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Published in:
IEEE Transactions on Medical Imaging

DOI:
10.1109/TMI.2013.2285500

Publication date:
2014

Document Version
Peer reviewed version

Link to publication in Heriot-Watt University Research Portal

Citation for published version (APA):
Quantitative comparison of reconstruction methods for intra-voxel fiber recovery from diffusion MRI

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Abstract—Validation is arguably the bottleneck in the diffusion MRI community. This paper evaluates and compares 20 algorithms for recovering the local intra-voxel fiber structure from diffusion MRI data and is based on the results of the “HARDI reconstruction challenge” organized in the context of the “ISBI 2012” conference. Evaluated methods encompass a mixture of classical techniques well-known in the literature such as Diffusion Tensor, Q-Ball and Diffusion Spectrum imaging, algorithms inspired by the recent theory of compressed sensing and also brand new approaches proposed for the first time at this contest. To quantitatively compare the methods under controlled conditions, two datasets with known ground-truth were synthetically generated and two main criteria were used to evaluate the quality of the reconstructions in every voxel: correct assessment of the number of fiber populations and angular accuracy in their orientation. This comparative study investigates the behavior of each algorithm with varying experimental conditions and highlights strengths and weaknesses of each approach.

This information can be useful not only for enhancing current algorithms and develop the next generation of reconstruction methods, but also to assist physicians in the choice of the most adequate technique for their studies.

Index Terms—diffusion MRI, local reconstruction, quantitative comparison, validation, synthetic data

I. INTRODUCTION

DIFFUSION MRI (dMRI) is a powerful magnetic resonance imaging modality which is sensitive to the Brownian motion of water molecules in living tissues. This peculiar feature is exceptionally effective in the brain as it permits to investigate the structural wiring of the nervous system in vivo and non-invasively. In fact, in the white matter, these microscopic random movements are highly hindered by the coherent organization in bundles of the neuronal tissue [1] and it is possible to infer the architecture of the neuronal connections by properly characterizing the distribution of the displacements. The richer and more detailed this description is, the more accurate any subsequent global connectivity analysis will be. In fact, all fiber-tracking algorithms [2], [3] (see [4] and references therein for a more complete list) rely on a suitable characterization of the diffusion process within each voxel as they attempt to estimate the most likely fiber tracts connecting different regions of the brain.

The first approach to model the anisotropic nature of the diffusion process in biological tissues has been Diffusion Tensor Imaging [5] (DTI), which is based on the Gaussian assumption on the displacements. Despite providing useful diagnostic information with very fast acquisitions, DTI is unable to resolve multiple fiber populations within a voxel. To overcome this limitation, Diffusion Spectrum Imaging [6] (DSI) has been proposed. DSI is a model-free technique exclusively based on the Fourier relationship [7] between the signal acquired in a Cartesian grid and the 3D displacement spectrum, also known as the Ensemble Average Propagator (EAP). DSI provides probably the most detailed description of diffusion within each voxel, but it requires hundreds of measurements which are not compatible with the normal acquisition time admitted in clinical protocols. To this aim, a large number of alternative reconstruction techniques have been developed to recover both the angular and radial component of the diffusion process. Among them, to mention a few, we can cite the models based on multiple [8]–[11] or higher-order tensors [12]–[14],

* indicates the corresponding author
the Q-Ball Imaging (QBI) [15] and its variant computed in constant solid angle [16], [17], the Persistent Angular Structure MRI (PAS-MRI) [18], the Composite Hindered And Restricted Model of Diffusion (CHARMED) [19], the Diffusion Orientation Transform (DOT) [20], [21], the Hybrid Diffusion Imaging (HIDY) [22], the Generalized Q-sampling Imaging (GQI) [23] and all the algorithms based on spherical deconvolution [24]–[29]. Multi-compartment models have been also proposed in the literature to explain the measured dMRI signal more accurately; see [30] for a detailed survey. All these techniques are characterized by fitting advanced and more descriptive models of diffusion (e.g. intra- and extra-axonal space) to get access to important micro-structure parameters such as axonal diameter and density. Lately, the advent of compressed sensing [31], [32] inspired new approaches to solve the reconstruction problem whose main goal is to drastically reduce the acquisition time [33]–[39]. This list is not exhaustive and many other algorithms exist in the literature; see [40]–[42] for a more comprehensive list of available techniques. All these methods differ in a great deal of aspects, like for instance the acquisition scheme adopted (number and position of the diffusion gradients), the targeted feature of the diffusion process to estimate, e.g. EAP, diffusion Orientation Distribution Function (ODF) or Fiber Orientation Distribution (FOD), and the assumptions made in the reconstruction. A direct comparison among techniques is hence very difficult.

Another source of confusion when comparing the approaches proposed in the literature stems from the different choices of the framework used to validate the reconstructions. Validation frameworks can be divided into three categories: biological phantoms, physical phantoms and numerical simulations. In the first case, the fiber configurations estimated from dMRI are compared with the results of histological or tracer-injection studies [43]–[46], either in vivo or ex vivo, performed on the same biological sample. In particular, an emerging technique called three-dimensional Polarized Light Imaging (3D-PLI) [47] is capable of providing the directionality of the axons in postmortem human brains at ultra-high resolution, i.e. sub-millimeter scale. All these techniques offer exquisite insights of the brain structure but they are very invasive and, more importantly, allow to investigate only a few specific axonal tracts or sample slices at once. On the other hand, with physical phantoms [48]–[57] it is possible to build specific arrangements of synthetic fibers that mimic bending, crossing and kissing configurations usually observed in the real brain anatomy and thus validate the algorithms in more complex scenarios. However, the construction of a realistic phantom is an extremely challenging task as the dimension and the complexity of the fiber architecture which is possible to achieve with current technologies are orders of magnitude different from the real nervous system. Moreover, all publicly available phantoms have normally been scanned only a limited number of times, hence providing images corresponding to a very limited number of sampling schemes and with only an intrinsic acquisition noise. Therefore they are not suitable for testing the performances of an algorithm in different scenarios. By contrast with the two previous validation frameworks, numerical simulations offer a simple and versatile way to assess the performances of a reconstruction algorithm with a broader set of experimental conditions, e.g. signal models [8], [58], fiber configurations, sampling schemes and noise levels. Interestingly, Tournier et al. [48] have shown that numerical simulations performed using these models agree well with physical phantoms but, as already mentioned, they represent an over-simplification of what happens in the white matter. Even though the level of complexity they offer is way too simplistic with respect to the real architecture of the nervous system, they actually constitute the validation framework of choice for the majority of the algorithms proposed in the literature, as in [8], [17], [20], [23], [27], [28], [33], [34], [37], [59], [60] to mention a few. Unfortunately, when a new algorithm is proposed, the performances are normally assessed with ad-hoc synthetic data and evaluation criteria, and comparing different approaches can be difficult. This lack of consensus on the validation framework is particularly crucial in a clinical perspective, as the availability of a comprehensive comparison of the most common techniques might help clinicians in the choice of the most adequate method for a specific application.

With these considerations in mind, the aim of this work is to provide all researchers in this field with a common validation framework to assess the performances of their own algorithms for the intra-voxel fiber recovery and fairly compare their results against other approaches under controlled conditions. It is not an intention of this work to determine the most suitable technique to adopt in a clinical study, as the choice highly depends both on the interests (e.g. research question or intra-voxel feature to extract) and possibilities (e.g. available scan time) of the study itself. Nor it is likely that an algorithm outperforms all the others in all possible scenarios. This study should be intended as a comprehensive set of analyses reporting strengths and weaknesses of a large selection of state-of-the-art techniques in a wide range of experimental conditions. The focus is on the angular component of the recovered diffusion profiles. Because of the large number of techniques proposed in the literature to this end, the evaluation presented in this work is not exhaustive. Nonetheless, to date and to the best of our knowledge it represents the comparison study with the largest number of different reconstruction techniques and the most heterogeneous set of experimental conditions.

The rest of the manuscript is organized as follows. In section II we present the challenge, describing in detail its organization, the ground-truth synthetic data and the evaluation criteria used, and the reconstruction methods that were considered in the study. The results of the comparison are presented and discussed in section III. We conclude this paper with general considerations and some take-home messages.

II. Materials and methods

In this section we describe in detail the organization of the contest, the reconstruction methods considered in this study and the evaluation framework that we used to quantitatively compare all the algorithms, i.e. synthetic datasets with known ground-truth and evaluation criteria. More information can be found on the challenge website\footnote{hardi.epfl.ch/static/events/2012_JSBI}. 
A. Context organization

The diffusion MRI reconstruction challenge was organized in the context of the 9th IEEE International Symposium on Biomedical Imaging (ISBI) conference, which was held in Barcelona (Spain) in May 2012. The purpose of this contest was to evaluate and compare different reconstruction methods for recovering the intra-voxel fiber structure, i.e., number and orientation of the fibers populations present in each voxel, from dMRI acquisitions, using the same ground-truth data and under controlled conditions.

Training data was released prior to the contest in order for the contestants to practice with the data format, test their methods and identify the setting providing the best results, e.g., acquisition protocol and specific parameters of the algorithm. For the final evaluation of the performances of each method, though, another set of testing data was used. In order to grant the maximum flexibility for the acquisition scheme required by each reconstruction algorithm, while still keeping the ground-truth of the testing data unknown to the contestants, we adopted the following single-blind procedure for the generation of the dMRI signal. Each participant had the possibility to probe the signal corresponding to the ground-truth dataset only once by submitting to the organizers an acquisition scheme specifying the sampling coordinates in q-space, i.e., direction and strength of the diffusion gradients. The minimum and maximum number of samples which was possible to probe was fixed to 6 and 257, respectively, corresponding to standard DTI and DSI acquisitions on one hemisphere, assuming central symmetry in the diffusion signal. There were no restrictions, however, on the position of the samples in the q-space. The signal was then numerically simulated in every voxel of the dataset by the organizers (see next section) according to the gradient list received from each team. This is the only information the contestants knew about the common dataset. For teams participating with more than one method, data sharing between the algorithms was not allowed. At this point, the participants could apply their proposed reconstruction algorithms on the data received from the organizers in order to recover the underlying fiber configuration. Any combination of algorithms, signal processing techniques and available software was allowed, with the only constraint that the outcomes of the proposed methodology must be reproducible.

B. Synthetic data

1) Signal simulation: To simulate the dMRI signal intensity in a voxel with $M$ fiber populations and corresponding to the application of a diffusion-sensitizing gradient we adopted the classical Gaussian Mixture Model proposed by [8]:

$$S(q) = S_0 \sum_{i=1}^{M} f_i \exp \left( -b u^T D_{(i)} u \right),$$  

(1)

where $q$ is the coordinate vector in q-space accounting for the direction and strength of the diffusion gradient, $u \in S^2$ is a unit vector in 3D with $q = qu = |q|u$, $b = 4 \pi^2 q^2 t$ is the b-value corresponding to $q$ and diffusion time $t$ and $S_0$ is the signal without diffusion weighting (i.e., $b = 0$). Each fiber population can be characterized by a tensor $D_{(i)}$ and a non-negative volume fraction $f_i$ such that $\sum_{i=1}^{M} f_i = 1$. Each $D_{(i)}$ can be expressed as a rotated version of a tensor $D = \text{diag}(\lambda_1, \lambda_2, \lambda_3)$ as $R^T D R$, $R$ being the rotation matrix rotating the main axis of $D$ to the direction of a given fiber population. Diagonal elements of $D$ are the diffusivities along the main axis of the fiber ($\lambda_1$) and in the plane perpendicular to it ($\lambda_2, \lambda_3$). In this contest, the diffusivities were generated from the following ranges typically observed in the human brain [60]: $\lambda_1 \in [1, 2] \times 10^{-3}$ mm$^2$/s and $\lambda_2 = \lambda_3 \in [0.1, 0.6] \times 10^{-3}$ mm$^2$/s. Also, we assumed $S_0 = 1$ without loss of generality.

2) Noise simulation: To be consistent with previous studies presenting new reconstruction methods, in this work the signal $S$ has been corrupted by Rician noise [61] as follows:

$$S_{\text{noisy}} = \sqrt{(S + \eta_1)^2 + \eta_2^2}$$

(2)

with $\eta_1, \eta_2 \sim \mathcal{N}(0, \sigma^2)$ and $\sigma = S_0/\text{SNR}$ controls the level of the noise, which is reported here as the signal-to-noise ratio (SNR) on the $S_0$ image. In our simulations the only source of artifact is Rician noise and the performances of the reconstruction methods have been evaluated as function of its magnitude for three different noise levels, i.e., SNR = 10, 20, 30. The study of the quality of the reconstructions as function of other realistic imaging artifacts such as eddy currents, $B_0$-field inhomogeneity and subject motion, was beyond the scope of this work.

3) Phantoms: In order to adequately evaluate the performances of each algorithm with different experimental conditions we created two separate synthetic phantoms.

![Structured field synthetic dataset](image)

Fig. 1. **Structured field synthetic dataset.** The 5 different fiber bundles (A-E) are reported separately. In each voxel, the directions of the fiber populations are color-coded depending on their orientation (x-axis, y-axis, z-axis). Subplot F shows a representative slice with the corresponding ODFs computed analytically from the equation (1) as in [15], [60].

The **structured field** dataset (hereafter SF) consists of a $16 \times 16 \times 5$ volume attempting to simulate a realistic 3D configuration of tracts occurring in reality. As shown in Fig. 1, this dataset comprises five different fiber bundles which give rise to non-planar configurations of bending, crossing and kissing tracts. All fiber tracts are characterized by having a fractional anisotropy between 0.75 and 0.90. The diffusion properties have been generated from the diffusivity ranges previously described and were kept constant along the full
trajectory of each bundle. The dataset contains the following proportion of voxel configurations: 35% with a single fiber population, 35% with two and 30% with three. In voxels with multiple fiber populations, then, the crossing angles between the fiber directions were distributed as follows: 27% smaller than 30°, 35% in the range 31°–60° and 38% above 61°. The purpose of this dataset was to test the average performance of each algorithm with plausible fiber configurations. Moreover, we used this dataset to investigate the ability of each algorithm to exploit additional information from neighboring voxels for possibly improving the reconstruction of the fiber configuration at a given position.

On the other hand, the isolated voxels dataset (hereafter IV) consists of a set of 9100 independent voxels with no spatial coherence among them. Each voxel contains a given configuration of 2 fiber populations crossing at specific angles, with 100 repetitions of every given configuration. The diffusion properties of each population have been randomly generated by means of uniform distributions in the ranges previously described, but the two fiber populations do not necessarily share the same values. As for the SF dataset, the fractional anisotropy was maintained between 0.75 and 0.90. For completeness, the analysis of the performance of each reconstruction algorithm is provided in the full range of crossing angles, 0°–90°, as in [17], [38], [48]. The purpose of this dataset is to explore more in detail the behavior of each approach as some experimental conditions change.

C. Evaluation criteria

To assess the quality of the reconstructions we have focused our attention on two main criteria: (i) correct assessment of the number of fiber populations present in each voxel and (ii) angular accuracy in their orientation. We have purposely adopted evaluation metrics widely used in the literature, e.g. [26], [29], [36]–[38], [62], [63], with the aim to simplify and standardize the comparison of our results with previously reported findings. To compute these metrics we have compared the fiber populations (also referred to as fiber compartments in this work) of the ground-truth with the directions estimated by each reconstruction algorithm. There was not consensus, though, in the strategy to estimate these directions: some teams used peak-finding procedures on the recovered ODFs while others extracted them directly in the reconstruction problem (e.g. DTI), some used a threshold to filter out small peaks and some other did not. More information on each method’s strategy can be found on the challenge website.

In order to assess the correct estimation of the number of fiber compartments we employed the success rate, hereafter reported as SR, which is defined as the proportion of voxels in which a reconstruction algorithm estimates correctly the right number of fiber populations. To understand the reasons of an incorrect assessment, we also used the following two quantities, \( n^+ \) and \( n^- \), which explicitly count the number of over-estimated and under-estimated fiber populations in a voxel, respectively. Furthermore, in order to distinguish between the estimated fiber populations that are close enough to the real ones, i.e. true positives, and those which are clearly spurious, i.e. false positives, we recomputed the previous metrics using a tolerance cone as suggested in [26], [63]. These three additional measures, namely \( SR_+, n^+_\theta \) and \( n^-\theta \), are defined as before but an estimated direction is considered “resolved” only if it falls within a tolerance cone around a real fiber population in the voxel. In this work the tolerance was set to 20°. Noteworthy, in this study we have employed both versions of these metrics, i.e. with and without tolerance cone, as this provided more insight into the reconstructions.

The angular accuracy in the orientation of the estimated fiber compartments was assessed by means of the average error (in degree) between the estimated fiber directions and the true ones present in a voxel:

\[
\bar{\theta} = \frac{180}{\pi} \arccos(|d_{\text{true}} \cdot d_{\text{estimated}}|),
\]

where the unitary vectors \( d_{\text{true}} \) and \( d_{\text{estimated}} \) are, respectively, a true fiber population in the voxel and the closest of the estimated directions; then \( \cdot \) denotes the dot product. The final value in the voxel is the average computed by first matching the directions estimated by the algorithm to the ground-truth in order to minimize the error.

D. Reconstruction methods

Table I gives an overview of the 20 algorithms considered in this work. For more details about each reconstruction method (not reported here for conciseness), please refer to the proceedings of the workshop on the challenge website. In addition to the 13 submissions received during the contest, in this study we also considered 7 additional well-established reconstruction algorithms with the aim to have a more comprehensive comparison. To prevent unfair comparisons among methods using so different acquisition protocols, the algorithms have been grouped into four different classes depending on the characteristics of the sampling scheme adopted.

The DTI-like class includes approaches based on clinical acquisition protocols with less than 15 samples and maximum b-value equal to 1000 s/mm². In addition to the classical Diffusion Tensor Imaging [5] (DTI), this class comprises a novel approach named DTI\_neigh [64] which aims at improving the reconstructions by means of contextual enhancement of the spherical diffusion functions based on the notion of context.

In the HARDI-like class we grouped those methods based on acquisition schemes with one or multiple shells using a maximum b-value of 3000 s/mm². Six well established techniques have been included: Diffusion Orientation Transform [20] (DOT), analytical Q-Ball Imaging [59] (QBI), Q-Ball Imaging with Spherical Wavelet Transform [66] (QBI\_SWT), Q-Ball Imaging in Constant Solid Angle [17] (QBI\_CSA), Non-negativity Constrained Super-resolved Spherical Deconvolution [27] (CSD) and the Sharpening Deconvolution Transform [29] (SDT). In addition, two novel algorithms have been considered: STD, a spherical deconvolution approach based on Symmetric Tensor Decomposition, and SPIRAL, an innovative technique based on spiral sampling which exploits the periodic pattern of the signal in the gradient direction domain for denoising and estimating the intra-voxel fiber configuration.
### TABLE 1

**Summary of the reconstruction algorithms evaluated in this study.** The methods have been grouped into four different classes, depending on the sampling scheme adopted: DTI-like (violet), SPARSE-like (green), HARDI-like (yellow-red) and DSI-like (blue). The colors used here will be used coherently throughout the manuscript to identify each algorithm.

<table>
<thead>
<tr>
<th>Method</th>
<th>Acquisition</th>
<th>Brief Description of the Algorithm</th>
<th>Spatial Filter</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>DTI</strong></td>
<td>one-shell b = 1000 6 samples</td>
<td>Diffusion Tensor Imaging (DTI) [5]</td>
<td>none</td>
</tr>
<tr>
<td><strong>DTI_\text{smooth}</strong></td>
<td>one-shell b = 1000 12 samples</td>
<td>Contextual enhancing of spherical diffusion functions for improving DTI reconstructions, based on the notion of context to extrapolate additional information about the underlying fiber structure [64]</td>
<td>regularization</td>
</tr>
<tr>
<td><strong>L2-L1-DL</strong></td>
<td>one-shell b = 2000 15 samples</td>
<td>A parametric dictionary is learnt from a set of training diffusion data, providing a highly sparse representation of the diffusion signal and enabling to analytically recover important diffusion features as the ODF with a small number of measurements [39]</td>
<td>none</td>
</tr>
<tr>
<td><strong>L1-L1</strong></td>
<td>one-shell b = 750 24 samples</td>
<td>Sparse reconstruction using the $\ell_1$ norm both in the data fidelity term and the sparsity regularization with a Gaussian diffusion profile dictionary</td>
<td>none</td>
</tr>
<tr>
<td><strong>L2-L1-TV</strong></td>
<td>one-shell b = 1200 30 samples</td>
<td>Sparse reconstruction with combined q-space and k-space with Total Variation regularization in the spatial domain [65]</td>
<td>regularization</td>
</tr>
<tr>
<td><strong>L2-L2</strong></td>
<td>one-shell b = 3333 37 samples</td>
<td>Multi-tensor fitting guided by ODF estimation, to obtain the true diffusivities corresponding to each different fiber population in the voxel. In the estimation process, the number, orientations and relative intensities of fiber populations were assumed to be known and equal to the values determined from the ODF maxima computed as in [21]</td>
<td>smoothing</td>
</tr>
<tr>
<td><strong>NLR-L2</strong></td>
<td>one-shell b = 1500 48 samples</td>
<td>Diffusion Basis Functions + selective spatial smoothing based on FA and diffusion orientation similarity [33]</td>
<td>smoothing</td>
</tr>
<tr>
<td><strong>L2-L1-TGV</strong></td>
<td>multi-shell $b \in {1500,2500}$ 64 samples</td>
<td>$\ell_1$-based constrained spherical deconvolution with Total Generalized Variation regularization optimized by a first-order primal-dual approach</td>
<td>regularization</td>
</tr>
<tr>
<td><strong>DOT</strong></td>
<td>one-shell b = 3000 60 samples</td>
<td>Diffusion Orientation Transform (DOT) [20] with the following parameters: Spherical Harmonics (SH) order $\ell_{\text{max}} = 8$ and EAP evaluated at radius $R = 15 , \mu m$</td>
<td>none</td>
</tr>
<tr>
<td><strong>QBI</strong></td>
<td>one-shell b = 3000 60 samples</td>
<td>Analytical Q-Ball Imaging (QBI) [59] with the following parameters: SH order $\ell_{\text{max}} = 6$ and regularization parameter $\lambda = 0.006$</td>
<td>none</td>
</tr>
<tr>
<td><strong>QBI_\text{SWT}</strong></td>
<td>one-shell b = 3000 60 samples</td>
<td>Q-Ball imaging with Spherical Wavelet Transform [66] with the following parameters: SH order $\ell_{\text{max}} = 8$</td>
<td>none</td>
</tr>
<tr>
<td><strong>QBI_\text{CSA}</strong></td>
<td>one-shell b = 3000 60 samples</td>
<td>Q-Ball imaging in Constant Solid Angle [17] with the following parameters: SH order $\ell_{\text{max}} = 6$ and regularization parameter $\lambda = 0.006$</td>
<td>none</td>
</tr>
<tr>
<td><strong>CSD</strong></td>
<td>one-shell b = 3000 60 samples</td>
<td>Non-negativity constrained super-resolved spherical deconvolution (CSD) [27] with the following parameters: SH order $\ell_{\text{max}} = 8$ and diffusion profile of $[1.7, 0.3, 0.3] \times 10^{-3} , \text{mm}^2 / s$</td>
<td>none</td>
</tr>
<tr>
<td><strong>SDT</strong></td>
<td>one-shell b = 3000 60 samples</td>
<td>Sharping Deconvolution Transform [29] with the following parameters: SH order $\ell_{\text{max}} = 6$ and diffusion profile of $[1.7, 0.3, 0.3] \times 10^{-3} , \text{mm}^2 / s$</td>
<td>none</td>
</tr>
<tr>
<td><strong>STD</strong></td>
<td>one-shell b = 3000 64 samples</td>
<td>Spherical deconvolution via Symmetric Tensor Decomposition [67]</td>
<td>none</td>
</tr>
<tr>
<td><strong>SPIRAL</strong></td>
<td>spiral b = 1200 82 samples</td>
<td>Gradient directions were set on a spherical spiral curve resulting in voxel-wise diffusion signal periodicity. The periodic pattern of the signal in the gradient direction domain was exploited for de-noising and fiber orientation estimation</td>
<td>none</td>
</tr>
<tr>
<td><strong>DSI</strong></td>
<td>cartesian $b \leq 8000$ 257 samples</td>
<td>Diffusion Spectrum Imaging (DSI) [6] with the following parameters: zero-padded $35 \times 35 \times 35$ grid for the EAP reconstruction, Haning filtering before discrete Fourier transformation and ODF integration over the full radius range</td>
<td>none</td>
</tr>
<tr>
<td><strong>DSI_\text{LR}</strong></td>
<td>cartesian $b \leq 8000$ 257 samples</td>
<td>General framework based on Lucy-Richardson deconvolution to rectify the EAP reconstructed with DSI, taking into account the blurring introduced in the Fourier inversion due to the discrete and finite support of q-space measurements [68]</td>
<td>none</td>
</tr>
<tr>
<td><strong>DSI_\text{SWT}</strong></td>
<td>cartesian $b \leq 8000$ 257 samples</td>
<td>Multi-resolution approach based on 3D Stationary Wavelet Transform (3D-SWT) to de-noise the propagator reconstructed with DSI</td>
<td>none</td>
</tr>
<tr>
<td><strong>QBI_\text{Q}</strong></td>
<td>cartesian $b \leq 8000$ 257 samples</td>
<td>Extension of the Generalized Q-sampling Imaging [23] to analytically reconstruct the ODF with solid angle consideration. As it is expressed as a direct linear transform of the raw signal, it is faster to compute than DSI</td>
<td>none</td>
</tr>
</tbody>
</table>
The DSI-like class encompasses reconstruction methods based on q-space Cartesian sampling and maximum b-value of 8000 s/mm². The standard Diffusion Spectrum Imaging [6] (DSI) is studied in comparison to three additional methods that aim at improving its known weaknesses. First, the method DSI_{LR} [68] employs the Lucy-Richardson deconvolution to rectify the EAP reconstructed taking into account the blurring effect introduced in the Fourier inversion by standard DSI due to the discrete and finite support of q-space measurements. The second algorithm, DSI_{SQT}, is a multi-resolution approach based on the 3D Stationary Wavelet Transform (3D-SWT) whose aim is de-noising the propagator reconstructed with DSI to obtain sharper ODFs. At last, GQI$_2$ extends the Generalized Q-sampling Imaging [23] to analytically reconstruct the ODF with solid angle consideration. Furthermore, as GQI$_2$ is formulated as an efficient linear transform of the raw signal, it is sensibly faster than standard DSI to reconstruct the EAP and, consequently, the ODF.

In the last class, that we term SPARSE-like, we have evaluated six different dictionary-based reconstruction approaches inspired by the recent theory of compressed sensing [31], [32]. These methods are generally formulated as convex optimization problems and exploit various sorts of priors. NN-L2 [33] and L2-L2-L1 [21], for example, represent two extensions to already published works and the corresponding formulations are based on $\ell_2$ norm priors. On the other hand, the remaining four algorithms are all based on the $\ell_1$ norm for promoting sparsity. Notably, these approaches were proposed for the first time at the contest. L1-L1 is based on a Gaussian dictionary estimated directly from the acquired diffusion data and exploits the $\ell_1$ norm both in the data and the sparsity regularization terms; this $\ell_1-\ell_1$ formulation had never been used in local diffusion estimation before. L2-L1-DL uses the more common $\ell_2-\ell_1$ paradigm of the reconstruction problem [31], [32], but implements an efficient strategy to learn a parametric dictionary from a set of training data. Such estimated dictionary enables highly sparse representations of the diffusion signal and allows to analytically estimate important diffusion features, such as EAP and ODF, with a very limited number of measurements [39]. The last two approaches, L2-L1-TV [65] and L2-L1-TGV, both use the same $\ell_2-\ell_1$ problem formulation but they implement two different spatial regularization schemes based on Total Variation and Total Generalized Variation, respectively. It should be noted that in case of the IV dataset, for which no spatial coherence exists among voxels, these two algorithms do not make use of these spatial regularization schemes. In this scenario they can be considered two different implementations, but conceptually equivalent, of a reconstruction algorithm based on the classical $\ell_2-\ell_1$ paradigm and no spatial regularization, as for instance [34], [38]. Hence, we can assume that we have also evaluated in this comparison, for free, an additional reconstruction algorithm, which might be called L2-L1 for consistency, whose performances can always be assumed equivalent to those of L2-L1-TV and L2-L1-TGV on the IV phantom.

In contrast to many previous studies, the comparison provided in this work is as fair as possible as the majority of the reconstructions were tested using the original implementations provided by their own developers. We used in-house implementations for the additional 7 classical techniques we evaluated, with the exception of DTI and CSD whose reconstructions were computed using FSL [9] and MRTRIX [69], respectively.

As last remark, we point out a few notes concerning some algorithms and their corresponding results. As the method DTI_{neigh} needs to have access to the neighborhood of a voxel to work properly, only the reconstructions relative to the SF dataset were submitted. For this reason, this algorithm was evaluated only for this phantom whereas its performances on the IV dataset can be assumed, as a reference, as good as those of DTI. On the other hand, the method SPIRAL internally fixed a priori the number of directions to extract and it always reconstructed two fibers. As this inevitably biases all the corresponding evaluations, the reader should keep in mind this fact throughout the manuscript.

III. RESULTS AND DISCUSSION

The reader is recommended to keep the Table I at his fingertips while examining the following results. In fact, two methods might have obtained similar results but using radically different acquisitions schemes, which makes a direct comparison very difficult. To this aim, as already said, the algorithms have been grouped into four separate sampling classes and, to further simplify the reading, are listed in Table I and in all the plots with increasing number of samples.

A. Overall reconstruction quality

To provide the reader with a first impression of the algorithms with plausible fiber configurations likely to happen in real human brain data, we assessed the overall quality of the reconstructions in every voxel of the SF dataset in a scenario with a mild level of noise. Fig. 2 reports the summary of the two metrics SR and $\bar{\theta}$ corresponding to the simulations performed with a SNR=30. The robustness to noise of each algorithm will be investigated in detail later in the paper.

The first result to meet the eye is the expected inability of DTI to distinguish multiple fiber populations in a voxel, as clearly reflected in the lowest SR among all methods and high $\bar{\theta}$ values. Also the enhancement proposed by DTI_{neigh} did not seem capable of overcoming this intrinsic limitation of DTI. DTI_{neigh} can extrapolate additional information about the underlying fiber structure in a voxel (e.g. infer a crossing) by looking at adjacent voxels, and the best results are obtained when in the neighborhood there are single fiber profiles pointing at that voxel. Hence, the compact configuration of bundles present in the SF dataset might have limited its performance.

Despite the wide variety of sampling schemes employed, all remaining approaches were characterized by moderate results in the estimation of the number of fiber populations, with SR always between 50% and 70%, but with very good performances in terms of angular accuracy, with $\bar{\theta}$ always below 10°. At the extreme side of the sampling panorama, DSI-like methods showed reconstructions with the topmost quality (highest SR and lowest $\bar{\theta}$ values) and remarkably stable (fewer outliers, i.e. red dots). These approaches are
A. Number of fiber populations

Fig. 3 presents the detailed performance of each method with respect to the estimation of the number of fiber populations in a voxel. The average values of the three metrics $SR$, $n^+$ and $n^-$ are reported as a function of the crossing angle, where the angle between the two fiber populations was increased from $0^\circ$ to $90^\circ$. Results refer to the numerical simulations performed on the TV dataset with a SNR=30.

At a first glance, it is apparent that the main source of inaccuracy for all reconstruction methods is mainly due to under-estimation other than over-estimation. Over-estimated directions are rather mild, with $n^+$ always below 0.20–0.25, and happen mostly at high rather than low crossing angles. On average, fiber compartments started to be underestimated at $60^\circ$ and a plateau was reached at about $30^\circ$. Above and below these thresholds all algorithms performed essentially the same. Surely DTI always under-estimates one of the two fiber populations and $n^-$ is always 1. In the mid-range of crossing angles, DSI-like approaches behaved very well ($52\% < SR < 85\%$), while the algorithms in the HARDI-like class, especially D0T and all those based on Q-Ball Imaging, appeared to suffer more ($SR \approx 25\%–30\%$) and the effects of under-estimation started at higher crossing angles. Concerning spherical deconvolution methods, we can see that results are quite heterogeneous. CSD was found to be the method suffering the most from under-estimation in the HARDI-like class ($SR \approx 20\%$), while SDT appeared moderately more robust ($SR \approx 45\%$). STD, on the contrary, performed very well and exhibited the best success rate among all methods ($SR \approx 88\%$), which is linked to a remarkably low rate of under-estimation ($n^- \approx 0.05$). This method implemented an iterative heuristic to accurately assess the number of compartments in each voxel: a configuration with one fiber population was first reconstructed, then with two etc, and finally the configuration best fitting the data was retained. This procedure, in combination with spherical deconvolution, appeared particularly effective and was able to reliably resolve two fiber populations down to a crossing angle of $30^\circ–35^\circ$.

B. Number of fiber populations

Fig. 3 presents the detailed performance of each method with respect to the estimation of the number of fiber populations in a voxel. The average values of the three metrics $SR$, $n^+$ and $n^-$ are reported as a function of the crossing angle, where the angle between the two fiber populations was increased from $0^\circ$ to $90^\circ$. Results refer to the numerical simulations performed on the TV dataset with a SNR=30.

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characterized by a very dense acquisition of the signal (257 samples) and the price to pay is a very long scan time which is normally not suitable in a clinical environment.

A sensible reduction in the number of samples (64 to 82) is possible with HARDI-like techniques. However, the reconstructions looked very unstable and presented many outliers which inevitably deteriorated the average performance. The algorithm SPIRAL showed particularly poor scores, but these are compatible with the previous remark on the number of estimated orientations fixed a priori. As an exception the STD method, which is based on symmetric tensor decomposition, obtained very interesting performances rivaling with those of DSI-like, but using only a quarter of the samples. In particular, these high performances seem to be driven especially by its superior ability to correctly identify the right number of fiber populations in each voxel, as reflected by the highest success rate among all methods ($SR \approx 70\%$).

SPARSE-like methods not only made it possible to further reduce the number of samples (15 to 64) but, interestingly, these algorithms exposed reconstructions with quality comparable to those of DSI-like approaches. In this case, however, despite the fact that $SR$ values were in line with all other methods, the orientation of the fiber populations turned out to be as accurate as more demanding and complex techniques. Considering that SPARSE-like methods only require between 6% and 25% of the number of samples needed by DSI-like ones, these results can be considered a remarkable achievement which opens the way to attain the quality of HARDI/DSI acquisitions at the price of DTI.

For sake of completeness, during the contest we had also compared the reconstructions with respect to the accuracy in the estimation of the ODF. However, not all the algorithms directly recover an ODF; Fig. 5 shows some representative diffusion profiles estimated by each reconstruction method. Consequently, we decided not to report the results of this analysis in the manuscript as this evaluation did not add any informative value to the comparison. The interested reader can find more information on the challenge website.
with $SR \approx 90\%$ at $35^\circ$.

In light of the significant under-sampling rates they permit, SPARSE-like methods exhibited impressive performances. Overall, all algorithms performed at least as good as HARDI-like approaches, with the scores of NN-L2, L1-L1 and L2-L1-TGV competing even with those of DSI ($SR \approx 65-70\%$). Remarkably, L1-L1 and L2-L1-TGV showed the lowest number of under-estimated compartments at low crossing angles, even better than DSI-like methods. L1-L1 appeared to have some difficulties at large crossing angles, where it showed a higher number of over-estimated compartments. This behavior is most probably linked to the $\ell_1$ norm used as goodness measure for the data term in the optimization procedure. Overall, though, these methods performed as good as DSI-like ones but, and it is important to emphasize this aspect, they required only 9%-25% of the samples.

Interesting observations can be made by repeating the previous analysis and considering a tolerance cone around each ground-truth fiber population, i.e. $SR_\perp$, $n^+_\perp$ and $n^-_\perp$ metrics. Comparing Fig. 4 with Fig. 3 we confirm what previously noted, in that differences among algorithms are restricted only to the mid-range of crossing angles, see $SR_\perp$ plot in $30^\circ$–$60^\circ$, and that the main source of inaccuracy is caused by under-estimation ($n^-_\perp$ plot). However, in this case both $n^+_\perp$ and $n^-_\perp$ scores reach much higher values than before, respectively up to 0.7 and 1.7. As we spotted that the tendency is to under-estimate and $n^-$ is at worst 1, this means that the single estimated direction must be outside both tolerance cones around the true fiber orientations: this behavior is in fact captured both as missing either of them (increased $n^-_\perp$ values) and also as recovering a spurious one (increased $n^+_\perp$ values). DTI is the archetype of this situation: it obtained in fact the highest $n^+_\perp$ score, almost 1 for crossing angles above $50^\circ$, which is clearly counter-intuitive for the nature of the
technique, but in accordance with the estimation of one single direction which lies somewhere between the two real ones. As before, the method STD achieved superb performances, with $SR_\angle$ always above 90% for all crossing angles. Also the performances of SPIRAL seemed almost independent from the crossing angle but, in comparison, its scores were rather modest ($SR_\angle \approx 70\%$), once more probably caused by the hard-constraint set internally. Curiously, L1-L1 and L2-L1-TGV turned out to have the lowest $SR_\angle$ scores in the crossing range $0^\circ$–$30^\circ$ even though, in the same range, they showed the highest $SR$, which was mainly driven by a reduced rate of under-estimation, $n^-$, and no over-estimation, $n^+$. However, both algorithms showed also a higher $n^\pm$ rate, $\approx 0.2$, which is an indication that one of the two directions estimated is not close to the corresponding true one and is considered spurious.

We conclude this section by noting two drawbacks of this latter set of metrics, i.e. $SR_\angle$, $n^+\angle$ and $n^-\angle$. First, for configurations with crossing angle less than $40^\circ$ these indices become ambiguous, as in this case the tolerance cones of the two fiber populations unavoidably overlap. However, above $40^\circ$ they essentially describe the same trends. Second, they are very dependent on the used threshold, $20^\circ$ in this study. For these reasons and for compactness of the exposition, we decided to report in the remaining only the measures without tolerance cone, i.e. $SR$, $n^+$ and $n^-$, as they appeared to us more intuitive and sufficiently informative for the scope of this work. All evaluations are available online.

C. Angular accuracy

Fig. 6 compares all reconstruction methods with respect to the accuracy in the orientation of the estimated fiber populations, $\tilde{\theta}$, as a function of the crossing angle. Results refer to the simulations performed with a SNR=30.

DTI is easily recognizable. The fact that $\tilde{\theta}$ is approximately
Fig. 5.  Estimated diffusion features. One representative diffusion profile as reconstructed by each algorithm (e.g. ODF or FOD) is shown for four different crossing configurations ($90^\circ$, $60^\circ$, $45^\circ$, $30^\circ$). Results correspond to the reconstructions on the IV dataset with a SNR=30. As these data were not available for method DTI$_{neigh}$, the corresponding profiles are not shown.
half of the angle separating the two ground-truth directions, together with the observations in the previous paragraph, is compatible with the reconstruction of a single orientation lying in the middle of them and, in practice, it identifies a boundary under which the technique is unable to resolve multiple directions. This limit was reached by all the methods as the crossing angle between the two fiber populations decreased: it happened at 48° for the CSD algorithm while the limit was 25° in the case of STD and DSI_Lk.

Despite quite heterogeneous results obtained concerning the estimation of the number of fiber populations, these variations translated only to fairly small differences between methods with regard to the orientation of the estimated directions. All tested algorithms exposed very high angular accuracy both at low and high crossing angles ($\bar{\theta} \approx 3°–7°$), while in the mid-range the performances quickly deteriorated. The largest difference between algorithms was observed at a crossing angle of 48°, where $\bar{\theta}$ varied between 4° in case of both STD and DSI_Lk and 21° for CSD. The observed tendency of the L1-L1 algorithm to over-estimate compartments at high crossing angles had quite a severe impact on the angular accuracy of the corresponding reconstructions ($\bar{\theta} \approx 13°$), three times higher than the average angular error of all the other algorithms ($\bar{\theta} \approx 4°$). At low crossing angles, on the other hand, the $\bar{\theta}$ values for both L1-L1 and L2-L1-TGV were higher than the average angular error of all other methods, even thought they previously had shown the best under-estimation behavior ($n^- \approx 0.8$) and over-estimation was almost absent ($n^+ \approx 0$). Inspecting the corresponding reconstructions we could observe that both algorithms normally recovered one fiber lying halfway the two true peaks like all others algorithms but, additionally, they reconstructed very often a spurious fiber direction (captured in $n^+$ plots). This extra compartment explains both the reduced under-estimation rate exposed by these algorithms and, as it was normally orientated randomly, the drop in the average angular accuracy of the reconstructions.

Special reference needs to be made about the algorithm SPIRAL, as the angular error appeared to be in the average range but almost independent from the crossing angle. The reasons for this trend are not clear, but it might be a side effect of the procedure used to recover their orientations. The authors used in fact the 82 sampling directions both to represent the signal and estimate the fiber directions, and thus the granularity of this grid (the points are separated by about 20°) might have hampered the performances of their method.

D. Robustness to noise

Fig. 7 compares the algorithms with respect to the sensitivity to noise of the corresponding reconstructions, reported in the plots as the variation of the three metrics $\bar{\theta}$, $n^+$ and $n^-$ as the acquisition SNR decreases. The performances obtained at a SNR=30 are taken as reference and the variation of the scores is reported for SNR=20 and SNR=10, which resemble more closely the typical SNR in real dMRI data. Positive values mean a degradation of the performances.

As expected, the angular resolution of the reconstructions gradually deteriorated as the noise increased, with no clear differences among the algorithms. Yet, methods with higher number of samples seemed to be in general more stable. DTI and SPIRAL appeared the most robust to noise ($\Delta_{\bar{\theta}} < 4°$) while DSI the most susceptible, especially at SNR=10 where it degraded abruptly showing a $\Delta_{\bar{\theta}} \approx 21°$. However, it is worth noting how the enhancements implemented in DSI_Lk and DSI_SWT allowed a significant boost in the robustness to noise in methods based on Cartesian signal sampling, turning standard DSI from the most susceptible to noise method to the most robust. Also CSD showed very stable performances, appearing to suffer only at high crossing angles, despite no significant worsening in under- and over-estimation rates.

In general, all methods showed the highest susceptibility to noise and, correspondingly, the major drop in the performances at medium-high crossing angles (boxes and upward triangles), with drops in the angular accuracy $\Delta_{\bar{\theta}}$ of the reconstructions up to 8°–11°. In particular, this degradation seemed driven mostly by over-estimation, as easily visible in the $\Delta_{n^+}$ plot. This observation is in accordance with the tendency of all methods, previously observed in Fig. 3, to recover spurious compartments at high, more than at low, crossing angles. On the other hand, high noise levels induced an opposite behavior in algorithms using highly under-sampled acquisition schemes, like L2-L1-DL, L1-L1 and L2-L1-TV, with less than 30 samples. In fact, these methods did not show an increased rate of over-estimated compartments but, on the contrary, an higher number
Fig. 7. Overall performances as a function of the noise level. For each method, the variations of the estimated number, \( n^+ \) and \( n^- \), and orientation, \( \theta \), of the fiber populations are reported as the noise level changes. The first mark of every bar corresponds to the difference between the scores at SNR=30 and those at SNR=20, while the second mark between SNR=30 and SNR=10. Results refer to the simulations performed on the IV datasets and, as usual, are reported separately for three crossing angle ranges: 0\(^\circ\)–30\(^\circ\) (downward triangles), 31\(^\circ\)–60\(^\circ\) (squares) and 61\(^\circ\)–90\(^\circ\) (upward triangles).

The method STD showed the same tendency to under-estimate more with higher noise levels and it resulted the most prone to under-estimation at SNR=10. Interestingly, the ability of STD to under-estimate less than all other algorithms in the medium range of crossing angles was the reason behind its exquisite performances (\( n^- \) plot in Fig. 3). STD performed very well with low noise, but unfortunately this robustness to missing fibers rapidly degraded as the noise increased. Curiously, the \( \Delta n^- \) plot in Fig. 7 shows that most of the algorithms got some benefit (\( \Delta n^- < 0 \)) from higher noise levels at low-medium crossing angles (boxes and downward triangles), with DSI being the most “beneficial”. However, this strange behavior is a side-effect of over-estimation. In fact, at SNR=30, all methods were shown to suffer heavily from under-estimation for crossing angles below 60\(^\circ\) (see \( n^- \) plot in Fig. 3). Higher noise levels caused an increased rate of spurious fiber compartments recovered, which not only were clearly captured by the positive values in the \( \Delta n^+ \) plot, but also by negative \( \Delta n^- \) values.

The observed tendency to recover spurious compartments as the noise increases can be explained, in part, by the fact that all the algorithms considered in this work are based on the assumption that the distribution of the noise follows a Gaussian distribution with zero mean. However, the noise corrupting the data follows a Rician distribution and the deviation from Gaussianity is particularly evident at low SNR values [61]. Rician noise produces, on average, a net increase in the signal which is different along every gradient direction. As a result, reconstructions assuming zero-mean Gaussian noise tend to produce small spurious fibers to explain this discrepancy.

As last remark, the results obtained in our study are strictly valid only for single-channel or SENSE multi-channel acquisitions [70] and they should not be extrapolated to other parallel techniques such as GRAPPA [71], where the noise follows a noncentral-\( \chi \) distribution. Please note however that nearly all reconstruction methods proposed to date were tested only on Rician-corrupted data, i.e. using the same assumption, and thus our results are consistent with previous literature.

Fig. 8. Effects of using information from neighboring voxels on the quality of reconstructions. For each method, the distribution of the average angular errors \( \bar{\theta} \) obtained on the IV phantom (left bar) are compared with those corresponding to SF (right bar). Results are reported for two noise levels: SNR=30 (top) and SNR=10 (bottom).

E. Spatial neighborhood

Fig. 8 investigates the effect of using additional information from neighboring voxels on the quality of reconstructions. For each method, we compared the distributions of the average angular errors \( \bar{\theta} \) obtained on the IV (left bar) and SF (right bar) phantoms. To be sensitive to this effect only, we considered a
subset of voxels with two fiber populations from both datasets and we matched the distributions of the crossing angles. We compared the results corresponding to low and high noise levels, respectively with SNR=30 (top) and SNR=10 (bottom).

The algorithms reported in Table I to exploit additional information from neighboring voxels in some form, either performing spatial smoothing on the DWI images or employing a regularization scheme, are clearly identifiable. For these methods, in fact, the angular accuracy of the reconstructions clearly benefitted (smaller " errors) from the spatial coherence among voxels present in the SF phantom (right bar in each method’s results) if compared to the results obtained on the IV dataset (left bar), which indeed consists of un-correlated voxels. These de-noising approaches appeared definitely more effective on data corrupted with higher noise levels, as apparent in the plot corresponding to SNR=10, where improvements in the angular accuracy up to 7◦–8◦ can be appreciated. On the other hand, with high quality data the regularization-based approaches appeared more effective than those using spatial smoothing. In this case, in fact, these latter did not seem to give any benefit (see performances of L2-L2 and MN-L2 at SNR=30) while methods using spatial regularization schemes, i.e. L2-L1-TV and L2-L1-TGV, still provided a 3◦–4◦ improvement in the average angular accuracy.

Surprisingly, even though we did not expect any significant variation for those methods not exploiting any spatial coherence in the data, a curious behavior can still be observed in the figure. In fact, some of the approaches in the HARDI-like class showed a performance reduction when recovering the intra-voxels structure in the SF dataset, which is sort of counter-intuitive. One possible speculation for this odd behavior can be attributable to different settings for the parameters used in the two datasets. As the structure of the SF dataset did include voxels with more than two crossing fibers (even though these configurations were excluded from the evaluation), more tolerant thresholds have been probably used in this case.

F. Signal-to-noise ratio and voxel-size

Because of the assumption on $S_0$ we made in section II-B, in our experiments we have implicitly considered a constant echo-time (TE) for all acquisition protocols, hence neglecting the additional dependence of the signal on the damping factor $\exp(-TE/T2)$, where $T2$ is the spin-spin relaxation time of the tissue. However, in clinical systems, increasing the b-value inevitably implies a longer TE which, in turn, negatively impacts the effective SNR of the acquisitions. To obtain images with adequate SNR, thus, clinical protocols typically adapt the spatial resolution and use a rather large voxel-size for high b-value sequences such as DSI, e.g. $2.5 \times 2.5 \times 2.5$ mm$^3$, whereas DTI is normally acquired with $1 \times 1 \times 1$ mm$^3$ resolution.

Fig. 9 shows a further comparison between the reconstructions performed on the original data and those repeated after adjusting the noise level to account for the specific TE of each sequence. For compactness the results are reported only for the most popular techniques, such as DTI, CSD and DSI. The new adjusted SNRs reported in Table II have been computed taking the protocol with the longest TE as reference and assuming the same spatial resolution, yielding reasonable values commonly observed in real datasets. Realistic TE values have been extracted from a clinical 3T scanner with maximum gradient strength of 40 mT/m, and the T2 in the white matter was assumed equal to 80 ms as found in [72].

<table>
<thead>
<tr>
<th>b-value (mm$^2$/s)</th>
<th>SF</th>
<th>IV</th>
</tr>
</thead>
<tbody>
<tr>
<td>80</td>
<td>83.6</td>
<td>79.5</td>
</tr>
<tr>
<td>84</td>
<td>76.6</td>
<td>72.9</td>
</tr>
<tr>
<td>87</td>
<td>66.8</td>
<td>59.7</td>
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<td>91</td>
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<td>107</td>
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<td>114</td>
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<tr>
<td>118</td>
<td>50.0</td>
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</tr>
<tr>
<td>162</td>
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</table>

The performances of DSI and DSI$_LR$ are clearly constant as their SNR remained fixed. Also the success rate of DTI did not change, but this is due to the intrinsic limitation of the model. As previously reported in Fig. 7, all methods can provide better reconstructions when working on images with higher quality, but they are characterized by different levels of robustness to the noise. At a noise level typically observed in real acquisitions (upper plots), in fact, no significant improvement can be noted for any of the methods when increasing the effective SNR to account for shorter TE, both in terms of SR and . With very low image quality (bottom plots) the methods with shorter TE ameliorate the reconstructions, even though clearly the same trend of performances is preserved across methods. Hence, for the scope of this study, the effect of each sequence echo-time on the effective SNR of the data did not seem to introduce significant bias in the results.
G. Dependence on the signal model

During the contest some concerns arose with respect to the Gaussian Mixture Model (1) that was used to simulate the dMRI signal in place of the more precise expression for particles diffusing in a restricted cylindrical geometry given in [58]. Both models have been largely used in the literature for validation of diffusion estimators, as well as the use of physical phantom. As already pointed out in the introduction, all these models are undoubtedly simplifications of the actual biophysical processes occurring in white matter. In fact, for a more accurate synthetic simulation, it should be taken into account the complex axonal structure, the membrane permeability, the bi-exponential behavior of the signal with respect to the b-value magnitude, the existence of slow and fast pools etc. In the particular case of the contest, however, the main concern was that our choice might have favored the model-based reconstruction algorithms, such as those in the SPARSE-like class, using implicitly or explicitly the same Gaussian assumption.

To investigate any possible influence of this choice on the reconstructions considered in our study, we simulated the dMRI signal attenuation corresponding to a single fiber configuration using both expressions: (i) Gaussian Mixture Model [8] and (ii) Soderman’s Model [58]. Fig. 10 compares the signal decay as predicted by the two models, named MultiTensor and Soderman in the plot, as a function of the b-value. For the latter, we used the following parameters as reported in previous studies [20], [34]: \( L = 5 \text{ mm}\), \( \rho = 5 \mu \text{m}\), \( D_0 = 2.02 \times 10^{-3} \text{ mm}^2/\text{s}\), but we set \( \Delta = 80 \text{ ms}\) and \( \delta = 35 \text{ ms}\) to resemble an acquisition with maximum b-value of 4000 s/mm$^2$ on a clinical 3T scanner with a maximal gradient strength of 40 mT/m [68]. To simulate the signal with the model (1) and match the diffusion pattern, a single tensor was fitted to the data and the estimated parameters were used for the simulation. The diffusivities along the three main axes were found to be: \( \lambda_1 = 2.3 \times 10^{-3} \text{ mm}^2/\text{s}\) and \( \lambda_{2,3} = 0.1 \times 10^{-3} \text{ mm}^2/\text{s}\).

![Fig. 10. dMRI signal attenuation of a single fiber configuration simulated using two different models. The decays predicted by the MultiTensor [8] (dashed lines) and the Soderman [58] (solid lines) models are plotted as a function of the b-value. Results are reported at four different angles between the diffusion gradient and the fiber population (0°, 30°, 60°, 90°).](image)

As we can see, the signal decays predicted by the two models, dashed lines for MultiTensor and solid for Soderman, appear almost indistinguishable for b-values up to 3000 s/mm$^2$, with differences becoming appreciable (note the logarithmic scale) above 4000 s/mm$^2$, especially when the diffusion gradient is parallel to the orientation of the fiber (red curves). Interestingly, 16 of the 20 algorithms evaluated in this study employed acquisition schemes with a maximum b-value of about 3000 s/mm$^2$. Furthermore, the remaining 4 methods belonged to the DSI-like class and were all characterized by model-free approaches for the reconstruction of the intra-voxel fiber configuration. In light of these arguments, we genuinely believe that for the scope of this study the choice of the model (1) that we used in our numerical simulations for the generation of the diffusion signal did not favor those model-based approaches which made use of the same Gaussian assumption and, for this reason, did not bias the comparison.

H. General observations

It is always difficult to draw general conclusions from comparative studies without going into the details of every single algorithm evaluated. This is particularly true in this work as the evaluated approaches differ in a great deal of aspects including acquisition scheme, assumptions made in the modeling (i.e. model-free and model-based approaches were included), numerical algorithms employed in the estimation process, descriptors used to characterize the intra-voxel structure (i.e. diffusion ODF or FOD), as well as some heuristic parameters such as the threshold used to select the main ODF local maxima and the regularization parameters. For this reason, during the “HARDI reconstruction challenge” we decided not to provide a final ranking based on scores to highlight a winner. This decision was mainly motivated due to the lack of a fair quantitative model to integrate all the different aspects involving each method.

In contrast, in order to help clinicians in the choice of the most adequate technique for a specific application according to their interests and possibilities, we summarize here the main lessons we have learned from this study:

- None of the methods was found to outperform the others in every experimental condition. On the contrary, results suggest that a combination of techniques might be the way to obtain optimal reconstructions.
- Overall, DSI$_{LR}$ appeared the most accurate and stable (i.e. fewer outliers), but the long scan time highly restricts its applicability in clinical environments.
- Accurate reconstructions (comparable to those obtained from HARDI/DSI) at the price of DTI acquisitions are becoming possible thanks to SPARSE-like approaches, which in general showed very interesting performances.
- With high quality data the main source of inaccuracy for all the algorithms seemed to be caused by under-estimation. On the other hand, for low/medium SNR acquisitions the degradation of the performances was mainly driven by over-estimation.
- Noise removal procedures, either in the form of image denoising or spatial regularization, are highly recommended as results showed they can improve the performances up to 30–40%. We did not observe any substantial difference in the efficacy of the two approaches; nevertheless, spatial regularization appeared to be effective also in low noise regimes, while merely denoising the images did not.
It is important to remark that the results presented in this work are based on numerical simulations and, as previously mentioned, there is no guarantee that these observations hold true to the same extent also in case of real data.

IV. CONCLUSION

We have quantitatively evaluated and compared a large selection of algorithms for recovering the local intra-voxel fiber structure from diffusion MRI acquisitions. Considered methods included a mixture of classical and well-established techniques as well as novel approaches recently proposed in the literature, in particular those based on compressed sensing theory. Despite this study has been conducted on synthetic dMRI data, results yet allowed to spot strengths and weaknesses of each approach and to draw interesting considerations common to all the algorithms. We believe that this information might be used both to improve current reconstruction methods and also as a guidance for physicians to identify the most adequate technique for their clinical studies.

ACKNOWLEDGEMENTS

This work benefited from the precious help of all the contest participants, who provided not only the reconstructions that were evaluated but also assistance and valuable comments during the writing of the present manuscript.

REFERENCES


