Sparse Solution of Fiber Orientation Distribution Function by Diffusion Decomposition

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Abstract

Fiber orientation is the key information in diffusion tractography. Several deconvolution methods have been proposed to obtain fiber orientations by estimating a fiber orientation distribution function (ODF). However, the L_2 regularization used in deconvolution often leads to false fibers that compromise the specificity of the results. To address this problem, we propose a method called diffusion decomposition, which obtains a sparse solution of fiber ODF by decomposing the diffusion ODF obtained from q-ball imaging (QBI), diffusion spectrum imaging (DSI), or generalized q-sampling imaging (GQI). A simulation study, a phantom study, and an in-vivo study were conducted to examine the performance of diffusion decomposition. The simulation study showed that diffusion decomposition was more accurate than both constrained spherical deconvolution and ball-and-sticks model. The phantom study showed that the angular error of diffusion decomposition was significantly lower than that of constrained spherical deconvolution at 30° crossing and ball-and-sticks model at 60° crossing. The in-vivo study showed that diffusion decomposition can be applied to QBI, DSI, or GQI, and the resolved fiber orientations were consistent regardless of the diffusion sampling schemes and diffusion reconstruction methods. The performance of diffusion decomposition was further demonstrated by resolving crossing fibers on a 30-direction QBI dataset and a 40-direction DSI dataset. In conclusion, diffusion decomposition can improve angular resolution and resolve crossing fibers in datasets with low SNR and substantially reduced number of diffusion encoding directions. These advantages may be valuable for human connectome studies and clinical research.

Introduction

Crossing fiber problem is still under active research in the field of diffusion MRI, and a method that offers accurate fiber orientation is the cornerstone of human connectome studies since it can facilitate fiber tracking and provide better mapping of neuronal connections [1,2]. To resolve crossing fibers, methods that make use of the orientation distribution function (ODF) have been widely used [3]. Tuch [4] proposed q-ball imaging (QBI) to estimate the diffusion ODF (dODF) by applying Funk-Radon transform to high angular resolution diffusion images [HARDI] [5]. Wedeen et al. [6] proposed diffusion spectrum imaging (DSI), which obtains ODF by applying inverse Fourier transform to diffusion MR signals and calculating the second moment along the radial distribution of the transformed signals. To further extend the applicability, Yeh et al. [7] proposed generalized q-sampling imaging (GQI), which can be applied to a variety of diffusion sampling schemes to obtain dODFs, and the results are consistent with those from QBI and DSI. Although these dODF methods have been used to determine fiber orientations, their accuracy is limited by the blurred contour of dODF. This problem is demonstrated in Fig. 1, an example of two fiber populations crossed at right angles (Fig. 1A). The fiber orientations can be resolved by the peak orientations of the dODF (Fig. 1B), but the blurred contour of the dODF may fail to resolve crossing fibers if the crossing angle is sufficiently small.

To better characterize fiber distribution, studies have used spherical deconvolution to obtain fiber ODF (fODF) and to measure the orientation distribution of fiber volume fractions [8,9]. Spherical deconvolution estimated the signal pattern of a single fiber bundle and applied deconvolution to the spherical harmonics of diffusion MR signals to calculate IODF. An example of IODF is shown in Fig. 1C, where IODF presents a contour sharper than dODF’s and achieves better angular resolution. This advantage makes IODF a useful way to resolve crossing fibers for diffusion MRI fiber tracking. Further studies have proposed improved methods to address the problem of partial volume effect [10,11], which can corrupt IODFs and produce false peaks. Moreover, different computational approaches have been proposed to sharpen IODFs, enforce non-negativity, or achieve more robust IODFs estimation [12,13,14,15,16,17]. Although IODF has been shown to be sensitive to crossing fibers, its specificity has been called into question [11,18]. IODF often has baseline fluctuation that may give rise to false peaks, as illustrated in the zoom-in figure of Fig. 1C. This fluctuation problem is often handled by applying an ODF filter, but the filter can indiscriminately eliminate less salient fibers and decrease the sensitivity of IODF. Besides the baseline fluctuation problem, L_2 regularization also introduces...
Diffusion Decomposition

Figure 1. An example of two crossing fibers at right angles. (A) The layout of the crossing fibers. (B) The corresponding diffusion ODF, showing the orientation distribution of the water diffusion due to the right angle crossing environment. (C) The corresponding fiber ODF obtained from diffusion deconvolution has non-zero values in most of the orientations, as shown in the blue colored fluctuation around the origin. This baseline fluctuation often causes false identification of crossing fibers. (D) The actual fiber ODF has zero values at most of the orientations and non-zero values only at the fiber orientations, suggesting that the fiber ODF is sparse.

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Materials and Methods

Diffusion model

We modeled a dODF using the mixed diffusion model [11]: a dODF can be viewed as a linear combination of multiple dODF components, meaning that the diffusion signal of a dODF is the summation of the diffusion signals from each constituted fiber components. Specifically, the overall dODF $\Psi_d$ is composed of a background isotropic component $f_0 e$ and a series of component dODFs, $\psi_1, \psi_2, ..., \psi_n$, each of which represents a fiber population oriented at a unique direction.

$$\Psi_d = f_0 e + f_1 \psi_1 + f_2 \psi_2 + ... + f_n \psi_n$$  (1)

where $f_0$ is the volume fraction of the background isotropic diffusion, and $e$ is an all-one vector modeling the isotropic diffusion. The coefficients $f_1, f_2, ..., f_n$ are the fiber volume fractions for each of the component dODFs. In our implementation, we further scaled the sum of the volume fractions (i.e. $\sum_{i=0}^{n} f_i$) to the spin density because the total volume is in fact proportional to the amount of the spins. The same scaling was also used in diffusion deconvolution to avoid the partial volume effect from the gray matter [11]. Furthermore, each dODF in Eq. (1) was represented by a vector with finite dimensions. Specifically, the sampling orientations of a dODF were defined by the vertices of an 8-fold tessellated icosahedron, which has a total number of 642 orientations. Since the dODF is assumed to be symmetric about the origin, a dODF can be represented by a 321-dimension vector, resulting in an angular resolution of around $8^\circ$. These 321 unique orientations were also used to assign the fiber orientations of the component dODFs, resulting in 321 component dODFs ($n = 321$). Using this setting, the orientation distribution of the volume fractions, $f_1, f_2, ..., f_n$, constitute the IODF that we aimed to estimate.

To solve Eq. (1), we first estimated the dODF, $\Psi_d$, using standard procedures of QBI, DSI, or GQI, and modeled the component dODFs with a putative common characteristic dODF that described the diffusion profile of a single fiber population. We selected the characteristic dODF from the voxel having the highest anisotropy, a strategy similar to deconvolution methods [8,26]. To calculate each component dODF from the characteristic dODF, we smoothed the characteristic dODF as described in our previous work [11]. The following is the formula for estimating the $i$-th component dODF:

$$\Psi_i(v) = z \sum_{j} \psi_j(u) \exp \left( - \frac{\{ \cos^{-1} \langle \tilde{v}, \tilde{u}_i \rangle \} - \cos^{-1} \langle \tilde{v}, \tilde{u}_j \rangle \}^2}{2\sigma^2} \right)$$  (2)

where $\psi_i$ is the $i$-th component dODF, and $\psi_j$ is the common characteristic dODF. $\tilde{v}$ and $\tilde{u}_i$ are the fiber orientations of $\psi_i$ and $\psi_j$, respectively. The operator $\langle \cdot, \cdot \rangle$ calculates the inner product of two vectors, $\sigma$ is the standard deviation for the Gaussian radial basis kernel, and $z$ is a normalizing factor to ensure that the component dODF integrates to one. The summation iterates through all sampling directions, $\tilde{u}_i$, and $z = 9^\circ$ was used as that in our previous work.

LASSO estimator

Without loss of generality, the regression equation in Eq. (1) can be reformulated by normalizing the vector of each component dODF in a way that the mean is zero, and the length of the vector is one.

$$\Psi_d = f_0 \psi_1 + f_1 \psi_2 + ... + f_n \psi_n$$

$$\sum_{i} \Psi_i(u) = 0, \sum_{i} \Psi_i^2(u) = 1$$  (3)
Table 1. Diffusion Decomposition Algorithm.

| Input | Comments |
|-------|----------|
| $\Psi_d$ | dODF |
| $\Psi_1, \Psi_2, \ldots, \Psi_n$ | Component dODF |
| $\Psi_d$ | Normalized dODF* |
| $\Psi_1, \Psi_2, \ldots, \Psi_n$ | Normalized component dODF |
| $\epsilon$ | Decomposition fraction |

| Output | Comments |
|-------|----------|
| $\nu$ | Volume fraction of the isotropic background diffusion |
| $\nu_1, \nu_2, \ldots, \nu_n$ | Volume fractions of the component dODFs |

| Algorithm | Comments |
|----------|----------|
| 1. while $\max_i \langle \Psi_i, \Psi_i \rangle > 0$ and $|A| < m$ | Repeat 1.1, 1.2, and 1.3 as long as the condition holds. |
| 1.1. $k = \arg \max_i \langle \Psi_i, \Psi_i \rangle$ | Selection: Select the most correlated component dODF |
| 1.2. $\Psi_d = \Psi_d - \epsilon \langle \Psi_d, \Psi_k \rangle \Psi_k$ | Decomposition: Decompose a small volume from the dODF |
| 1.3. $\lambda = \lambda_k$ | Record the selected component dODFs |
| 2. $\Psi_d = \Psi_d + \sum_i \nu_i \Psi_i$ | Solve volume fractions by ordinary least squares |
| 3. if $\min_i \langle \Psi_i, \Psi_i \rangle < 0$ then $A \leftarrow A \setminus \arg \min_i \langle \Psi_i, \Psi_i \rangle$ and go to 2 | If any of the volume fractions is negative, set the most negative one to zero and remove it from set $A$ and solve the regression again. |
| 4. if $\nu_i < 0$ then $\nu_i \leftarrow 0$ and go to 2 | If the volume fraction of background diffusion is negative, force it to zero and solve the regression again. |

*The normalized dODF is calculated by $\Psi_i \leftarrow (\Psi_i - \bar{\Psi_i}) / \| \Psi_i - \bar{\Psi_i} \|$. |

One should note that the isotropic component is removed since the mean is shifted to zero, and the coefficients are rescaled since the corresponding component dODFs are normalized. The sparse solution for the coefficients (i.e. the scaled volume fractions) can be estimated by the LASSO estimator, which uses $L_1$ regularization to promote sparsity.

$$
\min \left\| \Psi_d - \sum_{i=1}^n \nu_i \Psi_i \right\| \text{s.t. } \sum_{i=1}^n |\nu_i| \leq s
$$

The regularization is controlled by $s$, and it has been shown that the solutions for different levels of $s$ can be obtained by iteratively including different regressors (component dODFs) in the solution set [10,22,23]. This leads to our diffusion decomposition algorithm, which is based on the forward stagewise method [23] and further considers the nonnegative constraint of fODF.

**Diffusion decomposition**

Table 1 shows the diffusion decomposition algorithm, which includes the following steps (the C++ source codes of diffusion decomposition are available to the public at http://dsi-studio.labsolver.org). First of all, selection, the algorithm finds the component dODF that is most correlated with the dODF.

$$
k = \arg \max_i \langle \Psi_i, \Psi_i \rangle
$$

where $\langle \Psi_i, \Psi_i \rangle$ is the inner product of the normalized dODF and a normalized component dODF. One should note that in Eq. (5) diffusion decomposition considers only the positive correlation due to the nonnegative constraint of the solution. The next step is decomposition, which removes a small volume of $\Psi_k$ from $\Psi_d$.

$$
\Psi_d \leftarrow \Psi_d - \epsilon \langle \Psi_d, \Psi_k \rangle \Psi_k
$$

where $\epsilon$ is the decomposition fraction. These two steps, selection and decomposition, are recursively iterated until the maximum correlation is less than or equal to zero, or a total of $m$ unique component dODFs are selected. The selected component dODFs in the selection-decomposition recursion are recorded in a component set, $A$, whereas those not selected have zero fiber volume fractions.

Table 2. Summary of simulation and phantom studies.

| b-value (s/mm$^2$) | directions | SNR | crossing angles | simulation/experiment settings |
|---------------------|-------------|-----|----------------|-----------------------------|
| Simulation 1        | 1500        | 160 | 6.8            | 18° to 90°* Two tensors and an isotropic component |
| Simulation 2        | 1500        | 55  | 2.3            | 18° to 90°* Two tensors and an isotropic component |
| Phantom             | 4000        | 160 | 4.6            | 30° and 60° Two sets of capillary tubes immersed in water |

*divided into 40 steps.  
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One should note that the same component dODF can be selected more than once, but the component set only includes unique elements. After the selection-decomposition recursion, the algorithm proceeds to estimation, in which we reformulated Eq. (1) as follow.

\[ \Psi_d + f_0 e + \sum_{i \in A} f_i \Psi_i \]  

(7)

One should note that in Eq. 7 the original dODFs are used instead of the normalized dODFs, and the solution for the volume fractions can be obtained by ordinary linear regression. To enforce the nonnegative constraint, if any of the volume fractions is negative, the most negative volume fraction is forced to zero, and then the equation is solved again until all volume fractions are nonnegative. The resulting fiber volume fractions \( f_1, f_2, \ldots, f_n \) constitute the solution of the fODF.

Parameter settings and computation efficiency

The computation efficiency of diffusion decomposition algorithm is mainly determined by the decomposition fraction, and it is usually set to a sufficiently small value to achieve a good estimation [22,23]. However, this setting may burden the computation since the initial decompositions are always repeated on the same component dODF until another component dODF becomes dominant. To reduce the number of repetition, we can decompose a larger fraction that just allows the second component to dominate, as implemented in the least angle regression algorithm [22]. Following this paradigm, we can derive the algorithm for solving this large decomposition fraction.

\[ \min_{\Psi_1 \neq \Psi_i} \frac{\langle \Psi_d, \Psi_i - \Psi_i \rangle}{\langle \Psi_d, \Psi_i \rangle \langle \Psi_i, \Psi_i - \Psi_i \rangle} \]  

(8)

where \( \Psi_i \) is the most correlated component dODF selected in the first step of the algorithm, and the minimum searches all other

### Table 3. Correlation analysis on estimated fiber volume fraction.

| FA  | CSD  | Ball-and-sticks | Decomposition |
|-----|------|-----------------|---------------|
| 0.4 | 0.7783 | 0.8836 | 0.8109 |
| 0.5 | 0.8018 | 0.8984 | 0.8325 |
| 0.6 | 0.8226 | 0.9108 | 0.8461 |
| 0.7 | 0.8458 | 0.9215 | 0.8649 |

Figure 2. The average angular deviation of constrained spherical deconvolution (CSD), ball-and-sticks model, and diffusion decomposition applied to 160 diffusion weighted images with SNR of 6.8. The performance is examined under different fractional anisotropy, crossing angles, and fiber volume fractions. Diffusion decomposition outperforms CSD and ball-and-sticks model in smaller crossing angles.

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components dODFs ($\Psi_i, i = 1, 2, \ldots, n, i \neq k$) to find the least decomposition fraction required to allow another component dODF to dominate. In this study, we used Eq. 8 only for the first decomposition, and then the decomposition fraction was set to a fixed value thereafter (e.g. 0.05).

The maximum size of the component set, $A$, may also affect the computation efficiency, and a suitable value can be determined by a priori information that human brain has limited number of crossing fiber populations stacked within a voxel. For most of the white matter area, the number of crossing fibers is no more than 3 [27,28], and in this study, we used a maximum size of 10 to get a balance between detection ability and computation time.

**Simulation study**

We simulated crossing fibers to evaluate the performance of diffusion decomposition and to compare it with that of CSD and ball-and-sticks model. A multiple-tensor model was used to obtain the diffusion signals of crossing fibers. As summarized in Table 2, the model has a component of isotropic diffusion and two fiber populations [5,29].

$$S(b,v) = S(0)[f_0 \exp\{-b v^T D_0 v\} + f_1 \exp\{-b v^T D_1 v\} + f_2 \exp\{-b v^T D_2 v\}]$$

(12)

where $S(0)$ is the baseline signal without diffusion encoding ($b_0$), $b$ and $v$ are the b-value and the unit vector of the applied diffusion gradient, respectively. $f_0$ is the volume fraction of the isotropic background diffusion. $f_1$ and $f_2$ are the volume fractions of the two fiber populations. $D_0$, $D_1$, and $D_2$ are the diffusion tensor matrices for these three diffusion components ($D_0$ is a multiple of the identity matrix). The inclusion of the isotropic background diffusion results in partial volume effect, which is a more challenging condition for obtaining fODFs. We simulated this model with a variety of parameter combinations, including fiber volume fraction, crossing angle, and FA. In the simulation study, the $f_0$ was set to 0.2, and $f_1$ were assigned from 0.50 to 0.90 ($1 - f_0$), divided into 40 steps of size 0.01. The remaining volume was occupied by $f_2$. The crossing angles ranged from 18° to 90°, divided into 40 steps of size 1.8°. FA values for both of the simulated fibers were 0.4, 0.5, 0.6, and 0.7. Two signal-to-noise (SNR) settings were simulated. The first SNR setting was simulated with Rician noise [30] added to 160 diffusion weighted images (HARDI, b-value = 1,500 s/mm$^2$) and a $b_0$ image. The mean diffusivity was $1.0 \times 10^{-2}$ mm$^2$/s. The SNR of the $b_0$ image was 40, and the average SNR of the 160 diffusion weighted images was 6.8. The second SNR setting was simulated with Rician noise added to 55 diffusion weighted images (HARDI, b-value = 1,500 s/mm$^2$) and a $b_0$ image. The mean diffusivity was $0.5 \times 10^{-2}$ mm$^2$/s. The SNR of the $b_0$ image was 20, and the average SNR of the 55 diffusion weighted images was 2.3. To
average the angular deviation, each parameter combination (fiber volume fractions, crossing angle, and FA) was simulated independently for a total of 100 trials. The source codes for generating the simulation images are publicly available at https://github.com/frankych/DSI-Studio.

The ball-and-sticks model was calculated using the bedpostx program in the FMRIB’s Diffusion Toolkit v2.0 package (http://fsl.fmrib.ox.ac.uk/). Rician noise model was used, and the grid points within a sphere in the q-space, maximum b-value = 4000 s/mm², and NEX = 4. T₂-weighted images were also acquired to measure the exact crossing angles of the phantom. For crossing angles originally arranged at 30° and 60°, the measured crossing angles on the T₂-weighted images were 31.48° and 63.59°, respectively. The angle was measured by the crossing angles of placeholders on the T₂-weighted images. Similar to the simulation study, CSD, ball-and-sticks model, and diffusion decomposition were conducted using the same parameter settings. The angular deviation was calculated using the resolved fiber orientations and the actual fiber orientations shown on the T₂-weighted images.

In-vivo study

Diffusion MRI was acquired from a 25-year-old healthy male participant on a 3T MRI system (TIM Trio, Siemens, Erlangen, Germany). The scan was performed using a 12-channel head coil and a single-shot twice-refocused echo planar imaging (EPI) diffusion pulse sequence. On the same subject, shell (for QBI), grid (for DSI), and two-shell (for GQI) sampling schemes were acquired consecutively using the same spatial parameters: field of view = 240 mm × 240 mm, matrix size = 96 × 96, slice thickness = 2.5 mm (no gap), the number of the slices = 40 covering the cerebral cortex, resulting in isotropic voxel size of 2.5 mm. The grid scheme was acquired by 252 diffusion gradient directions, b-value = 4000 sec/mm², and TR/TE = 7200 ms/133 ms. The shell scheme was acquired by 202 diffusion gradient directions at the grid points within a sphere in the q-space, maximum b-value = 4000 sec/mm², and TR/TE = 7200 ms/144 ms. The two-shell scheme was acquired by two DTI scans based on the built-in DTI b-table. One DTI dataset had 64 diffusion-weighted images (DWI) acquired by b-value = 3000 sec/mm², and TR/TE = 6300 ms/121 ms. Another DTI dataset had 30 gradient directions, b-value = 1500 sec/mm², and TR/TE = 5500 ms/6300 ms/121 ms. Diffusion decomposition and diffusion deconvolution were conducted using the same dataset to facilitate comparison. Diffusion deconvolution was conducted using a regularization parameter of 7, a value recommended for in-vivo data [11], whereas diffusion decomposition was conducted using the same setting as the phantom study.

Ethics Statement

Data were analyzed anonymously, and written informed consent was waived because the study fulfills the following requirements. The study has the lowest risk. The risk to the studied subjects does not exceed the possible risks of people who do not participate in
Sensitivity and Specificity Test

We applied diffusion decomposition to datasets with reduced number of diffusion encoding directions (termed reduced datasets hereafter) and compared the results with datasets acquired by full diffusion encoding directions (termed full datasets hereafter). The performance of diffusion decomposition was quantified in terms of the angular error and was compared with that of diffusion deconvolution [11], which can be equally applied to both shell and grid sampling schemes for comparison. For the shell scheme, the reduced dataset was the 30-direction dataset acquired with $b$-value = 1500 mm/s, and the full dataset was the 252-direction shell dataset acquired with $b$-value = 4000 mm/s. For the grid scheme, the reduced dataset was the 40 lowest $b$-values acquisitions of the full 202-direction grid dataset (maximum $b$-value = 2300 sec/mm$^2$). The fODFs of the full datasets (both shell and grid) were calculated by diffusion deconvolution with a regularization parameter of 7, a value recommended for in-vivo data [11], whereas the fODFs of the reduced datasets were calculated separately by diffusion deconvolution and diffusion decomposition for comparison. Since the performance may be affected by different parameter settings, the reduced dataset was processed under 5 different parameters for each method. For diffusion deconvolution, we used decomposition fractions of 0.01, 0.02, 0.05, 0.1, and 0.2. For diffusion deconvolution, the regularization parameters were set to 1, 2, 4, 8, and 16. For both reconstructions, the fiber orientations were determined by peak orientations on the fODFs.

To calculate the angular error related to sensitivity, we selected each fiber in the full dataset and calculated its angular error with respect to a corresponding fiber in the reduced dataset. If multiple fibers were resolved in the reduced dataset, the one that had the least angular error was chosen. If no fiber was resolved, an angular error of 180 degrees was assigned.

Figure 5. The angular error of constrained spherical deconvolution (CSD), ball-and-sticks model, and diffusion decomposition applied to phantom data. The diffusion decomposition shows significantly better accuracy than CSD at 30° crossing and ball-and-sticks model at 60° crossing (p-value < 0.001). doi:10.1371/journal.pone.0075747.g005

Figure 6. The fiber ODFs obtained by applying diffusion deconvolution and diffusion decomposition to QBI, DSI, and GQI. Each of the methods reconstructs diffusion ODF from shell, grid, and two-shell sampling schemes, respectively. Both diffusion deconvolution and diffusion decomposition can be equally applied to QBI, DSI, and GQI to improve their ability to resolve crossing fibers, but the fiber ODFs obtained from deconvolution shows blunt lobes with fluctuating baseline, whereas those from decomposition show sharp spikes with clean baseline, suggesting that diffusion decomposition can achieve better sensitivity and specificity in resolving crossing fibers. doi:10.1371/journal.pone.0075747.g006

Figure 7. The fiber ODFs obtained from diffusion decomposition applied to the 30-direction QBI, 252-direction QBI, 40-direction DSI, and 202-direction DSI. Even in datasets with substantially reduced number of diffusion encoding directions, diffusion decomposition can still depict correct crossing fibers formed by the corpus callosum and corticospinal tract. doi:10.1371/journal.pone.0075747.g007

The study. Waiving the prior consent does not affect the rights and interests of the studied subjects. (File # 1010265083, Ministry of Health, Executive Yuan, Taiwan). The research procedures were also approved by the Institutional Review Board of National Taiwan University Hospital.
Specificity was examined conversely. We selected each fiber in the reduced dataset and calculated its angular error with respect to a corresponding fiber in the full dataset. If multiple fibers were resolved in the full dataset, the one that had the least angular error was chosen. If no fiber was resolved, an angular error of 45° was assigned. To exclude gray matter in the analysis, the angular error was calculated by fibers in white matter area defined by applying a threshold to the quantitative anisotropy (QA) [7] of the full dataset. The exact threshold values were adjusted by comparing the extent of the threshold with the gray-white matter junction to achieve the best match of the coverage in the white matter.

Results

Simulation study

Figure 2 shows the average angular deviation of CSD, ball-and-sticks model, and diffusion decomposition applied to 160 diffusion weighted images with SNR of 6.8. All methods were applied to different combinations of crossing angle, fiber volume fractions, and FA. As shown in the figure, CSD performs well with crossing fiber greater than 45°, but its angular deviation increases in smaller crossing angles due to the false peaks in fODFs. This result demonstrates a typical specificity problem of CSD in smaller crossing angles. The noise in the diffusion signals can give rise to false peaks and introduce substantial angular deviation. In addition to the noise in the diffusion MRI, the additional isotropic component in our simulation model may corrupt the CSD results and lead to poor performance. The results of ball-and-sticks model show a similar pattern with CSD, but the accuracy is better in smaller crossing angles. Lastly, diffusion decomposition outperforms CSD and ball-and-sticks model in angles less than 45°, whereas in angles greater than 45°, its performance resembles the ball-and-sticks model. This feature can be attributed to the sparsity feature of the decomposition that offers good specificity while still retaining the sensitivity to crossing fibers.

Table 3 shows the result of a correlation analysis conducted to examine the accuracy of the fiber volume fractions provided by CSD, ball-and-sticks model, and diffusion decomposition. The analysis was conducted using the simulation dataset with 160 diffusion weighted images (SNR = 6.8). The correlation coefficients between the measured and simulated values are reported against different FA values. All methods provide good correlation (r > 0.7), and higher FA values lead to higher correlation coefficients. Among these methods, ball-and-sticks model provides

Figure 8. The boxplot showing the distribution of the angular error related to sensitivity. The results from (A) 30-direction QBI and (B) 40-direction DSI are shown in the figure. Diffusion deconvolution and diffusion decomposition was examined by 5 different parameters. The result shows that the decomposition approach presents lower values in the medians and upper quartiles than those of diffusion deconvolution, suggesting that diffusion decomposition is more sensitive than deconvolution.

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Figure 9. The boxplot showing the distribution of the angular error related to the specificity. The results from (A) 30-direction QBI and (B) 40-direction DSI are shown in the figure. Diffusion deconvolution and diffusion decomposition was examined by 5 different parameters. The result shows that the decomposition approach presents lower values in the medians and upper quartiles than those of diffusion deconvolution, suggesting that the result obtained from diffusion decomposition is more specific.

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the best estimation of the fiber volume fraction, whereas diffusion decomposition slightly outperforms CSD.

Figure 3 shows the average angular deviation of CSD, ball-and-sticks model, and diffusion decomposition applied to 55 diffusion weighted images with SNR of 2.3. All methods were applied to different combinations of crossing angle, fiber volume fractions, and FA. As shown in the figure, the CSD shows higher angular deviation in smaller crossing angles due to the false peaks, demonstrating its problem of specificity. By contrast, the results of the ball-and-sticks model show better accuracy in smaller crossing angles but not in larger crossing angles. The error may be due to model incompatibility (the ball-and-sticks model is a simplified model for multiple tensors), and the incompatibility error can be enhanced in low SNR conditions. Lastly, diffusion decomposition showed substantial better performance than CSD and ball-and-sticks model. The overall results suggest that diffusion decomposition can be applied to a diffusion dataset with 1) substantially reduced number of diffusion sampling and 2) low SNR.

Phantom study

Figure 4 shows the results of CSD, ball-and-sticks model, and diffusion decomposition applied to 30° and 60° crossing phantoms. The voxels at the center of the phantom are selected for visualization. All three methods are able to resolve crossing fibers at 60° crossing, however, ball-and-sticks model present a slight tilting of the fiber orientations. A possible cause to this error can be due to the discrepancy between the ball-and-sticks model and the actual diffusion pattern in the phantom. CSD and diffusion decomposition uses a consistent diffusion model (the response function was estimated from the phantom), and thus they do not have this drawback. At 30° crossing, CSD resolves one fiber orientation with several false fibers crossing at the horizontal direction. Ball-and-sticks model, despite the discrepancy of the diffusion model, is able to resolve two fibers. Diffusion decomposition method also resolves two crossing fibers and shows a clean baseline—positive values at fiber orientations and zeros elsewhere, underlining the sparsity feature of the decomposition algorithm.

This is in contrast to CSD, which has fluctuation around the origin that may give rise to false fibers and compromise its specificity.

Figure 5 shows the angular error of CSD, ball-and-sticks model, and diffusion decomposition applied to 30° and 60° crossing phantom. At 60° crossing, both CSD and diffusion decomposition show angular deviations lower than 1°, while ball-and-sticks model has a significant increase (p < 0.001) in angular deviation. At 30° crossing, ball-and-sticks model and diffusion decomposition shows similar angular deviations around 1°, while CSD has a significant increase (p < 0.001) in the angular error due to false fibers resolved at the horizontal directions.

In-vivo study

Figure 6 shows the fODFs obtained by applying diffusion deconvolution and diffusion decomposition to the fODFs acquired by DSI, QBI, and GQI. The figure is presented at the coronal view of centrum semiovale. Both diffusion deconvolution and diffusion decomposition are applicable to different q-space imaging methods and different diffusion sampling schemes, and the resulting fODFs show consistent fiber orientations. However, the fODFs estimated by the decomposition method show sharp spikes with a clean baseline owing to the sparsity feature. In contrast, the fODFs estimated by the deconvolution method show blunt lobes with a fluctuated baseline. The above comparison suggests that diffusion decomposition can achieve better angular resolution and specificity in the in-vivo study.

Figure 7 shows the fODFs obtained from diffusion decomposition applied to the 30-direction QBI, 252-direction QBI, 40-direction DSI, and 202-direction DSI. The figure presents the centrum semiovale in the coronal view. As shown in Fig. 7, diffusion decomposition can reveal crossing fibers formed by the corpus callosum (horizontal fibers) and corticospinal tracts (vertical fibers) using reduced sampling schemes, and the crossing fiber orientations are consistent with those of the full schemes. The specificity of diffusion decomposition can be appreciated in the mid corpus callosum, where it presents single fiber population with correct orientation and is free from false fibers. The above results suggest that diffusion decomposition can be applied to a reduced dataset to provide correct information about fiber orientations.

Figure 8 shows the boxplot of the angular error related to sensitivity. Figure 8A shows the results obtained from the 30-direction QBI, whereas Figure 8B shows the results obtained from the 40-direction DSI. Both diffusion deconvolution and diffusion decomposition were tested using 5 different parameters to compare their performance. In both QBI and DSI, diffusion decomposition significantly reduced the angular error compared to CSD.

Table 4. Mean angular error of diffusion deconvolution and diffusion decomposition in 30-direction QBI and 40-direction DSI.

| Sensitivity test | Dataset | Results under different parameters* |
|-----------------|--------|----------------------------------|
| Diffusion deconvolution | 30-dir QBI | 20.19° 20.60° 20.77° 21.02° 22.12° |
|                  | 40-dir DSI | 18.36° 17.98° 17.73° 17.84° 18.33° |
| Diffusion decomposition | 30-dir QBI | 16.39° 16.29° 16.08° 15.90° 15.68° |
|                  | 40-dir DSI | 16.11° 16.04° 15.80° 15.45° 14.87° |
| Specificity test | Diffusion deconvolution | 30-dir QBI | 49.02° 48.33° 47.20° 44.67° 38.75° |
|                  | 40-dir DSI | 47.12° 46.62° 49.54° 49.69° 48.61° |
| Diffusion decomposition | 30-dir QBI | 33.35° 33.44° 33.73° 34.02° 34.51° |
|                  | 40-dir DSI | 28.73° 28.86° 29.23° 29.85° 30.89° |

*Diffusion deconvolution was conducted by regularization parameters of 1, 2, 4, 8, and 16. Diffusion decomposition was conducted by decomposition fractions of 0.01, 0.02, 0.05, 0.1, and 0.2.

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**Figure 10. Fiber volume mapping versus FA mapping. (A) The mapping of the total fiber volume calculated from diffusion decomposition and (B) the mapping of the fractional anisotropy (FA) calculated from the diffusion tensor analysis applied to the same data. The FA mapping shows decreased values in the centrum semiovale due to the crossing fibers in this region (annotated), whereas the total fiber volume mapping shows a relatively homogeneous intensity throughout the white matter. This suggests that the fiber volume can provide a better gray-white matter separation and facilitate further investigation into structural integrity.**

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decomposition shows lower values in medians and upper quartiles regardless of the parameter settings applied, suggesting that diffusion decomposition is more sensitive to crossing.

Figure 9 shows the boxplot of the angular error related to specificity. Figure 9A shows the results obtained from the 30-direction QBI, whereas Figure 9B shows the results obtained from the 40-direction DSI. In both datasets, diffusion decomposition shows substantially lower values in medians and upper quartiles, regardless of the parameter settings. This specificity study suggests that the fibers resolved by diffusion decomposition are more specific than those by diffusion convolution.

Table 4 lists the mean angular error of diffusion deconvolution and diffusion decomposition in the sensitivity and specificity tests. In the sensitivity test, the mean angular errors of diffusion deconvolution are around 20° in 30-direction QBI and 18° in 40-direction DSI, whereas those of the decomposition method are around 16° in both 30-direction QBI and 40-direction DSI. The decomposition method has an average of 4.87° less angular error in 30-direction QBI and 2.39° less angular error in 40-direction DSI. The difference is statistically significant (p-value < 0.001). Similarly, the specificity test demonstrates consistent conclusion but with a more substantial difference. The mean angular errors of diffusion deconvolution range from 38° to 49° in 30-direction QBI and 47° to 49° in 40-direction DSI, whereas those of the decomposition method are around 33° and 29° in 30-direction QBI and 29° in 40-direction DSI. The decomposition method has an average of 11.78° less angular error in 30-direction QBI and 19.20° less angular error in 40-direction DSI. The difference is statistically significant (p-value < 0.001). In terms of sensitivity and specificity, diffusion decomposition achieved significantly better performance than diffusion deconvolution.

Figure 10 shows the mapping of the total fiber volume fraction derived from diffusion decomposition (Fig. 10A) and the mapping of FA (Fig. 10B). The total fiber volume fraction was calculated by summing all fiber volume fractions in our mixed diffusion model ($\beta_0$ excluded), whereas the FA was calculated from the tensor analysis. Both images were obtained from the same 202-direction DSI dataset in an axial view at the level of the corpus callosum. The intensities of the images were scaled by their relative contrast so that the maximum value corresponded to the maximum intensity. Although the total fiber volume fraction mapping shows a pattern similar to the FA mapping, decreased values of FA can be observed in the regions where the corpus callosum crosses the corticospinal tracts (indicated by arrows). By contrast, the total fiber volume fraction mapping shows a relative homogeneous distribution of the intensity throughout the white matter. The above comparison suggests that the total fiber volume fraction is less affected by crossing fibers and may serve as an index specific to fiber volume fraction of white matter tracts.

**Discussion**

This paper proposes a sparse iODF estimation method called diffusion decomposition, which obtains iODF by decomposing the dODF acquired from DSI, QBI, or GQI. The simulation study shows that diffusion decomposition offers better accuracy than CSD and ball-and-sticks model in the dataset with substantially reduced number of diffusion sampling directions. The phantom study shows that the iODF of diffusion decomposition presents the sparsity feature—zero values at most orientations and positive values only at fiber orientations. This is in contrast to the iODF of CSD, which presents fluctuation at the baseline and often requires filtering to eliminate false fibers. Further quantitative analysis shows that diffusion decomposition presents significantly lower angular error than that of CSD at 30° crossing or ball-and-sticks model at 60° crossing. Similar conclusion can be drawn from the results of the in-vivo study where diffusion decomposition outperforms diffusion deconvolution owing to the sparsity feature of the solution.

From the perspective of computation, diffusion decomposition seems to be a hybrid of the model-fitting (e.g. ball-and-sticks model) and deconvolution approach. The performance in resolving crossing fibers is improved by promoting sparsity. Compared with model-fitting approach such as ball-and-sticks model, an interesting finding in our study is that while diffusion decomposition presents similar performance in both simulation and phantom study, ball-and-sticks model seems to perform worse in the phantom at 60° crossing than it does in the simulation. One possible reason for this underperformance may be due to the poor fitting of ball-and-sticks with the actual diffusion model in the phantom study. It is possible that the diffusion model is more complicated in real world scenario, and ball-and-sticks may not be adequate for fitting its signal responses. Compared with deconvolution, the outperformance of diffusion decomposition can be summarized in two aspects: sensitivity and specificity. The increased sensitivity can be attributed to the spiky iODF obtained by $L_1$ regularization, whereas the iODF obtained by $L_2$ regularization still retains part of the dODF blurring that leads to less sensitive results. The second aspect of the outperformance—specificity—is even more substantial. While the angular error related to sensitivity is improved by ~3° in the reduced datasets, almost negligible in terms of the ODF angular resolution, the angular error related to specificity is improved by ~15°. The substantial improvement in the specificity can be expected. As we have pointed out in our previous deconvolution study [11], the deconvolution method does not guarantee a good specificity although it has been shown to achieve a good sensitivity to crossing fibers. Higher smoothing or ODF filtering is often required to remove false fibers, but less salient fibers will also be removed indiscriminately. By contrast, diffusion decomposition offers a better specificity owing to the sparsity nature of $L_1$ regularization. The clean baseline of the sparse iODFs can avoid false identification of the crossing fibers. One should note that this outperformance does not come without a price. Diffusion decomposition sacrifices smoothness for sparsity to obtain better resolving power. Consequently, it cannot portray fiber dispersion due to the sparsity assumption. Nonetheless, since the fiber dispersion phenomenon is more obvious in low spatial resolution images, it may be possible to mitigate this limitation by acquiring diffusion images at a higher spatial resolution. Further study may be needed to fully investigate this limitation under different resolutions.

There are several other methods that have promoted sparsity in the spherical harmonics, whereas we promoted it in the ODF space. The advantage of our approach is that a sparse iODF can be easily represented by an ODF vector but not by spherical harmonics. This can be readily examined by calculating the spherical harmonics of a delta function (i.e. the iODF of a single fiber). It results in infinite non-zero responses of spherical harmonics, meaning that one cannot use a finite order of spherical harmonics to represent a single-fiber iODF. Methods based on spherical harmonics inevitably truncate higher responses of the spherical harmonics, and their estimated iODF becomes a smoothed version of the true iODF. Consequently, enforcing sparsity in the ODF space may achieve better accuracy.

In addition to the outperformance, another noteworthy feature of diffusion decomposition is its ability to resolve crossing fibers based on 30-direction QBI and 40-direction DSI. This feature is
particularly important since QBI and DSI have long been criticized for lengthy scanning time that limits their applications in clinical study. As shown in an optimization study [31], QBI and DSI typically requires maximal diffusion sensitivity about 3000 to 4000 s/mm² and more than 200 diffusion encoding directions, resulting in a scanning time of approximately 30 min. In this study, the 30-direction QBI was acquired using the built-in DTI sequence in the clinical scanner, which is readily available for the most of the MRI facilities. The 40-direction DSI needs only a maximum b-value of 2300 sec/mm², a setting that greatly reduces the demand for the gradient performance. For either 30-direction QBI or 40-direction DSI, the scanning time of the diffusion acquisition is approximately 5 minutes, making it a realistic option for inclusion in clinical studies. Nevertheless, one should note that the reduction of the number of gradient directions still has its shortcoming. Although diffusion decomposition can resolve crossing fibers in the white matter, we observe that its sensitivity is compromised at the gray-white matter junction. This could be due to the low SNR of diffusion MR signals in the region, and there seems to be a trade-off between the SNR and the number of diffusion gradient directions.

Lastly, diffusion decomposition has other features that are also worth mentioning. 1) It is equally applicable to grid, single-shell, and multi-shell encoding schemes. This makes the algorithm a convenient tool for data with various combinations of b-values and diffusion encoding schemes. 2) The fiber volume fraction is assigned to a specific fiber orientation, and thus it can be used as an index to quantify structural connectivity for human connectome study. The total fiber volume fraction derived from the decomposition method can also be used to delineate the gray-white matter junction and may be used to examine structural integrity.

There are drawbacks in diffusion decomposition. Similar to deconvolution methods, diffusion decomposition uses a putative common characteristic dODF to model the single fiber direction compartment despite the fact that the characteristic dODF varies across different regions in the white matter. As shown in our previous study, the dODFs obtained from the corpus callosum, cingulum bundle, and corticospinal tracts have different profiles [11]. Applying a universal characteristic ODF to fit all scenarios may introduce additional errors, and this drawback has been reported to decrease the specificity of spherical deconvolution [10]. Although this problem was not rigorously examined in this study, we believe that our method is also susceptible to this drawback as long as a universal characteristic dODF is used. Possible improvement may be achieved by using multiple characteristic dODFs instead of a common dODF or using an unsupervised clustering approach to characterize the component dODFs. Furthermore, the 642-direction ODF has a limited angular resolution around 8 degrees, and diffusion decomposition therefore has an inherent error due to limited resolution of an ODF. Lastly, a fixed value of decomposition volume was used for all voxels in our study. In practice, it is possible to optimize this value for each voxel using cross-validation, and this is a potential future work to further improve the performance of diffusion decomposition.

In conclusion, diffusion decomposition has a particular role in diffusion datasets with substantially reduced number of diffusion encoding directions, making it highly feasible for clinical studies that allows a limited scanning time. The decomposition algorithm provides a sparse solution of IODF to improve the ability in resolving crossing fibers and to avoid false fibers as encountered in diffusion deconvolution. The algorithm can be applied to diffusion data and to facilitate further fiber tracking algorithms. These features may be valuable for future human connectome study to define brain connectivity.

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Author Contributions
Conceived and designed the experiments: FY WT. Performed the experiments: FY WT. Analyzed the data: FY WT. Contributed reagents/materials/analysis tools: FY WT. Wrote the paper: FY WT.

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