Spherical deconvolution with tissue-specific response functions and multi-shell diffusion MRI to estimate multiple fiber orientation distributions (mFODs)

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A B S T R A C T

In diffusion MRI, spherical deconvolution approaches can estimate local white matter (WM) fiber orientation distributions (FOD) which can be used to produce fiber tractography reconstructions. The applicability of spherical deconvolution to gray matter (GM), however, is still limited, despite its critical role as start/endpoint of WM fiber pathways. The advent of multi-shell diffusion MRI data offers additional contrast to model the GM signal but, to date, only isotropic models have been applied to GM. Evidence from both histology and high-resolution diffusion MRI studies suggests a marked anisotropic character of the diffusion process in GM, which could be exploited to improve the description of the cortical organization. In this study, we investigated whether performing spherical deconvolution with tissue specific models of both WM and GM can improve the characterization of the latter while retaining state-of-the-art performances in WM. To this end, we developed a framework able to simultaneously accommodate multiple anisotropic response functions to estimate multiple, tissue-specific, fiber orientation distributions (mFODs). As proof of principle, we used the diffusion kurtosis imaging model to represent the WM signal, and the neurite orientation dispersion and density imaging (NODDI) model to represent the GM signal. The feasibility of the proposed approach is shown with numerical simulations and with data from the Human Connectome Project (HCP). The performance of our method is compared to the current state of the art, multi-shell constrained spherical deconvolution (MSCSD). The simulations show that with our new method we can accurately estimate a mixture of two FODs at SNR≥50. With HCP data, the proposed method was able to reconstruct both tangentially and radially oriented FODs in GM, and performed comparatively well to MSCSD in computing FODs in WM. When performing fiber tractography, the trajectories reconstructed with mFODs reached the cortex with more spatial continuity and for a longer distance as compared to MSCSD and allowed to reconstruct short trajectories tangential to the cortical folding. In conclusion, we demonstrated that our proposed method allows to perform spherical deconvolution of multiple anisotropic response functions, specifically improving the performances of spherical deconvolution in GM tissue.

1. Introduction

Diffusion Magnetic Resonance Imaging (dMRI) is a non-invasive technique sensitive to the microscopic motion (diffusion process) of water molecules (Einstein, 1905). In biologic tissues, the diffusion process is influenced by the presence of biologic membranes and macromolecules (Walter and Hope, 1971), which can hinder and/or restrict the molecular random walk in both isotropic and anisotropic fashions. In 1994, Basser et al. (1994) proposed the diffusion tensor imaging (DTI) framework to characterize the anisotropy of hindered diffusion processes. Later, it was shown that when the diffusion process exhibits a sufficient degree of anisotropy, its principal direction can be computed (Basser et al., 1994) and eventually tracked (Jones et al., 1999), ultimately leading to what is nowadays known as fiber tractography (Mori et al., 1999; Mori et al., 2005) of the brain white matter (WM).

DTI-based fiber tractography is efficient in terms of acquisition and computation but suffers from a number of limitations. Firstly, the technique fails when multiple brain pathways coexist in the same spatial location (Tuch et al., 2002), which has been estimated to be the case in over 90% of the human WM (Jeursen et al., 2013) and which is not likely to be solved by technological improvements as higher imaging resolutions (Schilling et al., 2017). Secondly, DTI-based fiber...
tractography only applies to data acquired within the Gaussian diffusion regime, and thus cannot take advantage of the strong diffusion weightings which are typical for high angular resolution diffusion imaging methods (Xie et al., 2015). Due to these considerations, DTI is not likely the method of choice to study a heterogeneous structure as gray matter (GM), which is characterized by low fractional anisotropy (FA) and, hence, high uncertainty of the principal diffusion direction.

Spherical deconvolution (Tournier et al., 2004; Dell’acqua et al., 2005) is one of the most popular methods to reconstruct fiber orientation distributions (FOD) and to perform fiber tractography in WM. Conversely, fiber tractography based on spherical deconvolution methods has found very limited application to investigate the organization of the cortical GM, despite its pivotal role as endpoint of most axonal bundles. This is most likely due to three unsolved challenges which limit the performance of spherical deconvolution in GM. Firstly, the presence of superficial WM next to the inner cortical surface limits the transition of the streamlines from the deep WM to the cortical GM, and vice versa (Reveley et al., 2015). Secondly, the fiber orientations of the superficial WM are typically tangent to the cortical surface, causing the premature termination of streamlines to and from the cortical banks and sulci, which results in the well-known gyral bias towards the cortical crowns (Schilling et al., 2018). Thirdly, the FODs derived in the cortex are noisy and exhibit spurious peaks, preventing reliable propagation of the streamlines in the cortical ribbon. Importantly, these limitations in fiber tractography propagate to other analysis techniques making use of fiber reconstructions as, among others, structural connectivity approaches (Hagmann et al., 2008), cytorachitectonical analyses (Bastiani and Roebroek, 2015), and pre-surgical planning (Mormina et al., 2016).

In this work, we focus on the third challenge, and propose a novel framework to improve the reliability of the fiber orientations determined with spherical deconvolution in the cortical gray matter. Of note, previous works have proposed geometric heuristics to determine the fiber orientation in the cortex based on the cortical normal (Tozer et al., 2012) or on the vector field from the inner to the outer cortex (Si-Onge et al., 2018). Although these methods have the merit to work with commonly available datasets, they oversimplify the way axons traverse the cortex (Kleinnijenhuis et al., 2013), and do not allow to take into account the simultaneous presence of tangential and perpendicular fiber orientations in the cortex (Kleinnijenhuis et al., 2013; Leuze et al., 2014), which is region- and layer-specific.

The advent of specialized techniques for diffusion imaging, such as ultra-strong gradients (McNab et al., 2013) and simultaneous multislice (Feinberg and Setsompop, 2013), are rapidly increasing the imaging quality and resolution achievable in dMRI. This allows to appreciate new details of the human cortex, as recently reviewed by Assaf (Assaf, 2018). In ex vivo studies (Kleinnijenhuis et al., 2013; Leuze et al., 2014; Aggarwal et al., 2015), it has been consistently shown that the diffusion process observed in the cortex has a large anisotropic component perpendicular to the cortical folding, but also that tangential directions can be observed, for instance, in the central sulcus (McNab et al., 2013). In particular, the anisotropy of the cortex has been observed with ex vivo MRI in animals, as well as in the human brain, both ex vivo (Leuze et al., 2014; McNab et al., 2009; Miller et al., 2012) and in vivo (McNab et al., 2013; Anwander et al., 2010). Such high-quality data acquisitions may soon be routinely available for in vivo applications, offering unprecedented research possibilities (Setsompop et al., 2018) in fiber tractography of GM, but a dedicated framework has not yet been investigated.

Multi-tissue spherical deconvolution approaches (Jeuring et al., 2014) have been suggested to further increase the accuracy of FOD reconstructions in WM by accounting for partial volume effects, e.g. in spatial locations where WM overlaps with GM or cerebrospinal fluid (CSF). Although GM and CSF are merely considered nuisance factors (Dell’acqua et al., 2010), such framework highlights the feasibility of explicitly accounting for multiple tissues in spherical deconvolution approaches. By separating multiple partial volume effects, multi-tissue approaches have shown improved FOD reconstructions in adults (Jeuring et al., 2014), neonates (Pietsch et al., 2019) and in patients with Parkinson’s disease (Kamagata et al., 2018). In current state-of-the-art multi-shell spherical deconvolution, only WM is assumed to be anisotropic, while GM is regarded as an isotropic tissue. To estimate the WM FOD, a WM fiber model – or response function – is estimated from voxels with a single fiber population, which are typically located in large bundles (Tournier et al., 2007) such as the corpus callosum or the corticospinal tract. The response function is then assumed to be valid throughout the brain. While the validity of this representation has been questioned even in WM itself (Schilling et al., 2019), it is highly unlikely that it can adequately represent the anisotropic signature of the cortex, which in addition to axons also contains dendrites and cell bodies (Jespersen et al., 2012; Zhang et al., 2012; Palombo et al., 2020; Lampinen et al., 2019).

In this study, we investigate whether the spherical deconvolution approach can be revisited to consider the different signal characteristics of WM and GM by explicitly modeling both signal mixtures simultaneously. This novel framework allows to reconstruct multiple FODs (mFOD) and is ultimately aimed at improving the FOD estimation and fiber tractography of GM while retaining state-of-the-art performances in WM. In our proposed method, we disregard the existence of a single response function for the brain, and investigate whether it is feasible to simultaneously estimate multiple anisotropic FODs corresponding to tissue-specific response functions. We also evaluate whether this improves the fiber orientation characterization in GM, and fiber tractography in GM as compared to existing spherical deconvolution methods. The mFOD method is in principle also applicable to deep GM structures such as the thalamus, which is often of interest in fiber tractography applications due to its hub role in several fiber bundles (Mallier et al., 2019).

In this work, however, we focus on the study on the cortical GM and only show proof-of-concept of the applicability of mFOD to deep GM, which should be investigated in more detail in future work. Preliminary results of this work have been presented at the 2019 ISMRM meeting in Montreal, Canada (De Luca et al., 2019).

2. Theory and methods

The following paragraphs present the mFOD framework and explain the qualitative and quantitative analyses that we performed using both simulations and in-vivo data.

2.1. Generalized Richardson Lucy (GRL)

In previous work (Guo et al., 2018; Guo et al., 2020), we introduced a multi-shell spherical deconvolution framework named Generalized Richardson Lucy (GRL) to account for multiple tissue types, such as WM, GM and cerebrospinal-fluid (CSF), as shown in Eq. (1).

\[
S_{\text{mFOD}} = f_{\text{DM}} \begin{bmatrix} H_{\text{WM},1}^1 & \cdots & H_{\text{WM},M}^1 \\ \vdots & \ddots & \vdots \\ H_{\text{WM},1}^n & \cdots & H_{\text{WM},M}^n \end{bmatrix} \odot \text{FOD}_{\text{WM}} + f_{\text{GM}} \begin{bmatrix} H_{\text{GM}}^1 \\ \vdots \\ H_{\text{GM}}^n \end{bmatrix} + f_{\text{CSF}} \begin{bmatrix} H_{\text{CSF}}^1 \\ \vdots \\ H_{\text{CSF}}^n \end{bmatrix}
\] (1)

In Eq. (1), \( S \) is the signal collected at \( n \) diffusion weightings, \( S_{\text{mFOD}} \) is the non-diffusion-weighted signal, the \( H \) matrices represent the signal model associated to WM, GM and CSF, \( f \) is the associated signal factor, and \( \text{FOD}_{\text{WM}} \) is the fiber orientation distribution associated to WM. The columns of the deconvolution matrix shown in Eq. (1) represent the possible solution of the spherical deconvolution on the unit sphere, resulting in a total number of \( k + 2 \) columns, as GM and CSF are both considered isotropic.

2.2. Multiple fiber orientation distributions (mFOD)

In the mFOD framework, we redesigned the deconvolution matrix to allow for an arbitrary number of independent anisotropic deconvolution
models. Considering solving the spherical deconvolution problem once more for WM, GM and CSF, but now accounting for anisotropic diffusion in both WM and GM, we can write:

\[
\begin{bmatrix}
F_{\text{WM}}^{H} & \cdots & F_{\text{WM,k}}^{H} & \cdots & F_{\text{GM}}^{H} \\
\vdots & \ddots & \vdots & \ddots & \vdots \\
F_{\text{GM},1}^{H} & \cdots & F_{\text{GM,k}}^{H} & \cdots & F_{\text{GM}}^{H}
\end{bmatrix}
\]

(2)

Eq. (2) can be solved using the iterative least-squares minimizer previously introduced for GRL but replacing the Richardson-Lucy deconvolution scheme with a regularized non-negative least-squares solver (De Luca et al., 2018), producing multiple independent FODs.

2.3. Signal models (response function)

To underline the model independence of the mFOD framework, the H-matrices of WM, GM and CSF, shown in Eq. (2), were evaluated with three different models. The columns of the H-matrices are generated with instances of a specific model, e.g. the signals are generated with a predefined set of parameters and then projected along a number of spatial directions defined on the unit sphere. In other works, the H-matrix is referred to as deconvolution matrix, or response function or kernel. \( H_{\text{CSF}} \) was generated using the ADC model with the typical diffusion coefficient of free water at 37°C (3.0 × 10⁻³ mm²/s). The columns of \( H_{\text{WM}} \) and \( H_{\text{GM}} \) were generated using the DKI model (Jensen et al., 2005) with isotropic kurtosis (De Luca et al., 2017), and the NODDI model (Zhang et al., 2012), respectively, both in simulations and with in-vivo data. For both models, the deconvolution matrix included 300 possible solutions equally distributed on the unit-sphere, which results in an angular resolution of about 4°. The DKI model was chosen as it can represent the anisotropy of the WM signal while also considering non-Gaussian diffusion effects. The NODDI model has been shown to characterize the GM microstructure better than tensor models (Kleimijenhuis, 2014), and allows to take into account the larger presence of cell bodies in GM as compared to WM. It should be noted that mFOD is a generic framework that could be used in combination with models other than DKI and NODDI, such as geometric models (Peri et al., 2014) or data driven representations (Tax et al., 2014). The combination of the simplified DKI and NODDI models should thus not be regarded necessarily as the optimal choice, but rather as a proof of concept. The parameters of the models were determined as explained in the following paragraphs.

2.4. Simulations

The feasibility of simultaneously computing multiple anisotropic FODs corresponding to different models was evaluated numerically. \( H_{\text{WM}} \) was generated with the DKI model by setting the eigenvalues of the diffusion tensor as \((1.7, 0.3, 0.3) \times 10^{-3} \text{mm}^2/\text{s}\), and isotropic kurtosis equal to 0.6. \( H_{\text{GM}} \) was generated with the NODDI model by simulating values of intra to extra-cellular ratio 0.4, parallel diffusivity \( 1.7 \times 10^{-3} \text{mm}^2/\text{s} \), concentration parameter of the Watson distribution equal to 1.0, and no partial volume of free-water was considered as the mFOD approach explicitly accounts for it. The parameters used for generating \( H_{\text{WM}} \) and \( H_{\text{GM}} \) matched the average values observed with in-vivo data.

Two sets of simulations were generated using the 3-shells HCP acquisition protocol. Simulation I aimed at evaluating the robustness to noise of the mFOD approach, whereas simulation II characterized its ability to separate partial volume effects for a given SNR level.

2.4.1. Simulation I

To investigate the feasibility of mFOD, we generated three sets of partial volume effects of WM and GM, to simulate the transition from deep WM to GM. The first configuration (a) simulates the partial volume of 1 WM-like fiber and CSF, which in-vivo can be observed in the deep WM, for instance in proximity of the ventricles. In this simulation, \( f_{\text{WM}} \) was set equal to 0.8, and \( f_{\text{GM}} \) equal to 0.2. The second configuration (b) simulates the partial volume of 1 GM-like fiber and CSF with signal fractions \( f_{\text{GM}} = 0.8, f_{\text{CSF}} = 0.2 \), and is meant to reproduce the setting observed in-vivo in the outer cortex. The third configuration (c) simulates the partial volume between 2 WM-like fibers with crossing angle 45°, and 1 GM-like fiber crossing the first WM-like fiber with angles 30°- 60°- 90°, respectively, to mimic the transition of a WM fiber to GM through the superficial WM. For this configuration, the signal fractions \( f_{\text{WM}} \) and \( f_{\text{GM}} \) were both set equal to 0.5. The fourth configuration (d) simulates the crossing of a WM-like and a GM-like fiber, which simplifies what may be observed in-vivo after the superficial WM is crossed. In this case, the simulation was repeated for crossing angles in the range 10° - 90° with signal fractions \( f_{\text{WM}} \) and \( f_{\text{GM}} \) both equal to 0.5. The fifth configuration (e) is representative of a voxel containing mostly GM. For this configuration, we used the same settings of the previous simulation but adjusted the values of the signal fractions to \( f_{\text{WM}} = 0.2 \) and \( f_{\text{GM}} = 0.8 \). The effect of noise was simulated by adding 1000 Rician realizations in correspondence of SNR levels 10, 20, 30, 40, 50, 60, 70, and 150. For each SNR level, we evaluated the error between the estimated and the simulated signal fractions, as well as the angular error between the simulated and the estimated FOD directions.

Additional simulations were performed to understand the performance of mFOD as function of the employed signal model and of the acquisition protocol and can be found in the Supplementary Material.

2.4.2. Simulation II

Three partial volume configurations were generated in analogy with simulation 1 – configuration e with a crossing angle equal to 75°, but increasing the ratio between the two simulated tissues from 0 to 1 with step 0.1. Rician noise was added by considering 1000 realizations at SNR 50. The error between the simulated and the estimated signal fractions was determined.

2.5. In-vivo data processing

Two subjects from the pre-processed dataset of the Human Connectome Project (HCP) (McNab et al., 2013; Sotirropoulos et al., 2013) were randomly chosen and included in this study. The datasets included a T1-weighted image acquired at a resolution of 0.7 mm isotropic, and a dMRI dataset acquired at 1.25 mm isotropic. The diffusion datasets consisted of 18 b = 0 s/mm² volumes in addition to 270 volumes acquired by sampling diffusion weightings b = 1000, 2000, and 3000 s/mm² along 90 directions each. One of the two subjects was chosen to showcase the main results of the mFOD approach, using the native dMRI space (1.25 mm) as analysis space. Generalizability of the methods was then shown on the second subject, using the T1 resolution (0.7 mm) to highlight some features of the mFOD method. The T1-weighted data of the first subject was rigidly registered to the pre-processed dMRI data. Then, the tissue type segmentations of both datasets in WM, GM and CSF were derived with FSL FAST (Zhang et al., 2001). The SNR of the two datasets was determined by computing the average and the standard deviation of the signals at b = 0 s/mm² in two regions of interest manually placed in deep WM and in the parietal cortex, respectively. The average SNR at b = 0 s/mm² of the first dataset was 20 ± 4 in WM and 32 ± 9 in GM. The average SNR at b = 0 s/mm² of the second dataset was 20 ± 5 in WM and 27 ± 8 in GM.

The dMRI data was fit with both the “MultiShell Constrained Spherical Deconvolution” (MSCD) (Jeuring et al., 2014) (only data at 1.25 mm) and the mFOD approaches using custom implementations in MATLAB R2018b (The Mathworks Inc.) and functions from the ExploreDTI toolbox (Leemans et al., 2009). The code of mFOD is freely available online as part of the “MRIToolkit” toolbox for MATLAB (https://github.com/delucaal/MRIToolkit). MSCD was initialized as previously suggested independently on each dataset with spherical harmonics
of order 8 by using the tissue maps derived from the T1-weighted image. For the generation of the H-matrices used for the mFOD approach, a whole brain DKI and NODDI fit were performed with the dMRI data of the two HCP datasets at 1.25 mm resolution. For the WM H-matrix, the eigenvalues of the diffusion tensor and the isotropic kurtosis were averaged within a white matter mask defined by fractional anisotropy values above 0.7. This resulted in eigenvalues [1.3, 0.5, 0.5] \times 10^{-3} mm^2/s and isotropic kurtosis value 0.64. For the H-matrix of GM, the NODDI metrics were averaged within the GM segmentation obtained with the T1-weighted image, which resulted in intra to extra-cellular ratio 0.4, parallel diffusivity 1.7 \times 10^{-3} mm^2/s and Watson’s concentration parameter (\lambda) equal to 1. In this work, identical H-matrices were used for the mFOD analysis of both datasets as proof-of-concept. More generally, however, we recommend optimizing the signal models on a per-subject basis, especially when studying subjects with large lifespan or in patients. A graphical representation of the H-matrices (response functions) used for the analysis of in-vivo data is shown in Fig. 1 for the data shell at b = 3000 s/mm^2. The figure shows that the H-matrix corresponding to mFOD1 is pancake-like and appears similar to the response function commonly used in constrained spherical deconvolution. The NODDI-based H-matrix used for mFOD2 is also clearly anisotropic and assumes a zeppelin-like shape.

2.6. FODs and signal fractions

The MSCSD fit produced signal fraction maps associated to WM, GM, and CSF, and one FOD describing WM orientations. The fit of the mFOD approach resulted in the same number of fractional maps but two FODs, mFOD1 describing WM geometry, and mFOD2 describing GM geometry, respectively.

The signal fraction maps computed with MSCSD and mFOD were quantitatively compared by evaluating their Pearson correlations. To this end, f_{WM} and f_{GM} were evaluated, respectively, in WM and GM masks derived from the T1-weighted image. Next, the signal fraction maps from both mFOD and MSCSD were visually compared to those obtained from the T1-weighted volume. Further, the FODs computed with MSCSD and the mFOD framework were compared in selected areas of the cortex and of deep GM, to determine whether mFOD2 provided additional information in GM as compared to mFOD1 and MSCSD.

Fig. 1. A graphical representation of the H-matrices (response functions) used in mFOD in correspondence of data at b = 3000 s/mm^2. The white matter (WM) signal is represented with the diffusion kurtosis imaging (DKI) model and shows a highly anisotropic profile. The NODDI model is used to represent the anisotropic diffusion signal in gray matter (GM). The corresponding H-matrix results in an anisotropic – zeppelin-like – 3D profile.

The FODs computed with the mFOD method can be either analyzed separately, or merged to combine accurate geometrical information in both WM and GM. As a proof of concept, we combined the two FODs by considering their normalized sum weighted by the respective signal fractions (mFOD-W). One of the difficulties when validating in-vivo dMRI techniques is the lack of ground truth. MSCSD can be regarded as a gold standard in WM, but debatably not in GM. In a first analysis, we investigated to which extent the principal diffusion directions of the mFOD-W agree with those of MSCSD in WM, GM and at the WM/GM interface. A voxel was assigned to the latter class when its probability of belonging to both WM and GM was non-zero in the T1-weighted segmentation. Per voxel, up to three peaks were identified for both FODs, then sorted per descending amplitude values. Subsequently, the dyadic angle between each mFOD-W peak and the closest MSCSD peak was determined.

If discrepancies are observed in GM, it remains impossible to ultimately judge which method is the most accurate, but comparisons with a third method could provide additional interpretations. Anatomical knowledge as well as previous MRI studies (Setsompop et al., 2018) suggest that the dominant diffusion directions in GM should be perpendicular to the cortical folding, with exceptions such as the primary visual cortex (Kleinnijenhuis et al., 2013). We therefore derived the normal vectors to the cortical folding based on the structural tensor (Kleinnijenhuis et al., 2012). In short, a 3D volume was initialized with 1-values in correspondence of the GM/WM and GM/CSF interfaces, as derived from the T1-weighted segmentation. The linear interpolation field between the two surfaces was derived, and the corresponding vectors computed by computing the X-Y-Z first order derivatives of the volume. Subsequently, we determined the minimum dyadic angle between the normal vectors and any peak orientation determined with the MSCSD and mFOD-W results, respectively.

2.7. Fiber tractography

2.7.1. Tracking parameters

We investigated fiber tractography results obtained with MSCSD, mFOD1, mFOD2, and mFOD-W, with a particular focus on GM areas. Deterministic fiber tractography was performed in ExploreDTI by using identical parameters for all methods: FOD threshold 0.1 (default), minimum maximum fiber length equal to 20/500 mm, angle threshold 45°, and step-size 0.6 mm. Fiber tractography was also performed in an axonal region of interest overlapping the central sulcus in a single slice, and using FOD thresholds of 0.1 and 0.3, minimum fiber lengths of 10 and 50 mm, and step-size of 0.35 mm.

Firstly, FODs computed with the MSCSD and mFOD-W approaches were computed in a frontal region and color encoded with a custom colormap assigning red, yellow and blue to pathways traversing WM, GM and CSF, respectively. Subsequently, whole-brain tractography of all FODs was performed by seeding all voxels with an FOD value above the default threshold. The fiber bundles were color encoded both with the conventional RGB color scheme (Pajevic and Pierpaoli, 1999) and the above-mentioned tissue-type color scheme, to convey both directional information and specificity of the FODs to the different tissue types.

Fiber pathways obtained with MSCSD and mFOD in a 3.75 mm wide sagittal slab were color encoded by normalized distance between the GM/WM interface and the WM/CSF interface – as obtained from the T1-weighted segmentation – and also directly overlaid, to appreciate differences in GM tracking distance and density between the compared methods.

Subsequently, we evaluated the number of tract pathways generated with MSCSD and mFOD-W, which is informative given that the seed-points were kept constant and that a deterministic tracking algorithm was used. Additionally, we counted the fraction of plausible tract pathways, i.e., trajectories having at least one endpoint in gray matter, and the spatial coverage of tractography within GM when considering only start/endpoints of tracks. To qualitatively evaluate whether mFOD-W
could benefit fiber tractography of deep gray matter regions, we performed deterministic tractography of both mFOD-WS and MSCSD in an axial sub-region of the thalamus.

To qualitatively assess whether the mFOD framework can reconstruct previously reported fibers tangential to the cortex (McNab et al., 2013), we investigated the fiber tractography results with the mFOD-WS approach. Fiber tractography was seeded in a single axial slice, then compared to the cortical surface reconstructed with the CAT12 toolbox (Seiger et al., 2018) for MATLAB.

3. Results

3.1. Simulations

Figs. 2 and 3 show the signal fractions estimated with the mFOD approach in simulation I, where different mixtures of WM, GM and CSF simulate 5 configurations at multiple SNR Levels. Fig. 2 highlights the results of configurations I, II, and V with crossing-angle 90°, which are the main focus of this work. The figure shows that at SNR level equal or greater than 20, the mFOD method disentangles mixtures of WM and CSF (configuration a) with signal fraction errors below 2.2% and angular errors below 3°. The separation of GM from both CSF (configuration b) and WM (configuration e) is more challenging and requires higher SNR to achieve an error below 10%. In configuration II, the signal fraction errors at SNR 50 are about 9%, 11% and 2%, for WM, GM and CSF, respectively, and the angular error on the GM FOD is 8°. In configuration V, when WM and GM-like fibers cross with angle equal to 90°, the corresponding signal fraction errors at SNR 50 are 7%, 10% and 2%, and the angular errors on the WM and GM FODs are 4 and 11°, respectively.

Fig. 3 shows the results of simulation I for configurations c-d-e as function of the crossing angle between WM and GM for SNR levels. In general, the crossing angle between the WM-like and the GM-like fibers seems to marginally affect the signal fraction estimation. At SNR 50, the min-max signal fraction errors of WM, GM and CSF in configuration c are 7% (60°) – 7.1% (30°), 8.8% (45°) – 9.2% (30°) and 1.9% – 1.9%, respectively. The min-max signal fraction errors at SNR 50 for configuration IV are 8.2% (60°) – 8.9% (10°) for f_{WM}, 10.5% (90°) – 11.2% (10°) for f_{GM}, 2.2% (90°) – 2.4% (10°) for f_{CSF}. For configuration e, the minimum signal fraction error was observed with a 90° crossing configuration, 7.8% for f_{WM}, 10.0% for f_{GM} and 2.1% for f_{CSF}, whereas the worst error was observed in correspondence of the 10° crossing, 8.4% for f_{WM}, 10.6% for f_{CSF} and 2.2% for f_{CSF}.

When the SNR level is above 30, the angular error of the WM FOD is below 10° in configuration c and d independently from the crossing angle, and in configuration e for crossing angles of 45° or above. Achieving a good estimate of the GM FOD at the GM/WM interface is challenging. In configuration c and d, the angular error of the GM FOD is greater than 10° at SNR < 70, independently from the crossing angle with WM. More reliable estimates of the GM FOD can be achieved in configuration e: at SNR 50, an angular error of about 9.6° is observed for crossing angles of 45° or greater.

In Supplementary Figure S1, the simulation of configuration d with crossing angle 45° was repeated with different values of the NODDI parameter κ. The figure shows that at SNR 50 it is possible to estimate the GM FOD with angular errors equal to 8° and 5.6° in correspondence of κ = 2 and κ = 3, respectively, which are plausible settings in-vivo at the WM/GM interface. The simulation also shows that the angular resolution of the GM FOD improves with larger values of κ even if a value κ = 1 is used to generate the mFOD signals.

The effect of the acquisition protocol on the performance of mFOD is investigated in Supplementary Figure S2 and S3. Halving the number of gradient directions as compared to the HCP protocol worsens the effective SNR of the data, resulting in larger spread of the signal fraction estimates. The angular error of the WM FOD is minimally affected by the halving of the gradients, whereas slightly worse performances can be observed for the GM FOD. In Supplementary Figure S3, we investigated whether the distribution of the gradients on the outer shell and its diffusion weighting affects the performance of mFOD. Among the tested scenarios, only increasing the maximum diffusion weighting to b = 4000 s/mm² appreciably affects the performance of mFOD, suggesting a better estimation of the GM FOD (~21% angular error at SNR 50) and reduced uncertainty in the signal fractions estimates for larger maximum diffusion weights.

Fig. 4 shows the signal fractions estimated in simulation II when linearly varying the simulated WM, GM and CSF fractions at SNR 50. For...
Fig. 3. Results of simulation I. Five configurations of WM and GM partial volume simulating a) a WM-like fiber with CSF partial volume (20%), b) a GM-like fiber with CSF partial volume (20%), c) two crossing WM-like fibers and a GM-like fiber, d) a WM-like and a GM-like crossing fibers with equal volume fraction, and e) a WM-like and a GM-like crossing fibers with volume ratios 0.2/0.8, respectively. For each simulation, the top plot shows the 25th – 75th percentile (shaded area) and the median (solid line) of the signal fractions estimated with the mFOD approach for different SNR levels. The bottom plot shows the angular error of the main direction of the WM and GM FODs as compared to the ground truth value. The solid line represents the median value, whereas the error bars the 25th and 75th percentile. The simulations were performed using anisotropic response functions for WM and GM, and an isotropic response function for CSF. The green dotted line corresponds to an angular error equal to 20°. To properly detect the main peak orientation associated to GM, a high SNR (≥50) is required.
all the cases, the estimated average signal fractions show excellent agreement with the simulated values. The maximum errors observed for WM, GM and CSF are 13%, 20% and 4%, respectively. The Pearson correlation coefficient between the simulated and estimated signal fractions have values between 0.99 and 1.

3.2. HCP data

3.2.1. Signal fractions

The WM, GM and CSF signal fractions estimated with the MSCSD and mFOD methods, as well as those derived from the segmentation of the T1-weighted image are shown for an example axial slice in Fig. 5. The maps estimated with the mFOD method exhibit excellent agreement with both MSCSD and the T1-derived segmentation, highlighting the adequacy of the NODDI model and of the chosen parameters to represent the in-vivo signal in GM. The Pearson correlation coefficient between mFOD and MSCSD were 0.96 and 0.95 for $f_{WM}$ and $f_{GM}$, respectively. Notice that the WM fractional maps estimated with both the MSCSD and mFOD methods have non-zero values in some GM areas, in contrast to the T1-derived segmentation.

3.2.2. FODs

The FODs estimated with the MSCSD and mFOD methods in two cortical areas are shown in Figs. 6 and 7 for an example axial slice and an example coronal slice, respectively. The WM fiber orientations estimated with the mFOD method (mFOD$_1$) show remarkable similarity to the orientations estimated with MSCSD. Both FODs have very sharp and comparable orientations in the vicinity of WM and tend to become smaller towards deeper cortical GM. In contrast, the GM fiber orientations with the mFOD method (mFOD$_2$) show dominant and anatomically plausible peaks in most GM voxel, with orientations mostly perpendicular to that of the underlying WM. When mFOD$_1$ and mFOD$_2$ are linearly combined (mFOD-WS), a continuous vector field from WM to deep cortical GM is revealed.

Fig. 8 shows the FODs determined with mFOD-WS and MSCSD in an axial slice of the thalamus. The FODs derived with the two methods appear overall similar at visual inspection, but those determined with mFOD-WS appear sharper. A closer look (zoomed pane) reveals that the FODs computed with mFOD-WS have less angular uncertainty than those derived with MSCSD, and a better resolution of the peaks can be observed (white arrows). In the ROI shown in Fig. 8, the average
Fig. 5. An example axial slice of the signal fractions estimated with the MSCSD and mFOD methods in correspondence of different response functions as compared to the tissue segmentation obtained from the structural T1 image. Estimates with the MSCSD and mFOD approaches are remarkably similar and have high anatomical correspondence with the tissue types that they are designed to represent.

Fig. 6. Fiber orientation distributions (FODs) obtained with MSCSD and the mFOD method in an axial slice. The middle and the right columns show increasing zoom levels to highlight details of FODs. The mFOD1 and mFOD2 refer to the FODs computed when using response functions meant to represent WM and GM, respectively. The mFOD-WS is the sum of mFOD1 and mFOD2 weighted by their corresponding signal fractions. The mFOD2 is non-zero only in GM and its proximity, and shows dominant orientations also in the deeper cortex, in contrast to both MSCSD and mFOD1. The mFOD-WS combines the information content of mFOD1 in WM with that of mFOD2 in GM.

Fig. 7. Fiber orientation distributions (FODs) obtained with MSCSD and the mFOD method in a coronal slice. The middle and the right columns show increasing zoom levels to highlight how the FODs change in vicinity of GM. The mFOD1 and mFOD2 refer to the FODs computed when using response functions meant to represent WM and GM, respectively. The mFOD2 is non-zero only in proximity of GM, and shows dominant orientations also in the deeper cortex, in contrast to both MSCSD and mFOD1. The mFOD1 provides directional information in strong agreement with MSCSD but occasionally shows more peaks.

The number of peaks detected with MSCSD is 2.31 ± 0.88 as compared to 3.54 ± 1.03 with mFOD-WS. The average angle deviation between the main direction derived with mFOD-WS and MSCSD is 4.7 ± 4.5°.

Fig. 9 shows the histogram of the peak deviations observed between MSCSD and mFOD-WS in WM, GM and at the WM/interface, as well as the angle between the FODs and the normal to the cortical folding as derived from the T1-weighted image with a structural tensor approach. When comparing the first peak direction of MSCSD and mFOD-WS in WM, we observe that more than 70% of the WM voxels have an angular deviation smaller or equal than 10°. For the second and third peak we observed similar results, but also the presence of clusters of voxels with angle differences equal to 55° and 90° between the two methods. In GM and at the WM/GM interface, the angular deviation between the two methods is larger, up to 29°, and the peak of the distributions are centered at about 0.1° and 5°, respectively. When looking at the angle with the cortex normal, the histogram of both peaks is centered at 20°, but the mFOD-WS method results on average in a smaller angular deviation than MSCSD.

3.3. Fiber tractography

Fiber tractography results in a frontal lobe region computed with the MSCSD and mFOD-WS methods are shown in Fig. 10. The fiber pathways are color encoded according to the tissue type they are spatially traversing, and show that the mFOD-WS method generates more dense tractography results in GM. Further, the fiber bundles reconstructed with the mFOD-WS approach cover the GM more uniformly and traverse the tissue with a more plausible shape and to a greater extent.
Subsequently, we performed the whole-brain fiber tractography with the MSCSD and mFOD methods, both individually (mFOD₁, mFOD₂) and after the weighted sum procedure (mFOD-WS). The tracking with mFOD₁ produces a whole brain reconstruction that is largely comparable to that of MSCSD, suggesting preservation of the WM tracking performance. Conversely, the tracking of mFOD₂ results mostly in cortical reconstructions and short U-fibers, in-line with the GM-specific nature of this FOD. For pathways reconstructed with the mFOD-WS approach, 42.1% have their end points in GM, as compared to 34.1% when performing tractography with MSCSD. This is qualitatively visible in the tissue-type encoded reconstruction in Fig. 11, which shows that trajectories reconstructed with the mFOD-WS method cover the GM more extensively as compared to MSCSD. Note, however, that this also results in introducing extra spurious fibers.

An example sagittal slab of the trajectories reconstructed with MSCSD and the mFOD-WS approach was obtained by cropping the whole brain fiber tractography, as shown in Fig. 12. In the first column, the trajectories are color encoded according to their distance from the WM/GM interface (blue) to the GM/CSF interface (red). It can be seen that the fiber bundles reconstructed with the mFOD-WS method consistently travel a longer distance within the cortical GM than with MSCSD, in-line with observations in Fig. 10. Further, the frontal, parietal and occipital side of the cortex appear to be better reconstructed with the mFOD-WS method.

While the mFOD framework has been primarily developed to target the cortical GM, deep GM regions as the thalamus have a central role in brain connectivity. In Fig. 13, we investigated the application of fiber tractography with mFOD-WS and MSCSD using a seed region covering part of the thalamus in one single axial slice. The fiber tractography obtained with both methods reveal the existence of thalamic – temporal and thalamic-sensory motor connections. With mFOD-WS, however, we also observe a remarkably larger number of streamlines reaching the brainstem, and the occipital cortex. Additionally, mFOD-WS results in longer reconstructions of the fiber tracts from the thalamus to both the sensory-motor and the frontal/pre-frontal cortex, in agreement with previous population studies (Cox et al., 2016).

Fiber tractography of the mFOD-WS method using the HCP data is shown in Fig. 14 for different choices of tracking criteria. Good coverage of the cortical structure is observed for all settings, even when enforcing a minimum length of 50 mm, ensuring that most of the reconstructions originating in GM can reach the WM. When using an FOD threshold equal to 0.1, more spurious reconstructions are revealed as compared to a more restrictive threshold (0.3), but a denser tract reconstruction in the cortical ribbon and more extensive coverage of the contralateral hemisphere can be observed. Allowing for trajectories as short as 10 mm, results in a very dense reconstruction of the cortical gyri, showing short bundles that seem to travel tangentially to GM, as well as many short radial reconstructions.

4. Discussion

In this work, we have introduced the mFOD framework to perform spherical deconvolution of multiple anisotropic response functions. With simulations and HCP data, we have shown that it is possible to reconstruct two anisotropic FODs, specific to WM and GM, while also accounting for the CSF partial volume effect. Further, we have shown that the mFOD approach improves the reconstruction of FODs in GM as compared to existing state-of-the-art methods, and that this results in improved fiber tractography reconstruction in regions near or in the GM.

Fig. 8. Fiber orientation distributions (FODs) obtained with the mFOD method and MSCSD in an axial slice of the thalamus. The right column zooms in a 3 × 3 neighborhood to better compare the visualized FODs. In general, the FODs in mFOD-WS appear sharper in the whole thalamus as compared to MSCSD. When looking into the zoomed region, the FODs in mFOD-WS appear sharper and appear to resolve complex configurations better than MSCSD (white arrows).

Fig. 9. The first three histograms show the normalized frequency (probability) of the angle between each mFOD-WS peak and the closest MSCSD peak in WM, GM and at the WM/GM interface, respectively. The right histogram shows the probability of the angle difference between the normal direction to the GM/WM surface and the main direction with MSCSD or mFOD-WS. With regards to the left plots, the first peak of two methods has very small angle differences in WM, within 10° in over 70% of voxels, whereas larger differences are observed in GM and at the WM/GM border. Larger differences were observed for the second and the third peak, with a subset of the voxels showing an angle of 60° between the two methods. mFOD-WS resulted in lower angle deviation with the surface normal (derived with the structural tensor technique) as compared to MSCSD.
4.1. Feasibility of mFOD

The reconstruction of multiple anisotropic FODs is a complicated, numerical challenge that requires the inversion of a matrix whose rank (number of independent rows/columns) is much lower than the number of possible solutions (FOD amplitudes on the unit sphere), a recurring challenge in deconvolution problems. Previous studies, as Tournier et al. (2007) and Jeurissen et al. (2014) used an L2 regularization term enforcing continuity (smoothness) in the FOD amplitudes. In this work, we used a non-negative least-squares solver with L2 regularization encapsulated in an iterative solver (Guo et al., 2020). Taking into consideration that the mFOD formulation (Eq. (2)) hosts multiple FODs, this might be suboptimal and introduce discontinuities in the solution. Nevertheless, our results suggest that the mFOD approach does not propagate smoothness between multiple FODs, likely due to the attenuation of small perturbations imposed by the median operator (Guo et al., 2018). Indeed, the results of simulations, shown in Figs. 2, 3 and 4, suggest that this approach allows to effectively disentangle the partial volume effects between mixtures of isotropic \( (n = 1) \) and anisotropic \( (n = 2) \) signal mixtures, if sufficient SNR is provided. In particular, the partial volume effects between WM and CSF are the easiest to disentangle, and an SNR equal or greater than 20 is sufficient to obtain accurate separation of signal fractions and minimal angular error on the WM FODs. In contrast, the separation of GM from both WM and CSF is a harder task, probably as a result of the limited anisotropy of this tissue type. In Fig. 4, we observe that the GM estimation shows a small but consistent offset from the simulated values. This is very likely caused by the above-mentioned median operator in the deconvolution framework (GRL) on which mFOD is built. Such operator is designed to be used in combination with deconvolution matrices containing highly anisotropic signal basis. When spherical deconvolution is performed with a deconvolution matrix including moderately anisotropic elements (e.g. generated with the NODDI model with \( \kappa = 1 \)), this is likely to produce some genuine small amplitudes which are discarded by the median filtering, causing a small but consistent bias in the \( f_{GM} \) estimate. Of note, in our simulations we did not compare the performance of mFOD to MSCSD, because such method is data-driven, e.g. it estimates its deconvolution matrix from the data, which is not feasible in a single-voxel simulation. Furthermore, we believe that such comparison would not be completely fair, as by design the MSCSD method lacks the ability to fit the anisotropic GM-like signal that we introduce in the simulated signals.

If an SNR equal or above 50 is achieved, the mFOD framework can preserve the ratios between different tissue types and can estimate their mixtures with minimal biases (Figs. 2 and 3). While an SNR of about 50 might be difficult to achieve in clinical settings, accurate pre-processing and denoising (Veraart et al., 2016), or even deep-learning based reconstructions (Ropers et al., 2017) might mitigate this need. According to Fig. 3, the crossing angle between a WM bundle entering into the cortex and the cortical folding itself is likely to have an impact on the performance of mFOD. Generally, white matter bundles travel to their cortical endpoint perpendicularly (Budde and Annese, 2013) to the cortical folding, with exceptions such as the primary visual cortex (Kleinnijenhuis et al., 2013) or the U-fibers adjacent to the cortex (Schilling et al., 2018). If the direction of the axons leaving the cortex (to the WM) would not match that of intra-cortical axons, the results of the simulations suggest that the GM FOD would be estimated with an angular error between 15 (SNR 30) and 10° (SNR 50), which are likely to perturbate the tracking trajectory. In practice, very high SNR is likely required to reconstruct fiber pathways tangential to the cortical folding, but less critical to detect radial trajectories. In light of the insights provided by simulation I, we conclude that mFOD can estimate the GM FOD with reasonable angular error in the cortex itself (configuration V in Fig. 3) and in the case of limited partial volume with a single WM fiber (configuration e in Fig. 3). The performance of mFOD at separ-
Fig. 11. Whole-brain tracking with MSCSD and the mFOD methods. The first row shows the fiber pathways color encoded with the conventional color scheme, whereas the second row shows the same tracking color encoded by traversed tissue type (subsampling with factor 8). MSCSD and mFOD$_1$ provide similar tracking results with the latter showing better characterization of the cerebellar cortex, as shown by the more homogeneous red coloring in the tissue type encoded tractography. The tracking with mFOD$_2$ resulted exclusively in cortical and short (U-fibers) connections. With mFOD-WS, a more complete coverage of the GM is achieved as compared to both mFOD$_1$ and MSCSD.

Fig. 12. A sagittal slab (red line in the bottom left picture) of the fiber tractography results obtained with MSCSD and the mFOD-WS method. The first column shows the fiber pathways color encoded according to their normalized distance from the GM/WM and the GM/CSF interfaces. The distance was determined in 3D as flow (linear interpolation field) between the two surfaces using the same method we previously used to determine the cortex normal. The central and right column show the same sagittal slab color encoded by reconstruction method, and its zoom in a cortical region. The WM-GM flow color encoding shows that a deeper cortical tracking was achieved with the mFOD-WS method as compared to MSCSD, especially in the frontal and occipital areas, as well as a more contiguous coverage of the cortex (white arrows). When looking at the tissue type encoded images, mFOD-WS resulted in a more extensive and contiguous reconstruction of the cortical folding, and the tracts traveled deeper into the cortical GM, although a larger number of spurious reconstructions can also be observed.

mFOD-WS is also confirmed by the in-vivo results, as the signal fraction map associated to GM is zero-valued in all non-GM voxels and has values of 1 in almost all GM voxels, which suggests excellent separation of the two tissue types. In proximity of the WM/GM interface, however, the possible presence of an additional crossing with the superficial U-fibers is likely to generate unreliable estimates (configuration c of Fig. 3), suggesting that further research is needed to address the challenge of crossing the superficial WM (Reveley et al., 2015). On the other hand, it should be noted that the above result corresponds to a GM-like signal generated with the NODDI parameter with $\kappa = 1$, which results in low signal anisotropy. While such value seems appropriate in the cortex, i.e., when $f_{GM} = 1$, at the WM/GM interface, higher values of $\kappa$ are typically observed. According to Supplementary Figure S2, with a higher value of $\kappa$, the angular error of the GM FOD drops drastically.
Fiber tractography of MSCSD and of the mFOD-WS method with a seed ROI located the Thalamus in one single axial slice. The streamlines reconstructed with mFOD-WS reached a larger extent of the cortex, especially in the occipital and frontal cortex. While the connection between the Thalamus and the sensory motor cortex was observed with both MSCSD and mFOD-WS, streamlines reconstructed with the latter branched more plausibly into the cortex (red tracts). When looking at the thalamic-frontal projection (yellow tracts), the fiber reconstructions with mFOD-WS covered a remarkably larger extent of the frontal cortex as compared to those obtained with MSCSD.

4.2. GM modeling

While recent work has introduced the reconstruction of multiple WM FODs (Pietsch et al., 2019) to account for inadequacies in WM modeling in infants, the mFOD framework has the unique characteristic of reconstructing multiple FODs that are specific of remarkably different tissue types as WM and GM, as shown in Figs. 6 and 7. The figure shows that WM fiber orientations estimated with mFOD (mFOD$_1$) and MSCSD are very similar in WM, and that both exhibit very small values in GM. In contrast, the GM fiber orientations estimated with mFOD$_2$ show consistent and well-defined diffusion directions for a large extent of the cortical GM. The performance improvement over MSCSD of the mFOD framework in GM is explained by considering the improved tissue modeling. Approaches as MSCSD essentially describe the WM and GM mixture as the sum of an anisotropic and an isotropic signal, similarly to the ball-and-stick model (Behrens et al., 2003), where the anisotropic signal is modelled with properties extracted from single fiber population WM (Tournier et al., 2007; Tax et al., 2014; Dell’Acqua et al., 2007). Therefore, in practice, by applying any constrained spherical deconvolution method in GM, one assumes the axonal component of GM to produce a similar signal to that of a WM bundle without crossing fibers. However, assuming the axons and dendrites of the cortex to be identical to the axons in the corpus callosum or other major WM bundles seems a stretch given their different diffusion signatures (Westin et al., 2016). In this work, we represent the GM signal with a special case of the NODDI model (Zhang et al., 2012), which represents a first approach to take into account additional effects as tortuosity (Sen and Basser, 2005) and intra-to-extra neurite signal ratios into the deconvolution problem.

4.3. How to use multiple FODs

One of the open challenges in the mFOD framework regards how to combine the multiple FODs computed by the method. In this work, we adopted a pragmatic approach and considered their weighted sum (mFOD-WS). This choice produces convincing and contiguous FODs, as shown in Figs. 6 and 7. However, this might be sub-optimal for fiber tracking approaches, as it might blur the peaks detected in individual FODs. Future work is needed to establish whether dynamically tracking the locally larger FOD, or dynamically choosing the FODs most aligned to the tracking direction might represent better alternatives. Additionally, sparsity constraints as L1 or L0 norm (Daducci et al., 2014) could be taken into consideration to improve the FOD separation and minimize overfit. Nevertheless, when we compared the main peaks of mFOD-WS to those obtained with MSCSD (Fig. 9), we observed remarkable similarity between the two in WM. Further, the average angular error between the two techniques was below 10° in over 70% of WM voxels, suggesting that the mFOD-WS approach does not affect the performance of spherical deconvolution in WM. In GM, the mFOD-WS approach resulted in smaller angular deviations with respect to the cortex normal as compared to MSCSD. While the cortex normal might not always be representative of the direction with which axons are organized in the cortex, they represent an accepted assumption (Tozer et al., 2012), and might serve as a comparison term between the two methods. The average angular deviation between the cortex normal and mFOD-WS is 20°, which is in-line with the angular error predicted in case of WM and GM mixtures at SNR 50 (Fig. 3).

4.4. Improved fiber tractography in GM

Fiber tractography is one of the main applications of spherical deconvolution techniques. Here, we used a deterministic approach to facilitate the interpretation of pathway reconstructions. As shown in Fig. 10, we obtained a higher percentage of tract pathways reaching a GM region with the mFOD-WS approach than with MSCSD given identical tracking parameters. Furthermore, the fiber pathways reconstructed with the mFOD-WS method covered the cortical structure more extensively, and their geometrical configuration appeared more similar to previous reports of high resolution cortical dMRI (Setsompop et al., 2018). While a similar tractography result might also be achievable with MSCSD by tuning the FOD threshold (Aerts et al., 2019), this is likely to introduce not only false positives in WM, but also to worsen the tracking in GM itself due to a larger number of spurious peaks (Parker et al., 2013). One important application of fiber tractography is the study of deep GM structures, due to their central role as hubs for many brain connections. In Fig. 8, we investigated the performance of mFOD-WS in the thalamus. We observe that the use of mFOD-WS seems also beneficial in deep GM, as it provides FOD estimates that are sharper and better resolved than those provided by MSCSD, which is desirable in fiber tractography applications. Furthermore, in the thalamus, we found a small but consistent angular bias of about 5° between the main directions estimated with the two methods, and a higher number of detected peaks with mFOD-WS, 3.5 vs 2.3. When performing fiber tractography from the thalamus (Fig. 13), the fiber tracts reconstructed with mFOD-WS covered a larger extent of the brain as compared to the fiber tracts reconstructed with MSCSD, including previously reported connections to the occipital, sensory-motor and frontal cortices, although a larger number of spurious fibers was also observed. Furthermore, the reconstruction of the thalamic-frontal tract with mFOD-WS appears more in line with previous reports.

When performing whole-brain tractography, the use of the mFOD-WS approach resulted in more tract pathways having a start or endpoint in the GM (+8%), and generally in more extensive GM coverage, as shown in Fig. 11. This was even more apparent when looking at a sagittal slab (Fig. 12), which showed how fiber bundles could be followed in GM up to the GM/CSF interface, at the price of potentially more spurious reconstructions as compared to MSCSD. In practice, mFOD could be beneficial to applications such as pre-surgical planning (Mormina et al., 2016) or structural connectivity studies (Beare et al., 2017), where propagating fiber pathways from the deep WM into the cortex is crucial. Ex-
isting fiber tractography methods do not generally define the seed points in the cortex, but rather at the GM/WM interface or in the adjacent WM, potentially introducing a number of false positives (Maier-Hein et al., 2017). For similar reasons, connectivity studies are often performed by first seeding in WM or at the WM/GM interface, then filtering the trajectories with anatomical constraints (Smith et al., 2012), and finally assigning the fiber bundles to the closest GM node after spatial dilation, potentially introducing uncertainty in the assignment (Wei et al., 2017).

To overcome these limitations and reconstruct complete fiber pathways to/from the cortical start/endpoints, 4 major steps are required: i) the estimation of reliable FODs in WM, ii) crossing the superficial WM at the WM/GM interface, iii) overcoming the gyral bias in fiber tractography and iv) the estimation of reliable FODs in GM. The mFOD framework is a first step towards the solution of this comprehensive challenge, as it allows to robustly determine the FODs in cortical areas. This problem has also been investigated in previous studies and has been addressed with computational approaches based on the geometry of the cortex itself. While such methods have the advantage of not requiring elaborate diffusion acquisition schemes as mFOD, they simplify the structure of the cortex, enforcing fiber tractography to proceed radially to the cortex, which does not take into account the potential presence of tangential fibers or the adverse effects related to disease or abnormal development of the cortex organization. Taking into account the results shown in Fig. 14, it seems possible with mFOD to also effectively reconstruct some crossing bundles in the GM itself, where radial projections cross previously reported tangential fibers (Kleinnijenhuis et al., 2013; Leuze et al., 2014; J. A McNab et al., 2013). Fig. 14 also highlights the importance of the FOD threshold parameter in fiber tractography. When the FOD threshold is lowered, a larger number of fiber tracts is reconstructed. These fiber reconstructions are also generally longer and can become erratic when the FOD is not well-defined, such as in the outer cortex where the partial volume with CSF dominates the signal. Increasing the value of the FOD threshold reduces the number of spurious fibers, but likely also the number of genuine pathways that are discarded (Maier-Hein et al., 2017). The accurate estimation of FODs at the WM/GM interface is more challenging than for GM itself according to the results of simulation I (Fig. 3). In our in-vivo experiments with mFOD, fiber tractography generally reached the cortex, but this might be more challenging in datasets with lower SNR. In that case, the mFOD approach might be further enhanced by taking into account geometric frameworks such as the “Surface Enhanced Tractography” (St-Onge et al., 2018) or the “Structure Tensor Informed Fiber Tractography” (Kleinnijenhuis et al., 2012), or with hybrid acquisition strategies aimed to better resolve the FODs at the WM/GM interface (Fan et al., 2017). In this manuscript, we have employed a deterministic tractography method to focus on the FOD estimation step. However, mFOD could also be used in combination with probabilistic fiber tractography (Tournier et al., 2010), multi-peak tractography (Descoteaux et al., 2009) or even global tractography (Christiaens et al., 2015), as shown in Supplementary Material Figure S4.

4.5. Methodological considerations

The models used here to describe WM and GM properties were chosen from current popular methods, but their combination might not be the optimal representations to estimate multiple FODs. The DKI model, here used in its simplified version with isotropic kurtosis (De Luca et al., 2017), efficiently represents non-Gaussian signals with a limited number of parameters but is sub-optimal to account for direction dependent signal restrictions. Alternatively, geometric representations of the WM signal (Ferizi et al., 2014), as well as data driven methods (Tax et al., 2014) could also be explored. Furthermore, in this work we focused on the modeling of the cortical GM, but mFOD can theoretically be used also with multiple WM response functions to account, for instance, for differently myelinated WM in infants (Pietsch et al., 2019), or for other anisotropic tissues such as lesioned WM (Ban et al., 2019) or malignancies (Aerts et al., 2019). Note that the diffusion approaches suggested for WM (DKI) and GM (NODDI) could be further adjusted or even replaced by other more specific tissue models to further improve the mFOD framework. In name of clarity, we fixed the parameters of the

![Fiber tractography of mFOD-WS from an axial seed region](image_url)
NODDI model to the average values observed in GM, but these values could be suboptimal at voxel level. In Supplementary Figure S1, we have briefly investigated the impact of mis-estimating the NODDI parameters in mFOD, observing that overestimating the value of $\kappa$ in mFOD results in worse angular errors in the GM FOD but minimal or no effects on the signal fraction estimates. Conversely, underestimating the value of $\kappa$ has marginal effects on the angular error, but biases the signal fraction estimates.

The dMRI acquisition protocol is expected to have a major influence on the performance of the mFOD framework. Here, however, we aimed to show a proof-of-concept of the feasibility of the mFOD approach, and therefore relied on the high-quality HCP data. Nevertheless, in the Supplementary Material we briefly investigated with simulations the impact on the performance of mFOD of specific changes to the acquisition protocol. Supplementary Figure S3 shows that halving the acquisition protocol with respect to the HCP gradient schemes would minimally affect the estimation of the WM FOD and of the signal fractions, but would worsen the precision of the estimates, as expected. The reduced acquisition protocol negatively impacts also the angular error of the GM FOD, increasing the error of about 5° for the 90° WM-GM crossing simulation. In Supplementary Figure S2, we investigated how different distributions of HCP-like multi-shell diffusion weightings would impact the mFOD performance. We observe that acquiring more directions in the outer shell or reducing its diffusion weighting to $b = 2500\text{s/mm}^2$ does not improve nor worsen the angular error of the GM FODs. Conversely, increasing the maximum diffusion weighting to $b = 4000\text{s/mm}^2$ seems to improve the estimation of the GM FOD, in line with recent reports using ultra-high diffusion weightings (Palombo et al., 2020; Tax et al., 2020).

In Fig. 9, we used the cortex normal direction as a comparison for FODs computed with both MSCD and mFOD. While it is reasonable to assume that most axonal projections will traverse the cortex with a perpendicular trajectory, this is not always the case. Furthermore, the method employed to compute the normal trajectory is not accurate when the sulci are separated by less than 2 voxels and should therefore be interpreted cautiously. The performance of the mFOD framework has been shown for two subjects and, although promising, should be further evaluated on larger cohorts and with more clinically achievable protocols. Finally, while we have shown that the mFOD framework improves the performance of spherical deconvolution and fiber tractography in GM, it remains to be shown to which extent such improvements will benefit clinical studies and structural connectivity investigations.

In conclusion, we have presented the mFOD framework to perform tissue-specific spherical deconvolution which can simultaneously reconstruct multiple distributions of fiber orientations per voxel. By integrating specific WM and GM models, we have shown that the mFOD approach can improve the performance of spherical deconvolution in GM while retaining the performance of state-of-the-art methods designed for WM.

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**Supplementary materials**

Supplementary material associated with this article can be found, in the online version, at doi:10.1016/j.neuroimage.2020.117206.

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