these few cases, is reduced. The proposed framework, however, 720 established accurate correspondences for the remaining samples in the AD 730 and MCI groups across both WM ROIs. The high deviations from the mean 731 fibre orientation errors in the corpus callosum for these groups (Table 6) 732 are thus attributed to the outlier subjects identified from the corresponding 733 histograms (Fig. 6). Similarly, for the cingulum, the high standard devia-734 tions observed for the AD group are attributed to the two subjects mentioned 735 above. However, no apparent outliers were identified in the MCI group based 736 on the registration errors and, by extension, the mean FA and fibre orien-737 tation errors reported in Tables 4 and 3, are low and consistent with their 738 corresponding histogram plots (Fig. 7 and Fig. 6). 739

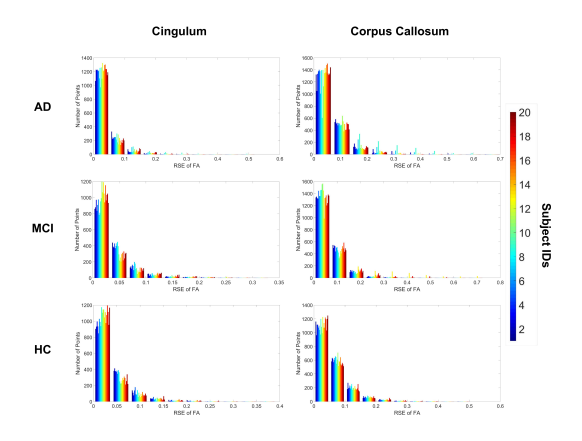


Figure 7: Histograms of root-squared-error (RSE) of FA for each subject in AD, MCI and HC groups, evaluated between established correspondences and ground truth voxels.

The foregoing results suggest the proposed framework established valid correspondences for both WM ROIs across all subjects in the HC group and for the majority of cases in the AD and MCI groups. This is indicative of the ability of the proposed HdMM to approximate the joint *PDF* of positions, fibre orientations and FA values across multiple subjects.

745 3.3. Group Comparisons

The ability of the proposed framework to identify significant differences between patient groups was assessed by comparing each pair of patient groups in terms of the variation in FA. These results were compared with those obtained from the widely used TBSS approach. Un-paired two-sample t-tests, assuming equal variances, were performed to compare FA values at corresponding spatial positions between patient groups. The procedure proposed in (Benjamini and Yekutieli, 2001) was used to correct for multiple comparisons by controlling the false discovery rate (FDR) for the set of hypothesis
tests. The desired FDR was fixed at 1% for all experiments. However, no
statistically significant reduction in FA was identified between any of the
groups, using the proposed approach, TBSS and VBM.

Interquartile ranges (IQRs) for the mean FA values estimated using each 757 approach were also evaluated to provide a quantitative means of comparing 758 the range of estimated FA values for both WM ROIs. This measure is adopted 759 as it provides a robust means of assessing dispersion in data. IQRs are 760 summarised in Table 8 for both WM ROIs, from which we infer that all 761 three methods do indeed show similarities in the range of estimated mean 762 FA values, for the corpus callosum. Conversely, for the cingulum, while 763 VBM and the proposed approach show similar IQRs, the ranges estimated 764 for TBSS are lower. This is because TBSS models the central skeleton of 765 the ROI, and there is substantial variation in FA between the center and 766 peripheral regions of cingulum region. Consequently, the variation in mean 767 FA values in the skeleton voxels is lower in comparison to the entire ROI (as 768 modelled by VBM and HdMM). 769

Corpus Calle			losum:	Cingulum:			
Method	IQR of mean FA			IQR of mean FA			
	AD	MCI	HC	AD	MCI	HC	
HdMM	0.24	0.24	0.24	0.17	0.16	0.16	
TBSS	0.20	0.21	0.21	0.08	0.08	0.09	
VBM	0.21	0.21	0.21	0.14	0.13	0.14	
	HdMM	Method IQR AD HdMM 0.24 TBSS 0.20	Method IQR of me AD MCI HdMM 0.24 0.24 TBSS 0.20 0.21	AD MCI HC HdMM 0.24 0.24 0.24 TBSS 0.20 0.21 0.21	Method IQR IQR AD MCI HC AD HdMM 0.24 0.24 0.24 0.17 TBSS 0.20 0.21 0.21 0.08	Method IQR of means IQR of means AD MCI HC AD MCI HdMM 0.24 0.24 0.24 0.17 0.16 TBSS 0.20 0.21 0.21 0.08 0.08	

Table 8: Interquartile ranges for mean FA values estimated using each approach for both WM ROIs.

As discussed previously, the primary advantage of the proposed HdMM 770 framework is its ability to model fibre orientations and facilitate their compar-771 ison across multiple subjects, which is not offered by conventional approaches 772 such as TBSS and VBM. Furthermore, the proposed method does not require 773 extraction of fibre trajectories using tractography in order to model fibre ori-774 entations as it operates directly on the raw DTI-derived eigenvectors, unlike 775 state-of-the-art approaches such as those proposed in (Garyfallidis et al., 776 2015) and (ODonnell et al., 2017). Inter-group statistical comparisons of 777 the angular deviation in fibre orientations, relative to study-specific mean 778

template, were also conducted. Here, the angular deviation of the modelpredicted fibre orientations at each spatial correspondence was first evaluated relative to the corresponding mean fibre orientation (for patients from all groups), as the minimum arc length between unit vectors. Subsequently, these deviations were compared between each pair of patient groups, while correcting for multiple comparisons using FDR. However, as with the FA analyses, no statistically significant differences were identified.

The proposed HdMM for the joint registration and clustering of data com-786 prising positions, orientations and scalar-valued features (such as FA) shows 787 promise for statistical analysis of diffusion derived measures across multiple 788 subjects and patient populations. Although the inter-group statistical com-789 parisons conducted to analyse the variation in FA and fibre orientations re-790 vealed no significant differences between patient groups, our results matched 791 those obtained using TBSS and VBM, in the case of the former. This may 792 be due to the underlying nature of the data as the samples used throughout 793 this study were part of the prospective cohort of the VPH-DARE@IT project. 794 Consequently, it is possible that no significant differences in FA and fibre ori-795 entation exist in the WM ROIs considered, between the subjects assigned to 796 the AD, MCI and HC groups. However, we believe the proposed approach 797 still holds merit due to the flexibility it affords, as: (a) it enables analysis of 798 various scalar-valued diffusion measures (although just FA was considered in 799 this study), similar to existing approaches such as TBSS and VBM; and (b) 800 also permits analysis of local fibre orientation, defined by primary diffusion 801 axes, a capability not afforded by existing techniques. Although approaches 802 based on clustering of fibre trajectories enable such analyses, they require 803 diffusion-tractography derived fibres to do so. The present work ameliorates 804 this need and acts directly on the raw eigenvector images. Additionally, our 805 approach is not restricted to a specific anatomical region or analysing voxel-806 wise (or structured grid) data and may be employed to jointly register and 807 cluster unstructured data as well. 808

A current limitation of the proposed approach is it only enables anal-809 ysis of DTI data generated using a single tensor model. However, the pro-810 posed HdMM framework could be imbued with greater flexibility by replacing 811 the Watson distributions with the Kent or the general 8-parameter Fisher-812 Bingham distribution, to model multi-fibre (or crossing fibre) regions by fit-813 ting to orientation distribution functions obtained from high angular diffusion 814 images. Extensions to the Von-Mises-Fisher mixture model for example, have 815 been proposed previously to accommodate antipodal symmetry and model 816

⁸¹⁷ diffusion ODFs (McGraw et al., 2006).

The sensitivity and discriminative capacity of the proposed framework 818 in comparison to existing approaches requires further investigation and val-819 idation, which will be the subject of future work. Natural extensions to the 820 proposed framework include whole WM volume analysis across multiple sub-821 jects, WM parcellation, and automatic region-of-interest analysis, to name 822 a few. As discussed previously, the proposed approach can be employed to 823 analyse the entire WM volume across subjects, i.e. *a priori* definition of ROIs 824 is not required, though the computational burden at present is substantial. 825 Such an approach naturally leads to the unsupervised parcellation of WM 826 into distinct clusters defined by the centroids of the HdMM, across multiple 827 subjects. This in turn provides a mechanism for automatic ROI-type analy-828 ses, as the generated clusters for each subject will correspond to similar WM 829 regions in terms of spatial position, fibre orientation and FA (or some other 830 scalar measure of interest). Furthermore, by employing a suitable prior/atlas 831 containing pre-defined labels for WM tracts of interest, the presented frame-832 work could be employed for automatic tractography segmentation (similar 833 to (O'Donnell and Westin, 2007)). The proposed approach can also be em-834 ployed to track and identify localised changes in WM over time for a single 835 subject, resulting from the progression of neuro-degenerative disorders such 836 as dementia, for example. Although WM changes in the brain were consid-837 ered in this study, the generic nature of the proposed framework permits its 838 application to other organs exhibiting tissue anisotropy, such as cardiac dif-839 fusion data, and modelling bone micro-architecture. Additionally, it can be 840 employed for a variety of other applications, such as vessel centerlines-based 841 image registration, as demonstrated by our recent study (Bayer et al., 2018). 842

4. Conclusions

In this study, a Watson-distribution based hybrid mixture model was pre-844 sented for jointly registering and clustering DTI-derived data from multiple 845 subjects and patient populations. This approach was shown to model the 846 observed fibre orientations and FA values accurately for all subjects within 847 the HC group, for both of the studied WM ROIs, namely, the cingulum and 848 corpus callosum. Registration to subjects in AD and MCI groups was suc-849 cessful for the majority of cases, with two in the former and one in the latter 850 resulting in high registration errors, due to significant pathology induced 851 changes in these cases. Group comparisons of FA values in the WM ROIs 852

using the proposed approach showed no statistically significant reductions in 853 FA between the AD, MCI and HC groups, as with TBSS and VBM. Similarly, 854 no significant variations in fibre orientation were identified between patient 855 groups. However, the proposed method has potential for use in a variety of 856 applications involving statistical analysis of diffusion data. Its generic and 857 flexible nature make it well suited to a variety of other computer vision and 858 medical image analysis tasks, such as: point set registration with the inte-859 gration of surface normals, vessel-based image registration, joint registration 860 and clustering of geometries with associated velocity fields (estimated from 861 computational fluid dynamic simulations for example) and texture mapping, 862 to name a few. The fidelity and extensibility of the proposed framework is 863 thus compelling as a general tool for multi-dimensional medical image anal-864 ysis. 865

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874 Appendix

⁸⁷⁵ M-step update equations for the Student's t-distribution parameters in ⁸⁷⁶ the HdMM and rigid registration parameters at the $(t + 1)^{\text{th}}$ EM-iteration, ⁸⁷⁷ discussed in section 2.6, are derived by maximizing the complete data log-⁸⁷⁸ likelihood $Q(\Theta_p^{t+1}, \mathbb{T}^{t+1} | \Theta_p^t, \mathbb{T}^t)$ with respect to each parameter as follows:

• Estimation of TMM centroids $\boldsymbol{\mu}_j$ at the $(t+1)^{\text{th}}$ EM-iteration:

$$Q(\Theta_p^{t+1}, \mathbb{T}^{t+1} | \Theta_p^t, \mathbb{T}^t) = -\frac{1}{2} \sum_{k,i,j} P_{kij}^{\star t} \Delta_{kij} + O.T.$$
(8a)

$$\Delta_{kij} = \frac{(\mathbf{x}_{ki} - s_k \mathbf{R}_k \boldsymbol{\mu}_j - \mathbf{t}_k)^T (\mathbf{x}_{ki} - s_k \mathbf{R}_k \boldsymbol{\mu}_j - \mathbf{t}_k)}{\sigma^2}$$
(8b)

O.T. summarizes terms in Q independent of $\pmb{\mu}_j.$

$$<\partial Q, \partial \boldsymbol{\mu}_{j}>=\left[-\frac{1}{2}\sum_{k,i}P_{kij}^{\star}\Delta_{kij}^{\boldsymbol{\mu}_{j}+\partial\boldsymbol{\mu}_{j}}\right]-\left[-\frac{1}{2}\sum_{k,i}P_{kij}^{\star}\Delta_{kij}^{\boldsymbol{\mu}_{j}}\right]$$
(9a)

$$\langle \partial Q, \partial \boldsymbol{\mu}_j \rangle = \sum_{k,i} P_{kij}^{\star} [(\mathbf{x}_{ki} - s_k \mathbf{R}_k \boldsymbol{\mu}_j - \mathbf{t}_k)^T s_k \mathbf{R}_k] \partial \boldsymbol{\mu}_j$$
(9b)

$$\langle \partial Q, \partial \boldsymbol{\mu}_j \rangle = 0 \implies \sum_{k,i} P_{kij}^{\star} [(\mathbf{x}_{ki} - s_k \mathbf{R}_k \boldsymbol{\mu}_j - \mathbf{t}_k)^T s_k \mathbf{R}_k] = 0$$
 (9c)

$$\sum_{k,i} P_{kij}^{\star} s_k \mathbf{R}_k^T (\mathbf{x}_{ki} - \mathbf{t}_k) = \sum_{k,i} P_{kij}^{\star} s_k \mathbf{R}_k^T \mathbf{R}_k s_k \boldsymbol{\mu}_j$$
(9d)

$$\boldsymbol{\mu}_{j} = \frac{\sum\limits_{k,i} P_{kij}^{\star} s_{k}^{-1} \mathbf{R}^{T} (\mathbf{x}_{ki} - \mathbf{t}_{k})}{\sum\limits_{k,i} P_{kij}^{\star}}$$
(9e)

• Estimation of model variance σ^2 :

$$\frac{\partial Q}{\partial \sigma^2} = \frac{\partial \sum_{k,i,j} \left[-\frac{P_{kij}}{2} \left[\log(\sigma^6)\right] - \frac{P_{kij}^*}{2} \left[\Delta_{kij}\right]\right]}{\partial \sigma^2} = 0$$
(10a)

$$\implies \sum_{k,i,j} -P_{kij}\frac{3}{\sigma} + P_{kij}^{\star}\frac{(\mathbf{x}_{ki} - s_k\mathbf{R}_k\boldsymbol{\mu}_j - \mathbf{t}_k)^T(\mathbf{x}_{ki} - s_k\mathbf{R}_k\boldsymbol{\mu}_j - \mathbf{t}_k)}{\sigma^3} = 0$$
(10b)

$$P^{2} = \frac{\sum_{k,i,j} P^{\star}_{kij} (\mathbf{x}_{ki} - s_{k} \mathbf{R}_{k} \boldsymbol{\mu}_{j} - \mathbf{t}_{k})^{T} (\mathbf{x}_{ki} - s_{k} \mathbf{R}_{k} \boldsymbol{\mu}_{j} - \mathbf{t}_{k})}{(100)}$$
(100)

$$\sigma^2 = \frac{n_{it,j}}{3\sum_{kij} P_{kij}}$$
(10c)

• Estimation of translation \mathbf{t}_k :

$$\langle \partial Q, \partial \mathbf{t}_k \rangle = \left[-\frac{1}{2} \sum_{i,j} P_{kij}^{\star} \Delta_{kij}^{\mathbf{t}_k + \partial \mathbf{t}_k} \right] - \left[-\frac{1}{2} \sum_{i,j} P_{kij}^{\star} \Delta_{kij}^{\mathbf{t}_k} \right]$$
(11a)

$$\langle \partial Q, \partial \mathbf{t}_k \rangle = \sum_{i,j} P_{kij}^{\star} [(\mathbf{x}_{ki} - s_k \mathbf{R}_k \boldsymbol{\mu}_j - \mathbf{t}_k)^T] \partial \mathbf{t}_k$$
(11b)

$$\langle \partial Q, \partial \mathbf{t}_k \rangle = 0 \implies \sum_{i,j} P_{kij}^{\star} (\mathbf{x}_{ki} - s_k \mathbf{R}_k \boldsymbol{\mu}_j)^T = \sum_{i,j} P_{kij}^{\star} \mathbf{t}_k^T$$
 (11c)

$$\mathbf{t}_{k} = \frac{\sum_{i,j} P_{kij}^{\star} \mathbf{x}_{ki}}{\sum_{i,j} P_{kij}^{\star}} - s_{k} \mathbf{R}_{k} \frac{\sum_{i,j} P_{kij}^{\star} \boldsymbol{\mu}_{j}}{\sum_{i,j} P_{kij}^{\star}}$$
(11d)

Setting the first term as \mathbf{d}_k and the second term as \mathbf{m}_k we get:

$$\mathbf{t}_k = \mathbf{d}_k - s_k \mathbf{R}_k \mathbf{m}_k \tag{11e}$$

• Estimation of strictly orthogonal rotation \mathbf{R}_k : Using the lemma outlined in (Myronenko and Song, 2010), the optimal rotation matrix maximises $\operatorname{tr}(\mathbf{C}_k^T \mathbf{R}_k)$ where \mathbf{C}_k represents a real covariance matrix (refer to equation 12d).

$$\tilde{\mathbf{x}}_{ki} = \mathbf{x}_{ki} - \mathbf{d}_k, \tilde{\mathbf{m}}_{kj} = \boldsymbol{\mu}_j - \mathbf{m}_k$$
(12a)

Using equations (11e) and (12a) we get:

$$Q(\Theta_p^{t+1}, \mathbb{T}^{t+1} | \Theta_p^t, \mathbb{T}^t) \propto \sum_{i,j} P_{kij}^{\star t}(\tilde{\mathbf{x}}_{ki}^T \mathbf{R}_k \tilde{\mathbf{m}}_{kj})$$
(12b)

$$Q(\Theta_p^{t+1}, \mathbb{T}^{t+1} | \Theta_p^t, \mathbb{T}^t) \propto \sum_{i,j} P_{kij}^{\star t} \operatorname{tr}[\tilde{\mathbf{m}}_{kj} \tilde{\mathbf{x}}_{ki}^T \mathbf{R}_k]$$
(12c)

As equation (12c) must be maximised with respect to \mathbf{R}_k ,

$$\mathbf{C}_{k} = \sum_{i,j} P_{kij}^{\star} \tilde{\mathbf{x}}_{ki} \tilde{\mathbf{m}}_{kj}^{T}$$
(12d)

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- $\mathbf{R}_k = \mathbf{U}\mathbf{S}\mathbf{V}^T$, where \mathbf{U}, \mathbf{V} are unitary matrices computed by singular value decomposition of \mathbf{C}_k and $\mathbf{S} = diag(1, 1, det(\mathbf{U}\mathbf{V}^T))$ is a diagonal matrix that prevents reflections.
- Estimation of scaling s_k :

$$\frac{\partial Q}{\partial s_k} = -\frac{1}{2} \frac{\partial \sum_{i,j} P_{kij}^{\star} \Delta_{kij}}{\partial s_k} = 0$$
(13a)

$$\sum_{i,j} P_{kij}^{\star} \frac{(\tilde{\mathbf{x}}_{ki} - s_k \mathbf{R}_k \tilde{\mathbf{m}}_{kj})^T (\mathbf{R}_k \tilde{\mathbf{m}}_{kj})}{\sigma^2} = 0$$
(13b)

$$\sum_{i,j} P_{kij}^{\star}[(\mathbf{\tilde{x}}_{ki})^T (\mathbf{R}_k \mathbf{\tilde{m}}_{kj})] = s_k \sum_{i,j} P_{kij}^{\star}[\mathbf{\tilde{m}}_{kj}^T \mathbf{R}_k^T \mathbf{R}_k \mathbf{\tilde{m}}_{kj}]$$
(13c)

$$s_k = \frac{\operatorname{tr}[\tilde{\mathbf{m}}_{kj}\tilde{\mathbf{x}}_{ki}^T]\mathbf{R}_k}{\operatorname{tr}[\tilde{\mathbf{m}}_{kj}\tilde{\mathbf{m}}_{kj}^T]} = \frac{\operatorname{tr}[\mathbf{C}_k^T\mathbf{R}_k]}{\operatorname{tr}[\tilde{\mathbf{m}}_{kj}\tilde{\mathbf{m}}_{kj}^T]}$$
(13d)

• Estimation of degrees of freedom ν_i :

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$$Q(\Theta_{p}^{t+1}, \mathbb{T}^{t+1} | \Theta_{p}^{t}, \mathbb{T}^{t}) = \sum_{k,i,j} P_{kij}^{t} [-\log \Gamma(\frac{\nu_{j}}{2}) + \frac{1}{2} \nu_{j} \log(\frac{\nu_{j}}{2}) + \frac{\nu_{j}}{2} [\log(U_{kij}^{t}) - U_{kij}^{t} + \Psi(\frac{\nu_{j} + D}{2}) - \log(\frac{\nu_{j}^{t} + D}{2})]] + O.T.$$
(14a)

O.T. summarizes terms in Q independent of ν_j .

$$\frac{\partial Q}{\partial \nu_j} = -\Psi(\frac{\nu_j}{2}) + \log(\frac{\nu_j}{2}) + 1 + \frac{1}{\sum_{k,i} P_{kij}^t} \sum_{k,i} P_{kij}^t (\log(U_{kij}^t) - U_{kij}^t) + \Psi(\frac{\nu_j^t + D}{2}) - \log(\frac{\nu_j^t + D}{2}) = 0$$
(14b)

Equation (14b) is solved using Newton's method to estimate the degrees 882 of freedom ν_j . 883

• Derivations for the M-step updates (refer to equations 3c - 3e) of the mean fibre orientation \mathbf{m}_{j}^{d} and fibre concentration κ_{j} parameters asso-885 ciated with Watson distributions in the HdMM, presented in section 2.4, are derived by maximizing the complete data log-likelihood Q (refer to equation 15a), with respect to each model parameter as follows: (Here $M(\kappa_i)$ denotes the Kummer function). 889

$$Q(\Theta_n^{t+1}|\Theta_n^t) = \sum_{k=1}^K \sum_{i=1}^{N_k} \sum_{j=1}^M P_{kij} \log p(\pm \mathbf{n}_{ki} | \mathbf{m}_j^d, \kappa_j) + \lambda_j (1 - \mathbf{m}_j^{d^T} \mathbf{m}_j^d)$$
(15a)

$$\langle \partial Q, \partial \mathbf{m}_{j}^{d} \rangle = 0 \implies \lambda_{j} \mathbf{m}_{j}^{d} = \kappa_{j} \sum_{k=1}^{K} \sum_{i=1}^{N_{k}} P_{kij}(\mathbf{n}_{ki}^{T} \mathbf{m}_{j}^{d}) \mathbf{n}_{ki}$$
 (15b)

$$\langle \partial Q, \partial \kappa_j \rangle = 0 \implies \frac{M'(\kappa_j)}{M(\kappa_j)} \sum_{k=1}^K \sum_{i=1}^{N_k} P_{kij} = \sum_{k=1}^K \sum_{i=1}^{N_k} P_{kij} (\mathbf{n}_{ki}^T \mathbf{m}_j^d)^2$$
(15c)

$$\mathbf{m}_{j}^{d^{T}}\mathbf{m}_{j}^{d} = 1 \implies \lambda_{j} = \kappa_{j} || \sum_{k=1}^{K} \sum_{i=1}^{N_{k}} P_{kij}(\mathbf{n}_{ki}^{T}\mathbf{m}_{j}^{d})\mathbf{n}_{ki} ||$$
(15d)

Substituting equation (15d) in (15b) results in a non-linear equation (16), which is solved numerically by fixed-point iteration.

$$\mathbf{m}_{j}^{d} = \frac{\sum_{k=1}^{K} \sum_{i=1}^{N_{k}} P_{kij}(\mathbf{n}_{ki}^{T}\mathbf{m}_{j}^{d})\mathbf{n}_{ki}}{\left|\left|\sum_{k=1}^{K} \sum_{i=1}^{N_{k}} P_{kij}(\mathbf{n}_{ki}^{T}\mathbf{m}_{j}^{d})\mathbf{n}_{ki}\right|\right|}$$
(16)

Based on equation (15c), the ratio of the derivative of the Kummer function to the function itself, is expressed as shown in equation (17a). This ratio may be expressed as a continued fraction, as shown in equation (17b). Consequently, using equations (17a) and (17b), the concentration parameters κ_j can be approximated as shown in equation (17d), by solving the linear equation (17c) (similarly to (Bijral et al., 2007)).

$$\frac{M'(\kappa_j)}{M(\kappa_j)} = \frac{\sum_{k=1}^{K} \sum_{i=1}^{N_k} P_{kij} (\mathbf{n}_{ki}^T \mathbf{m}_j^d)^2}{\sum_{k=1}^{K} \sum_{i=1}^{N_k} P_{kij}}$$
(17a)

$$\frac{\kappa_j M'(\kappa_j)}{M(\kappa_j)} = \frac{\kappa_j/2}{(D/2) - \kappa_j + \frac{(3/2)\kappa_j}{(\frac{D}{2} + 1) - \kappa_j + \dots}}$$
(17b)

$$\frac{\kappa_j M'(\kappa_j)}{M(\kappa_j)} \approx \frac{\kappa_j/2}{(D/2) - \kappa_j + \frac{\kappa_j M'(\kappa_j)}{M(\kappa_j)}}$$
(17c)

$$\kappa_j \approx \frac{1}{2} \left[\frac{1 - \frac{M'(\kappa_j)}{M(\kappa_j)} D}{\left(\frac{M'(\kappa_j)}{M(\kappa_j)}\right)^2 - \frac{M'(\kappa_j)}{M(\kappa_j)}} \right]$$
(17d)

• The mean-squared distance (MSD) metric (refer to equation (18)) is used to assess registration errors in terms of spatial position. MSD values were evaluated between the correspondences established following registration of the (study-specific) mean template, and the corresponding original hybrid point sets (i.e. between the estimated correspondences and the voxel centroids defining the WM ROIs). In equation (18) $\mathbf{d}_{\min}(A, B)$ denotes the minimum Euclidean distance between each point in sample A and sample B.

$$MSD = mean(mean(\mathbf{d_{min}}(A, B)), mean(\mathbf{d_{min}}(B, A)))$$
(18)

• The "model-predicted" values for FA (\hat{f}_{kj}) and fibre orientation $(\hat{\mathbf{n}}_{kj})$ estimated at each established spatial correspondence, for each patient, are weighted averages of the neighbouring voxels in their original DTI-derived images (original hybrid point sets), where the weights are defined by the estimated posterior probabilities following non-rigid registration of the study-specific mean template to each sample. These values were estimated for FA and fibre orientation as described by equations 19a and 19b, respectively.

$$\hat{f}_{kj} = \sum_{i=1}^{N_k} \frac{P_{kij} f_{ki}}{\sum_l P_{klj}}$$
 (19a)

$$\hat{\mathbf{n}}_{kj} = \sum_{i=1}^{N_k} \frac{P_{kij} \mathbf{n}_{ki}}{\sum_l P_{klj}}$$
(19b)

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