

How to measure functional connectivity using resting-state fMRI? A comprehensive empirical exploration of different connectivity metrics

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ABSTRACT

Background: Functional connectivity in the context of functional magnetic resonance imaging is typically quantified by Pearson's or partial correlation between regional time series of the blood oxygenation level dependent signal. However, a recent interdisciplinary methodological work proposes >230 different metrics to measure similarity between different types of time series.

Objective: Hence, we systematically evaluated how the results of typical research approaches in functional neuroimaging vary depending on the functional connectivity metric of choice. We further explored which metrics most accurately detect presumed reductions in connectivity related to age and malignant brain tumors, aiming to initiate a debate on the best approaches for assessing brain connectivity in functional neuroimaging research.

Methods: We addressed both research questions using four independent neuroimaging datasets, comprising multimodal data from a total of 1187 individuals. We analyzed resting-state functional sequences to calculate functional connectivity using 20 representative metrics from four distinct mathematical domains. We further used T1- and T2-weighted images to compute regional brain volumes, diffusion-weighted imaging data to build structural connectomes, and pseudo-continuous arterial spin labeling to measure regional brain perfusion.

Results: First, our findings demonstrate that the results of typical functional neuroimaging approaches differ fundamentally depending on the functional connectivity metric of choice. Second, we show that correlational and distance metrics are most appropriate to cover reductions in connectivity linked to aging. In this context, partial correlation performs worse than other correlational metrics. Third, our findings suggest that the FC metric of choice depends on the utilized scanning parameters, the regions of interest, and the individual investigated. Lastly, beyond the major objective of this study, we provide evidence in favor of brain perfusion measured via pseudo-continuous arterial spin labeling as a robust neural entity mirroring age-related neural and cognitive decline.

Conclusion: Our empirical evaluation supports a recent theoretical functional connectivity framework. Future functional imaging studies need to comprehensively define the study-specific theoretical property of interest, the methodological property to assess the theoretical property, and the confounding property that may bias the conclusions.

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1. Introduction

Over the past twenty years, the use of connectivity-based methods has played a leading role in characterizing of both the normal brain organization and alterations due to various brain disorders (van den Heuvel and Sporns, 2019). In the context of resting-state functional magnetic resonance imaging (rs-fMRI), functional connectivity (FC) reflects the statistical interdependence of the blood-oxygenation-level-dependent (BOLD) signal of two or more brain regions during rest (Biswal et al., 1995; Fox and Raichle, 2007; van den Heuvel and Hulshoff Pol, 2010). Given an appropriate denoising strategy, the BOLD signal results from changes in cerebral blood flow, volume and oxygenation and is interpreted as an indirect measure of neural activity (Fox and Raichle, 2007; Logothetis and Wandell, 2004). Hence, FC refers to the degree of similarity of neural activation between different brain regions (Aertsen et al., 1989; Friston et al., 1993). The higher the statistical similarity between the neural activity of different brain regions is, the stronger both regions are assumed to be functionally connected (Biswal et al., 1995; van den Heuvel and Hulshoff Pol, 2010).

Studies that utilize FC as an outcome of interest often apply one of the following approaches: First, based on typically occurring FC patterns, the human brain is divided in several functional brain networks such as the default-mode network (Laird et al., 2011; Yeo et al., 2011). Second, these brain networks can also be regarded as macroscale gradients that distribute along different spatial axes (Margulies et al., 2016) and represent developmental markers from adolescence to adulthood (Dong et al., 2021). Third, many studies attempt to relate certain FC patterns to behavioral domains such as cognitive functioning, aiming to clarify the contribution of certain brain regions to observable human behavior. For instance, FC between the hippocampus and the middle frontal gyrus has been associated with multiple cognitive processes such as working memory (Sigurdsson and Duvanci, 2015). Fourth, to identify robust biomarkers of pathological conditions, many approaches examine alterations in certain FC patterns by comparing clinical samples to matched healthy controls (Brandl et al., 2019; Li et al., 2019; Sha et al., 2019). For example, aberrant FC between certain brain networks has been demonstrated in multiple neurological (Tang et al., 2022) and psychiatric conditions (Sha et al., 2019).

Importantly, most studies that follow one of the four approaches use Pearson's correlation or partial correlation to quantify FC. Since the correlation coefficient only covers the linear relationship between two BOLD time series, other metrics aiming to assess FC in rs-fMRI research have been proposed in recent years (Bobadilla-Suarez et al., 2020; Honari et al., 2021; Mahadevan et al., 2021; Mohanty et al., 2020; Phillips et al., 2022; Savva et al., 2019; Wu et al., 2021). Thereby, exploratory evidence demonstrates that the type of FC metric affects the resulting functional brain configuration regarding the number and size of the extracted brain networks and the regions assigned to each network (Mohanty et al., 2020). A recent interdisciplinary work assigns >230 metrics that quantify similarity between different types of time series to the following six categories: basic measures, distance measures, spectral measures, information-theoretic measures, causal measures, and miscellaneous measures (Cliff et al., 2023). These similarity metrics are computed based on various distinct approaches and thus differ in their mathematical properties such as dis-/similarity, directionality, directness, domain, or linearity (Cliff et al., 2023).

With regard to the vast amount of applicable and mathematically different FC metrics, it remains unknown if the abovementioned common approaches in FC research provide stable results independent of the actual FC metric of choice. Therefore, in the first part, we examine the impact of different FC metrics on several common FC-based outcomes in rs-fMRI research. In particular, we assess how FC strength within the default-mode network depends on the actual FC metric of choice. We further inspect the composition of macroscale gradients across different FC metrics. Moreover, we investigate if the association between hippocampal-frontal FC and cognitive functioning remains stable across

all FC metrics. Finally, we study if FC between certain brain networks in patients with schizophrenia differs from healthy controls independent of the chosen FC metric.

Irrespective of potential variations across FC metrics, the above-mentioned and widely used statistical definition of FC fails to differentiate between the theoretical property of interest and the methodological approach to assess this property (Reid et al., 2019). In other words: In many cases, it remains unclear what kind of biological interaction between two brain regions (theoretical property) is covered, when computing the purely statistical correlation between of two BOLD time series (methodological approach). Nowadays, FC has become an inherent entity by itself, although the biological underpinnings are not clarified sufficiently (Reid et al., 2019). This issue is of particular interest in clinically heterogeneous populations without strong and obvious neurological indications such as psychiatric samples (Gratton et al., 2022).

In this work, we assume that FC, on the biological level, reflects the ability of two or more brain regions to synchronize their neural activation patterns, facilitating an efficient information flow between them (theoretical property), whereas the choice of the FC metric is only the method to quantify this ability of signal synchronization (methodological approach). Hence, studying factors that are likely to reduce the ability of different brain regions to synchronize their neural activity reflects an empirical approach to investigate which FC metric is most appropriate metrics to measure connectivity in the human brain.

One such factor is normal aging, which is accompanied by a decline of global and regional grey matter volumes revealed through large-scale normative modelling approaches (Bethlehem et al., 2022; Ge et al., 2024). In addition to ageing, another factor of interest is a severe neurological condition such as a malignant tumor. Malignant high-grade brain tumors are more likely to invade the surrounding healthy brain tissue and thus can lead to disruptions in structural connectivity (Manan et al., 2023). Given the interrelation between structural connectivity and FC (Liu et al., 2023), we assume that such age- and tumor related deteriorations in grey and white matter tissue are likely to impede the brain's ability to effectively synchronize its neural activity between different regions leading to a decline in global FC throughout the later life. Indeed, a recent large-scale normative modelling approach reveals reductions of global FC starting in the late fourth decade of life (Sun et al., 2024). In the following, we use terms such as "reduction of connectivity" to describe the age- and tumor-related disturbed ability of neural signal synchronization.

Hence, in the second part, we aim to determine which FC metric is most suitable to display the presumed reduction of connectivity related to age and malignant brain tumors. We further study with which FC metric associations between age-related reductions of connectivity and cognitive decline are found.

Through our in-depth multimodal and transdiagnostic investigations, we contribute to a broader field of neuroimaging research to better understand if brain connectivity can be quantified in empirical data using metrics other than Pearson's or partial correlation.

2. Methods

2.1. Study samples

The current study was based on four independent datasets acquired at different sites. All research procedures were performed in compliance with relevant laws and institutional guidelines and have been approved by the respective institutional committees. We analyzed behavioral and neuroimaging data from the Mind-Brain-Body dataset from the Max-Planck-Institute Leipzig in Germany (MBB, DOI:10.18112/openneuro.ds000221.v1.0.0, reference number and date of approval: 154/13-ff, 10/02/2013) (Babayan et al., 2019), the Clinical Deep Phenotyping cohort from the Department of Psychiatry and Psychotherapy of the Ludwig-Maximilians-University Hospital Munich in Germany (CDP,

reference number and date of approval: 20–528, 08/28/2020) (Krčmář et al., 2023), the aging cohort of the Human Connectome Project (HCP-Aging, reference number and date of approval: 0925-0667, 03/14/2025) (Harms et al., 2018), and from four patients with malignant tumors from the Brain Tumor Connectomics data from the Ghent University Hospital in Belgium (BTC, DOI:10.18112/openneuro.ds001226.v4.0.0, reference number and date of approval: 2013/881, 03/03/2014) (Aerts et al., 2020; 2018). The CDP and MBB datasets were used to address the first aim of this study, namely to examine the impact of different FC metrics on several common FC-based outcomes in rs-fMRI research. The HCP-Aging cohort, the MBB dataset, and the BTC data were utilized to achieve the study's second objective, targeting the capability of different FC metrics to capture age- and tumor-related reductions of connectivity. Table S1 provides an overview of the sample characteristics. Ages and sexes of the four subjects from the BTC dataset are shown in Figs. 6 and 7.

2.2. MRI data acquisition

The MBB dataset was measured at a Siemens 3T Magnetom Verio scanner with a 32-channel head coil, applying a magnetization-prepared two rapid acquisition gradient echoes (MP2-RAGE) sequence, a T2-weighted sequence, a resting-state functional echo-planar-imaging (EPI) sequence, and a diffusion-weighted imaging (DWI) sequence. A detailed description of the scanning protocols is provided elsewhere (Babayan et al., 2019). The imaging protocol of the CDP study was based on the HCP protocol (Harms et al., 2018). It contained a T1-weighted magnetization-prepared rapid acquisition gradient echo (MP-RAGE) sequence, a T2-weighted sampling perfection with application-optimized contrasts using different flip angle evolution (T2-SPACE), a resting-state EPI sequence and a DWI sequence conducted at a Siemens 3T Magnetom Prisma scanner with a 32-channel head coil (for details see Krčmář et al. (2023)). The HCP-Aging protocol comprised a T1-weighted MP-RAGE sequence, T2-weighted SPACE sequence, a resting-state EPI sequence, a DWI sequence, and a pseudo continuous Arterial Spin Labeling (PCASL) sequence acquired at a Siemens 3T Magnetom Prisma scanner with a 32-channel head coil. Details are provided in Harms et al. (2018). For the BTC dataset, a T1-weighted MP-RAGE sequence, a resting-state functional EPI sequence, and a multi-shell High Angular Resolution Diffusion Imaging (HARDI) sequence were acquired in a Siemens 3T Magnetom Trio MRI scanner using a 32-channel head coil. The entire scanning protocol is described elsewhere (Aerts et al., 2020; 2018). Table S2 provides an overview of the scanning parameters.

2.3. Multimodal MRI data processing

T1- and T2-weighted images of the MBB and HCP-Aging datasets were processed using FreeSurfer v7.2 (Dale et al., 1999; Fischl et al., 2002, 1999). DWI sequences of the MBB, HCP-Aging, and BTC datasets were processed using functions from MRtrix3 v3.0.3 (Tournier et al., 2019), FSL v6.0 (Jenkinson et al., 2012; Smith et al., 2004), FreeSurfer v7.2 (Dale et al., 1999; Fischl et al., 2002; 1999), AFNI v22.1.09 (Cox, 1996; Cox and Hyde, 1997) and ANTS v2.3.5 (Avants et al., 2009). PCASL images of the HCP-Aging dataset were processed using Oxford-ASL from FSL v6.0 (Chappell et al., 2009). fMRIPrep v22.1.1 (Esteban et al., 2019) was utilized to preprocess the resting-state functional MRI images of all datasets. After removal of the first ten volumes and subsequent smoothing (FWHM = 6 mm), the global signal, cerebrospinal fluid signal, white matter signal, and the extracted noise components from Automatic Removal Of Motion Artifacts based on independent component analysis (ICA-AROMA) were regressed from BOLD time series using the *clean_img* function from Nilearn. The denoised BOLD time series were extracted using the *maskers* module from Nilearn. Subject-specific FC was computed for 20 different metrics using the python-based *pyspi* module (Cliff et al., 2023).

Details on the preprocessing steps performed for each MRI modality and dataset and the applied quality control procedures are described in the supplemental information.

2.4. Selection of FC metrics

A recent interdisciplinary study proposed >230 metrics from distinct categories to quantify dis-/similarity between different types of time series and released the python package *pyspi* to compute these metrics (Cliff et al., 2023). We selected 20 metrics from four mathematical categories already used to quantify FC in the context of resting-state fMRI research. We used Pearson's correlation, partial correlation, Spearman's rank correlation, Kendall's tau, and cross-correlation as **correlational metrics**. As **distance metrics**, Euclidean distance, city-block distance, cosine distance, constrained dynamic time warping using the Itakura parallelogram, and constrained dynamic time warping using the Sakoe-Chiba band were computed. Distance metrics were multiplied by minus one prior to data analysis to convert them to similarity measures. Coherence magnitude, phase coherence, phase-locking value, phase-slope index, and spectral Granger causality were assessed as **metrics in the frequency domain**, while mutual information with either a gaussian, kernel-based, or Kraskov-Stögbauer-Grassberger density estimation and transfer entropy with either a gaussian or a Kraskov-Stögbauer-Grassberger density estimation served as **metrics from information theory**. The supplemental information provides a detailed description of these metrics with the respective formulas.

2.5. Multimodal neuroimaging outcomes and cognitive assessments

As outlined previously, the first objective of this study was to investigate the impact of the selected FC metric on several common FC-based approaches in rs-fMRI research. We considered the following four commonly applied approaches: Investigation of default-mode network connectivity, assessment of macroscale gradients, examination of the association between hippocampal-frontal FC and cognition, and identification of FC-based biomarkers for schizophrenia based on a case-control comparison. Details on how these approaches were addressed are provided in the supplemental information.

Concerning the second aim of this study, addressing the sensitivity of different FC metrics regarding age- and tumor-related decline in connectivity, we used the HCP-Aging cohort and the MBB dataset for aging effects, and the BTC dataset to study the impact of malignant tumors. Prior to comparing the FC metrics in terms of their ability to capture age- and tumor-related reductions in connectivity, we first intended to demonstrate that multimodal neural decline was observable in our particular cohorts using other MRI modalities.

With regard to aging effects on neural and cognitive outcomes, we considered T1- and T2-weighted structural MRI data and DWI data, and PCASL data. Structural MRI and DWI were available for the HCP-Aging and MBB datasets, whereas PCASL data was only provided by the HCP-Aging project. Global cognitive functioning in the HCP-Aging cohort was assessed by the NIH toolbox (Hodes et al., 2013).

We used the outputs from FreeSurfer v7.2 gained from the structural MRI data to summarize cortical and subcortical brain volumes based on subcortical parcellations from FreeSurfer and the DKT atlas. We first extracted the regional volumes of several previously defined hub regions in the brain (precuneus, posterior cingulate gyrus, anterior cingulate gyrus, insula, superior frontal gyrus, the pallidum, putamen, thalamus, hippocampus) (Oldham and Fornito, 2019; van den Heuvel and Sporns, 2013). Volumes were z-standardized and averaged to a score indicating the mean volume in these hub regions. We focused on these hub regions because they are characterized by their particularly pronounced connectivity with all other brain regions (Oldham and Fornito, 2019; van den Heuvel and Sporns, 2013).

Considering the DWI data, structural connectivity between subcortical and cortical regions defined by subcortical parcellations from

FreeSurfer and the DKT atlas was estimated as the number of white matter tracts standardized by the size of the respective regions using the `tck2connectome` function from MRtrix3 (Tournier et al., 2019). We extracted the structural connectivity values between the hub regions, z-standardized the values, and averaged them to a global structural connectivity score between all hubs.

In case of the PCASL data, Oxford ASL provided partial volume-corrected measures of the cerebral blood flow for all subcortical and cortical regions specified in the subcortical parcellations from FreeSurfer and the DKT atlas. We extracted the measures of the cerebral blood flow for each hub region, performed a z-standardization, and averaged the values to calculate the mean perfusion across all hubs.

Finally, based on rs-fMRI data, BOLD timeseries of all hub regions defined by the subcortical parcellations of FreeSurfer and the DKT atlas were extracted. FC between all hub regions was calculated for each metric and subject. FC values between all hub regions were z-standardized across all subjects separately for each metric and then averaged across functional connections, resulting in one subject-specific value of mean FC between all hub regions per metric.

We considered DWI and the rs-fMRI data from the BTC dataset to identify biologically plausible disruptions of connectivity related to malignant tumors.

Based on the processed DWI data, we computed structural connectivity between all regions from the multimodal Brainnetome atlas (Fan et al., 2016) using the `tck2connectome` function from MRtrix3 (Tournier et al., 2019). For each patient with a malignant tumor, we extracted structural connectivity values between regions close to the specific tumor location and between the same regions from the contralateral hemisphere. Structural connectivity values were z-standardized and averaged to a subject-specific structural connectivity score between regions that were located close to the individual malignant tumor versus a structural connectivity score between the same regions from the contralateral hemisphere. After extracting the BOLD timeseries of all regions close to the tumor and the same regions in the contralateral hemisphere, the same strategy was applied for all FC metrics computed from rs-fMRI data.

Figs. 1 and 2 illustrate the processing strategies to address the first and second research question, respectively, and Table S5 provides an overview of all neuroimaging outcomes of interest.

2.6. Statistical data analysis

R v4.2.2 was used for statistical data analysis. With regard to the first aim of this study, we evaluated both FC strength within the default-mode network and the composition of macroscale gradients in the MBB dataset descriptively. To investigate if the association between hippocampal-frontal FC and cognitive functioning in the healthy subjects of the CDP cohort remains stable independent of the chosen FC metric, we computed one Bayesian multiple linear regression for each hippocampal-frontal connection and each FC metric with the BACS composite score as dependent variable and hippocampal-frontal FC, age, and sex as predictors. Aiming to evaluate if FC between common brain networks differs between patients with schizophrenia and healthy controls in the CDP cohort independent of the utilized FC metric, we calculated one Bayesian multiple linear regression for each inter-network connection and each FC metric with inter-network FC as dependent variable and group (schizophrenia, healthy controls), age, and sex as predictors.

Regarding the second objective of this study, we explored the association between aging and multimodal neural and cognitive decline in the HCP-Aging and MBB datasets. We used Bayesian multiple linear regressions with either mean volume, mean perfusion, mean structural connectivity, and mean functional connectivity (for all 20 metrics) of the hub regions as dependent variables and age and sex as predictors. Note that we used age as a binary predictor in the case of the MBB dataset (younger adults between 20 and 35 years versus older adults between 55 and 80 years), because only age categories instead of actual ages of participants were published for this dataset. Moreover, we calculated Bayesian multiple linear regressions with the cognitive composite score from the NIH toolbox as dependent variable and either mean volume, mean perfusion, mean structural connectivity, and mean functional connectivity (for all 20 metrics) of the hub regions and sex as predictors.

1st Research Question: Effects of choice of FC metric on results in typical functional neuroimaging studies

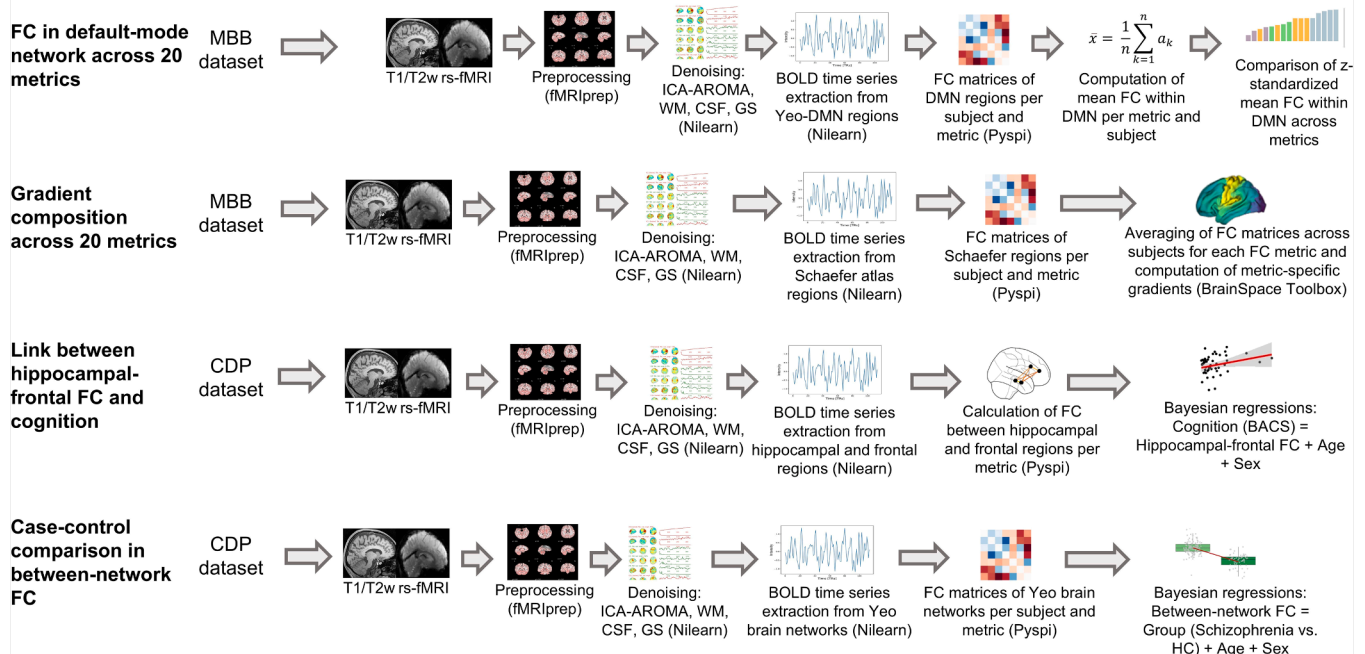


Fig. 1. Multimodal processing strategy for the first research question.

This figure illustrates the processing strategies to address the first research question of this study.

2nd Research Question: Sensitivity of FC metrics regarding disruptions of connectivity

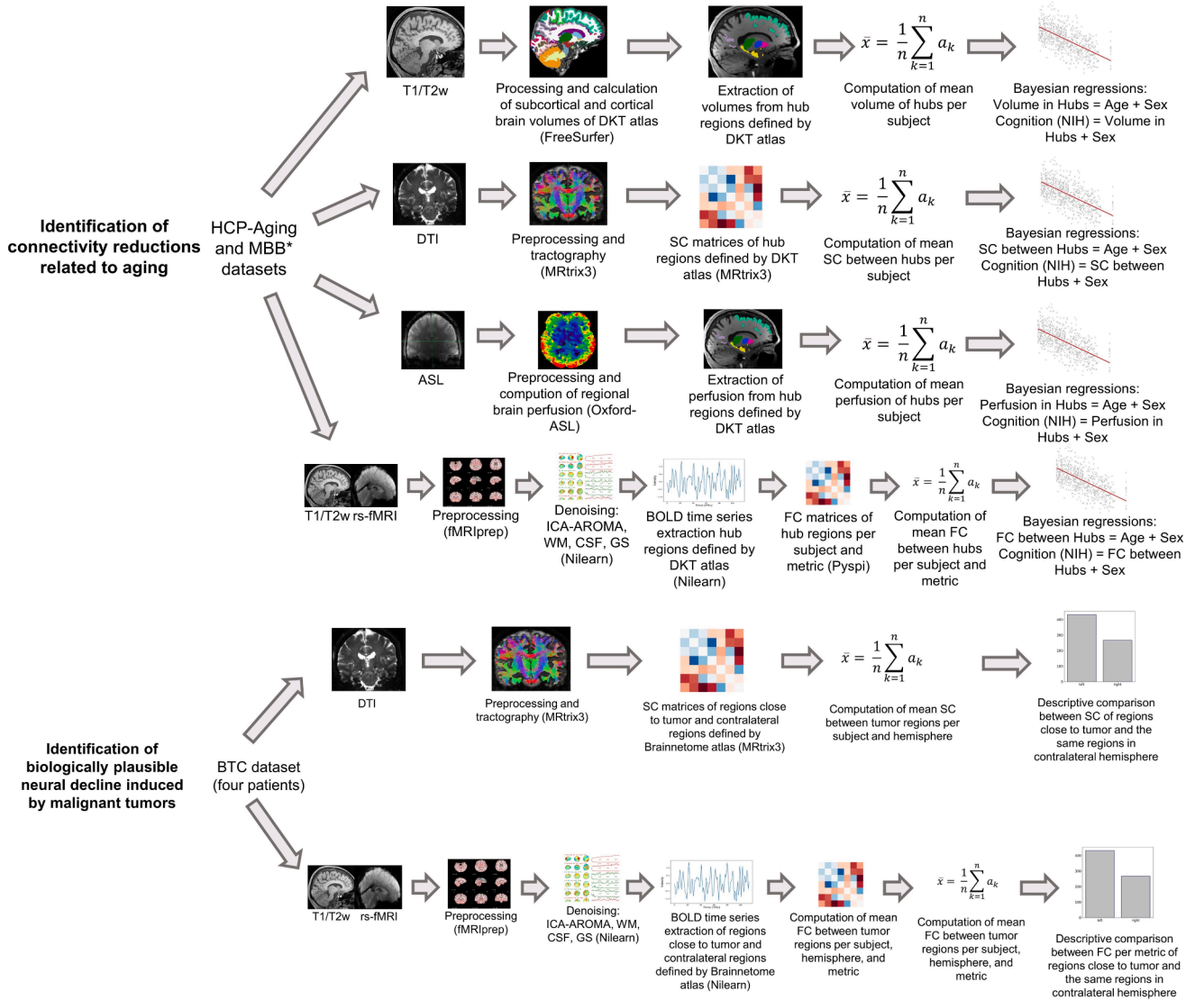


Fig. 2. Multimodal processing strategy for the second research question

This figure illustrates the processing strategies to address the first research question of this study. *, in case of the MBB dataset no PCASL data were acquired and neural outcomes were not linked to cognitive performance.

Structural connectivity and metric-specific FC between regions close to the malignant tumor and the same regions from the contralateral hemisphere were compared descriptively.

The main test statistics of interest from the Bayesian multiple linear regression computed with the *brms* package (Bürkner, 2017) was Jeffrey's default Bayes Factor (BF_{10}), representing a continuous, relative measure of evidence the data is providing for the alternative hypothesis ($H_1: \beta_z \neq 0$) compared to the null hypothesis ($H_0: \beta_z = 0$) (Ly et al., 2016). A BF_{10} between one and three reflects anecdotal evidence for the alternative hypothesis, between three and ten is considered to be moderate evidence, between ten and 30 is labeled as strong evidence, between 30 and 100 is seen as very strong evidence, and a BF_{10} above 100 is decisive evidence for the alternative hypothesis (Lee and Wagenmakers, 2013).

3. Results

3.1. Default-mode network connectivity across FC metrics

We compared the absolute connectivity strength for all 20 FC metrics across both hemispheres. Based on the distance metrics, we obtained the highest absolute connectivity strength within the DMN followed by the correlational metrics, whereas the information-theory and frequency metrics showed the lowest. Among the correlational metrics, DMN connectivity was the lowest for partial correlation (Fig. S1).

3.2. Composition of macroscale functional gradients across FC metrics

Depending on the FC metric used, gradient patterns vary in their graphical representation, as indicated in Fig. S2. Especially, phase coherence, dynamic time warping with an Itakura constraint, spectral Granger causality, and transfer entropy with a Kraskov-Stögbauer-Grassberger density estimation do not show patterns related to the DMN as proposed in Margulies et al. (2016). In contrast, the remaining FC

metrics show similar patterns, most dominantly reflected by the correlational metrics. Importantly, the first and second gradients were inverted for some metrics compared to Margulies et al. (2016).

3.3. Association between hippocampal-frontal FC and cognition across FC metrics

Fig. 3A illustrates the β -coefficients and BF_{10} of the respective hippocampal-frontal connection extracted from the Bayesian multiple linear regression that assessed the association between hippocampal-frontal FC and cognition using the healthy controls from the CDP cohort. Results indicate that depending on the FC metric of choice a different number and different types of hippocampal-frontal connections show associations with cognitive functioning. For instance, when using Pearson's correlation, Spearman's correlation, Kendall's tau, Euclidean distance, Cityblock distance, Cosine distance or dynamic time warping with a Sakoe-Chiba band, higher FC between the left parahippocampal gyrus and the left caudal anterior cingulate gyrus is related to better cognitive performance. This finding is not reproduced in the case of partial correlation or any other FC metric, demonstrating that the choice of the FC metric affects the results and interpretations of brain-behavior associations.

3.4. Dysconnectivity patterns in schizophrenia across FC metrics

Fig. 3B shows the β -coefficients and BF_{10} of the predictor group extracted from the Bayesian multiple linear regression that covered the case-control differences in FC between brain networks based on patients with schizophrenia and healthy controls from the CDP study. Results reveal that depending on the FC metric of choice different numbers and types of dysconnectivity patterns are found. For example, when using partial correlation, coherence magnitude or phase-locking value, patients with schizophrenia reveal a hyperconnectivity between the visual network and the dorsal attention network. At the same time, this is not the case for any other FC metric. Hence, the choice of the FC metric impacts the results of disorder-related dysconnectivity patterns.

3.5. Sensitivity of FC metrics regarding age-related decline in connectivity and cognition

Figs. 4 and 5 visualize the associations between age and cognition and multimodal neural outcomes in the hub regions and the test statistics extracted from the respective Bayesian multiple linear regressions based on the HCP-Aging and the MBB cohort.

With regard to the HCP-Aging dataset (Fig. 4A–D), we found decisive evidence that the older the participants were, the lower their mean volume (Fig. 4A), structural connectivity (Fig. 4B), and perfusion (Fig. 4C) in the hub regions was, on average. The strongest effects were found for volume, followed by perfusion and structural connectivity. These findings demonstrate that multimodal age-related neural decline in the hub regions was present in the HCP-Aging cohort. Considering the rs-fMRI data (Fig. 4D), our findings reveal decisive evidence that FC between the hub regions decreases with age, when using Pearson's correlation, Spearman's correlation, Kendall's tau, cross correlation, Euclidean distance, or cosine distance as FC metric. Correspondingly, we found very strong evidence for this association with Cityblock distance and strong evidence when utilizing Gaussian mutual information. In the case of partial correlation and dynamic time warping with a Sakoe-Chiba band we only found anecdotal evidence, but strong and anecdotal evidence for phase coherence and gaussian transfer entropy towards a positive relation between age and FC in hub regions, respectively. Effect sizes were overall smaller than in the other MRI modalities.

With respect to the MBB dataset (Fig. 4E–G), we found decisive evidence that elderly, on average, had lower mean volume (Fig. 4E) and structural connectivity (Fig. 4F) in the hub regions than younger

participants. The effect for volume was stronger than the effect for structural connectivity. These findings demonstrate that multimodal age-related neural decline in the hub regions was also observable in the MBB cohort. Considering the rs-fMRI data (Fig. 4G), our findings demonstrate decisive evidence that the elderly show lower FC between the hub regions than younger participants, when using cross-correlation and dynamic time warping constrained by an Itakura parallelogram. We also obtained strong evidence for this group difference in the case of Pearson's correlation, Spearman's correlation, Kendall's tau, and cosine distance, but moderate and anecdotal evidence by Euclidean distance and partial correlation, respectively. Higher FC between the hub regions in elderly was evident when using dynamic time warping with a Sakoe-Chiba band, coherence magnitude, phase coherence, phase-locking value, spectral Granger causality, and both transfer entropy versions. Apart from dynamic time warping constrained by an Itakura parallelogram, effect sizes were smaller than in the other MRI modalities.

Referring back to the HCP-Aging cohort, we obtained very strong and decisive evidence that larger mean volumes (Fig. 5A) and higher mean perfusion (Fig. 5C) in the hub regions, respectively, were linked to better cognitive performance in the HCP-Aging cohort. The effect for perfusion was stronger than the effect for volume. We did not encounter such an association for structural connectivity between hub regions (Fig. 5B). These findings demonstrate that age-related cognitive decline in the hub regions was present for volumes and perfusion in the HCP-Aging cohort. Regarding the rs-fMRI data (Fig. 5D), only anecdotal evidence was observed, suggesting that higher FC between the hub regions was linked to better cognitive performance, when using Pearson's correlation, Kendall's tau, cosine distance, and dynamic time warping constrained with an Itakura parallelogram. Effect sizes were smaller than in structural MRI and PCASL.

3.6. Sensitivity of FC metrics regarding tumor-related decline of connectivity

Figs. 6 and 7 display structural connectivity and FC data from the BTC dataset for four patients with malignant tumors.

Our findings indicate that the average number of white matter tracts between regions close to the subject-specific tumor location was smaller compared to the same regions in the contralateral hemisphere for the first three subjects, whereas this was not the case for the healthy controls included in the BTC dataset. For the first subject (Fig. 6A), lower FC between regions close to the subject-specific tumor location compared to the same regions in the contralateral hemisphere was observed, when using Pearson's correlation, Spearman's correlation, Kendall's tau, Euclidean distance, Cityblock distance, cosine distance, dynamic time warping constrained by a Sakoe-Chiba band, or the phase-slope index as FC metrics. In case of the second subject (Fig. 6B), only kernel-based mutual information and both transfer entropy metrics revealed corresponding results. In contrast, correlational and distance metrics, as well as several frequency metrics indicated an effect in the opposite direction. For the third subject (Fig. 7A), no FC metric except phase coherence and dynamic time warping constrained by an Itakura parallelogram revealed clear differences in FC between regions close to the subject-specific tumor location, which was smaller compared to the same regions in the contralateral hemisphere. In case of the fourth subject (Fig. 7B), the average number of white matter tracts did differ substantially between both hemispheres in contrast to the healthy controls. Only phase coherence and spectral Granger causality showed a corresponding pattern.

4. Discussion

Based on four independent multimodal MRI data sets, our study first explored if FC metrics with distinct mathematical properties lead to different conclusions in the context of typical FC-based research questions. Furthermore, we evaluated whether FC metrics other than

Fig. 3. Brain-behavior associations and dysconnectivity patterns between cases and controls for 20 metrics

This figure displays the results from Bayesian multiple linear regression analyses, addressing the associations between hippocampal-frontal FC and cognitive functioning (A) and the FC differences between patients with schizophrenia and healthy controls (B) across 20 FC metrics. Each dot reflects one Bayesian multiple linear regression. (A) The β -coefficient of the respective hippocampal-frontal connection is shown on the x-axis and the BF10 of this predictor is illustrated on the y-axis. A positive β indicates a positive association between the particular functional connection and cognition, and a negative β reveals a negative association. (B) The β -coefficient of the predictor group (schizophrenia vs. controls) is shown on the x-axis and the BF10 of this predictor is illustrated on the y-axis. A positive β indicates a higher FC between the respective networks in patients with schizophrenia and a negative β reveals lower FC. The higher the BF10, the darker the color of the dots. Functional connections are labelled if the BF10 is higher than one. Pearson corr, Pearson's correlation; Partial corr, partial correlation; Spearman corr, Spearman's correlation; Kendall tau, Kendall's tau; Cross corr, cross correlation; Euclidean dist, Euclidean distance; Cityblock dist, Cityblock distance; Cosine dist, Cosine distance; DTW itakura, dynamic time warping constrained with Itakura parallelogram; DTW sakoe-chiba, dynamic time warping constrained with Sakoe-Chiba band; COH magnitude, coherence magnitude; COH phase, phase coherence; PLV, phase-locking value; PSI, phase slope index; SGC, spectral Granger causality; MI gaussian, mutual information with gaussian density estimation; MI kernel, mutual information with kernel-based density estimation; MI kraskov, mutual information with Kraskov-Stögbauer-Grassberger density estimation; TE gaussian, transfer entropy with gaussian density estimation; TE kraskov transfer entropy with Kraskov-Stögbauer-Grassberger density estimation; L, left hemisphere; R, right hemisphere; PHIG, parahippocampal gyrus; CMFG, caudal middle frontal gyrus; CACG, caudal anterior cingulate gyrus; RACG, rostral anterior cingulate gyrus; RMFG, rostral middle frontal gyrus; EC, entorhinal cortex; HI, hippocampus; VN, visual network, SMN, somatomotor network; DAN, dorsal attention network; VAN, ventral attention network; LN, limbic network; FPN, fronto-parietal network; DMN, default-mode network; H_0 , null hypothesis; H_1 , alternative hypothesis.

Pearson's or partial correlation are more appropriate for detecting reductions of connectivity related to aging or malignant brain tumors.

4.1. Influence of FC metrics on typical research approaches in functional neuroimaging

Our findings suggest that the selected FC metric affects the results and conclusions in several common FC-based research approaches, such as examining FC within the default-mode network, exploring the composition of macroscale gradients, investigating brain-behavior associations, and studying disorder-related dysconnectivity patterns. This lines with previous evidence demonstrating that distinct FC metrics computed within the same dataset lead to substantial differences in the resulting brain network configuration (Mohanty et al., 2020). Hence, the choice of FC metric represents an essential step in studies including FC as a relevant outcome, because the resulting conclusions on the particular research questions will be fundamentally different. With respect to the noticeable mathematical differences of potentially applicable FC metrics, the resulting variation in typical FC-based outcomes is not surprising. In particular, the FC metrics examined in this study differ in terms of their type (similarity or dissimilarity measure or both), directionality (non-directional, unidirectional, or bidirectional), directness (direct or non-direct measure), domain (time or frequency), and linearity assumption (linear or non-linear) (Bastos and Schoffelen, 2015; Cliff et al., 2023). For instance, partial correlation reflects both similarity and dissimilarity between two BOLD time series. It does not assume a direction of the association, considers the influence of other regions, and thus assesses the direct association between the two regions of interest. Moreover, partial correlation captures similarity in the time domain and includes a linearity assumption. The supplemental information provides an overview of the mathematical properties of all applied FC metrics.

4.2. Superiority of correlational and distance metrics in detecting age-related decline of connectivity

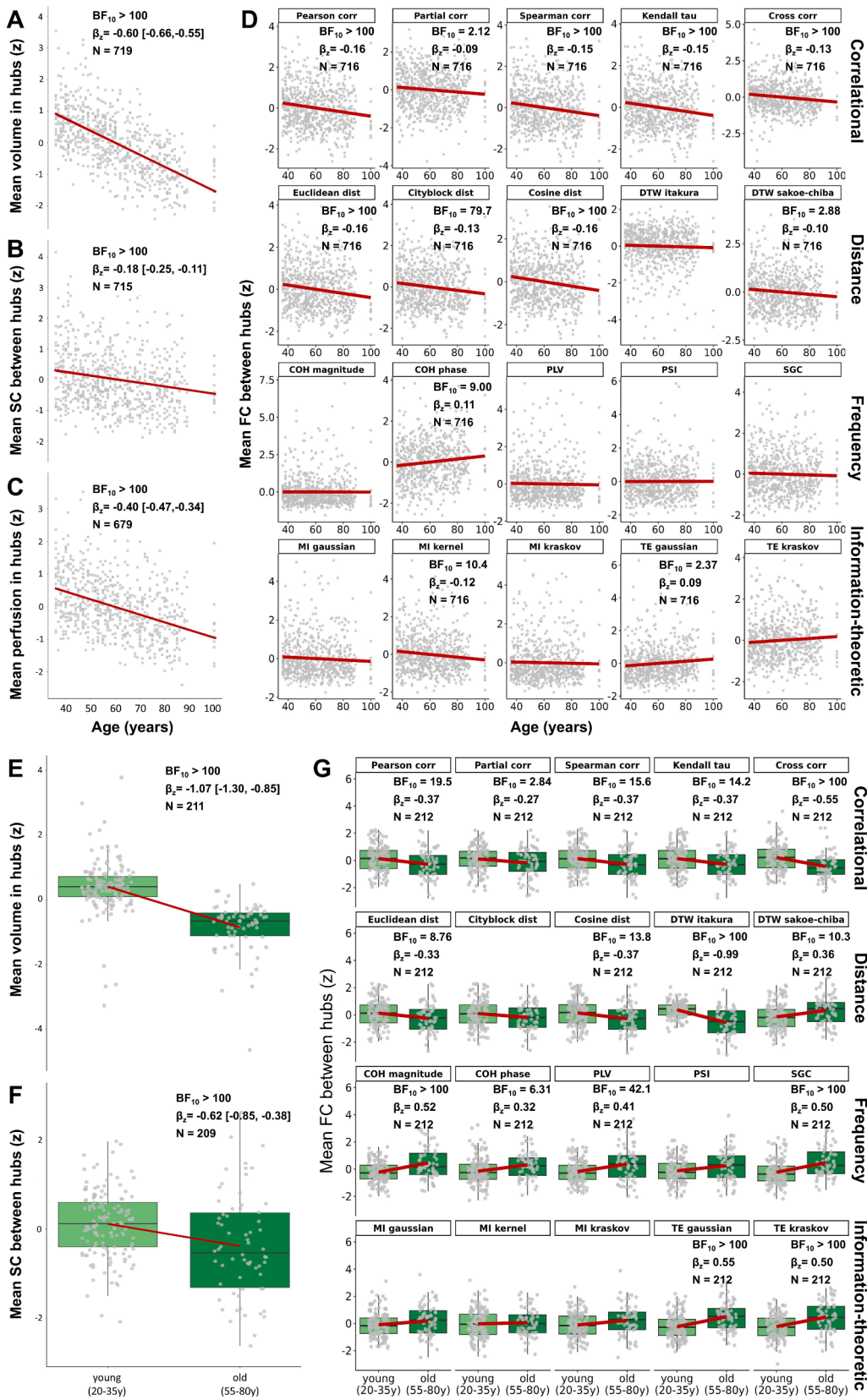
Apart from demonstrating the impact of different FC metrics on the results in common FC-based research approaches, our findings reveal that correlational and distance metrics most accurately capture connectivity decline. Although generally in line with previous simulation studies that also suggest the superiority of correlational measures in detecting specific alterations in brain networks (Smith et al., 2011), we observed that partial correlation only provided anecdotal evidence in favor of age-related connectivity decline, whereas other correlational and distance metrics indicated more robust associations. In contrast, partial correlation was found to be one of the most accurate FC metrics when tested in simulated BOLD time series (Smith et al., 2011) and is generally suggested to be less prone to confounds resulting from other

neural entities (Reid et al., 2019). Despite these convincing examinations and the underlying reasonable theoretical foundation, our findings suggest that using partial correlation should have some additional consideration. Here, we used a rather global measure of FC as the main outcome by averaging the subject-specific FC values per metric between all hub regions. If a certain brain region is highly connected to many other regions, it is more likely to influence FC between all other regions. When not controlling for its impact on other functional connections, this highly connected region will be weighted stronger than less connected regions in an average FC value used in this study. Given that we demonstrated an age-related decline of white matter tracts between hub regions, a summarizing FC that gives more weight to highly connected regions may be beneficial in covering decline of connectivity and cognition related to age. This emphasizes the role of directness as one important property of FC metrics.

However, directness is not the only important property, as FC metrics in the frequency domain and from information theory did not capture age-related decline of connectivity and cognition properly, despite also covering indirect associations between brain regions like most correlation and distance metrics. In the case of frequency metrics computed from BOLD time series, previous simulation studies raise concerns about their sensitivity regarding alterations in the underlying network architecture (Smith et al., 2011), which corresponds to our empirical findings. These concerns also apply to the directional metrics we examined, namely spectral Granger causality and transfer entropy, whose utility for BOLD time series has been questioned (Smith et al., 2011). Finally, mutual information as an FC metric that also covers non-linear associations between BOLD time series has been suggested to be a promising alternative to correlational metrics (Pereda et al., 2005), but our results do not support its use to assess age-related decline in connectivity and cognition. This again corresponds to evidence in simulated data (Smith et al., 2011) and is in line with theoretical considerations emphasizing that non-linear metrics of FC are not necessarily better than linear metrics, but rather address distinct aspects of the similarity between two signals (Pereda et al., 2005). In sum, our empirical examination supports the use of correlational and distance metrics in order to detect reductions of connectivity and impaired cognition related to age, while questioning the utility of partial correlation at least for summarizing FC scores across different regions.

4.3. Variability across datasets, brain regions, and individuals

In addition to this general tendency towards the superiority of correlational and distance metrics, we obtained dataset-specific effects that have also been reported previously (Bobadilla-Suarez et al., 2020). Specifically, when using dynamic time warping constrained by an Itakura parallelogram as FC metric, age-related connectivity decline was robustly identified in the MBB dataset, whereas no effect was observed



(caption on next page)

Fig. 4. Associations between age and multimodal neural outcomes for the HCP-Aging and MBB cohort

This figure depicts the associations between age and mean volume, structural connectivity, perfusion, and functional connectivity in the hub regions, as well as the test statistics of the respective Bayesian multiple linear regressions. Each dot represents a healthy subject from the HCP-Aging (A-D) and the MBB (E-G) cohort. The red regression line reflects the correlation between age and the particular neural outcome. A) Association between age and mean volume in hub regions with age displayed on the x-axis and mean volume on the y-axis. B) Association between age and mean structural connectivity between hub regions with age displayed on the x-axis and mean structural connectivity on the y-axis. C) Association between age and mean perfusion in hub regions with age displayed on the x-axis and mean perfusion on the y-axis. D) Association between age and mean FC between hub regions with age displayed on the x-axis and mean FC on the y-axis for each FC metric. E) Difference in mean volume in hub regions with the age groups displayed on the x-axis and mean volume on the y-axis. F) Difference in mean structural connectivity between hub regions with the age groups displayed on the x-axis and mean structural connectivity on the y-axis. G) Difference in mean FC between hub regions with the age groups displayed on the x-axis and mean FC on the y-axis for each FC metric. Pearson corr, Pearson's correlation; Partial corr, partial correlation; Spearman corr, Spearman's correlation; Kendall tau, Kendall's tau; Cross corr, cross correlation; Euclidean dist, Euclidean distance; Cityblock dist, Cityblock distance; Cosine dist, Cosine distance; DTW itakura, dynamic time warping constrained with Itakura parallelogram; DTW sakoe-chiba, dynamic time warping constrained with Sakoe-Chiba band; COH magnitude, coherence magnitude; COH phase, phase coherence; PLV, phase-locking value; PSI, phase slope index; SGC, spectral Granger causality; MI gaussian, mutual information with gaussian density estimation; MI kernel, mutual information with kernel-based density estimation; MI kraskov, mutual information with Kraskov-Stögbauer-Grassberger density estimation; TE gaussian, transfer entropy with gaussian density estimation; TE kraskov transfer entropy with Kraskov-Stögbauer-Grassberger density estimation; BF₁₀, Bayes factor of the predictor age; β_z , standardized beta coefficient of the predictor age and 95 % confidence interval; N, sample size considered in the respective analysis; SC, structural connectivity; FC, functional connectivity.

in the HCP-Aging cohort. The utilized MRI scanners and the underlying scanning parameters differed noticeably between both projects, as the MBB protocol used a repetition time of 1400 ms with 657 timepoints, while the single HCP-Aging sequences had a repetition time of 800 ms with 478 timepoints. Given that especially the sequence length and repetition time have been shown to affect the sensitivity of FC metrics towards network alterations (Smith et al., 2011; Wang et al., 2014), it appears plausible that the FC metric of choice, to a certain degree, depends on the acquired EPI sequence. Hence, our findings indicate that methodological considerations regarding the scanning parameters need to be considered, but it remains to be determined which FC metric suits best under which scanning conditions in empirical data.

In addition to the impact of the scanning sequence, we also observed inter-individual variation between four patients with malignant tumors from the BTC dataset. Particularly, in the case of the first patient (Fig. 6A), correlational and distance metrics were sensitive to white matter decline between regions close to the tumor. However, with regard to the other patients, this was not the case, as other FC metrics such as transfer entropy or kernel-based mutual information (2nd patient, Fig. 6B), phase coherence and dynamic time warping constrained by an Itakura parallelogram (3rd patient, Fig. 7A) or partial correlation and the frequency metrics (4th patient, Fig. 7B) mimicked the existing white matter decline in the ipsilateral compared to the contralateral hemisphere. While having been scanned under the same conditions, the four patients differ in terms of age, sex, tumor location, and tumor type (only 4th patient). Consequently, these descriptive observations could suggest that the choice of the appropriate FC metric may depend on the individuals characteristics or on the regions of interest.

4.4. Embedding current findings into a theoretical FC framework

Our empirical findings can be embedded in the theoretical FC framework proposed by Reid et al. (2019). They categorize properties essential for mechanistic inferences from FC data into three types: theoretical, methodological, and confounding properties. Theoretical properties refer to the characteristics of the neural connections or pathways investigated (e.g., directionality between two neural assemblies). Methodological properties comprise all methodological approaches to assess these theoretical properties (e.g., type of FC metric). Confounding properties consist of any factors that may induce bias (e.g., motion artifacts). Reid et al. (2019) emphasize the need for empirical validations of methods used in FC research and suggest that future FC studies should comprehensively describe and discuss the three properties of the framework in the context of their particular research question to refine their conclusions.

Our approach aimed for empirical validation of different FC metrics (methodological property) to explore deteriorations in connectivity related to age and malignant tumors (theoretical property). Importantly,

our findings serve as an empirical endorsement of the FC framework proposed by Reid et al. (2019), as indicated by the following examples: First, using partial correlation as FC metric eliminates the confounding effect of unmeasured neural activity (confounding property), but proves less effective than other FC metrics that do not consider directness in detecting global connectivity decline (theoretical property). Second, the dataset-specific effects described above reveal that the selection of FC metrics (methodological property) to assess age-related connectivity decline (theoretical property) may depend on the parameters of the underlying scanning protocol (methodological property). Lastly, the findings in patients with malignant tumors show that the choice of FC metric (methodological property) may also depend on the type and location of the brain regions of the individual between which neural impairments are assessed (theoretical property). These examples demonstrate that no single FC assessment is superior; the optimal approach depends on the theoretical property of interest, other methodological properties than the choice of the FC metric itself, and relevant confounding properties.

4.5. Brain perfusion measured by PCASL as a neural representation of age-related cognitive decline

Lastly, beyond the major scope of this study, we observed that volumetric and perfusion-based measures in the hub regions gained from structural MRI and PCASL are more stable in mirroring multimodal neural and cognitive decline than structural connectivity and FC outcomes assessed by DWI and rs-fMRI, respectively. This is particularly interesting with respect to PCASL since this sequence is not yet commonly used in clinical neuroimaging research. According to our findings, cerebral blood flow in the hub regions assessed by PCASL decreases substantially with age and shows the strongest associations with age-related cognitive decline. Notably, cognitive impairments reflect a robust transdiagnostic phenomenon in various fields such as psychiatry (Goodkind et al., 2015), but associations between cognitive functioning and the underlying neural processes are mostly small and often not reliably detected (Gratton et al., 2022; Marek et al., 2022). The robust correlation between reductions in the cerebral blood flow of central brain hubs and age-related cognitive decline identified in the current work can serve as a promising basis for future clinical neuroimaging studies that aim to identify the neural underpinnings of cognitive deficits. Thus, our results may leverage the use of PCASL to investigate neural correlates of behavior in human neuroimaging.

4.6. Study limitations and future directions

Our study navigates through multiple FC metrics with room for enhancement, while maintaining a solid foundation for future exploration. First, our empirical validation of FC requires knowledge about the

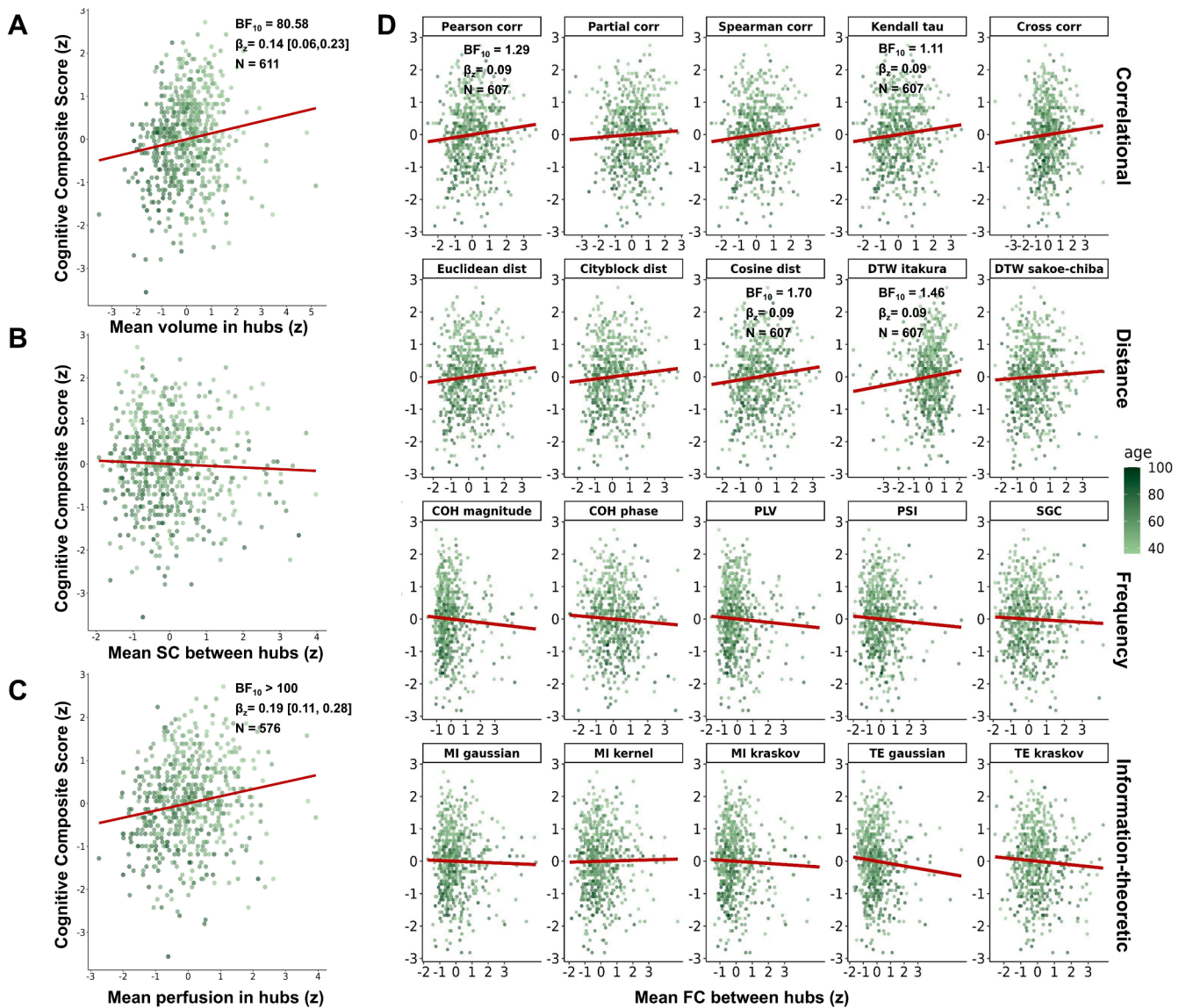


Fig. 5. Associations between multimodal neural outcomes and global cognition for the HCP-Aging cohort

This figure shows the associations between mean volume, structural connectivity, perfusion, and functional connectivity in the hub regions and the cognitive composite score, as well as the test statistics of the respective Bayesian multiple linear regressions. Each dot represents an individual subject from the HCP-Aging cohort. The red regression line reflects the correlation between the particular neural outcome and cognitive performance. A) Association between mean volume in hub regions and cognition with mean volume displayed on the x-axis and the cognitive composite score on the y-axis. B) Association between mean structural connectivity between hub regions and cognition with mean structural connectivity displayed on the x-axis and the cognitive composite score on the y-axis. C) Association between mean perfusion in hub regions and cognition with mean perfusion displayed on the x-axis and the cognitive composite score on the y-axis. D) Association between mean FC between hub regions and cognition with mean FC between hub regions displayed on the x-axis and the cognitive composite score on the y-axis. Pearson corr, Pearson's correlation; Partial corr, partial correlation; Spearman corr, Spearman's correlation; Kendall tau, Kendall's tau; Cross corr, cross correlation; Euclidean dist, Euclidean distance; Cityblock dist, Cityblock distance; Cosine dist, Cosine distance; DTW itakura, dynamic time warping constrained with Itakura parallelogram; DTW sakoe-chiba, dynamic time warping constrained with Sakoe-Chiba band; COH magnitude, coherence magnitude; COH phase, phase coherence; PLV, phase-locking value; PSI, phase slope index; SGC, spectral Granger causality; MI gaussian, mutual information with gaussian density estimation; MI kernel, mutual information with kernel-based density estimation; MI kraskov, mutual information with Kraskov-Stögbauer-Grassberger density estimation; TE gaussian, transfer entropy with gaussian density estimation; TE kraskov, transfer entropy with Kraskov-Stögbauer-Grassberger density estimation; BF_{10} , Bayes factor of the predictor age; β_z , standardized beta coefficient of the predictor age and 95 % confidence interval; N , sample size considered in the respective analysis; SC, structural connectivity; FC, functional connectivity.

ground truth of aberrant FC patterns in the brain (Reid et al., 2019). We assumed that ageing and tumors lead to a decline in the ability of the hub brain regions to synchronize their activation patterns and thus result in a reduction of efficient information flow (theoretical property). Hence, an appropriate functional connectivity metric (methodological property) should mirror this effect. However, especially in the case of brain tumors it may be possible that our presumed ground truth may not hold. Patterns of increased FC have also been observed in such patients

(Fox and King, 2018) and, more generally, may reflect a pathological rather than a healthy brain state in neurological disorders like Alzheimer's disease (Roemer-Cassiano et al., 2025). We used a multimodal approach to demonstrate that the mean volume, perfusion, and structural connectivity in the hub regions decrease with age among different datasets, assuming that this decline should also manifest in decreased FC between the hubs. This served as the ground truth presumed in this study. Given the known link brain structure and function (Liu et al.,

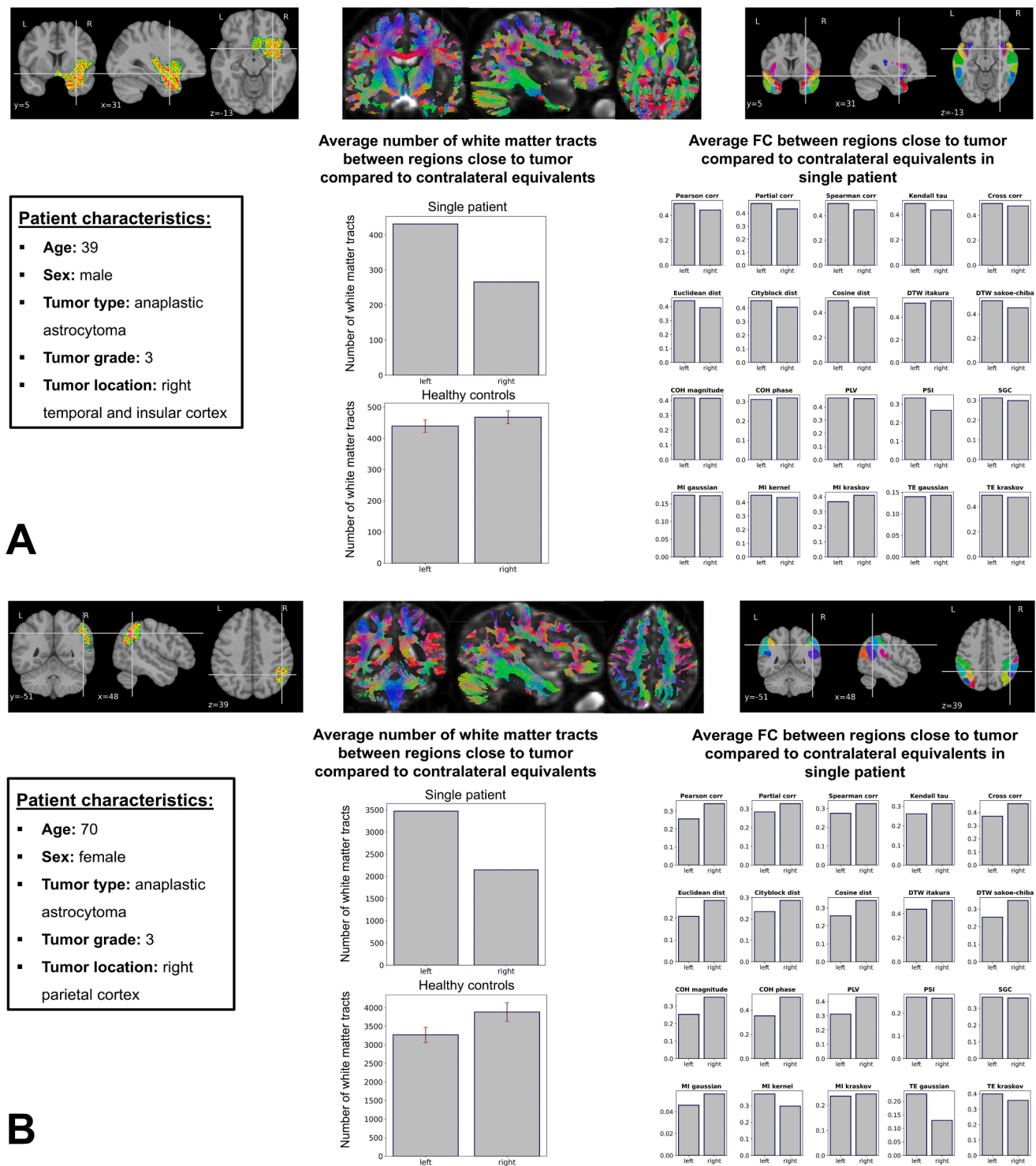


Fig. 6. Comparison of white matter tracts and FC between regions close to tumor and contralateral equivalents

This figure shows the average number of white matter tracts between regions close to the subject-specific tumor location compared to the same regions in the contralateral hemisphere for two subjects with malignant tumors and for respective healthy controls of the BTC dataset. In each panel (A-B), the tumor's location is shown on the top left, the respective tractography on the top middle, and the regions from the Brainnetome atlas used to compute the average number of white matter tracts and FC on the top right. Patient characteristics and the comparison in the average number of white matter tracts and FC between selected regions from both hemispheres are illustrated.

2023), this assumption stands on biologically plausible ground and is supported by recent large-scale normative modeling approach on the functional connectome across the life span (Sun et al., 2024). Nonetheless, the relationship between brain structure and function is not fully explicit due to complex multi-synaptic interactions (Zamani Esfahlani

et al., 2022). There may persist an uncertainty if age-related decreases in FC between the hub regions truly reflect a stable and replicable ground truth. It may be possible that different FC metrics address different theoretical properties in the brain. In other words: Distinct metrics may cover different aspects of neural interactions between brain regions.

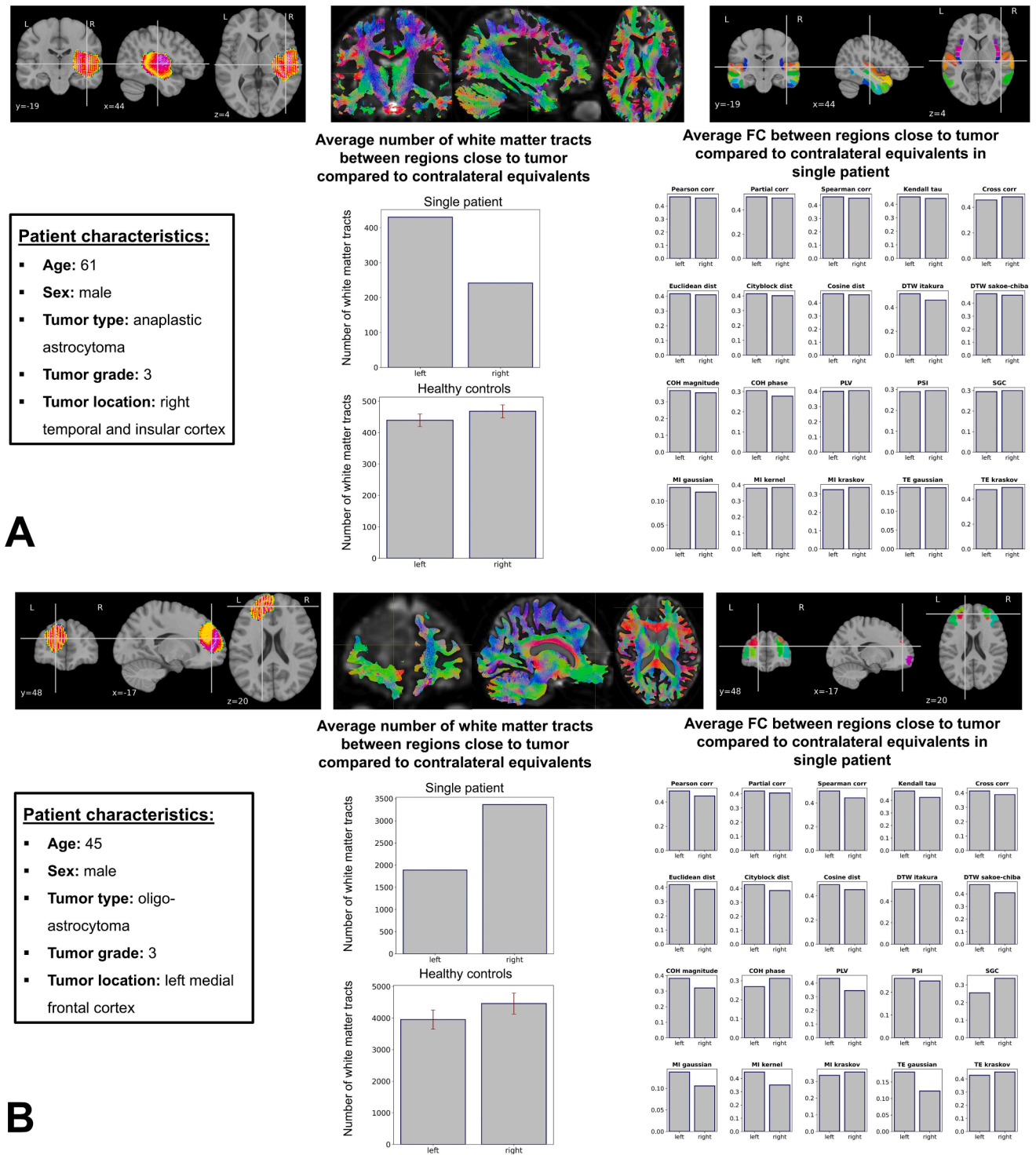


Fig. 7. Comparison of white matter tracts and FC between regions close to tumor and contralateral equivalents

This figure shows the average number of white matter tracts between regions close to the subject-specific tumor location compared to the same regions in the contralateral hemisphere for two subjects with malignant tumors and for respective healthy controls of the BTC dataset. In each panel (A-B), the tumor's location is shown on the top left, the respective tractography on the top middle, and the regions from the Brainnetome atlas used to compute the average number of white matter tracts and FC on the top right. Patient characteristics and the comparison in the average number of white matter tracts and FC between selected regions from both hemispheres are illustrated.

Some of these aspects may be affected by ageing and tumors, while others may not. In this case, a reduction of FC would not necessarily mean that the respective FC metric is more appropriate. This is important to consider when interpreting our current findings.

Based on this discussion, future examinations could combine

empirical factors that induce strong neural impairments with model-based simulated data to bolster biologically plausible assumptions and to decrease the uncertainty about the ground truth in FC research.

Secondly, we conclude that selecting a proper FC metric depends on the acquired EPI sequence and the type and location of the connections

of interest. While our data do not provide clear-cut guidelines for metric selection, future studies could evaluate how certain scanning parameters, such as the session length or repetition time, affect the utility of different FC metrics in capturing connectivity decline. This could involve leveraging respective biophysical models or simulating specific brain networks to test if distinct brain pathways require different FC metrics to evaluate the sensitivity of several FC metrics regarding known patterns of information flow within these networks.

Thirdly, our selection of 20 representative metrics from four domains, based on their previous application in fMRI research, opens the field for further inquiry time series (Cliff et al., 2023). Future investigations are encouraged to test the applicability of these yet-unutilized metrics, as it has been done in a recent comprehensive work comparing all metrics included in the pypsi package (Liu et al., 2024).

Fourth, we did not systematically evaluate if our results depend global signal regression, as it was not the primary focus of this work. However, future studies in this field should incorporate such analysis, given that the global signal regression is known to affect FC (Murphy and Fox, 2017).

Fifth, we did not have proper longitudinal data available to proof the test-retest reliability of different FC metrics over time. Future studies should address this issue to provide insights on the longitudinal stability of different FC metrics.

5. Conclusion

To conclude, we first provide empirical evidence that the utilized FC metric strongly affects the results in the context of typical fMRI research approaches. Secondly, our results demonstrate that correlational and distance metrics perform best in detecting age- and tumor-related decline in connectivity, while questioning the utility of partial correlation when summarizing FC scores across regions are used. Thirdly, we demonstrate that the sensitivity of FC metrics towards connectivity decline is influenced by the parameters of the acquired EPI sequence, may vary between individuals that underwent the same scanning sequence, and depends on the regions of interest. Lastly, our results emphasize the promising role of the cerebral blood flow measured by PCASL as a neural representation of aging and cognitive impairment. These empirical findings strongly support the considerations by Reid et al. (2019), which illustrate the urgent need to define the respective theoretical, methodological, and confounding properties more carefully in future FC-based studies.

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Availability of data, code, and material

Behavioral and imaging data of the HCP-Aging cohort are available through application at the Human Connectome Project, while data from the MBB and BTC cohort are freely available at OpenNeuro. Analysis commands, python scripts, and the singularity containers used will be published on a Github repository. Additional data can be made available upon request.

CRedit authorship contribution statement

Lukas Roell: Writing – review & editing, Writing – original draft, Visualization, Validation, Project administration, Methodology, Investigation, Formal analysis, Data curation, Conceptualization. **Stephan Wunderlich:** Writing – review & editing, Writing – original draft, Visualization, Validation, Methodology, Formal analysis, Conceptualization. **David Roell:** Writing – review & editing, Validation, Formal analysis. **Florian Raabe:** Writing – review & editing, Funding

acquisition. **Elias Wagner:** Writing – review & editing, Funding acquisition. **Zhuanghua Shi:** Writing – review & editing, Validation, Formal analysis. **Andrea Schmitt:** Writing – review & editing, Supervision, Funding acquisition. **Peter Falkai:** Writing – review & editing, Supervision, Funding acquisition. **Sophia Stoecklein:** Writing – review & editing, Supervision. **Daniel Keeser:** Writing – review & editing, Writing – original draft, Supervision, Methodology, Conceptualization.

Declaration of competing interest

PF is a co-editor of the German (DGPPN) schizophrenia treatment guidelines and a co-author of the WFSBP schizophrenia treatment guidelines; he is on the advisory boards and receives speaker fees from Janssen, Lundbeck, Otsuka, Servier, and Richter. **LR, SW, DR, EW, FR, ZS, AS, SS, and DK** report no conflicts of interest.

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The data of the MBB project was obtained from the OpenNeuro database (DOI:10.18112/openneuro.ds000221.v1.0.0) with the accession number ds000221. The project was partially supported by the Volkswagen Foundation (AZ.: 89 440).

The data of the BTC project was obtained from the OpenNeuro database (DOI:10.18112/openneuro.ds001226.v4.0.0) with the accession number ds001226. This project was supported by the Special Research Funds (BOF) of the University of Ghent (01MR0210 and 01J10715), as well as Grant P7/11 from the Interuniversity Attraction Poles Program of the Belgian Federal Government.

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Supplementary materials

Supplementary material associated with this article can be found, in the online version, at doi:10.1016/j.neuroimage.2025.121195.

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