Statistical significance in DTI group analyses: How the choice of the estimator can inflate effect sizes

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Abstract

Diffusion magnetic resonance imaging (dMRI) is one of the most prevalent methods to investigate the micro- and macrostructure of the human brain in vivo. Prior to any group analysis, dMRI data are generally processed to alleviate adverse effects of known artefacts such as signal drift, data noise and outliers, subject motion, and geometric distortions. These dMRI data processing steps are often combined in automated pipelines, such as the one of the Human Connectome Project (HCP). While improving the performance of processing tools has clearly shown its benefits at each individual step along the pipeline, it remains unclear whether – and to what degree – choices for specific user-defined parameter settings can affect the final outcome of group analyses. In this work, we demonstrate how making such a choice for a particular processing step of the pipeline drives the final outcome of a group study. More specifically, we performed a dMRI group analysis on gender using HCP data sets and compared the results obtained with two diffusion tensor imaging estimation methods: the widely used ordinary linear least squares (OLLS) and the more reliable iterative weighted linear least squares (IWLLS). Our results show that the effect sizes for group analyses are significantly smaller with IWLLS than with OLLS. While previous literature has demonstrated higher estimation reliability with IWLLS than with OLLS using simulations, this work now also shows how OLLS can produce a larger number of false positives than IWLLS in a typical group study. We therefore highly recommend using the IWLLS method. By raising awareness of how the choice of estimator can artificially inflate effect size and thus alter the final outcome, this work may contribute to improvement of the reliability and validity of dMRI group studies.

1 Introduction

Diffusion magnetic resonance imaging (dMRI) has been used extensively to study fundamental biological concepts (Assaf et al., 2019; Novikov et al., 2019), pathologies of the brain (Cercignani and Gandini Wheeler-Kingshott, 2019; Lunven et al., 2015; Phillips et al., 2016; Sabia et al., 2017), and the architectural configuration of white matter (WM) tracts (Catani et al., 2013; David et al., 2019; Thiebaut de Schotten et al., 2012). As dMRI became more commonly used, there was a need to improve its reliability for clinical applications (Eierud et al., 2014; Nir et al., 2013; Owen et al., 2013; Rudie et al., 2013; Schwarz et al., 2013). Methodological developments that contributed to this improvement are related to cardiac gating (Chang et al., 2005; Kozák et al., 2013), high-field MRI scanners (Moser et al., 2017), stronger and faster switching MR gradients (McNab et al., 2013; Setsompop et al., 2013), image reconstruction techniques (Lustig et al., 2007), diffusion model
Effect of DTI estimator on group study outcome

Processing tools are the key contributors in minimizing adverse effects of confounding factors on the final results. Despite the theoretical benefits of integrating novel methodological developments in the dMRI processing pipeline, there is no consensus on which settings or algorithms should be preferred for, for instance, a typical diffusion tensor imaging (DTI) study in which two groups of subjects (e.g., healthy controls vs. patients) are compared. This lack of agreement is reinforced by our limited understanding of whether a specific processing method has a significant contribution to the reliability of the subsequent group analysis in terms of outcome. In this context, one could state that, in practice, the added benefit of a particular data correction procedure is nullified if there are other data aspects with a much higher variability. As an example, the decrease in diffusion parameter estimation bias due to Gibbs ringing correction may be completely swamped by the high noise levels in low-SNR dMRI data, obviating the relevance of performing this processing step.

In general, the relative improvement of one processing step not only depends on the intrinsic quality of the data, but also on the performance of the other processing steps used in the dMRI pipeline. Correcting spatial misalignment across multiple diffusion-weighted images (DWIs) due to subject motion, for instance, may benefit from preceding denoising of these images. In addition, after the data has been corrected for artifacts, strategies to further analyze the data (e.g., using fiber tractography, histograms, ROIs, voxel-based approaches, or network graphs) may have a difference in sensitivity to the benefit of some of the individual processing steps and potentially generate differences in the final outcome of a group study.

While many steps in a dMRI processing pipeline can be considered as optional, for several diffusion approaches such as DTI or diffusion kurtosis imaging (DKI), there is the mandatory step of choosing the diffusion estimation method to obtain model parameters. Over the last decade, a plethora of such estimators have been used, including ordinary linear least squares (OLLS), non-linear least squares (NLLS), weighted linear least squares (WLLS), and their constrained, robust and conditional extensions, among others (Andersson, 2008; Chang et al., 2012, 2005; Collier et al., 2015; Jones and
Effect of DTI estimator on group study outcome

Basser, 2004; Koay et al., 2009; Kristoffersen, 2012, 2007; Salvador et al., 2005; Tax et al., 2015; Veraart et al., 2013b, 2011). Assuming that data outliers have been identified and removed, a specific version of the WLLS, iterative WLLS (IWLLS), shows high performance characteristics in terms of accuracy and precision and may even be preferred over advanced NLLS estimation methods (Veraart et al., 2013b). Yet, OLLS is still the most widely used estimation method and often defined as the default in common software tools (e.g., FSL – (Jenkinson et al., 2012)).

Similar to the other dMRI processing steps, one can also question the relevance of choosing a particular diffusion estimation approach. Does it really matter which estimator is used for the final outcome of a group study? In this work, we address this concern. More specifically, we performed a dMRI group analysis using Human Connectome Project (HCP) data sets and compared the results obtained with OLLS and IWLLS. To this end, and without loss of generality, we investigated gender related differences (Caeyenberghs and Leemans, 2014; Herting et al., 2012; Hsu et al., 2008; Ingalhalikar et al., 2014; Kanaan et al., 2012; Menzler et al., 2011; Núñez et al., 2017; Tyan et al., 2017; Westerhausen et al., 2003; Wierenga et al., 2017) to evaluate the potential differences in final outcomes using the two estimators. Preliminary results of this work were presented at the International Society for Magnetic Resonance in Medicine (ISMRM) meeting in Toronto, Canada (David et al., 2015).

2 Methods

2.1 Subject data and processing

Minimally preprocessed DWIs were collected from the HCP S500 release (Essen et al., 2012; Glasser et al., 2013). Briefly, the data consist six separate acquisitions of 90 DWIs acquired with diffusion weightings (b-values) equal to 1000/2000/3000 s/mm² and five, six or seven non-DWIs (b-value = 0 s/mm²). Every image was acquired with both left-to-right and right-to-left phase encoding directions; the voxel size was 1.25 mm isotropic. Susceptibility artifacts, eddy current induced distortions, and subject motion were corrected with the FSL tools taking into account any reorientations of the diffusion gradient orientations (Andersson et al., 2003; Jenkinson et al., 2012; Leemans and Jones, 2009; Sotiropoulos et al., 2013). All datasets were further processed with ExploreDTI version 4.8.6. (Leemans et al., 2009) using two different tensor estimation approaches: (a) OLLS (Basser et al., 1994) and (b) IWLLS (Veraart et al., 2013b). For this step, only the 90 DWIs with b-value of 1000 s/mm² and 9 non-DWIs per participant were selected for diffusion tensor estimation. In addition, we
also corrected for the gradient nonlinearities in the diffusion-weighted gradients during this estimation procedure (Bammer et al., 2003; Mesri et al., 2019; Sotiropoulos et al., 2013). Every participant for which all the 90 $b = 1000$ s/mm$^2$ images were available, and which was not listed among the participants with known anatomical anomalies or data quality issues, was included in the analysis. The complete list of the excluded participants can be found on the appropriate HCP wiki page (HCP, 2017). The final sample size is 409 participants, consisting of 244 females and 165 males.

### 2.2 Voxel-based analysis

For each subject, fractional anisotropy (FA) maps were calculated from the fitted tensors (using OLLS and IWLLS) and transformed to the Montreal Neurological Institute (MNI) template via the native-to-MNI warp files, provided by the HCP team (Fonov et al., 2011). Voxelwise statistical comparisons of FA between the male and female groups were performed using the permutation analysis of linear models (PALM) (Holmes et al., 1996; Nichols and Holmes, 2003; Winkler et al., 2014), a Matlab based open-source toolbox, version alpha104 with 10000 permutations. For all the tests (next section), calculations are based on nonparametric permutations as this approach was proven to be more efficient in producing fewer false positives than parametric methods (Eklund et al., 2016). Significance was determined at $p_{corr} < 0.05$ using family-wise error rate (FWER) adjustment to correct for multiple comparisons after applying threshold-free cluster enhancement (TFCE) (Smith and Nichols, 2009). Calculation speed was accelerated using the tail approximation (Winkler et al., 2016). A Dell server with 72 Intel Xeon E7-8870 v3 @ 2.10 GHz dual cores with 1 TB RAM was used for calculations.

### 2.3 Statistical tests

#### 2.3.1 Effect of tensor estimator

For each participant, there are two FA maps: one obtained from the diffusion tensor estimated with OLLS and one with IWLLS. In order to investigate the potential differences in FA (regardless of gender) between the OLLS and IWLLS pipelines, we used a paired two-sample t-test. This procedure tests whether there is a significant effect of using a different tensor estimation method on FA, without considering if the participant is female or male.
2.3.2 Effect of Gender

Differences in FA values between males and females (denoted as $FA_m$ and $FA_f$) were investigated using an unpaired two-sample t-test for the OLLS and IWLLS pipelines separately. A further correction was applied via the “-corrcon” option in PALM, which accounts for the multiple contrasts during the FWER correction.

2.3.3 Pipeline dependent gender differences

To test whether gender differences depend on the tensor estimation method, we performed a two-sample t-test on the gender, where the tested variable is the difference in FA, denoted as $\Delta FA$, between the IWLLS and OLLS pipelines:

$$\Delta FA = FA_{IWLLS} - FA_{OLLS}. \quad (1)$$

More specifically, we evaluated with this test whether the $\Delta FA$ values for males, denoted as $\Delta FA_m$, differ significantly from the $\Delta FA$ values for females, denoted as $\Delta FA_f$. Statistically, this procedure is the same as the interaction part of a two-group analysis of variance (ANOVA) test with two levels per participant. A significant effect means that the gender differences are solely driven by the choice of estimation method. Independent and symmetric errors were assumed to boost the statistical power of the test, by using the command “-ise” in PALM. Effect sizes and their distributions were analyzed in detail within the regions of significance.

2.3.4 Effect size

The practical significance of the findings was further evaluated by reporting effect sizes, as suggested by the American Statistical Association’s (ASA) recent statement on p-values: “A p-value, or statistical significance, does not measure the size of an effect or the importance of a result.” (Wasserstein and Lazar, 2016). Accordingly, we used Cohen’s $d$, a frequently applied effect size estimator. Furthermore, because Cohen’s $d$ is not a robust effect size measure to outliers, skewness, heavy-tails and the combinations of these factors, the shape differences between the voxelwise distributions of FA values were studied via the shift function (Rousselet et al., 2017). The 95% percentile confidence intervals for the decile differences were estimated with a bootstrap estimation (1000 samples), using the Harrell–Davis estimator (Wilcox, 2012), as implemented in the Matlab Robust Graphical Methods For Group Comparisons (matrogme) toolbox, version 0.0.9000 (Rousselet et al., 2017).
3 Results

3.1 Effect of tensor estimator

Fig. 1 shows the result for the paired t-test that investigates the difference in FA between the OLLS and IWLLS estimation methods. To further emphasize the differences, we show the effect size (with Cohen’s $d$) only for the voxels that were statistically significant after applying the multiple comparison correction procedure. The map shows that these differences are significant in the whole brain and are tissue-dependent. Larger effect sizes were revealed in the core WM, such as in the corpus callosum (CC), the corticospinal tract (CST), and the optic radiation (OR), where FA values are relatively high. Areas with lower FA values near the cortical and deep GM regions (thalamus, hippocampus, putamen, etc.) resulted in no or negligible differences, as expressed by the white areas in the image that indicate a near zero effect size. Overall, the IWLLS estimator results in significantly higher FA values in the vast majority of the WM compared to using OLLS.

The systematic deviation in FA between OLLS and IWLLS is further highlighted in Fig. 2, where the FA values are averaged across all 409 subjects. It is clear that for most of the WM voxels ($\sim$FA>0.2) the mean FA values are higher for the IWLLS estimator than for the OLLS estimator.
Fig. 1 Effect sizes (defined as Cohen’s $d$) are shown as color maps overlaid on regions with statistically significant differences in FA between using the IWLLS and OLLS estimators, presented in MNI space. Notice the different color scale magnitudes for the effect sizes. The reddish and blueish color bars reflect regions where $\Delta FA > 0$ and $\Delta FA < 0$, respectively (see Eq. 1). (Radiological view: left on the image is right in the brain and vice versa).
Fig. 2 Scatterplot of the ratios of the FA values from the IWLLS and OLLS estimators as a function of FA from the IWLLS estimator. Each point in the scatterplot represents the average FA value across all 409 subjects for each brain voxel in MNI space. If there was no systematic deviation between the OLLS and IWLLS estimators, the points should be located around the unity value, indicated by the red dashed line.
3.2 Effect of gender

Fig. 3 shows the result of the voxelwise two-sample t-tests for both the OLLS and the IWLLS estimator, indicating the regions where $\text{FA}_{f} > \text{FA}_{m}$ with $p_{\text{corr}} < 0.05$. The results of the opposite tests, that is, the regions where $\text{FA}_{m} > \text{FA}_{f}$ with $p_{\text{corr}} < 0.05$, are shown for both OLLS and IWLLS in Suppl. Fig. 1. Note that the overlap itself of the two tests does not necessarily indicate identical results. In addition, the lack of overlap is not indicative of a difference in outcome between the OLLS and IWLLS results. At this stage, the results merely illustrate that there is general agreement in spatial overlap of the regions that were deemed significant in terms of FA based gender differences.
Fig. 3 Results of the voxelwise analysis, indicating the regions where FA is significantly higher for females than males. Voxels colored in red and blue represent the regions where FA estimates were obtained with OLLS and IWLLS, respectively. The green voxels show their overlap, i.e., the regions where both OLLS and IWLLS reflect significantly higher FA values for females than for males. (Radiological view: left on the image is right in the brain and vice versa).
3.3 Pipeline dependent gender differences

Fig. 4 shows to which extent gender-based FA differences are driven by the choice of estimator (i.e., using OLLS or IWLLS). Overall, gender differences depend on the choice of estimator mainly in the following areas with $p_{corr} < 0.05$: parts of the CC and brainstem for $\Delta FA_m > \Delta FA_r$ and parts of the CST for $\Delta FA_r > \Delta FA_m$. To get a more detailed insight into the effect of estimation choice on the observed gender-based FA differences, we investigate the four possible scenarios ($FA_r > FA_m$ or $FA_m > FA_r$ in regions where $\Delta FA_m > \Delta FA_r$ or $\Delta FA_r > \Delta FA_m$) in the following subsections.
Fig. 4 Significance maps are shown for the interaction of estimator choice with gender-based FA differences. To enhance the contrast for significance, color-encoding is according to $-\log_{10}(p\text{-value})$ with minimum and maximum values of $-\log_{10}(0.05) \approx 1.3$ and $-\log_{10}(1/10000) = 4$ ($1/10000$ is the smallest achievable p-value with 10000 permutations), respectively. The difference in color encoding reflects how the choice of estimator can drive the gender-based FA difference in opposite directions, i.e., $\Delta \text{FA}_m > \Delta \text{FA}_f$ (red-to-yellow coloring) and $\Delta \text{FA}_f > \Delta \text{FA}_m$ (blue-to-green coloring). (Radiological view: left on the image is right in the brain and vice versa).
3.3.1 Scenario 1: $\text{FA}_f > \text{FA}_m$ in regions of $\Delta\text{FA}_m < \Delta\text{FA}_f$

Fig. 5 a) shows the area of investigation. The generality of the estimator-induced bias can be seen on Fig. 5 b), which shows the differences of the effect sizes as a function of OLLS-based effect sizes.

To get a better insight into the underlying effect of how estimator choice can drive gender-based FA differences, we explicitly show the data points of all participants for a single voxel. To showcase this effect, we performed a detailed analysis for the voxel in which the effect size of the $\text{FA}_f > \text{FA}_m$ test decreased the most, when the estimation was changed from OLLS to IWLLS (Fig. 6). MNI coordinates of this voxel, located in the midsagittal plane of the splenium, are: $x = 0; y = -38; z = 16$.

Figs. 6 a) and b) show the distribution of FA values from all subjects in the given voxel when using the OLLS ($\text{FA}_{\text{OLLSS}}$) and IWLLS ($\text{FA}_{\text{IWLLS}}$) estimators, respectively. The effect size is lower for IWLLS than for OLLS: Cohen’s $d$ decreased from 0.49 to 0.34. By investigating the $\text{FA}_{\text{IWLLS}} / \text{FA}_{\text{OLLSS}}$ ratios (Fig. 6 c)), it can be readily seen that $\text{FA}_m$ increased more than $\text{FA}_f$ when changing the estimator from IWLLS to OLLS. The $\text{FA}_m - \text{FA}_f$ difference is plotted for each decile with the bootstrapped confidence intervals as a function of male deciles, indicating that the increase in $\text{FA}_m$ was systematically larger than the increase in $\text{FA}_f$ by 0.5-2% due to this change (Fig. 6 d)). Note that if a confidence interval does not include zero, one may also conclude that said difference is significant between the changes of these ratios.
Fig. 5. a) The spatial distribution of the voxels in MNI space, where males have a significantly larger $\Delta FA$ than females and where $FA_f > FA_m$, regardless of whether the test was significant or not with any of the estimators. There were no voxels where the IWLLS-based $FA_f > FA_m$ test was significant, while the OLLS-based was not. b) Scatterplot of the difference in effect sizes between OLLS ($d_{OLLS}$) and IWLLS ($d_{IWLLS}$) based effect sizes as a function of $d_{OLLS}$. (Radiological view: left on the image is right in the brain and vice versa).
Fig. 6 The FA distribution for males (blue) and females (red) for OLLS (a) and IWLLS (b), respectively, in a voxel located in the corpus callosum (CC), where the effect size decreased the most from $d_{\text{OLL}} = 0.49$ to $d_{\text{IWLLS}} = 0.34$. c) The ratio of $\text{FA}_{\text{IWLLS}} / \text{FA}_{\text{OLL}}$ per gender, with the vertical lines indicating the deciles. d) The quantile differences between males and females for the ratios shown in panel c).
3.3.2 Scenario 2: FAf > FA m in regions of ΔFA f > ΔFA m

Fig. 7 a) shows the area of investigation. The generality of the estimator-induced bias can be seen on Fig. 7 b), which shows the differences of the effect sizes as a function of IWLLS-based effect sizes.

Fig. 8 shows the detailed analysis for the voxel in which the effect size of the FA f > FA m test increased the most, when the estimation was changed from OLLS to IWLLS. MNI coordinates of the voxel, located in the superior longitudinal fasciculus (SLF), are: x = 28; y = -20; z = 36. Figs. 8 a) and b) show the distribution of FA values from all subjects in the given voxel when using the OLLS (FA OLLS) and IWLLS (FA IWLLS) estimators, respectively. The effect size is higher for IWLLS than for OLLS: Cohen’s d increased from 0.19 to 0.27. Fig. 8 c) shows the FA IWLLS / FA OLLS ratios per gender, indicating that FA f increased more than FA m when changing the estimator from IWLLS to OLLS. Fig. 8 d) shows the shift function. The FA m - FA f difference is plotted for each decile with the bootstrapped confidence intervals as a function of male deciles, indicating that the increase in FA f over FA m was larger with 1-2%, except in the highest decile, where FA increased nearly at the same rate. Note that if a confidence interval does not include zero, one may also conclude that said difference is significant between the changes of these ratios.
Fig. 7 a) shows the spatial distribution of the voxels in MNI space, where females have a significantly larger ΔFA than males and where $\text{FA}_f > \text{FA}_m$, regardless of whether the test was significant or not with any of the estimators. There were no voxels where the OLLS-based $\text{FA}_f > \text{FA}_m$ test was significant, while the IWLLS-based was not. b) Scatterplot of the difference in effect sizes between OLLS ($d_{\text{OLLS}}$) and IWLLS ($d_{\text{IWLLS}}$) based effect sizes as a function of $d_{\text{IWLLS}}$. (Radiological view: left on the image is right in the brain and vice versa).
Fig. 8 The FA distribution for males (blue) and females (red) for OLLS (a) and IWLLS (b), respectively, in a voxel located in the superior longitudinal fasciculus (SLF), where the effect size increased the most from $d_{\text{OLL}} = 0.19$ to $d_{\text{IWLLS}} = 0.27$. c) The ratio of $F_{\text{IWLLS}} / F_{\text{OLL}}$ per gender, with the vertical lines indicating the deciles. d) The quantile differences between males and females for the ratios shown in panel c).
3.3.3 Scenario 3: $F_{A_m} > F_{A_f}$ in regions of $\Delta F_{A_m} > \Delta F_{A_f}$

Males have a smaller area where $F_{A_m} > F_{A_f}$, therefore the area where estimators could have any effect is also smaller compared to females. The area of investigation is located where $\Delta F_{A_m} > \Delta F_{A_f}$ is significant, as shown in Fig. 4, but within that region it is limited to voxels where $F_{A_m} > F_{A_f}$. Fig. 9 shows the differences of the effect sizes as a function of IWLLS-based effect sizes. For the sake of simplicity, the spatial distribution of the voxels in MNI space is not shown.

Fig. 9 Scatterplot of the difference in effect sizes between OLLS ($d_{OLLS}$) and IWLLS ($d_{IWLLS}$) based effect sizes as a function of $d_{IWLLS}$, where males have a significantly larger $\Delta F_A$ than females and where $F_{A_m} > F_{A_f}$. 
3.3.4 Scenario 4: $FA_m > FA_f$ in regions of $ΔFA_f > ΔFA_m$

The area of investigation is located where $ΔFA_f > ΔFA_m$ is significant, as shown in Fig. 4, but within that region is limited to voxels where $FA_m > FA_f$. Fig. 10 shows the differences of the effect sizes as a function of OLLS-based effect sizes. For the sake of simplicity, the spatial distribution of the voxels in MNI space is not shown.

Fig. 10 Scatterplot of the difference in effect sizes between OLLS ($d_{OLLS}$) and IWLLS ($d_{IWLLS}$) based effect sizes as a function of $d_{OLLS}$, where females have a significantly larger $ΔFA$ than males and where $FA_m > FA_f$. 
In this work, we investigated how making a different choice for a specific data processing step can affect the outcome in a typical DTI group study. More specifically, we performed a voxel-based analysis, comparing FA values between males and females using HCP data, and revealed that a higher effect size was obtained with the OLLS diffusion tensor estimator than with its IWLLS counterpart. If we consider that the IWLLS estimator has a higher accuracy, we can conclude that OLLS overestimates the observed FA based gender differences. With the majority of published DTI studies having used the OLLS estimator, it is not hard to imagine that the lack of general agreement in findings for several research topics (both in neuroscience and clinical applications) could also be partly attributed to the higher number false positives introduced by the OLLS estimator as compared with the IWLLS estimator. In the following paragraphs, we will discuss how our findings relate with what is known in functional MRI (fMRI) and we will place our results in the context of other dMRI studies.

The term ‘blobology’ (Poldrack, 2012) corresponds to the colorful patches, the ‘blobs’, of fMRI brain studies, summarizing the localization of the results after processing and statistical thresholding. The phrase reflects an inherent frustration within the neuroimaging community, partly due to the lack of effect size reports. In dMRI studies, unfortunately, effect sizes are rarely reported. Researchers often spend most of their efforts on reporting statistically significant results from the data, while the extent of these effects, which is highly complementary, is hardly considered.

With large databases like ADNI (n > 2000) (Mueller et al., 2005), ENIGMA (n > 10000) (Thompson et al., 2014), HCP (n = 1200), UK BioBank (final n = 100000) (Sudlow et al., 2015), or the Whitehall study (n = 6035) (Filippini et al., 2014), the challenges are shifting toward huge sample sizes to allow the detection of small effects, which otherwise could not be identified (Smith and Nichols, 2018). But even for group studies based on these cohorts, not properly processing the data according to best practices may still result in biases that will affect the reliability of the final outcome measures. Thompson et al. (Thompson et al., 2016) reached a similar conclusion in relation to the genome-connectome association in the ENIGMA project: “… Clearly, the ability to pursue such an approach on a large scale, within ENIGMA, depends on several factors: a working group, ENIGMA-DTI, was set up to assess its feasibility. First, unless diffusion-weighted MRI measures show greater genetic effect sizes than other traits assessed so far, there must be tens of thousands of DTI scans available from people with GWAS for such a study to be well powered …”. According to the ENIGMA-DTI
processing protocol (ENIGMA DTI protocol, 2018), the OLLS estimator is used via the FSL toolbox dtifit. In all of the aforementioned large-scale cohorts (ADNI, HCP, UK BioBank, Whitehall study), OLLS is also used which, in light of our findings, may adversely affect the reliability of the final outcome in a group study. Generally, lower-quality dMRI data in terms of effective SNR or CNR benefit more from using an estimator with better performance characteristics such as the IWLLS approach (Veraart et al., 2013a, 2013b). In this work, we used HCP data, which are among the highest quality data available in current large-scale cohorts (Bastiani et al., 2019). Given the lower number of DWIs, the lower SNR and CNR, and the higher amount of physiological artifacts in more conventional neuroimaging studies, especially in a clinical setting, one can expect even more inflated effect sizes by using the OLLS estimator than those observed in this work.

In this work, we carried out the voxelwise analysis with the Statistical Parametric Mapping (SPM) toolbox (Penny et al., 2007), rather than with another common approach, i.e. tract-based spatial statistics (TBSS) (Smith et al., 2006). While our results in this manuscript would be conceptually the same when using TBSS, confounds may arise from the skeletonization step, which may be different between the OLLS and the IWLLS. Differences in their local FA maxima could then affect statistical analysis and may further complicate interpretation of the outcome (Bach et al., 2014). Assuming the same skeleton could be provided for both datasets, e.g., via the overlap and the fusion of the skeletons, there is no reason to consider that the results presented in this work would be significantly different.

Researchers often justify the choices made for specific processing steps in their data processing pipeline by referring to previously peer-reviewed studies, which used the same settings or algorithms, despite the availability of more reliable alternatives. In addition, as OLLS generates an artificially higher effect size than IWLLS, it stimulates the positive bias in publications (Rothstein et al., 2006) and contributes to “the natural selection of bad science” (Smaldino and McElreath, 2016). To some extent, following the implementation of “registered reports” may mitigate this concern as the processing pipeline can be reviewed and scrutinized before starting the actual analysis (Nosek and Lakens, 2014).

In a recent review paper by Poldrack et al. (Poldrack et al., 2017) the lack of common consensus in processing and analysis was showcased for fMRI. With common fMRI software packages, it was shown that the number of possible analysis workflows can be as much as 69,120. For DTI, it is not hard to achieve the same order of magnitude for this number of workflows given the vast amount of
Effect of DTI estimator on group study outcome

options and parameter settings one can think of. In this work, we specifically investigated the effect of choosing between the OLLS and the IWLLS estimator on the outcome of the analysis, as using a diffusion tensor estimator is mandatory. Other processing steps, such as denoising and correcting for artifacts are not per se necessary (although highly recommended, of course) to continue with performing an actual group study. In this context, there may be several aspects of a typical processing or analysis workflow for DTI that may result in much larger effects than shown in this work.

Eklund et al. (Eklund et al., 2016) used resting-state fMRI to obtain “null data”, i.e., truly negative data, to test the false-positive ratios for task fMRI. Unfortunately, for DTI, such an experimental testing setup to evaluate statistical inferences related to methodological factors is not trivial. However, without loss of generality, in this work, we performed a standard group study on gender as the framework to evaluate the effect of using different diffusion tensor estimation approaches. We used HCP data because of the excellent data quality and the large number of subjects with proper male-female balance, thereby eliminating issues related to small sample size and low power during statistical inference (Button et al., 2013).

In this work, we did not opt for analyzing the “statistical” significance (i.e., p-values) of our findings, but rather considered the difference in effect sizes that can be observed. In a similar context, shifting the focus from p-values to effect sizes was also recently presented by Ritchie et al. (Ritchie et al., 2018). They compared volumes and DTI based metrics of cortical, subcortical, and WM regions between females and males from the UK BioBank for more than 5000 participants. The comparison of the right CST revealed that males have larger FA values than females, with a p-value of $4 \times 10^{-65}$ using Cohen’s $d = 0.54$. After adjusting for total brain volume, the values changed to $8 \times 10^{-12}$ with Cohen’s $d = 0.22$. While these p-values are indeed very significant, they do not contain any useful information. On the other hand, the effect size measures provide more practical information. That is, adding another 5000 or more participants to the analysis will not result in any meaningful change in terms of the effect size, as this investigation is already statistically well-powered, while the p-value would decrease further. For the same reason, i.e., avoiding under-powered study design, we used HCP data for our group comparison, allowing us to focus on the performance of the DTI estimators.

Despite the efforts of optimizing the dMRI processing pipeline, it is often not clear what the benefits are of new developments for group-based studies. In this work, however, we showed that the application of IWLLS should be preferred over the OLLS for diffusion tensor estimation. The current framework can be easily extended to examine effects of modifying other processing elements, but
also to investigate choices in algorithms and settings for specific analysis strategies, like tractography and connectomics, further improving the reliability and validity of future dMRI group studies.

5 Conflict of Interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Suppl. Fig. 1 Results of the voxelwise analysis, indicating the regions where the FA is significantly higher for males than for females. Voxels colored in red and blue represent the regions where the FA estimates were obtained with the OLLS and IWLLS estimators, respectively (only visible in a few voxels). The green voxels show their overlap, i.e., the regions where both OLLS and IWLLS reflect significantly higher FA values for males compared to females.