MASiVar: Multisite, Multiscanner, and Multisubject Acquisitions for Studying Variability in Diffusion Weighted Magnetic Resonance Imaging

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ABSTRACT

Purpose

Diffusion weighted imaging (DWI) allows investigators to identify microstructural differences between subjects, but variability due to session and scanner biases is still a challenge.

Methods

To investigate DWI variability, we present MASiVar, a multisite dataset consisting of 319 diffusion scans acquired at 3T from $b = 1000$ to $3000$ s/mm$^2$ across 14 healthy adults, 83 healthy children (5 to 8 years), three sites, and four scanners as a publicly available, preprocessed, and de-identified dataset. With the adult data, we demonstrate the capacity of MASiVar to simultaneously quantify the intrasession, intersession, interscanner, and intersubject variability of four common DWI processing approaches: (1) a tensor signal representation, (2) a multi-compartment neurite orientation dispersion and density model, (3) white matter bundle segmentation, and (4) structural connectomics. Respectively, we evaluate region-wise fractional anisotropy (FA), mean diffusivity, and principal eigenvector; region-wise cerebral spinal fluid volume fraction, intracellular volume fraction, and orientation dispersion index; bundle-wise shape, volume, length and FA; and connectome correlation and maximized modularity, global efficiency, and characteristic path length.

Results

We plot the variability in these measures at each level and find that variability generally increases with intrasession to intersession to interscanner to intersubject effects across all processing approaches and that sometimes interscanner variability can approach intersubject variability.

Conclusions

This study demonstrates the potential of MASiVar to investigate DWI variability across multiple levels and processing approaches simultaneously and suggests harmonization between scanners for multisite analyses is critical prior to inference of group differences on subjects.
INTRODUCTION

Diffusion weighted MRI imaging (DWI) is a noninvasive way of elucidating the brain’s microstructural makeup (1). Common modes of DWI analysis include representing the diffusion signal with tensors (2,3), representing biological tissues with multi-compartment models (4–6), identifying white matter bundles (7), and investigating the human structural connectome (8). These approaches form the basis for many studies including those investigating a wide range of neurological disorders including autism (9,10), diabetes (11,12), multiple sclerosis (13), and schizophrenia (14) as well as differences due to aging (15) and sex (16). These types of studies, however, rely on the identification of group differences with respect to an independent variable. Often this variable reflects whether the scanned subject has a particular disease, or the age or sex of the subject. Robust study design can control for additional subject-level confounders through age- and sex-matching and related approaches. However, one level of potential confounding in DWI studies that has not been thoroughly characterized is the variability of calculations due to differences within and between imaging sessions and scanners.

One particular reason for this is the difficulty in acquiring data configured to perform such a characterization. For instance, to quantify variation within a session, imaging sessions with repeated scans are needed. To quantify variation between sessions and between scanners, multiple imaging sessions on at least one scanner and at least one imaging session on multiple scanners are required, respectively. Last, to assess the session and scanner effects relative to the subject effect size, multiple scanned subjects are needed as well.

Another reason for this is the low number of properly configured publicly available datasets. Some of the few that exist that allow for investigations of DWI variability are the MASSIVE dataset (17), the Human Connectome Project (HCP) 3T dataset (18), the MICRA dataset (19), the SIMON dataset (20), and the multisite dataset published by Tong et al. (21). MASSIVE consists of one subject scanned repeatedly on one scanner (17); HCP consists of multiple subjects with multiple acquisitions per session all on one scanner (18); MICRA consists of multiple subjects scanned repeatedly on one scanner (19); SIMON consists of one subject scanned at over 70 sites (20), and the Tong dataset consists of multiple subjects each scanned on multiple scanners (21).

These difficulties have resulted in existing DWI variability studies that are largely limited in scope and that offer a fragmented view of the variability landscape (Table 1). Many of these studies each capture portions of the spectrum of effects due to session, scanner, and subject biases, but are unable to assess for all levels at once. In addition, most of the existing investigations each focus on one specific DWI processing approach and/or model and as such do not provide a holistic assessment of DWI variability. As such, the understanding of how one study’s variability estimates in tensor-based metrics between sessions might compare to another’s estimates of tractography biases
between scanners is not obvious, for instance. Thus, to bring the field toward a more global understanding of DWI variability, the release of additional publicly available datasets configured to characterize DWI variability and a global analysis of variability on multiple levels and across different processing approaches is needed.

To fill the first need, we propose MAsiVar, a multisite, multisite, and multisubject dataset able to characterize DWI variability due to session, scanner, and subject effects. To fill the second need, we demonstrate the potential of MAsiVar to characterize DWI variability by presenting a simultaneous quantification and comparison of these effects on four different common diffusion approaches, hypothesizing that variability increases with session, scanner, and subject effects.

**METHODS**

*Data acquisition*

MAsiVar consists of data acquired from 2016 to 2020 to study both DWI variability and other phenomena. As such, the data exist in four cohorts, designated I, II, III, and IV (Figure 1).

Cohort I consists of one healthy adult subject (male, age 25 years) with multiple imaging sessions on a 3T Philips Achieva scanner at site 1 (scanner A). This subject underwent three imaging sessions, one each consecutive day, and received two to three scans during each session (Figure 1). Each scan consisted of 96-direction acquisitions at b = 1000, 1500, 2000, 2500, and 3000 s/mm² (Table 2). These scans were acquired at 2.5mm isotropic resolution with an echo time (TE) and repetition time (TR) of TE / TR = 94ms / 2650ms.

Cohort II consists of five healthy adult subjects (3 male, 2 female, age 27 to 47 years) scanned for one to two sessions on each of three to four different scanners. Each subject underwent all sessions within one year. The scanners included scanner A, another 3T Philips Achieva scanner at site 1 (scanner B), a 3T General Electric Discovery MR750 scanner at site 2, and a 3T Siemens Skrya scanner at site 3 (Figure 1). For each imaging session, each subject received one scan, consisting of 96-direction acquisitions at b = 1000, 1500, 2000, 2500 (or 2465 at site 3 due to hardware limitations) s/mm² and a 30- or 32-direction acquisition at b = 1000 s/mm² (Table 2). The scans acquired on scanner B, at site 2, and at site 3, and all the 30- or 32-direction scans were acquired at 2.5mm isotropic resolution. On scanner A, one subject’s 96-direction acquisitions were also acquired at 2.5mm isotropic resolution while the remainder were acquired at 1.9mm by 1.9mm by 2.2mm (sagittal, coronal, and axial) resolution. For acquisitions on scanner A, the 2.5mm isotropic 96-direction scans were acquired with TE / TR = 90ms / 5200ms, while the other 96-direction acquisitions were acquired with TE / TR = 90ms / 5950ms, and TE / TR = 55ms /...
6127ms to 7309ms for the 32-direction acquisitions. For acquisitions on scanner B, the 96-direction scans were acquired with TE / TR = 90ms / 5800ms or 5900ms, while the 32-direction acquisitions were acquired with TE / TR = 55ms / 7022ms to 7069ms. For the 96-direction acquisitions acquired at site 2, TE / TR = 90ms / 5800ms or 5900ms, while the 32-direction acquisitions were acquired with a TE / TR of either 58ms / 7042ms or 59ms / 4286ms. All scans acquired at site 3 were acquired with TE / TR = 95ms / 6350ms. All sessions acquired on scanner A that contained scans of varying resolution were resampled to match the resolution of the 96-direction acquisitions prior to analysis.

Cohort III consists of 8 healthy adult subjects (4 male, 4 female, ages 21 to 31 years) scanned for one to six sessions on scanner B (Figure 1). Each subject underwent all sessions within one year. Each subject received one to two scans during each session, with each scan consisting of a 40-direction b = 1000 s/mm\(^2\) and a 56-direction b = 2000 s/mm\(^2\) acquisition (Table 2). The majority of these scans were acquired at 2.1mm by 2.1mm by 2.2mm (sagittal, coronal, and axial) resolution and TE / TR = 79ms / 2900ms, with a few acquired at 2.5mm isotropic resolution and TE / TR = 75ms / 3000ms.

Cohort IV consists of 83 healthy child subjects (48 male, 35 female, ages 5 to 8 years) scanned for one to two sessions on scanner B (Figure 1). For the subjects with multiple sessions, the sessions were longitudinally acquired, spaced approximately one year apart. As with Cohort III, during each session, each subject received one to two scans, with each scan consisting of a 40-direction b = 1000 s/mm\(^2\) and a 56-direction b = 2000 s/mm\(^2\) acquisition (Table 2). These scans were acquired at 2.1mm by 2.1mm by 2.2mm (sagittal, coronal, and axial) resolution with TE / TR = 79ms / 2900ms.

All acquisitions were phase encoded in the posterior to anterior direction (APP) and were acquired with one b = 0 s/mm\(^2\) volume each. Reverse phase encoded (APA) b = 0 s/mm\(^2\) volumes were also acquired for all scans in all cohorts except for those from one subject in cohort II at site 3. Most sessions also included a T1-weighted image for structural analysis or distortion correction (36). All images were deidentified and all scans were acquired only after informed consent under supervision of the project Institutional Review Board.

Data preprocessing

After acquisition, all scans in MASiVar were preprocessed and quality checked with the PreQual pipeline (37). In brief, all acquisitions per scan were denoised with the Marchenko-Pastur technique (38–40), intensity normalized, and distortion corrected. Distortion correction included susceptibility-induced distortion correction (41) using APA b = 0 s/mm\(^2\) volumes when available and the Synb0-DisCo deep learning framework (36) and associated T1 image...
when not, eddy current-induced distortion correction, intervolume motion correction, and slice-wise signal drop out imputation (42,43). The estimated volume-to-volume displacement corrected during preprocessing and signal-to-noise ratios of the scans are reported in Supporting Information Figure S1.

Overview of variability study

Using data acquired in adults, we sought to demonstrate the capacity of MASiVar to simultaneously investigate DWI variability due to

1. intrasession (scans acquired within the same session on the same scanner of the same subject),
2. intersession (scans acquired between different sessions on the same scanner of the same subject),
3. interscanner (scans acquired between different sessions on different scanners of the same subject), and
4. intersubject (scans acquired of different subjects in different sessions on the same scanner) effects.

We quantified these levels of effects in four common types of DWI analysis, including

1. a diffusion tensor imaging (DTI) signal representation,
2. a multi-compartment neurite orientation dispersion and density imaging (NODDI) model (4),
3. the RecoBundles white matter bundle segmentation technique (44), and
4. a connectomics representation with graph-based measures (45).

For DTI, we investigate variability in regional fractional anisotropy (FA), mean diffusivity (MD), and principal eigenvector (V1) measurements. For NODDI, we investigate variability in regional cerebrospinal fluid (CSF) volume fraction (cVF), intracellular volume fraction (iVF), and orientation dispersion index (ODI) measurements. For bundle segmentation, we investigate variability in bundle shape, volume, length, and FA. For connectomics we investigate whole connectome variability as well as that of the maximum modularity (MM), global efficiency (GE), and characteristic path length (CPL) graph measures (Figure 2).

Defining variability

Due to the nested nature of session, scanner, and subject effects, we wished to define variability for a given effect in a way that limited confounding from the other effects. For example, because sessions are necessarily nested in scanners, we wanted to be able to investigate interscanner effects without confounding from intrasession effects and vice versa. To do this, we elected to use a paired difference approach. By identifying pairs of images that should produce the same measurements, we can estimate variability by analyzing the differences between the scans in the pair. For example, when the same person is scanned once on two different scanners, interscanner variability can be quantified by computing the differences in measurements between the two images. Additionally, since a pair of scans can only satisfy the criteria for exactly one of the listed effects, computing variability within these pairs
reduces confounding by holding the other effects constant. As such, we identified all pairs of scans in cohorts I to III of MASIvar that satisfied the criteria for each of the four listed effects (Figure 3), resulting in 41 intrasession pairs, 188 intersession pairs, 53 interscanner pairs, and 80 intersubject pairs. Only cohort II was used for the intersubject pairings to reduce bias toward scanners A and B.

We quantify variability at a given level of variation and type of DWI measurement by “summarizing” the “differences” within the relevant pairs. For example, to compute the variability of intersession DTI FA measurements, we compute the “difference” in FA in each intersession scan pair and report a “summary” across all pairs. Due to the distinct properties of each type of DWI processing, the exact definition of “difference” (Figure 4) and “summary” (Figure 5) vary by measurement type and are detailed in the following sections.

**Variability in DTI and NODDI**

For the DTI approach, we extract the \( b = 1000 \text{ s/mm}^2 \) acquisition from each scan with the largest number of directions. We then calculate the diffusion tensor for each scan using an iteratively reweighted least squares approach implemented in MRtrix3 (46). The tensors are subsequently converted to FA, MD, and V1 representations of the data (47). These images are then deformably registered to the Montreal Neurological Institute (MNI) image space with the ANTs software package (48,49). From there, we identify the 48 regions of interest (ROIs) in each image defined by the Johns Hopkins white matter atlas (50–52) (Figure 2a).

For the NODDI approach, we extract the \( b = 1000 \text{ s/mm}^2 \) acquisition from each scan with the largest number of directions and the \( b = 2000 \text{ s/mm}^2 \) acquisition. We then fit the multicompartiment model with the UCL NODDI Toolbox as implemented in MATLAB (4). The models are subsequently converted to cVF, iVF, and ODI representations. These images are then deformably registered to MNI space with the ANTs software package. From there, we identify the 48 ROIs in each image defined by the Johns Hopkins white matter atlas (Figure 2b).

We perform the DTI and NODDI paired difference calculations on a regional basis in MNI space with voxel-wise correspondence between images. For a given region and level of variation, we calculate the difference within a pair of scans for FA, MD, cVF, iVF, and ODI as the median voxel-wise absolute percent difference. For V1, we define it as the median voxel-wise absolute angular difference in degrees (Figure 4a). Once the paired differences are computed for all pairs in all regions, we compute the regional medians and take the resulting distribution across regions to summarize a given level of variability (Figure 5a). Additionally, we compute the 95% confidence intervals for each of the regional medians with nonparametric bootstrapping of the scan pairs with 1000 iterations (Supporting Information Figures S2-S7) (53).
Variability in bundle segmentation

For the white matter segmentation approach, we extract the b = 2000 s/mm$^2$ acquisition from each scan. We calculate a whole-brain tractogram with DIPY of 2 million streamlines (54). We use the constrained spherical deconvolution model (55) with probabilistic local tracking with a maximum angle of 25°, a seeding criterion of FA > 0.3, and a stopping criterion of FA < 0.2. We extract 43 white matter bundles (Supporting Information Table S1) from each tractogram using the RecoBundles algorithm as implemented in DIPY. In short, each tractogram is registered to an MNI tractogram template and streamlines from each tractogram are assigned to bundles within the template (44). The length, volume, and FA of each bundle are then calculated. We calculate bundle length by calculating the median streamline length. We calculate volume by first converting each bundle to a tract density image representation. From there, a binary bundle mask is calculated by thresholding the tract density image at 5% of the 99$^{th}$ percentile density. Volume is calculated by multiplying the number of voxels in the mask by the volume of each voxel. FA is calculated by first converting the image to a tensor representation (46) and then to an FA representation (47). Each bundle’s binary mask is then applied to obtain the voxel-wise median FA value per bundle (Figure 2c).

Because streamline-wise and subsequent voxel-wise correspondence cannot be achieved with tractography and bundle segmentation, we compute paired differences differently than in the DTI and NODDI case. For a given bundle and level of variation, we calculate the paired difference of bundle shape with the Dice similarity index (56) between the tract density images from the two images. We define the difference of bundle volumes as the absolute percent difference. We calculate the difference of bundle FA as the absolute percent difference between the voxel-wise medians from each image and the difference of bundle length as the absolute percent difference between the streamline-wise medians from each image (Figure 4b). Similar to the DTI and NODDI analysis, once the paired differences are computed for all pairs in all bundles, we compute the bundle-wise medians and take the resulting distribution across bundles to summarize a given level of variability (Figure 5a). Additionally, we compute the 95% confidence intervals for each of the bundle-wise medians with nonparametric bootstrapping of the scan pairs with 1000 iterations (Supporting Information Figures S8-S11).

Variability in connectomics

For the connectomics approach, we extract the b = 2000 s/mm$^2$ acquisition from each scan. We then calculate a whole-brain tractogram with MRtrix3 (57). We first use the constrained spherical deconvolution model with probabilistic tracking with a maximum angle of 25°, a seeding criterion of FA > 0.3 and a stopping criterion of FA < 0.2 to calculate a 10 million streamline tractogram. The tractogram is then filtered with the SIFT approach to 2
We parcellate the brain into 96 cortical regions using the Harvard-Oxford cortical atlas (59–62) and compute a connectome where each edge represents the average streamline distance connecting the two nodes. The MM, GE, and CPL are then calculated from each connectome using the Brain Connectivity Toolbox as implemented in MATLAB (45) (Figure 2d).

To evaluate paired differences of connectomics, we characterize each pair of connectomes as both (1) a whole and (2) through scalar measures. First, we calculate the Pearson correlation between the connectomes within each image pair as an estimate for connectome agreement within a pair. Second, we calculate the percent absolute difference in MM, GE, and CPL (45) between the connectomes in the pair (Figure 4c). Unlike the DTI, NODDI, and bundle segmentation cases, we do not have multiple regions or bundles for the connectomics analysis with which we can obtain a distribution to summarize variability at a level. As a result, we instead compute a distribution for the median paired difference of each metric with nonparametric bootstrapping of the scan pairs with 1000 iterations (Figure 5b).

Comparing variability across levels

To compare variability across the intrasession, intersession, interscanner, and intersubject levels, we use non-parametric statistical tests for all four processing approaches with a significance level of 0.05 and report the uncorrected \( p \)-values. For the DTI, NODDI, and bundle segmentation analyses, we use the Wilcoxon signed-rank test for paired distributions, as all points in each distribution are measured from corresponding regions or bundles (63). For the connectomics analysis, we use the Wilcoxon rank-sum test for unpaired distributions, as the distributions were constructed from bootstrapping and thus do not have correspondence (63).

Comparing variability across processing approaches

Last, to obtain a more global understanding of the relative intrasubject (intrasession, intersession, and interscanner) effects across different processing approaches, we normalize these effects to intersubject variation for each approach. We calculate the ratios of each of the intrasubject differences to the corresponding intersubject estimates. For the DTI, NODDI, and bundle segmentation analysis, we compute this ratio on a region- or bundle-wise basis, and for the connectome analysis, we compute this ratio on the bootstrapped distributions. For the Dice and correlation similarity measures, we first subtract them from 1 to obtain dissimilarity estimates. With this design, ratios of <1, 1, and >1 indicate the intrasubject variation is less than, equal to, or greater than the intersubject variation, respectively.
RESULTS

Variability in DTI

As shown in Figure 6, we find that intrasession FA measurements vary around 8.3%, that intersession measurements vary around 9.1%, that interscanner measurements vary around 13.7%, and that intersubject measurements vary around 17.7%. We find the corresponding measurements in the MD case to be 3.6%, 4.9%, 8.6%, and 9.1% and for the V1 case to be 10.0°, 10.7°, 14.5°, and 20.2°, respectively. All these differences were statistically significant (p < 0.0005, Wilcoxon signed-rank test), except for the interscanner and intersubject comparison for MD (p < 0.05).

Variability in NODDI

As shown in Figure 7, we find that the intrasession cVF measurements vary around 47.3%, that intersession measurements vary around 55.1%, that interscanner measurements vary around 57.5%, and that intersubject measurements vary around 61.9%. We find the corresponding measurements in the iVF case to be 7.9%, 8.6%, 11.8%, and 12.1% and for the ODI case to be 11.7%, 13.5%, 19.1%, and 26.5%, respectively. All these differences were statistically significant (p < 0.0005, Wilcoxon signed-rank test), except for the interscanner and intersubject iVF comparison. We evaluated cVF only in white matter regions defined by the Johns Hopkins atlas and thus dealt with very low cVF values when calculating percent difference.

Variability in bundle segmentation

As shown in Figure 8, we find that intrasession bundles overlap at 0.61 Dice, that intersession bundles overlap at 0.61 Dice, that interscanner bundles overlap at 0.57 Dice, and that intersubject bundles overlap at 0.53 Dice. We find the measurements for the corresponding levels of variation in the bundle volume comparisons to be 11.4%, 10.2%, 12.8%, and 17.9%, in the FA case to be 1.7%, 1.9%, 5.0%, and 4.9%, and in the bundle length case to be 3.2%, 3.6%, 4.1%, and 8.0%, respectively. All these differences were statistically significant (p < 0.0005, Wilcoxon signed-rank test), except for the intrasession and intersession shape, volume, and length (p < 0.05) comparisons as well as the interscanner and intersubject FA comparison.

Variability in connectomics

As shown in Figure 9, we find that the intrasession connectomes correlate at 0.62, that the intersession connectomes correlate at 0.59, that the interscanner connectomes correlate at 0.59 and that the intersubject connectomes correlate
at 0.54. We find the measurements for the corresponding levels of variation in the MM case to be 7.2%, 8.2%, 9.5%, and 13.3%, in the GE case to be 1.0%, 1.6%, 6.1%, and 6.2%, and in the CPL case to be 0.8%, 1.5%, 6.2%, and 6.1%, respectively. All these differences were statistically significant ($p < 0.0005$, Wilcoxon rank-sum test) except for the interscanner and intersubject comparison of CPL.

Comparing variability across processing approaches

We plot the ratios of intrasubject to intersubject variability in Figure 10. Previous studies have demonstrated what we find to be good intrasession and intersession FA reproducibility with coefficients of variation (CoV) up to 3% as well as moderate interscanner reproducibility with CoV up to 8% (Table 1). Based on these findings, we draw thresholds to separate the FA intrasession and intersession ratios from the interscanner ratios and the FA interscanner ratios from the intersubject reference in order to compare the remaining metrics. As such, we take a ratio of <0.65 to represent good reproducibility, 0.65-0.95 to represent moderate, and >0.95 to represent poor reproducibility. With this heuristic, we find good to moderate reproducibility in most of the intrasession and intersession measures across all four approaches. One notable deviation from this trend is that intersession white matter cVF demonstrated moderate to poor reproducibility. Additionally, despite the thresholds drawn, we note that the intrasession and intersession variability estimates are all within an order of magnitude of the intersubject effects as designated by ratios between 0.1 and 1. Last, for most measures, we find moderate to poor interscanner reproducibility and that interscanner effects can introduce variability that is on par with that from intersubject effects. We find this trend is most striking for the GE and CPL analyses that demonstrated intrasession ratios of around 0.2 but interscanner ratios near 1. In summary, we find that consistently across all four DWI analysis approaches intrasubject variability is good to moderate but non-negligible compared to intersubject variability and that the size of interscanner effects can approach the size of intersubject effects.

DISCUSSION AND CONCLUSIONS

Here, we present, MASiVar, a dataset designed for investigation of DWI variability. Additionally, to demonstrate the capacity of MASiVar as a resource, we characterize intrasession, intersession, interscanner, and intersubject variability and plot the ratios of intrasubject to intersubject variability in four common diffusion processing approaches. In support of our hypothesis, we consistently find that variability increases with consideration of session, scanner, and subject effects. We also consistently find across all approaches that intrasubject variability accounts for a non-negligible portion of intersubject variability and that at times interscanner variability can approach intersubject variability. We interpret two primary conclusions from these results. The first is that MASiVar provides the field a resource to obtain an improved global understanding of session, scanner, and subject effects.
within and between different DWI processing approaches. Second, we interpret these results to mean that harmonization between scanners for multisite analyses should be carefully considered prior to inference of group differences on subjects.

The reproducibility of DWI analyses has received significant attention in the field, including the analysis of tensor representations (23–26), multi-compartment models (26, 27), tractography and bundle segmentation (28, 29), and connectomics (33, 35) (Table 1). Looking at the literature, we find similar trends between our results and those of prior studies that variation increases with session and scanner biases, although the numbers are not directly comparable due to different definitions of variability. However, review of the literature also demonstrates a fragmented picture of DWI variability. Previous studies have largely each primarily focused on one type of approach and one or two levels of variation. This coupled with the different definitions of variability and different study objectives have made it difficult to understand how the different effects relate to each other and how they affect a multitude of common DWI processing approaches. To the best of our knowledge, this study represents the first attempt and to characterize all four types of diffusion processing and all four levels of variation consistently and simultaneously. Thus, we hope that the dataset and study presented here will promote further investigation into a wide spectrum of DWI variability issues from a large pool of models to push the field toward a global understanding of the effects of session, scanner, and subject biases on different DWI measurements.

Of note, the bulk of existing studies used CoV to estimate variation and the intraclass correlation coefficient (ICC) to characterize the proportion of intersubject variation attributable to session or scanner effects. In contrast, we used an intrasubject and intersubject paired difference approach for the former and quantified their ratios for the latter. We chose these approaches to improve isolation of session and scanner effects. For instance, by pairing images, we remove other potential confounders like subject or acquisition effects. This approach also has the benefit of characterizing the relative sizes of intra- and intersubject variation without the necessary assumptions of traditional ICC, including not having additional confounders within each class, an assumption that is inherently violated when looking at sessions nested within scanners nested with subjects.

For this study, we chose popular software toolboxes to do all the analyses, parameter configurations that we were familiar with, and consistent similarity assessments that we found to be interpretable. However, we recognize that there are many other software options available to do similar tasks, each with a large number of different configurations, and a large number of ways to assess variability. For instance, there are different methods for fitting tensors (64–66), for identifying regions (61, 67–69) and bundles (70–73), for comparing bundles (74), and for configuring and representing connectomes (35, 45, 75, 76). Additionally, there are a number of other microstructural measures that can be characterized as well (19). Thus, the goal of the present study was not to provide an analysis
between different processing toolboxes or parameters, and since each approach was not necessarily optimized, we do not recommend thorough utilization of absolute reproducibility values presented here for any one processing approach. Instead, we aimed to contribute to a global understanding of DWI variability and its relative trends across the four processing approaches and across sessions, scanners, and subjects in a generally interpretable way that demonstrated the potential of the dataset. As such, we hope that the release of MASiVar will prompt other investigators in the field to optimize and further characterize differences between software tools and their parameters, different DWI processing and variability measures, and other potential confounders in DWI analysis.

In addition to the ability of MASiVar to serve as a utility for variability analysis, we note that the pediatric subjects in cohort IV present another unique resource for the field. The majority of the existing DWI datasets and studies for variability use adult subjects. Of existing pediatric datasets, many have focused on older age ranges. For example, the Adolescent Brain Cognitive Development project (77) and the Lifespan Human Connectome Project in Development (78) contain longitudinal DWI data acquired from children starting at age 9 and 10 through adolescence. Thus, to the best of our knowledge, MASiVar represents one of the first publicly available longitudinal DWI datasets of children prior to adolescence aged 5-8 years old and is further distinguished by its inclusion of repeated scans within each session. As a demonstration of the usefulness of cohort IV, we include a characterization of the longitudinal intersession variability in children with one year between sessions compared to the adult intersession variability computed above for all four processing approaches (Supporting Information Figure S12). We find that, as expected, the pediatric cohort exhibits increased variability for many measurements, supporting that the expected developmental changes in this age group likely introduce additional variability on top of session effects. We hope that investigators in developmental neuroscience and pediatric neurology will be able to take advantage of this resource for their work.

One limitation of the variability study is the differences in number of gradient directions between the different cohorts. Cohort III consists of a 40-direction b = 1000 s/mm² acquisition and a 56-direction b = 2000 s/mm² acquisition in contrast to the 96 directions for cohorts I and II. There is a potential effect that could be biasing the results. However, our study design focuses on paired scans and thus all variability measurements are between acquisitions of the same configuration, which should minimize this effect. In a similar vein, due to hardware limitations, the data collected at site 3 in cohort II was collected at a maximum shell of 2465 s/mm² as opposed to the 2500 s/mm² across the rest of MASiVar. This shell was not used for the present variability analysis, but this discrepancy should be noted on future studies using the dataset.

Last, we have made the MASiVar dataset publicly available at https://openneuro.org/datasets/ds003416 in Brain Imaging Data Structure (BIDS) format (79) with deidentified metadata and defaced images.
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69. Figley TD, Mortazavi Moghadam B, Bhullar N, Kornelsen J, Courtney SM, Figley CR. Probabilistic white matter atlases of human auditory, basal ganglia, language, precuneus, sensorimotor, visual and visuospatial


### Table 1. A survey of existing DWI variability estimates (cited with first author and publication year) against those presented in the present work (blue).

<table>
<thead>
<tr>
<th></th>
<th>Intrasection</th>
<th>Intersession</th>
<th>Interscanner</th>
<th>Intersubject</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>DTI</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FA</td>
<td>8.32% PD</td>
<td>9.07% PD</td>
<td>13.73% PD</td>
<td>17.66% PD</td>
</tr>
<tr>
<td></td>
<td>2% CoV (Farrell, 2010) (22)</td>
<td>1% CoV (Magnotta, 2012) (23)</td>
<td>3% CoV (Magnotta, 2012) (23)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>3% CoV (Palacios, 2017) (25)</td>
<td>0.5% CoV (Andica, 2020) (26)</td>
<td>0.90-0.99 ICC (Vollmar, 2010) (24)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>0.74-1.00 ICC (Andica, 2020) (26)</td>
<td>0.95-0.97 ICC (Koller, 2020) (19)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>3% CoV (Farrell, 2010) (22)</td>
<td>0.6-1% CoV (Koller, 2020) (19)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.93-0.97 ICC (Koller, 2020) (19)</td>
</tr>
<tr>
<td>MD</td>
<td>3.65% PD</td>
<td>4.92% PD</td>
<td>8.61% PD</td>
<td>9.06% PD</td>
</tr>
<tr>
<td></td>
<td>1% CoV (Farrell, 2010) (22)</td>
<td>1% CoV (Magnotta, 2012) (23)</td>
<td>2% CoV (Magnotta, 2012) (23)</td>
<td>6% CoV (Palacios, 2017) (25)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2% CoV (Palacios, 2017) (25)</td>
<td>0.2% CoV (Andica, 2020) (26)</td>
<td>3% CoV (Andica, 2020) (26)</td>
</tr>
<tr>
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<td></td>
<td>1% CoV (Farrell, 2010) (22)</td>
<td>0.5-1% CoV (Koller, 2020) (19)</td>
<td>0.94-0.96 ICC (Koller, 2020) (19)</td>
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<tr>
<td>V1</td>
<td>9.97°</td>
<td>10.72°</td>
<td>14.46°</td>
<td>20.19°</td>
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<tr>
<td></td>
<td></td>
<td>7-10° (Farrell, 2010) (22)</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>NODDI</strong></td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>cVF</td>
<td>47.32% PD</td>
<td>55.07% PD</td>
<td>57.47% PD</td>
<td>61.93% PD</td>
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<tr>
<td></td>
<td></td>
<td>15.9% CoV (Andica, 2020) (26)</td>
<td>15.0% CoV (Andica, 2020) (26)</td>
<td>0.013-0.545 ICC (Andica, 2020) (26)</td>
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<td>0.133-0.997 ICC (Andica, 2020) (26)</td>
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</tr>
<tr>
<td>iVF</td>
<td>7.92% PD</td>
<td>8.61% PD</td>
<td>11.76% PD</td>
<td>12.13% PD</td>
</tr>
<tr>
<td></td>
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<td>0.4% CoV (Andica, 2020) (26)</td>
<td>0.9% CoV (Andica, 2020) (26)</td>
<td>0.300-0.935 ICC (Andica, 2020) (26)</td>
</tr>
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<td>0.773-0.989 ICC (Andica, 2020) (26)</td>
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<tr>
<td>ODI</td>
<td>11.74% PD</td>
<td>13.51% PD</td>
<td>19.05% PD</td>
<td>26.48% PD</td>
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<td>0.181-0.962 ICC (Andica, 2020) (26)</td>
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<td></td>
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<td>0.789-0.998 ICC (Andica, 2020) (26)</td>
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<tr>
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<td>0.5% CoV (Tariq, 2013) (27)</td>
<td></td>
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<tr>
<td><strong>Bundle Segmentation</strong></td>
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<td></td>
<td></td>
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<tr>
<td>Shape</td>
<td>0.61 Dice</td>
<td>0.61 Dice</td>
<td>0.57 Dice</td>
<td>0.53 Dice</td>
</tr>
<tr>
<td></td>
<td>~0.67 Dice (Nath, 2020) (28)</td>
<td>~0.64 Dice (Nath, 2020) (28)</td>
<td>~0.58 Dice (Nath, 2020) (28)</td>
<td>~0.5-0.6 Dice (Schilling, 2020) (32)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.65-0.92 Dice (Besseling, 2012) (29)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.72 wDice (Cousineau, 2017) (30)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.71-0.87 wDice (Boukadi, 2019) (31)</td>
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<tr>
<td>Volume</td>
<td>11.38% PD</td>
<td>10.20% PD</td>
<td>12.76% PD</td>
<td>17.90% PD</td>
</tr>
<tr>
<td></td>
<td></td>
<td>3-22% CoV (Besseling, 2012) (29)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.53-0.96 ICC (Besseling, 2012) (29)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.41-0.83 ICC (Boukadi, 2019) (31)</td>
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<tr>
<td>FA</td>
<td>1.66% PD</td>
<td>1.91% PD</td>
<td>5.00% PD</td>
<td>4.93% PD</td>
</tr>
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<td></td>
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<td>1-4% CoV (Besseling, 2012) (29)</td>
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<tr>
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<td>0.65-0.94 ICC (Besseling, 2012) (29)</td>
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<td>0.62-0.89 ICC (Boukadi, 2019) (31)</td>
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<tr>
<td>Length</td>
<td>3.18% PD</td>
<td>3.57% PD</td>
<td>4.05% PD</td>
<td>8.04% PD</td>
</tr>
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<td></td>
<td></td>
<td>0.68-0.89 ICC (Boukadi, 2019) (31)</td>
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<tr>
<td><strong>Connectomics</strong></td>
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<tr>
<td>Correlation</td>
<td>0.62 PC</td>
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<td>0.59 PC</td>
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<td>0.59 PC (Prčkovska, 2016) (33)</td>
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<td>32.7-39.9% CD (Girard, 2015) (34)</td>
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<tr>
<td>MM</td>
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<td>GE</td>
<td>1.03% PD</td>
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<td>6.06% PD</td>
<td>6.24% PD</td>
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<tr>
<td></td>
<td>31% CoV (Roine, 2019) (35)</td>
<td>31% CoV (Roine, 2019) (35)</td>
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<tr>
<td></td>
<td>0.78 ICC (Roine, 2019) (35)</td>
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<tr>
<td>CPL</td>
<td>0.80% PD</td>
<td>1.53% PD</td>
<td>6.19% PD</td>
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</tr>
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<td>2% CoV (Roine, 2019) (35)</td>
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<td></td>
</tr>
<tr>
<td></td>
<td>0.77 ICC (Roine, 2019) (35)</td>
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</tbody>
</table>

PD: paired difference
PC: Pearson correlation
CoV: coefficient of variation
ICC: intraclass correlation coefficient
wDice: weighted Dice
CD: connectome distance, Σ(C1 - C2)/2, where C1 and C2 are matrices
Figure 1. Overview of the MASIVar dataset. This dataset consists of four cohorts. Cohort I consists of one adult subject scanned repeatedly on one scanner. This subject underwent three separate imaging sessions and acquired 3-4 scans per session. Cohort II consists of 5 adult subjects each scanned on 3-4 different scanners across 3 institutions. Each subject underwent 1-2 sessions on each scanner and had one scan acquired per session. Cohort III consists of 8 adult subjects all scanned on one scanner. Each subject underwent 1-6 sessions on the scanner and had two scans acquired per session. Cohort IV consists of 83 child subjects all scanned on one scanner. Each subject underwent 1-2 sessions on the scanner and had two scans acquired per session.
Table 2. Acquisitions acquired in each scan for the different MASiVar cohorts.

<table>
<thead>
<tr>
<th>Cohort</th>
<th>Shell (b-value)</th>
<th>Number of Directions</th>
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<td>96</td>
</tr>
<tr>
<td></td>
<td>1500</td>
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<td>2500</td>
<td>96</td>
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<tr>
<td></td>
<td>3000</td>
<td>96</td>
</tr>
<tr>
<td>II</td>
<td>1000</td>
<td>30 or 32</td>
</tr>
<tr>
<td></td>
<td>1000</td>
<td>96</td>
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<td>2000</td>
<td>96</td>
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<td></td>
<td>2465 or 2500</td>
<td>96</td>
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<td>III</td>
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<tr>
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<tr>
<td></td>
<td>2000</td>
<td>56</td>
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</table>
Figure 2. Outline of processing and measurements investigated presently in four common diffusion MRI analysis approaches. (A) We quantify variability in the tensor-based FA, MD, and V1 measurements in MNI space for each of the 48 Johns Hopkins white matter atlas regions. (B) We quantify variability in the NODDI-based cVF, iVF, and ODI measurements in MNI space for each of the 48 Johns Hopkins white matter atlas regions. (C) We quantify variability in bundle shape, volume, FA and length for 43 white matter bundles (Supporting Information Table S1) identified with the RecoBundles segmentation method. (D) We quantify variability in whole brain structural connectomes and the MM, GE, and CPL graph measures.
Figure 3. Example identification of scan pairs at the four levels of variation. The MASiVar dataset consists of scans that can be paired in order to satisfy intrasession, intersession, interscanner, and intersubject criteria. Each of these pairs represent scans that should produce the same measurements, thus quantification of differences within pairs provides an estimate of variability.
Figure 4. Defining variability within a pair of images. (A) Regional FA, MD, cVF, iVF, and ODI variability was defined as the median voxel-wise absolute percent difference. Absolute angular difference was used for V1. (B) Bundle-wise variability was defined as the Dice similarity for shape (1), as the absolute percent difference for volume (2), as the absolute percent difference in median voxel-wise FA for FA (3), and as the absolute percent difference in median streamline-wise length for length (4). (C) Whole connectome variability was defined as the Pearson correlation between connectivity matrices (1), and variability in the MM, GE, and CPL graph measures was defined with absolute percent difference (2-4).
**Figure 5.** Summarizing variability at a given level of variation. (A) DTI, NODDI, and bundle segmentation. After differences within all N image pairs for all M regions or bundles are computed, the regional or bundle-wise medians are taken as the final distribution of variability for a given level of variation. (B) For the connectomics analysis, multiple regions or bundles are not used. Instead, a bootstrapped distribution of medians for each metric is taken as the final distribution of variability.
Figure 6. Variability in DTI. Visualization of differences within intrasession, intersession, interscanner, and intersubject pairs across 48 Johns Hopkins white matter atlas regions illustrates increased variability with session, scanner, and subject effects. Statistical significance was determined with the Wilcoxon signed-rank test.
Figure 7. Variability in NODDI. Visualization of differences within intrasession, intersession, interscanner, and intersubject pairs across 48 Johns Hopkins white matter atlas regions consistently illustrates increased variability with session, scanner, and subject effects. Statistical significance was determined with the Wilcoxon signed-rank test.
Figure 8. Variability in bundle segmentation. Visualization of differences within inrasession, intersession, interscanner, and intersubject pairs across 43 white matter bundles identified with the RecoBundles algorithm (Supporting Information Table S1) consistently illustrates increased variability with session, scanner, and subject effects. Statistical significance was determined with the Wilcoxon signed-rank test.
Figure 9. Variability in connectomics. With the exception of the intersession and interscanner correlation comparison, visualization of differences within intrasession, intersession, interscanner, and intersubject pairs consistently illustrates increased variability with session, scanner, and subject effects. Statistical significance was determined with the Wilcoxon rank-sum test.
Figure 10. Comparing variability across processing approaches. The ratio of intrasubject to intersubject differences averaged across regions or bundles (DTI, NODDI, bundle segmentation) or bootstrapped distributions (connectomics) are shown with error bars denoting standard deviation. With the regional FA in the tensor-based model as a reference, the remaining DTI measurements as well as the NODDI, bundle segmentation, and connectomics measurements generally exhibit good (<0.65) to moderate (<0.95) intrasession and intersession variability, intrasubject variability that is non-negligible compared to intersubject variability, and interscanner effects that approach the size of intersubject effects.