Effects of Inaccurate Response Function Calibration on Characteristics of the Fiber Orientation Distribution in Diffusion MRI

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Abstract

2 Diffusion MRI of the brain enables to quantify white matter fiber orientations noninvasively. 3 Several approaches have been proposed to estimate such characteristics from diffusion MRI 4 data with spherical deconvolution being one of the most widely used methods. Constrained 5 spherical deconvolution requires to define – or derive from the data – a response function, 6 which is used to compute the fiber orientation distribution (FOD). This definition or derivation 7 is not unequivocal and can thus result in different characteristics of the response function which 8 are expected to affect the FOD computation and the subsequent fiber tracking. In this work, we explored the effects of inaccuracies in the shape and scaling factors of the response 9 10 function on the FOD characteristics. With simulations, we show that underestimation of the 11 shape factor in the response functions has a larger effect on the FOD peaks than overestimation of the shape factor, whereas the latter will cause more spurious peaks. 12 Moreover, crossing fiber populations with a smaller separation angle were more sensitive to 13 the response function inaccuracy than fiber populations with more orthogonal separation 14 15 angles. Furthermore, the FOD characteristics show deviations as a result of modified shape 16 and scaling factors of the response function. Results with the in vivo data demonstrate that the deviations of the FODs and spurious peaks can further deviate the termination of propagation 17 in fiber tracking. This work highlights the importance of proper definition of the response 18 19 function and how specific calibration factors can affect the FOD and fiber tractography results.

- 21 Keywords: Diffusion MRI; constrained spherical deconvolution (CSD); response function; fiber
- 22 orientation distribution (FOD); brain fiber tractography; apparent fiber density (AFD).

24 **1. Introduction**

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Diffusion MRI allows to characterize tissue microstructure in vivo and noninvasively by 26 measuring the anisotropic diffusion of water molecules [1,2]. Diffusion tensor imaging (DTI) [3] 27 28 is the most widely used model in clinical studies to relate the diffusion MRI signals to the diffusion characteristics of the underlying tissue. However, DTI is inadequate to estimate the 29 directional information in voxels containing crossing fibers [4,5]. A commonly used approach 30 31 to resolve more complex fiber configurations in the brain is spherical deconvolution (SD) [6-32 8]. SD also allows for the extraction of fiber population specific microstructural measures derived from the magnitudes of the fiber orientation distribution (FOD) functions, such as 33 apparent fiber density (AFD) [9] and hindrance modulated orientational anisotropy (HMOA) 34 [10]. 35

36 SD requires an appropriate response function as input to estimate the FOD [7]. The 37 response function, representing the diffusion signal for a single fiber population, is ideally 38 calibrated from the acquired diffusion MRI data [11,12]. In brief, for each subject, the voxels 39 containing only single fiber populations are localized, and an average of the diffusivity 40 characteristics within those voxels is used to represent the subject specific response function. 41 An inadequately chosen response function can affect the quantification of FOD characteristics 42 like AFD and HMOA, as well as the fiber tractography.

In order to compare inter-subject AFD, Raffelt and colleagues [9] chose a response 43 44 function common to all subjects to minimize the differences between subjects for voxel-wise AFD comparison. However, this may potentially result in a bias in the estimated FOD. 45 Specifically, the use of such a common response function for group-wise analysis may cause 46 47 biases in the FOD peak orientations for individual subjects. Therefore, whereas a common 48 response function is optimal for the comparison of AFD and HMOA in group studies [9], it is unclear whether this is also optimal for group-wise tractography studies because of the 49 potentially inaccurate FOD peak orientations and concomitant spurious FOD peaks. Intuitively, 50 the difference in response function characteristics across healthy subjects are not expected to 51

be large, as response functions are generally averaged from more than hundreds of voxels 52 that are supposed to contain single fiber populations [6,7,12]. This was partly demonstrated by 53 Jeurissen and colleagues [13], who studied the inter-subject response functions of 100 healthy 54 55 subjects from the Human Connectome Project (HCP) [14] and observed only subtle differences. Accordingly, it seems justified not to be too concerned about inter-subject 56 response function variability in healthy subjects, since either using averaged response 57 functions or individual response functions is not likely to affect the FOD profiles in the HCP 58 59 dataset. However, although the differences in the response functions of healthy subjects may 60 be small [13], this is likely not the case for subjects with some form of pathology. The intersubject signal deviations do raise concern for aging and diseased groups. 61

White matter degeneration, whether caused by aging or by a disease process, may substantially alter the response function. Hence, studying subjects of different ages with a common response function might introduce errors due to discrepancies in white matter characteristics. Therefore as the focus of this work, it is useful to investigate such differences in response functions and the resulting variations of the FOD. A thorough numerical evaluation focusing on the angular characteristics of FOD is needed to shed more light on this issue.

Previous studies have discussed the effect of improperly calibrated response functions 68 on the FOD characteristics and fiber tracking. Tournier [7] and Dell'Acqua [8] reasoned from a 69 70 mathematical point of view that a wrongly chosen response function would affect the 71 magnitudes of FOD peaks, thus also AFD and HMOA, but would leave their orientations 72 unaffected. Dell'Acqua and colleagues [8,10] investigated with simulations and in vivo data the effects of various response function changes on the FOD profiles, including variations in the 73 74 response function shape and scaling factor, as well as in axonal radius and in angle of crossing 75 pathways for the damped Richardson-Lucy (dRL) method. Their paper focused on the effect of the response function on FOD amplitudes and the sensitivity of HMOA to diffusivity changes 76 per fiber population, as compared to traditional metrics as fractional anisotropy (FA) and mean 77 78 diffusivity (MD). Parker [15] studied the FOD peak orientations and the existence of spurious 79 peaks in simulations as a function of the response function miscalibration for CSD and dRL.

80 The results of that study demonstrate that sharper response functions resulted in more spurious peaks in the FOD profiles, and that the mismatch of the calibrated-targeted response 81 82 functions introduced uncertainty on the main FOD peak orientations. However, in previous 83 work[15], the authors used the FA value as a metric to characterize the response functions, a strategy which is unable to describe the true axial and radial diffusivities in crossing fibers [16]. 84 Changes in FA entangle changes in the axial and radial diffusivities, so that the effects on 85 these two diffusivities could not be studied straightforwardly. Here we seek to disentangle 86 87 these effects and, complementing earlier studies [15,17], also aim to quantify both the effect on peak magnitude and angular deviation. 88

In this manuscript we studied how variations in the response function affect voxel-wise 89 FOD characteristics and fiber tracking. Changes in pathology are likely reflected in changes in 90 either the axial or the radial diffusivity, which in our study, is represented by the shape and 91 92 scale factor of the response function. Simulations were designed to explore the effects of the response function shape and scaling factor on the FOD properties, such as the number of FOD 93 94 peaks, their orientation (for tractography) and magnitude, and the AFD. Additionally, in vivo 95 data from the Human Connectome Project (HCP) were used to illustrate how the choice of the response function in CSD can affect the FOD quantification and fiber tracking. 96

98 2. Methods

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In Sections 2.1 and 2.2, we give a brief background on (constrained) spherical deconvolution methods to reconstruct the FOD. In Section 2.3 we outline the simulation experiments and introduce the shape and scaling factor that characterize the response function. Section 2.4 presents the parameter settings used in these simulations. In Section 2.5, the in vivo data experiments are described.

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106 2.1 Constrained Spherical Deconvolution

Recent studies showed that crossing fibers account for over 90% of white matter voxels 108 109 [4]. The DTI representation cannot resolve crossing fibers by design and thus provides nonspecific metrics in such voxels. Spherical deconvolution approaches [6-8,18,19] overcome this 110 limitation and allow for estimating the FOD for more complex fiber configurations, while 111 retaining reasonable computation and acquisition time compared with other methods [20-23]. 112 113 CSD assumes that the diffusion MRI signals can be expressed as the spherical 114 convolution of a fiber response function with the FODs in the spherical harmonics basis, thus also assuming the validity of the response function in all voxels. The response function 115 represents the diffusion-weighted signal of a single fiber population. Spherical harmonics form 116 117 a complete basis on the sphere. However, to fully reconstruct a signal on the sphere, the spherical harmonics should have infinite order, which is not possible in practice. In clinical 118 studies, signals with up to 60 gradient directions are generally acquired, limiting the order of 119 the spherical harmonics to 8, which we also adopted in this work. 120

The FODs are used to infer information on the orientation of the fiber pathways under the assumption that the FOD peak orientations coincide with the underlying fiber directions. To reconstruct the FOD, truncation of the spherical harmonics is needed, causing the so-called "ringing" effect on the FOD profiles, which introduces implausible negative values. In order to suppress the ringing effect and the sensitivity to noise, the regularization of FOD was proposed

[7,19,24] to improve the conditioning of the deconvolution problem, which is further referred to as constrained SD (i.e., CSD). In addition to directional information, the magnitudes of the FOD are used to compute additional metrics, such as AFD [9] and HMOA [10]. The accurate estimation of FOD peak directions and magnitudes is therefore essential for subsequent analysis.

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132 2.2 Shape and scaling of response functions

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The response function used in the CSD process can be either simulated or derived 134 135 directly from the data. Following the latter approach, which is more common, voxels that have 136 a high chance of containing single fiber populations are used to calibrate the response function. A straightforward approach to numerically implement the concept of a single fiber population 137 is to threshold, for instance, the fractional anisotropy (FA), above a pre-defined value. 138 139 However, the choice of FA threshold is not trivial and can cause inaccuracies in the response 140 function estimation [12]. A data-driven method using a recursive calibration framework was proposed to estimate the response function from the subject data in an unbiased way [12]. 141 This method estimates which voxels contain single fiber populations by iteratively excluding 142 143 voxels which do not have a single dominant orientation and updating the estimated response function. 144

The choice of the fiber response function has an impact on the peak directions and magnitudes of the FODs [10,15,19]. Theoretically, changes in the response function are directly reflected in the FOD estimation, but should affect only peak magnitudes while leaving their orientations untouched [6,10]. However, in practice, due to the low SNR level in diffusionweighted MRI data, the ill-posedness of inverse problems, and the regularization process, the effects of the choice of response function on the FODs become less obvious.

Parker et al. [15] investigated alterations of response function by changing its FA value. Here, we acknowledge that changing the FA affects both the scale and the shape of the response function. It is thus not straightforward to disentangle an FOD change into scale and shape effects. To this end, we decompose general changes in the response function into specific changes in shape and scale [8] and analyze their individual effects on the FOD characteristics (i.e., magnitude, the number of peaks, and peak orientations). The following sections describe how we can achieve such changes in shape and scale of the response functions in the simulated and in vivo diffusion MRI data experiments.

- 159
- 160 2.3 Simulation experiments
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162 2.3.1 Modeling of single fiber populations and response functions

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164 If the diffusivity *D* associated with the underlying fiber population is expressed by an 165 axially symmetric diffusion tensor, whose first eigenvector is in parallel with the *z*-axis in the 166 reference coordinate frame, then $D(\theta, \varphi)$ can be written as (Anderson 2005)

$$D_{(\theta,\varphi)} = [\sin\theta\cos\varphi \quad \sin\theta\sin\varphi \quad \cos\theta] \begin{bmatrix} \beta & 0 & 0\\ 0 & \beta & 0\\ 0 & 0 & \lambda \end{bmatrix} \begin{bmatrix} \sin\theta\cos\varphi \\ \sin\theta\sin\varphi \\ \cos\theta \end{bmatrix}, \tag{1}$$

where λ and β are the axial and the radial diffusivity of the single fiber population, (θ, φ) is the polar angle set between the fiber orientation and the applied gradient. Given the axial symmetry property of the diffusion tensor, Eq. (1) can be simplified as

$$D_{(\theta)} = \lambda \cos^2 \theta + \beta \sin^2 \theta = \alpha \cos^2 \theta + \beta,$$
 (2)

where $\alpha = \lambda - \beta$ is the absolute difference between the axial and radial diffusivity. For simplicity, if we assume that the signal $S(\theta, \varphi)$ from each fiber population is a function of $D(\theta, \varphi)$, then the diffusion-weighted signal *S* can then be rewritten as [3]

$$S_{(\theta,\varphi)} = S_0 e^{-bD_{(\theta,\varphi)}},\tag{3}$$

where S_0 is the non-diffusion-weighted signal and *b* is the b-value that represents the strength of diffusion weighting. Combining Eq. (1) – Eq. (3), the diffusion-weighted signals can be expressed as [18]

$$S(\theta) = S_0 e^{-b(\alpha \cos^2 \theta + \beta)} = S_0 K e^{-b\alpha \cos^2 \theta},$$
(4)

where $K = e^{-b\beta}$. Eq. (4) highlights the dependency of *S* on the shape factor α and the scaling factor *K*, following the definition in previous studies [8]. In this equation, the scaling factor *K* depends only on the radial diffusivity of the fiber response, representing the isotropic diffusion within the fiber population, whereas the shape factor α depends on the difference between the axial and radial diffusivities, representing the anisotropic diffusion within the fiber population.

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182 2.3.2 Modifying the shape and scaling factor of the response functions

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Since the response function *R* is intrinsically based on the shape and scaling of the fiber population diffusivities, *R* can be written in the same form as the signal of a fiber population imposed by the gradient at an elevation angle θ with the fiber orientation, which is identical to Eq. (4), i.e.,

$$R(\theta) = S_0 K e^{-b\alpha \cos^2 \theta}.$$
 (5)

According to Eq. (5), we can modify (i) the shape factor α of the response function, by varying only the axial diffusivity with a fixed radial diffusivity, to keep *K* constant; and (ii) the scaling factor *K* of the response function, by changing simultaneously the axial and radial diffusivity, to not alter the shape factor α . We can then study the effects of *R* on FOD characteristics, by selectively introducing a discrepancy into the shape or the scale of a simulated single fiber signal with respect to the response function.

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195 2.3.3 Modeling of multi-fiber populations

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We model the diffusion-weighted signal within a voxel as the sum of multiple compartments measured from each fiber population. Each compartment is assumed to share an identical response function, so the diffusion-weighted signals are depending only on the orientations of the fiber populations in the voxel and on data noise. We further assume that there is no exchange of water molecules between fiber populations, and that each single fiber

population can be represented by a signal $S_{i(\theta)}$ (where *i* denotes the *i*th fiber population). The signal S_{DW} generated by a crossing fiber configuration can then be described by

$$S_{DW} = \sum_{i=1}^{n} f_i S_{i(\theta)},\tag{6}$$

where f_i is the volume fraction of each fiber population, n is the total number of fiber populations intercrossing the voxel, and $i(\theta)$ is the angle between the applied gradient and the i^{th} fiber population. In our work, we focus on configurations of two crossing fiber populations, but the equations of generating the diffusion-weighted signals can also be extended to analyze the FOD characteristics for more than two fiber populations.

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210 2.3.4 Data analysis

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Amongst the SD frameworks, the CSD approach is implemented in several software 212 packages, such as MRtrix [25], Dipy [26] and ExploreDTI [27]. In this work, the FODs were 213 214 estimated with CSD as implemented in *ExploreDTI*. The FOD peak orientations, which are assumed to reflect the underlying fiber orientations [6], and the magnitudes of the FOD peaks, 215 were extracted using a Newton-Raphson gradient descent method [28]. All FOD peaks that 216 were smaller than an absolute threshold of 0.1 were regarded as contributions from noise and 217 218 thus discarded to reduce false positives [29]. All peaks were clustered to the nearest simulated peak directions, by using an angular threshold of 45° to determine whether or not two peaks 219 220 were belonging to the same fiber population. In case of simulating multiple fiber populations, 221 only the estimated FOD peaks closest to the simulated fiber populations were considered. For 222 each simulation, the mean and standard deviation of the following FOD metrics were 223 evaluated:

a. the average difference between the estimated and simulated number of FOD peaks;

b. the angular deviations between the estimated FOD peak orientation and the simulated fiberorientation;

227 c. the estimated separation angles in case of multiple fiber populations;

228 d. the FOD peak magnitudes in case of single fiber populations;

e. the percentage difference of the estimated AFD with respect to the AFD with the referenceresponse function.

The AFD computation was performed as the integral of the FOD magnitudes assigned to each peak, which in the literature is commonly referred to as "lobe". The calculation of the AFD is similar to what was used in a previous study [30], except that we use the gradients generated by the electromagnetic model [31] to segment the FODs for each fiber population instead of using gradients generated by an icosahedron model.

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237 2.4 Parameter settings

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239 We simulated different fiber configurations with a predefined *b*-value equal to 3000 s/ mm^2 , a set of 60 gradient directions [31], and $S_0 = 1$. Rician noise (1000 noise instances) was 240 241 added to the diffusion weighted signals to simulate SNR (with respect to S_0) levels of [50 40] 30 20 15 10]. In the first simulation, a single-fiber configuration was generated with the main 242 diffusion direction along the z-axis, setting $\alpha = 1.2 \times 10^{-3} \text{ mm}^2/\text{s}$ and K = 0.4 (i.e. $\beta \sim (0.3 \times 10^{-3} \text{ mm}^2/\text{s})$ 243 $10^{-3} \,\mathrm{mm^2/s}$)). In the second simulation, a second fiber population was rotated around the y-244 axis and combined with the single-fiber population generated in the first simulation to achieve 245 246 a separation angle ω . Here we simulated crossing fiber populations with separation angles ω = [90°, 75°, 60°, 55°, 50°, 45°, 40°]. 247

For both simulations, two sets of response functions were tested to achieve (a) different shape but the same scaling factors, by increasing α from 0.6 × 10⁻³ mm²/s to 1.8 × 10⁻³ mm²/s with steps of 0.1 × 10⁻³ mm²/s, while keeping *K* constant (Fig. 1a); and (b) the same shape but different scaling factors, by decreasing *K* from 0.7 to 0.3 with steps of 0.1, while keeping α constant (Fig. 1b).

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Fig. 1. The 2D projection of response functions obtained by changing (a) the shape factor α and (b) the scaling factor *K*. The shape factors are defined from 0.6 × 10⁻³ mm²/s to $1.8 \times 10^{-3} \text{ mm}^2/\text{s}$ in steps of $0.1 \times 10^{-3} \text{ mm}^2/\text{s}$. The scaling factors are varied from 0.7 to 0.3 in steps of 0.05.

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- 259 2.5 Peak clustering and angular threshold
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We clustered the peak directions to make sure that we are always comparing the angular deviations between the simulated fiber orientation and the FOD peak orientation most closely aligned to that orientation. Like in other studies [16,32,33] that compare axial and radial diffusion characteristics, we also included an angular threshold (e.g., $\cos(\theta) > 0.7$, which means approximately $\theta < 45^{\circ}$) to make sure the correct peaks were being extracted for further evaluations.

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268 2.6 In vivo data experiments

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Diffusion-weighted MRI data of a single HCP subject was further used to illustrate the 270 effects of ill-defined response functions on voxel-wise FOD characteristics and brain 271 tractography. In summary, diffusion-weighted images were acquired along 90 diffusion 272 gradient directions with a *b*-value of 3000 s/mm^2 in addition to 18 non-diffusion-weighted 273 images, and with an isotropic spatial resolution of 1.25 x 1.25 x 1.25 mm³. We performed CSD 274 based tractography in ExploreDTI with a step size of 1 mm, an FOD threshold of 0.1, an 275 276 angular threshold of 30°, and seeding points per 2mm x 2mm x 2mm across the whole brain. All the tracts were constructed with deterministic fiber tracking to facilitate data interpretation. 277

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279 2.6.1 Modeling the response function

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The reference response function for the in vivo dataset was represented by the diffusion tensor fit to the response function, as estimated with the recursive calibration approach [12]. Similar to the method described in Section 2.3.2, the diffusion tensor was used to model the changes in the shape and the scaling factor of the response functions. The shape factor α of

the response function was modified by +/- $[0.1 - 0.3 \times 10^{-3} \text{ mm}^2/\text{s}]$, while the scaling factor *K* was modified by +/- [0.1 - 0.2].

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288 2.6.2 Evaluation of in-vivo data

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In analogy with the simulations, we computed the voxel-wise difference in number of estimated FOD peaks, the angular deviations of the main orientation, and the percentage difference in AFD of the dominant fiber orientation, for all the estimated FODs. The comparisons of number of FOD peaks were computed for the whole brain, whereas the comparisons of angular deviation and AFD were only computed for voxels with FA > 0.2.

Individual white matter fiber bundles were extracted by using the regions of interest 295 (ROIs) as suggested by Wakana [34]. The segmented fiber pathways include parts of the 296 297 splenium of corpus callosum (sCC), the genu of corpus callosum (gCC), the cingulum (Cg), the uncinate fasciculus (UF), the corticospinal tract (CST), and the temporal part of the superior 298 longitudinal fasciculus (tSLF). The average FOD characteristics for each fiber bundle were 299 calculated. In addition, FOD characteristics of the response function were computed from (1) 300 301 the region with a single fiber population as identified during the recursive calibration step 302 (referred to as "SFP-mask"); and (2) the region with voxels for which FA > 0.2 (referred to as "FA-mask"). 303

304 **3. Results**

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- 306 3.1 FOD characteristics of single fiber populations
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Fig. 2 shows the effect of changing the shape factor and the scaling factor of the response function on the FOD characteristics in a single fiber population. At SNR < 20, the average number of spurious peaks increases when the shape factor increases, but only slightly increases when the scaling factor decreases (Fig. 2A). The angular deviation depends mainly on the SNR and is far less affected by changes in shape or scale factor of the response function

313 (Fig. 2B). By contrast, changes in peak magnitude (Fig. 2C) and the AFD (Fig. 2D) as a function 314 of shape and scaling factor of the response function are more pronounced than due to 315 differences in SNR level alone. Notice that the effect of changing the scaling factor (up to 316 \sim 60%) is roughly three times larger compared to changing the shape factor (up to \sim 20%).

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Fig. 2. Effect of simulating changes in the response function on FOD characteristics for 318 a single fiber configuration at different SNR levels. Shape factor α and the scaling factor 319 K of the response function (RF) are varied at different SNR levels to investigate (A) the 320 introduction of spurious peaks, i.e., the average difference between the estimated and 321 predefined number of FOD peaks; (B) the confidence interval (average ± standard error) of the 322 angular deviation of the primary FOD peak; (C) the percentage difference between the 323 amplitudes of the estimated FOD peak and the ground-truth FOD peak; and (D) the percentage 324 difference between the estimated AFD of the primary fiber population and the ground-truth 325 AFD. The dashed vertical lines represent the ground-truth values. 326

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328 3.2 Occurrence of spurious peaks

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Fig. 3 shows the average difference between the number of estimated and simulated 330 FOD peaks in relation to the shape (left) and the scaling (right) factor of the response functions 331 for different SNR levels. Overall, performing spherical deconvolution with sharper response 332 333 functions (i.e., higher values of the shape factor) generally introduces more spurious peaks. 334 On the other hand, CSD fails to extract all the simulated peaks from the estimated FODs when the response function shape factor has smaller values, in particular for separation angles 335 336 below 55°. With higher noise levels, more spurious peaks are introduced, especially for higher 337 values of the shape factor. Furthermore, adjusting the scaling factor has no significant effect on the estimated number of spurious peaks. While there are hardly any spurious peaks 338 introduced at the lower noise levels (SNR = 30 and 50), additional incorrect peaks can be 339 observed at the higher noise level (SNR = 10). 340

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Fig. 3. The average difference between the number of estimated and simulated FOD peaks as a function of shape (left) and scaling (right) factor of the response function (RF) at three SNR levels (different SNR for each row). Brighter yellow areas show a higher probability of introducing spurious peaks, whereas darker blue areas show a higher probability of merging the two simulated peaks into one peak. The dashed vertical lines indicate that the settings of the response function are identical to those used for generating the underlying signals. Notice that different scaling of the colorbars were used for better contrast.

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350 3.3 Angular deviation

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352 **3.3.1** The effect of the shape factor

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354 Fig. 4 shows the results of investigating the effect of the response function's shape 355 factor on the angular characteristics of FOD peaks at SNR = 50, 30 and 10 for crossing fiber configurations with different separation angles. At lower noise levels (SNR = 30 and 50), lower 356 357 values of the shape factor generally cause an underestimation of the separation angles, except 358 when the two simulated fiber populations are orthogonal to each other (i.e., 90°) (Fig. 4A). At 359 the higher noise level (i.e., SNR = 10), the bias in the estimated separation angle due to 360 changes in the shape factor is swamped by the noise itself, especially for lower separation 361 angles. From the observed angular deviations in Fig. 4B (the first peak) and Fig. 4C (the second peak) we can observe, in general, that for smaller simulated separation angles, the 362 adverse effects of changing the shape factor of the response function on the estimated FOD 363 angular characteristics are more pronounced. 364

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Fig. 4. Results of exploring the impact of response functions with different shape factor α on the FOD peaks for crossing fiber configurations simulated with separation angles ranging from 90° to 40°. Fig. 4A shows the estimated separation angles between the two primary peaks. Dashed horizontal lines indicate the simulated separation angles. Fig. 4B and Fig. 4C show the angular deviations between the estimated first (p1) and second (p2) FOD peaks and their corresponding simulated fiber orientations. Solid line interruptions occurred when one of the two peaks was not detected. The means of the estimated values are plotted with the standard error as the shaded areas. Dashed vertical lines are defined as in Fig. 3.

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375 3.3.2 The effect of the scaling factor

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Fig. 5 shows the angular deviations between the orientation of the estimated FOD 377 378 peaks and the simulated fiber orientations as a function of the scaling factor. Overall, crossing 379 fibers with separation angles smaller than 45° show larger angular deviations than those with 380 more orthogonal separation angles. In Fig. 5A, the estimated separation angles do not change 381 significantly as a function of the scaling factor of the response function. Nevertheless, smaller simulated separation angles result in a larger bias of the estimated separation angles. Fig. 5B 382 383 and Fig. 5C present the angular deviations of the first and second FOD peak, respectively. The angular deviations are not significantly affected by the scaling factor, but do depend on the 384 385 magnitude of the separation angles of the two fiber populations.

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Fig. 5. The effect of varying the scaling factor (K) of the response function on the FOD 387 peaks for crossing fiber configurations simulated with separation angles ranging from 388 **90**° to **40**°. Fig. 5A shows the estimated separation angles between the two primary peaks. 389 390 Dashed horizontal lines indicate the simulated separation angles. Fig. 5B and Fig. 5C show the angular deviations between the estimated first (p1) and second (p2) FOD peaks and the 391 corresponding simulated fiber orientations. Solid line interruptions occurred when one of the 392 393 two peaks was not detected. The means of the estimated values are plotted with the standard 394 error as the shaded areas. Dashed vertical lines are defined as in Fig. 3.

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396 3.4 AFD per fixel

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Fig. 6 shows the percentage difference of the AFD of the first and second fiber 398 399 population in relation to the response function shape factor (A, B) and scaling factor (C, D). In Fig. 6A, at SNR 50 and 30, the AFD started at a very high value when the shape factor is 400 401 smaller than 0.8, 1.0 and 1.4 x 10⁻³ mm²/s for the simulated separation angles of 55°, 50° and 45°, respectively. The AFD values converge to the AFD of the other separation angles as the 402 shape factor increases. As shown in the angular characteristics results (Fig. 4), when the 403 404 response function becomes sharper, the drop points of AFD for small separation angles 405 indicate the boundaries at which CSD is just able to separate the two fiber populations. In case of the 40° separation angle, only one FOD peak is obtained. The large difference in AFD for 406 small separation angles (45°-55°) with decreased shape factors can be a confounding factor in 407 inter-subject comparisons of AFD studies, which will be discussed further in Section 4.3. At 408 409 SNR 10, the AFD differences are more related to noise than to the shape of the response 410 function for smaller separation angles (below 60°). As for the second peak (Fig. 6B), the AFD 411 can change from -30% to 20% when the shape factor was modified from -50% to 50%, 412 respectively.

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Fig. 6. The percentage difference of the estimated AFD of the first peak (p1) and the second peak (p2) in relation to the response function shape factor α (A, B) and scaling factor *K* (C, D) at different SNR levels. The quick drop of the AFD difference while increasing the shape factor indicates when CSD was able to separate the two fiber populations. Dashed vertical lines are defined as in Fig. 3.

Fig. 6C and Fig. 6D show the percentage difference of the AFD of the first and second fiber population in relation to the scaling factor of the response function. In line with the simulation results for single fiber populations (Fig. 2D), AFD can change up to 80% due to the scaling factor changes for the second peak. For simulated separation angles of approximately 45°, AFD of the first fiber population can be over-estimated up to as much as 150%. For the

| 424 | other simulated separation angles, the AFD of the primary peak can vary from -40% to 70% at |
|-----|---|
| 425 | SNR = 50 and SNR = 30, irrespective of the simulated separation angles. Notice that the AFD |
| 426 | changes are not linearly related with changes in the scaling factor. |

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428 3.5 In vivo HCP data set

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430 3.5.1 FOD characteristics of white matter

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In this section, we present the effect of changing the shape and scaling factors of the 432 response function on FOD characteristics for an axial slice of the HCP data set. The difference 433 in number of FOD peaks per voxel is shown in Fig. 7. Differences are typically seen in areas 434 with partial volume effects and with mostly a peak number difference value of one. When the 435 difference in shape factor, denoted by $\Delta \alpha$, increases by 0.1 $\times 10^{-3}$ mm²/s to 0.3 $\times 10^{-3}$ 436 mm²/s, one can see the increase in occurrence of peak number deviations, such as, for 437 instance, in mid-sagittal regions of the corpus callosum. With the increase of difference in 438 scaling factor, denoted by ΔK , regions containing CSF showed higher peak number differences 439 than regions with white and gray matter. 440

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Fig. 7. The difference between the number of FOD peaks estimated with the tensorbased response function and the number of FOD peaks computed with the response function modified according to certain changes in scaling (ΔK) and shape ($\Delta \alpha$) factors. The background is an axial view of the FA map. The peak number difference mostly occurs in grey matter and CSF areas, and crossing fiber regions for white matter, as indicated by the colormap. In regions with single fiber populations (e.g., middle parts of the corpus callosum) spurious peaks are hardly present.

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450 Fig. 8 shows the angular difference between the primary FOD peak, computed with the 451 tensor-fit to the recursive calibrated response function, and the FOD peak obtained with the 452 modified shape and scale factors of the response function. In general, regions containing 453 crossing fibers are affected most when modifying the shape of response functions, with angular 454 deviations of the main FOD peak of up to 3°. Notice that the angular deviation is mostly affected 455 by changing the shape factor, rather than the scaling factor. In addition, while changing ΔK did 456 not affect the angular deviation, increasing the magnitude of $\Delta \alpha$ resulted in larger angular 457 deviations in the same locations.

458

Fig. 8. The angular deviations between the FOD peaks estimated with the tensor-fit of 459 460 the response function and the FOD peaks estimated with the response function 461 modified according to certain changes in scaling (ΔK) and shape ($\Delta \alpha$) factors. The 462 background is an axial view of the FA map and, for clarity, the angular deviations are shown only in regions where FA > 0.2. Most angular differences are in the range of $0-3^{\circ}$. Similar to the 463 results of spurious peaks shown in Fig. 7, angular deviations are larger in regions with crossing 464 fiber populations than regions with single fiber populations, such as the middle part of the 465 466 corpus callosum. Notice that the angular deviations are much higher with regard to shape factor changes than scaling factor changes. 467

468

Fig. 9 shows the voxel-wise AFD difference for the dominant fiber direction between 469 470 the FOD estimated using the tensor-fit to the recursive calibrated response function and the FOD obtained with the modified shape and scale factors of the response function for the HCP 471 data set. The AFD shows a very different pattern in relation to the shape factor changes 472 compared to scaling factor changes. The AFD differences are homogenous throughout the 473 474 brain when the scaling factor varies, while the outliers indicate the voxels where there are 475 potential geometrical differences in the estimated AFD from the reference, such as merging or spurious peaks. The AFD differences are up to 98% when the scaling factor K decreased by 476 0.2. When changing the shape factor with -0.3 $\times 10^{-3}$ mm²/s to 0.3 $\times 10^{-3}$ mm²/s, the highest 477 differences (around 6 to 8%) were observed in areas with a single-fiber population, such as 478

479 the corpus callosum. Notice that bigger changes of the shape factor α makes the AFD 480 difference more heterogeneous across the brain.

481

Fig. 9. The percentage difference of the apparent fiber density (AFD) between the FOD peaks estimated with the tensor-fit of the response function and the FOD peaks estimated with the response function modified according to certain changes in scaling (ΔK) and shape ($\Delta \alpha$) factors. The background is an axial view of the FA map and, for clarity, the AFD percentage differences are shown only in regions where FA > 0.2. Notice that the AFD difference stays homogenous with respect to the scaling factor changes, whereas it is heterogeneous when the shape factor changes.

489

490 3.5.2 Effect on fiber tractography

491

Fig. 10 shows the effect of changing the scaling and shape factors of the response function on the reconstruction of the pathways of the tSLF. The reference trajectories (shown in yellow) are computed with the recursive calibration method. While not much differences can be observed for the main part of the reconstructed tracts, changing the response function mainly affected the trajectories where the tSLF enters the frontal and temporal lobes (see enlarged regions in Fig. 10).

498

Fig. 10. The temporal part of the superior longitudinal fasciculus (tSLF) reconstructed 499 with the FODs estimated using the tensor-fit to the recursively calibrated response 500 501 function (yellow), and the tSLF from the same ROIs reconstructed with FODs estimated using the modified response functions. The other fiber bundles (shown in red, blue, cyan, 502 magenta, and green) indicate the effect of changing the scaling (ΔK) and shape ($\Delta \alpha$) factors 503 504 of the response function on the trajectory of the tSLF. Notice the subtle differences in how the fiber trajectories terminate in the temporal lobe (zoomed areas; the "+" and "-" indicate increase 505 and decrease in the scaling and shape factors, respectively). 506

507

Fig. 11 shows the FOD characteristics for the FA-mask, the SFP-mask, and the 508 509 extracted fiber bundles (gCC, sCC, CST, UF, Cg and tSLF). From all the three FOD characteristics (i.e., spurious peaks, angular deviations, and AFD percentage differences), we 510 can spot a similar trend for all the bundles and the masks with respect to the changes in the 511 shape and scaling factors of the response function. Overall, the UF has the highest average 512 number of spurious peaks. The lowest average angular deviations of the first FOD peak can 513 514 be seen for the SFP-mask. Furthermore, the alterations of the shape factor of the response function can cause angular deviations up to 6°, while the alterations of the scaling factor hardly 515 cause any angular differences in the masks or the selected fiber bundles (see the enlarged 516 plot). Finally, the differences in AFD are relatively homogenous across the extracted fiber 517 bundles and masks with as a function of changing the shape or the scaling factors. 518

519

Fig. 11. The average number of spurious peaks, the average angular deviations, and the 520 average percentage differences in AFD of the first fiber population for the FA-mask, the 521 SFP-mask, and the selected fiber bundles (shown on the right) when a modified 522 523 response function was used in comparison to the original tensor-fit to the recursive calibrated response function. The effect of the changes in the scaling (ΔK) and shape ($\Delta \alpha$) 524 factors of the response function on the selected fiber bundles are reflected in the different color 525 encoding. sCC = splenium of corpus callosum; gCC = genu of corpus callosum; Cg = cingulum; 526 UF = uncinate fasciculus; CST = corticospinal tract; tSLF = temporal part of superior 527 528 longitudinal fasciculus.

529 4. Discussion

530

531 In this work we investigated the effect of changing response function properties on the 532 FOD characteristics using numerical simulations and in vivo HCP data. In particular, we show 533 how miscalibration of the response function, as defined by adjusting the scaling and shape

factors, can introduce a bias in the orientation and magnitude of fiber population peaks. Our 534 findings demonstrate that CSD is prone to produce spurious FOD peaks in the presence of 535 miscalibrated response functions, especially in data with insufficient SNR levels. The 536 537 occurrence of such spurious peaks can also introduce inaccurate fiber pathway reconstructions with fiber tractography. Overall, in agreement with former studies, spurious 538 peaks are introduced due to overestimating the shape factor of the response function, while 539 underestimating the shape factor will result in lower angular resolution of the FOD lobes 540 541 [10,15]. Proper tuning of the response function is therefore necessary to achieve an optimal balance between increasing the angular resolution and minimizing the number of spurious 542 peaks, especially for smaller separation angles (i.e., below 60°) and at low SNR levels. Further, 543 AFD estimation can be influenced by the choice of response function, which will be discussed 544 545 in section 4.3.

546

547 4.1 Effect of shape and scaling factors with simulations

548

At SNR levels of 30 and 50, the FOD characteristics are consistently affected by the 549 choice of the response functions, while at SNR of 10, noise is the dominating factor that affects 550 551 the FOD properties (Fig. 3). In addition, more spurious peaks are observed at SNR of 10. At relatively high SNR levels, the shape factor of the response function has a greater impact on 552 the results than the scaling factor. In particular, using a sharper response function for 553 separation angles below 50° can potentially increase the angular resolution of CSD and can. 554 555 therefore, improve the estimation of the number of peaks (Fig. 3). The shape of the response function was reported to vary with axonal injury and brain maturation, whereas the scaling 556 factor was observed to change as result of demyelination, axonal diameters and axonal density 557 changes [10,35]. This implies that in brain regions affected by disease, applying CSD with a 558 559 response function determined by healthy white matter data can result in unreliable estimates 560 of FOD characteristics.

561

- 562 4.2 Effect of the separation angle between crossing fiber populations
- 563

The extent to which the FODs will be affected by the response function depends largely 564 565 on the separation angle between crossing fiber populations (Fig. 4). More orthogonally crossing fiber orientations are less sensitive to response function changes, as originally 566 suggested in the spherical deconvolution paper [6]. In voxels containing crossing fiber 567 configurations with smaller separation angles (e.g., below 60°), the average angular deviations 568 569 and their variance increase rapidly with lower shape factors of the response function. By contrast, a higher shape factor of the response function results in a smaller bias in the 570 571 computation of the FOD peak orientations than the underestimation of the shape factor (Fig. 4 572 and Fig. 5).

573

574 4.3 Effect of shape factor on AFD

575

For fiber populations with separation angles below 55°, CSD fails to estimate the correct 576 number of peaks when response functions with a lower shape factor are employed, leading to 577 artificially higher AFD values (Fig. 6). As FOD peaks merge together when the shape factor is 578 579 further decreased, the AFD becomes close to the integral of the total FOD amplitudes within the voxel. This is shown in Fig. 6 for simulated separation angles between 45° to 55°. For these 580 relatively small separation angles, the large AFD difference is caused by the limited angular 581 resolution of CSD with the simulated settings. Previous studies [36] reports AFD as a more 582 583 sensitive diffusion marker in traumatic brain injury than the traditional metrics. However, one should be aware that these changes in AFD in the presence of pathology could result from 584 global response function differences between subjects, rather than local diffusivities 585 alterations. 586

587

588 4.4 Effect of FOD angular deviations on fiber tracking

590 If the angular deviations of the FOD peaks are similar in the neighborhood voxels along the white matter pathways, accumulating effects on reconstructed fibers will be significant. By 591 592 contrast, the heterogonous angular deviations of the FOD peaks may only change the voxel-593 wise characteristics like AFD and number of fiber population peaks, the fiber pathways remains if the angular deviations of FOD was not big enough to end in different voxels in the trajectory. 594 Generally, fiber tractography results will not be severely affected in the main part of the fiber 595 bundles, but may show subtle differences at the edges (Fig. 10). In addition, the termination of 596 597 fiber pathways passing through crossing regions can be affected [12]. With the in vivo HCP data, only minor changes in the tSLF trajectories are detected when using the modified 598 response functions with different shape factors. Nevertheless, an inaccurate response function 599 will influence the FODs and subsequently fiber tractography results. 600

601

4.5 Limitations and future directions 602

603

The reference value of the shape and scaling factor of the simulated diffusion-weighted 604 605 signals match with the values in the corpus callosum as reported before. However, recent studies [37-40] indicated that the diffusivities of fiber bundles in the brain are not always the 606 607 same. There is not a full map of diffusivity characteristics of each white matter structure yet. Although our simulation study included the same configurations of crossing fiber bundles in a 608 voxel, in reality, the diffusivities of these crossing fibers may not be identical. 609

In this study, we showed tractography results of an HCP subject using the tensor-fit to 610 611 the recursively calibrated response function and modified response functions. In group studies between healthy subjects and patients with neural degradation diseases (e.g., Alzheimer's 612 disease), it would be useful to compare the alterations of response functions. If there is a 613 group-wise alteration of the shape and the scaling factor of the response functions, we should 614 615 first exclude the deviations of the diffusivities of the diseased group from the healthy subjects, 616 to ensure that FOD characteristics and fiber tractography changes are not the effects of the

response function alteration itself. Furthermore, we can separate the effects of disease onwhite matter fiber tracking from the effects of response functions used in the FOD estimation.

619 **5. Conclusion**

620

This study demonstrates with numerical simulations and in vivo HCP data that 621 decreasing the shape factor of the response function can cause large angular deviations of the 622 623 FOD peak orientations in crossing fibers. Sharper response functions are responsible for introducing spurious peaks, which can also confound subsequent tractography results. 624 Extremely low shape factors of the response function can cause significant angular deviations 625 and may complicate the interpretation in studies involving pathology. In addition, although 626 627 individual angular deviations of FOD peak orientations are small for single voxels at most 628 separation angles, the adverse effect can accumulate for brain tractography. Since smaller separation angles are more sensitive to changes of response function shape factors, future 629 work of inter-subject AFD and pathological groups should be aware of this possible 630 631 confounding factor when investigating brain structures with crossing fiber configurations.

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634 **References**

- Stejskal EO, Tanner JE. Spin diffusion measurements: Spin echoes in the presence of a time dependent field gradient. J Chem Phys. 1965;42(1):288–92.
- Le Bihan D, Breton E, Lallemand D, Grenier P, Cabanis E, Laval-Jeantet M. MR imaging of
 intravoxel incoherent motions: application to diffusion and perfusion in neurologic disorders.
 Radiology. 1986;161:401–7.
- Basser PJ, Mattiello J, LeBihan D. MR diffusion tensor spectroscopy and imaging. Biophys J
 [Internet]. 1994;66(1):259–67. Available from: http://dx.doi.org/10.1016/S0006 3495(94)80775-1
- 4. Jeurissen B, Leemans A, Tournier JD, Jones DK, Sijbers J. Investigating the prevalence of
 complex fiber configurations in white matter tissue with diffusion magnetic resonance
 imaging. Hum Brain Mapp. 2013;34(11):2747–66.
- 647 5. Behrens TEJ, Berg HJ, Jbabdi S, Rushworth MFS, Woolrich MW. Probabilistic diffusion
 648 tractography with multiple fibre orientations: What can we gain? Neuroimage. 2007;34:144–
 649 55.
- Tournier JD, Calamante F, Gadian DG, Connelly A. Direct estimation of the fiber orientation
 density function from diffusion-weighted MRI data using spherical deconvolution.
 Neuroimage. 2004;23(3):1176–85.
- 7. Tournier JD, Calamante F, Connelly A. Robust determination of the fibre orientation
 distribution in diffusion MRI: Non-negativity constrained super-resolved spherical
 deconvolution. Neuroimage. 2007;35:1459–72.
- Bell'Acqua F, Rizzo G, Scifo P, Clarke RA, Scotti G, Fazio F. A model-based deconvolution
 approach to solve fiber crossing in diffusion-weighted MR imaging. IEEE Trans Biomed Eng.
 2007;54(3):462–72.
- 859 9. Raffelt D, Tournier JD, Rose S, Ridgway GR, Henderson R, Crozier S, et al. Apparent Fibre
 bensity: A novel measure for the analysis of diffusion-weighted magnetic resonance images.
 Neuroimage [Internet]. 2012;59(4):3976–94. Available from:
 http://dx.doi.org/10.1016/j.neuroimage.2011.10.045
- 10. Dell'Acqua F, Simmons A, Williams SCR, Catani M. Can spherical deconvolution provide more
 information than fiber orientations? Hindrance modulated orientational anisotropy, a truetract specific index to characterize white matter diffusion. Hum Brain Mapp.
 2013;34(10):2464–83.
- 667 11. Dhollander T, Raffelt D, Connelly A. Unsupervised 3-tissue response function estimation from
 668 single-shell or multi-shell diffusion MR data without a co-registered T1 image. ISMRM Work
 669 Break Barriers Diffus MRI [Internet]. 2016;5. Available from:
- 670 https://www.researchgate.net/publication/307863133_Unsupervised_3-
- 671 tissue_response_function_estimation_from_single-shell_or_multi-
- 672 shell_diffusion_MR_data_without_a_co-registered_T1_image
- Tax CMW, Jeurissen B, Vos SB, Viergever M a., Leemans A. Recursive calibration of the fiber
 response function for spherical deconvolution of diffusion MRI data. Neuroimage [Internet].
 2014;86:67–80. Available from: http://dx.doi.org/10.1016/j.neuroimage.2013.07.067
- Jeurissen B, Sijbers J, Tournier J-D. Assessing inter-subject variability of white matter response
 functions used for constrained spherical deconvolution. In: ISMRM 23th Annual Meeting,

678 Toronto, Ontario, Canada. 2015. p. 2834. 679 14. Van Essen DC, Ugurbil K, Auerbach E, Barch D, Behrens TEJ, Bucholz R, et al. The Human Connectome Project: A data acquisition perspective. Neuroimage. 2012;62(4):2222–31. 680 681 15. Parker GD, Marshall D, Rosin PL, Drage N, Richmond S, Jones DK. A pitfall in the reconstruction of fibre ODFs using spherical deconvolution of diffusion MRI data. Neuroimage [Internet]. 682 683 2013;65:433–48. Available from: http://dx.doi.org/10.1016/j.neuroimage.2012.10.022 Wheeler-Kingshott CAM, Cercignani M. About "axial" and "radial" diffusivities. Magn Reson 684 16. 685 Med. 2009;61(5):1255-60. 686 17. Dell'Acqua F, Scifo P, Rizzo G, Catani M, Simmons A, Scotti G, et al. A modified damped 687 Richardson-Lucy algorithm to reduce isotropic background effects in spherical deconvolution. 688 Neuroimage [Internet]. 2010;49(2):1446–58. Available from: http://dx.doi.org/10.1016/j.neuroimage.2009.09.033 689 18. Anderson AW. Measurement of fiber orientation distributions using high angular resolution 690 691 diffusion imaging. Magn Reson Med. 2005;54:1194–206. 19. 692 Jian B, Vemuri BC. A unified computational framework for deconvolution to reconstruct 693 multiple fibers from diffusion weighted MRI. IEEE Trans Med Imaging. 2007;26(11):1464–71. 694 20. Descoteaux M, Angelino E, Fitzgibbons S, Deriche R. Regularized, fast, and robust analytical Q-695 ball imaging. Magn Reson Med. 2007;58:497-510. 696 21. Tuch DS. Q-ball imaging. Magn Reson Med. 2004;52(6):1358–72. 697 22. Jansons KM, Alexander DC. Persistent angular structure: new insights from diffusion magnetic resonance imaging data. Inverse Probl [Internet]. 2003;19(5):1031. Available from: 698 699 http://stacks.iop.org/0266-5611/19/i=5/a=303 700 23. Wedeen VJ, Hagmann P, Tseng WYI, Reese TG, Weisskoff RM. Mapping complex tissue 701 architecture with diffusion spectrum magnetic resonance imaging. Magn Reson Med. 702 2005;54(6):1377-86. 703 24. Ramirez-Manzanares A, Rivera M, Vemuri BC, Carney P, Mareei T. Diffusion basis functions 704 decomposition for estimating white matter intravoxel fiber geometry. IEEE Trans Med 705 Imaging. 2007;26(8):1091-102. 25. 706 Tournier J-D, Calamante F, Connelly A. MRtrix: Diffusion tractography in crossing fiber regions. 707 Int J Imaging Syst Technol [Internet]. 2012;22(1):53–66. Available from: http://dx.doi.org/10.1002/ima.22005 708 709 26. Garyfallidis E, Brett M, Amirbekian B, Rokem A, Van Der Walt S, Descoteaux M, et al. Dipy, a 710 library for the analysis of diffusion MRI data. Front Neuroinform. 2014;8:8. 711 27. Leemans A, Jeurissen B, Sijbers J, Jones D. ExploreDTI: a graphical toolbox for processing, 712 analyzing, and visualizing diffusion MR data. Proc 17th Sci Meet Int Soc Magn Reson Med 713 [Internet]. 2009;17(2):3537. Available from: http://www.mendeley.com/research/exploredti-714 a-graphical-toolbox-for-processing-analyzing-and-visualizing-diffusion-mrdata/%5Cnhttp://www.exploredti.com/ref/ExploreDTI ISMRM 2009.pdf 715 716 28. Jeurissen B, Leemans A, Jones DK, Tournier J-D, Sijbers J. Probabilistic fiber tracking using the 717 residual bootstrap with constrained spherical deconvolution. Hum Brain Mapp. 2011 718 Mar;32(3):461-79. 719 29. Jeurissen B, Leemans A, Tournier J-D, Jones DK, Sijbers J. Investigating the prevalence of

complex fiber configurations in white matter tissue with diffusion magnetic resonance

| 721 | | imaging. Hum Brain Mapp. 2013;34(11):2747–66. |
|---------------------------------|-----|---|
| 722 723 724 | 30. | Smith RE, Tournier JD, Calamante F, Connelly A. SIFT: Spherical-deconvolution informed filtering of tractograms. Neuroimage [Internet]. 2013;67:298–312. Available from: http://dx.doi.org/10.1016/j.neuroimage.2012.11.049 |
| 725 726 | 31. | Jones DK, Horsfield MA, Simmons A. Optimal strategies for measuring diffusion in anisotropic systems by magnetic resonance imaging. Magn Reson Med. 1999;42(3):515–25. |
| 727 728 729 | 32. | Jbabdi S, Behrens TEJ, Smith SM. Crossing fibres in tract-based spatial statistics. Neuroimage [Internet]. 2010;49(1):249–56. Available from: http://dx.doi.org/10.1016/j.neuroimage.2009.08.039 |
| 730 731 732 | 33. | Tax CMW, Westin CF, Dela Haije T, Fuster A, Viergever MA, Calabrese E, et al. Quantifying the brain's sheet structure with normalized convolution. Med Image Anal [Internet]. 2017;39:162–77. Available from: http://dx.doi.org/10.1016/j.media.2017.03.007 |
| 733 734 735 736 | 34. | Wakana S, Caprihan A, Panzenboeck MM, Fallon JH, Perry M, Gollub RL, et al. Reproducibility of quantitative tractography methods applied to cerebral white matter. Neuroimage [Internet]. 2007;36(3):630–44. Available from: http://dx.doi.org/10.1016/j.neuroimage.2007.02.049 |
| 737 738 | 35. | Feldman HM, Yeatman JD, Lee ES, Barde LHF, Gaman-Bean S. Diffusion tensor imaging: a review for pediatric researchers and clinicians. J Dev Behav Pediatr JDBP. 2010;31(4):346. |
| 739 740 741 | 36. | Wright DK, Johnston LA, Kershaw J, Ordidge R, O'Brien TJ, Shultz SR. Changes in apparent fiber density and track-weighted imaging metrics in white matter following experimental traumatic brain injury. J Neurotrauma. 2017;34(13):2109–18. |
| 742 743 744 745 746 | 37. | Tax CMW, Novikov DS, Garyfallidis E, Viergever MA, Descoteaux M, Leemans A. Localizing and Characterizing Single Fiber Populations Throughout the Brain. Proc 23rd Annu Meet ISMRM, Toronto, Canada [Internet]. 2015;59(6):473. Available from: http://scil.dinf.usherbrooke.ca/wp-content/papers/tax-etal-ismrm15a.pdf VN - readcube.com |
| 747 748 749 | 38. | Novikov DS, Jespersen SN, Kiselev VG, Fieremans E. Quantifying brain microstructure with diffusion MRI: Theory and parameter estimation. ArxivOrg [Internet]. 2016;1–38. Available from: http://arxiv.org/abs/1612.02059 |
| 750 751 | 39. | Jespersen SN, Kroenke CD, Østergaard L, Ackerman JJH, Yablonskiy DA. Modeling dendrite density from magnetic resonance diffusion measurements. Neuroimage. 2007;34(4):1473–86. |
| 752 753 | 40. | Kroenke CD, Ackerman JJH, Yablonskiy DA. On the nature of the NAA diffusion attenuated MR signal in the central nervous system. Magn Reson Med. 2004;52(5):1052–9. |
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| 755 | | |



Fig_1











Difference in number of FOD peaks



Angular deviations of FOD peaks



Difference in AFD (%)

Difference in AFD (%)



https://doi.org/10.1101/760546tSLF when applying modified response functions





 $\Delta K = -0.2$

 $\Delta K = 0.2$



K = 0.4 $\alpha = 1.8 \ge 10^{-3} \text{ mm}^2/\text{s}$





 $\Delta \alpha = -0.1 \ge 10^{-3} \text{ mm}^2/\text{s}$ $\Delta \alpha = 0.1 \ge 10^{-3} \text{ mm}^2/\text{s}$



 $\Delta \alpha = -0.2 \ge 10^{-3} \text{ mm}^2/\text{s}$ $\Delta \alpha = 0.2 \ge 10^{-3} \text{ mm}^2/\text{s}$



 $\Delta \alpha = -0.3 \ge 10^{-3} \text{ mm}^2/\text{s}$ $\Delta \alpha = 0.3 \ge 10^{-3} \text{ mm}^2/\text{s}$





Average number of spurious peaks