Integrating temporal and spatial scales: human structural network motifs across age and region of interest size

Christoph Echtermeyer1,2, Cheol E. Han3, Anna Rotarska-Jagiela4, Harald Mohr5, Peter J. Uhlhaas4 and Marcus Kaiser2,3,6*

1 Institute for Chemistry and Biology of the Marine Environment, Carl von Ossietzky University, Oldenburg, Germany
2 School of Computing Science, Newcastle University, Newcastle-upon-Tyne, UK
3 Institute of Neuroscience, The Medical School, Newcastle University, Newcastle-upon-Tyne, UK
4 Department of Neurophysiology, Max-Planck Institute of Brain Research, Frankfurt a. M., Germany
5 Goethe University, Institute of Psychology, Frankfurt a. M., Germany
6 Institute of Neuroscience, The Medical School, Newcastle University, Newcastle-upon-Tyne, UK

Edited by:
Olaf Sporns, Indiana University, USA
Reviewed by:
Alex Fornito, University of Melbourne, Australia
Lucas Antiqueira, University of Sao Paulo, Brazil
*Correspondence:
Marcus Kaiser, School of Computing Science, Newcastle University, Claremont Tower, Newcastle-upon-Tyne NE1 7RU, UK. e-mail: m.kaiser@newcastle.ac.uk

Human brain networks can be characterized at different temporal or spatial scales given by the age of the subject or the spatial resolution of the neuroimaging method. Integration of data across scales can only be successful if the combined networks show a similar architecture. One way to compare networks is to look at spatial features, based on fiber length, and topological features of individual nodes where outlier nodes form single node motifs whose frequency yields a fingerprint of the network. Here, we observe how characteristic single node motifs change over age (12–23 years) and network size (414, 813, and 1615 nodes) for diffusion tensor imaging structural connectivity in healthy human subjects. First, we find the number and diversity of motifs in a network to be strongly correlated. Second, comparing different scales, the number and diversity of motifs varied across the temporal (subject age) and spatial (network resolution) scale: certain motifs might only occur at one spatial scale or for a certain age range. Third, regions of interest which show one motif at a lower resolution may show a range of motifs at a higher resolution which may or may not include the original motif at the lower resolution. Therefore, both the type and localization of motifs differ for different spatial resolutions. Our results also indicate that spatial resolution has a higher effect on topological measures whereas spatial measures, based on fiber lengths, remain more comparable between resolutions. Therefore, spatial resolution is crucial when comparing characteristic node fingerprints given by topological and spatial network features. As node motifs are based on topological and spatial properties of brain connectivity networks, these conclusions are also relevant to other studies using connectome analysis.

Keywords: network analysis, network motifs, structural connectivity, human

1 INTRODUCTION

The set of connections in the brain can be described as the connectome (Sporns et al., 2005). Connectome data is currently becoming available at different levels of the structural organization: from neuronal networks of connections between neurons to fiber-tract networks between brain regions. These data promise to give valuable new insights, but analyses integrating different data sets are challenging. One problem of data integration across labs is that the raw data might show different voxel resolutions due to different magnetic field strengths of the magnetic resonance imaging (MRI) scanner. In addition, different labs might use different parcelation routines or brain atlases. This leads to different spatial resolutions and consequently to different structural connectivity networks (eventually with different numbers of nodes) (Bassett et al., 2010; Fornito et al., 2010; Hagmann et al., 2010). Another problem of data integration is caused by age differences between subjects. There is a certain evidence that characteristics of the connectome vary over age (Fair et al., 2009; Uhlhaas et al., 2009; Fan et al., 2010; Hagmann et al., 2010). Given a network analysis result of two studies, say concerning small-world features (Watts and Strogatz, 1998), are differences due to the study population's differences in connectivity or are they due to differences in age or network resolution? In this article, we observe how topological and spatial network features change across age and network resolution for structural connectivity, based on diffusion tensor imaging (DTI), in healthy human subjects. Using single node motifs, we also apply a network mapping technique, which yields a compact and easily comparable representation of complex network structures.

1.1 SCALES OF A HIERARCHICAL ORGANIZATION

Hierarchy is a central feature in the organization of complex biological systems and particularly the structure and function of neural networks (Kaiser and Hilgetag, 2010; Kaiser et al., 2010). In neural networks, the term hierarchy may be understood in several different ways and can apply to topological, spatial, temporal as well as functional properties. These features can be observed at different scales or resolutions.

1.1.1 Temporal scales

Network analysis is often presented as the analysis of a network at a given time, but many networks change both over time and with temporal resolution. Structural connectivity, given by anatomical
connections between neurons or brain regions, may change due to activity-dependent plasticity (Butz et al., 2009). Functional connectivity, given by correlations in the activity patterns of network nodes, can change for different tasks for a subject or due to different system states (e.g., awake vs. asleep). In our case, connectivity might also change over longer time-scales, as part of brain development.

Recovered networks can appear differently depending on temporal resolution. At a fine-grained temporal resolution of a few hundred milliseconds, cortical tissue can show distinct oscillations, e.g., in the α, γ, or δ frequency bands (Roopun et al., 2008). For more coarse-grained time-resolutions with sample lengths of several seconds or minutes such distinct frequency peaks disappear and the frequency distribution displays a 1/θ power-law behavior instead (Buzsaki, 2006).

Here, we observe temporal scale in terms of snapshots of structural connectivity at different points in time. The human brain undergoes large structural developments until the age of around 20–25 years. These developments coincide with large functional changes during child development and during the teenage years. They further coincide with changes in functional connectivity (Uhlhaas et al., 2009). Such changes during brain network development are not only of interest for the study of healthy subjects, but also for many psychiatric disorders like schizophrenia (Uhlhaas and Singer, 2010), which have an onset around the time when the brain network matures (age 18–25 years). We therefore test the potential of single node motifs to integrate information across temporal network “snapshots”.

1.1.2 Spatial scales
The network structure not only depends on the time of a “snapshot” but also on the spatial resolution. One can distinguish the micro- and macro-connectome as the connectivity between individual neurons and connections between brain regions, respectively (DeFelipe, 2010). In this study of structural connectivity we focus on the macro-connectome level based on MRI data [see Seung (2009) for more information about neuronal connectivity instead].

The macro-connectome can be analyzed at different levels of spatial resolution and thus different levels of brain network organization. With a low spatial resolution, it corresponds to connectivity between brain areas where network nodes correspond to brain regions. The resulting networks show features of modular and small-world networks (Hilgetag et al., 2000; Sporns et al., 2000) and typically consist of up to 100 nodes per cortical hemisphere for the primate brain. At the mesoscale, connectivity between regions of interest (ROIs) of the same size, e.g., 1 cm² cortical surface area, can be studied. Such networks consist of around 1000 nodes for the human brain (Hagmann et al., 2008). With even higher resolution, the microscale of the macro-connectome could be studied, i.e., connections between cortical columns (Mountcastle, 1997). However, such networks with nodes representing one cortical column each would consist of 1,250,000 nodes for the human brain (both hemispheres, not including subcortical structures) and identifying columns with diameters ranging from 200 μm to 1 mm (Hubel et al., 1977; Horton and Adams, 2005) in humans is (currently) beyond the reach of standard MRI. Such extremely high resolutions might only be achieved with higher magnetic fields or extremely high magnetic fields in post mortem studies.

We therefore study the global and regional scale of the macro-connectome. Importantly, this level allows us to assess whether motifs which are characteristic for a certain cortical region are also characteristic for all parts of that cortical area or whether a region displays a variety of motifs at higher spatial resolutions. Also, changes in spatial resolution affect spatial measurements, which form part of the motif detection. It is often interesting to observe how many connections go to nearby targets and how many extend over a long distance, potentially linking different components of the neural network. This can be readily observed using a histogram of the connection lengths of a network. It is known that the probability that two neurons are connected decays almost exponentially with distance (Hellwig, 2000; Schüz et al., 2005; Kaiser et al., 2009): connections over a long distance are less likely than short-distance connections. In different organisms, ranging from neuronal connectivity in C. elegans and layers in the rat visual cortex to fiber-tract connectivity in the macaque, the actual distribution was approximated best by a Gamma probability density function (Kaiser et al., 2009). Looking at different spatial resolutions within humans, we would expect that the proportion of short-distance connections relative to long-distance connections increases when the spatial resolution is increased. We will therefore observe how the length distributions change over different spatial scales.

1.2 SINGLE NODE MOTIFS
Networks can be analyzed following different concepts: to investigate a single network, for example, analysis can focus on its nodes, comparing them through connectional fingerprints based on the absolute values of node properties (e.g., node degree or local clustering coefficient; Passingham et al., 2002). On a more abstract level, nodes of brain networks can be regarded with respect to their different organization levels. These approaches look at one network of a particular subject, but networks can also be compared across different subjects – or different networks might exist for the same subject across time. However, mapping between networks becomes difficult if the number of nodes and edges varies, such that direct comparisons (i.e., link-to-link or node-to-node) between structures are ruled out. In contrast to such methods of cross-comparison, the approach used in this paper is less sensitive to these changes: although a network with a slightly higher number of nodes could show additional motifs, most nodes are non-outliers making such changes less likely. Similarly, a network with slightly more edges might show absolute changes in network features without affecting the outliers of the feature distribution leading to similar single node motifs. Again, these comparisons of the number and classes of motifs become possible for non-identical numbers of nodes or edges but only if the numbers do not differ too much. These motif-classes are significantly different from each other and they allow to classify network nodes at the same level even if they appear in different networks.

The concept of node motifs – a combination of local network features – complements that of network motifs, which are specific connectivity patterns that have been used to characterize networks before (Milo et al., 2002). An example for a node motif are highly connected nodes or hubs that affect spreading phenomena and can be important components of a network. More complex, multi-dimensional node motifs, which are characterized by multiple features in combination, specify nodes more comprehensively. With
these more precise descriptions, new kinds of motifs can be formulated and Costa et al. (2009) presented a routine for their detection and specification. We proposed first improvements and procedures for automating parameter choices for this method (Echtermeyer et al., 2011), such that large numbers of networks can be processed and compared to each other. This enhanced technique is applied to brain networks in this paper.

### 1.3 Comparing Motif Counts Across Networks

The usefulness of mapping brain networks (which itself are abstract representations) to motif-classes is demonstrated for two scenarios: first, regional connectomes of 53 children (aged 12–23 years) are compared to show how the fingerprint of a DTI network can change over time. Second, we test the effect of different connectome-resolutions on the resulting network-fingerprint. Therefore, DTI networks at different levels of organization were generated by changing the level of granularity from 414 ROIs to up to 1615 ROIs.

Comparing networks from the same subject but at different granularities has so far been complicated by the critical dependence of standard approaches on the resolution of the network (Antiqueira et al., 2010). We show how our novel node fingerprints can be used to compare different levels of connectome organization.

Note that comparing motifs has several advantages over comparing network features. Features, such as the small-world measures of clustering coefficient and characteristic path length, critically depend on the edge density of a network. Therefore, changes in these measures could be either due to changes in network organization (e.g., modularity) or due to a difference in the number of connections. To identify the changes related to network organization, the edge density needs to be constant when comparing networks. However, this leads to a severe problem: when transforming raw data of connection strengths between nodes to binary data, each network needs a different threshold for establishing a connection (Rubinov and Sporns, 2010). To avoid the problem of either having different edge densities or different thresholds, researchers increasingly use weighted networks without thresholding (Bullmore and Bassett, 2011). For our analysis, we expect that differences in edge density are not only due to inter-subject variability, as in other studies, but are more heavily influenced by changes across different temporal or spatial scales. Therefore, we observe how network measures, and as a result node motifs, change over scales. These changes can be due to changes in organization or edge density.

Although a detailed analysis is outside the scope of this paper, we show the relation between both scales in Figure 1. Whereas edge density decreases with network size, there are no significant changes between age groups when looking at networks with the same spatial resolution. However, we would like to point out that (a) motifs change over time and can therefore be used to find characteristic changes in the developing brain and (b) motifs change over spatial scales, therefore comparisons between studies are only meaningful when the same (or a comparable) spatial resolution has been used.

### 2 Materials and Methods

#### 2.1 Recruiting and Data Recording

We recruited 53 participants (age between 12 and 23 years) from local community and high schools. The study received ethics committee approval and written informed consents were obtained from the participants’ parents for under-age participants, or from themselves if older than 18 years after study procedures were described. Participants were screened for a history of psychiatric and neurological disorders and current drug abuse using ophthalmological assessment including monocular and binocular visual acuity. Participants were grouped into four age-categories (Table 1).

A 3-Tesla scanner (Siemens Trio, Erlangen, Germany) was used at the Brain Imaging Centre (BIC), Frankfurt am Main, Germany. We recorded T1 weighted MRI (voxel-size 1 mm × 1 mm × 1 mm) and Diffusion Tensor MRI (voxel-size 2 mm × 2 mm × 2 mm). To minimize head motion, we used tightly padded clamps on the eight-channel head coil. The T1 weighted MRI was recorded with the following parameters: 176 slices, field of view (FoV) 256 mm, repetition time (TR) 2250 ms, and echo time (TE) 2.6 ms. Three Diffusion Tensor Images per participant were recorded with the following parameters: 60 slices, FoV 192 mm, TR 7600 ms, TE 91 ms, 60 directions b-vectors with b-factor of 1000 and 10 b0-images.

### 2.2 Data Processing and Network Construction

We used FreeSurfer to obtain surface meshes of the boundary between gray matter and white matter from T1 anatomical brain images1, and Diffusion toolkit along with TrackVis (Wang et al., 2007) to obtain stream-line tractography from diffusion tensor images of connection strengths between nodes to binary data, each network needs a different threshold for establishing a connection (Rubinov and Sporns, 2010). To avoid the problem of either having different edge densities or different thresholds, researchers increasingly use weighted networks without thresholding (Bullmore and Bassett, 2011). For our analysis, we expect that differences in edge density are not only due to inter-subject variability, as in other studies, but are more heavily influenced by changes across different temporal or spatial scales. Therefore, we observe how network measures, and as a result node motifs, change over scales. These changes can be due to changes in organization or edge density.

Although a detailed analysis is outside the scope of this paper, we show the relation between both scales in Figure 1. Whereas edge density decreases with network size, there are no significant changes between age groups when looking at networks with the same spatial resolution. However, we would like to point out that (a) motifs change over time and can therefore be used to find characteristic changes in the developing brain and (b) motifs change over spatial scales, therefore comparisons between studies are only meaningful when the same (or a comparable) spatial resolution has been used.

#### Table 1 | Ranges of age-categories.

| Category | Age range (years) | Number of subjects |
|----------|------------------|--------------------|
| 1        | 12–14            | 9                  |
| 2        | 15–17            | 20                 |
| 3        | 18–20            | 16                 |
| 4        | 21–23            | 8                  |
images. We registered surface meshes into brains of the diffusion
tensor images to extract networks. FreeSurfer provides parcelation
of anatomical regions of cortices (33 for each hemisphere) based
on the Deskian atlas (Fischl et al., 2004b; Desikan et al., 2006) and
subcortices (Fischl et al., 2002, 2004a). We subdivided those ana-
tomical regions of cortices into many of similar surface area using
our own algorithm. It is known that the size of each ROI affects
network connectivity and various network measures (Hagmann
et al., 2007; Zalesky et al., 2010). The core of our algorithm is an
expectation–maximization (EM) algorithm. First, we choose the
expected number of subdivisions (400, 800, and 1600) for cortical
regions (we do not subdivide subcortical areas) and decided the
number of subdivisions based on the ratio of each surface area
over whole surface area in the average subject. Then we subdivide
each anatomical cortical region to the assigned number of subdivi-
sions using the EM algorithm, minimizing variance of subdivisions’
surface area sizes. The ratio of variance to mean does not exceed
10% in the average subject. In each subject, this ratio was higher
(400: 414.78 ± 140.06 mm², 33.77%; 799: 207.65 ± 77.51 mm²,
37.33%; 1601: 103.63 ± 44.33 mm², 42.78%, statistics of cortical
ROIs only) because the size of each subject’s anatomical region var-
ied (2468.06 ± 1813.10 mm², 73.46%, which represents individual
subjects’ regions were 12 ± 30% bigger than the average subject’s;
ratio of variance to mean is 26.49%). Another useful statistic is the
ratio of interquartile range to median: larger values represent higher
dispersion. The values (Table 2) are smaller than those determined
by Fornito et al. (2010).

We projected this new parcelation template into each sub-
ject’s surface. Using this projection procedure we kept topologi-
cal consistency across all subjects. We also included a few selected
subcortical areas in the ROI list: nucleus accumbens, Amygdala,
Caudate, Hippocampus, Pallidum, Putamen, and Thalamus. Thus
the actual number of ROIs (414, 813, and 1615, respectively) is
slightly larger than specified. These three numbers of ROIs on the
brain and resulting networks are shown in Figure 2. Before further
processing the diffusion tensor images, we serialized three diffusion
tensor images and b-vectors for each participant; thus, a collated
image of each subject has 30 b0-images and 180 diffusion images.
The eddy current was corrected through FSL2. We used Diffusion
Toolkit along with TrackVis (Wang et al., 2007) with fiber assign-
ment by continuous tracking (FACT) algorithm (Mori et al., 1999)
and 35° of angle threshold. After registering the surface meshes
(FreeSurfer3) of each subject to his/her DTI, we generated cortical
volume ROIs, which are voxels in the gray matter. Also the selected
subcortical areas (see above) were registered to DTIs and used as
ROIs. Then, we used the UMCP toolbox4 to obtain connectivity
matrices from the defined and registered ROIs and tractography.
This counts the number of fibers from a tractography from the
Diffusion toolkit, between all pairs of defined ROIs’ gray matter;
the fiber counts between all ROI-pairs yield the weight-matrix. We
also computed the average fiber lengths between ROIs (if there is
no connection between a pair, the length is set to zero). The fiber
length is based on the actual three-dimensional trajectory of the
fiber tract and may be larger than the Euclidean distance between
connected nodes. Note that long fiber length usually also corre-
sponds to a large Euclidean distance between ROIs: earlier studies
in fiber tracts in primates have shown that 85% of the fibers go
in a straight line or are only slightly curved (Hilgetag and Barbas,
2006). However, only a fully bended (U-turn) fiber could result in
a long fiber-tract connection to spatially nearby nodes.

Diffusion tensor imaging fiber-tracking yielded weighted matrices
where weights indicate the probability of connections between
corresponding ROIs. Non-zero weights have been interpreted as an
indication for a link when constructing the brain networks. The
resulting adjacency matrices were similarly sparse for each network
resolution (Figure 1, average link-densities 7.5, 4.0, and 2.0% for
networks of 414, 813, and 1615 nodes, respectively).

2.3 NETWORK ANALYSIS
Single node motifs were identified from the constructed networks
by applying a technique presented by Costa et al. (2009) with addi-
tions in order to choose parameters automatically (Echtermeyer

Table 2 | Region of interest (ROI) surface area (mm²) statistics with ratio of
interquartile range to median.

| Parcelation | Median | Interquartile range | Ratio |
|-------------|--------|---------------------|-------|
| aparc       | 1935   | 2811                | 1.45  |
| 414         | 395    | 183                 | 0.46  |
| 813         | 195    | 101                 | 0.52  |
| 1615        | 95     | 56                  | 0.59  |

1http://www.fmrib.ox.ac.uk/fsl/
2http://surfer.nmr.mgh.harvard.edu/
et al., 2011). The method yields a global characterization of the network based on its local properties. Here, each node is quantified by nine different local measures, which are motivated in the next paragraph. We chose such a large number of local measures, some of which share similarities, to ensure that as many network features as possible are considered. Redundancies, where two features are highly correlated, are removed during the node-motif detection, which is explained further below.

A node’s degree is its number of connections to other nodes; dividing this by the number of all links in the network yields the normalized node degree $K$. The average over all neighbors’ degrees is called the normalized average degree $r$. (Nodes that are directly linked to each other are called neighbors.) Degrees of a node’s immediate neighbors can further be taken into account by their respective coefficient of variation $cv$. Connectivity among neighbors of a node is quantified by the clustering coefficient $cc$, which reflects how many of all possible connections between neighbors actually exist (Watts and Strogatz, 1998; Kaiser et al., 2007a). The hierarchical clustering coefficient of level two $cc_2$ extends this concept to connections between neighbors’ neighbors (Costa and Silva, 2006). To what degree a node’s neighbors connect to the same target is quantified by the locality index $loc$, which is based on the matching index (e.g., Kaiser and Hilgetag, 2004a). The six measures mentioned so far quantify topological aspects of the network, while the next three measures also take into account spatial features. The average connection length $acl$ for a node is the average length of fiber tracts to its neighbors and the furthest away neighbor determines the maximum connection length $mcl$. The average over connection lengths to nodes two steps away, i.e., neighbors’ neighbors, is called the average indirect reach $air$. Further details and measures can be found in the literature (Albert and Barabási, 2002; Newman, 2003; Newman et al., 2006; Costa et al., 2007b).

The motif-detection workflow starts by applying the above listed network measures (Figure 3): local network measures are calculated for all nodes of the network, which yields a nine-dimensional characterization (feature vector) for each node. This high-dimensional feature vector is compacted to two dimensions (Step 1) in order to estimate a probability distribution (Step 3), which is used to identify nodes with uncommon features (Step 4). All nodes are grouped (Step 5) to form high-dimensional motif-regions (Step 6), which are eventually joined, if too close to each other (Step 7). The number of nodes in each motif-region yields a fingerprint of the network (Step 8).

**FIGURE 3 | Figure from Echtermeyer et al. (2011).** Illustration of network analysis workflow: high-dimensional characterization of all network nodes through local network measures $\mu_i$ (Step 1) is compacted to two dimensions (Step 2) in order to estimate a probability distribution (Step 3), which is used to identify nodes with uncommon features (Step 4). All nodes are grouped (Step 5) to form high-dimensional motif-regions (Step 6), which are eventually joined, if too close to each other (Step 7). The number of nodes in each motif-region yields a fingerprint of the network (Step 8).
representation is simplified by exploiting similarities between different measures (Costa et al., 2007b): using principal component analysis (PCA; Johnson and Wichern, 2007, Chapter 8), correlation between measures is removed and the feature vectors are reduced to two dimensions (PCA-plane). The position on the PCA-plane characterizes the nodes and allows for comparisons: nodes close to each other share similar features whereas well separated ones differ. The next step is to estimate how likely specific combinations of features are. This is done by smoothing over points in the PCA-plane using the Parzen window approach (Parzen, 1962; Duda et al., 2001, Chapter 4.3) with width of the smoothing kernel scaled according to the SD along the corresponding principal component axis (Echtermeyer et al., 2011). Thereby, each node is assigned a probability, which is used to distinguish those with common and rare features. The $w$ many nodes with uncommon properties, termed outlier nodes, were determined as follows. Given all nodes’ probabilities $p = (p_k)_{k = 1}^\ldots^w$ (sorted increasingly), the respective mean $\overline{p}$, and SD $\sigma(p)$, the number of singular nodes $w$ is chosen as

$$w = \text{arg max}_{k, p_k \in \overline{p} - \sigma(p)升降} p_{k+1} - p_k,$$  \hspace{1cm} (1)

or $w = 0$, if all probabilities are greater than $\overline{p} - \sigma(p)升降$. Nodes corresponding to the smallest $w$ probabilities are the identified outliers.

Next, the outlier nodes were assigned to clusters each of which represents a different motif-group, i.e., a set of features. This was done by first centering equally sized ellipses on each point and determining their overlap. Maximal sets of nodes whose ellipses form a connected area yield both the number $k$ of motif-groups as well as the groups themselves. The above mentioned mechanisms have been evaluated and discussed in detail in an earlier publication (Echtermeyer et al., 2011). Motif-groups correspond to nine-dimensional motif-regions in feature space, such that all nodes within a motif-group are similar with respect to the local network measures used in the beginning. In the final step, motif-regions from all networks were reduced to the final nine motif-regions by subsequently joining the closest two of them. The pairwise distances have been determined with a modified Mahalanobis measure (Mahalanobis, 1936). The number of nodes that express a particular motif yields a fingerprint of the network, which characterizes the network and allows for comparisons (Costa et al., 2009), e.g., by considering the motif-diversity, which is the number of unique motifs $k$ that were identified. Another possibility is to compare the number of outlier nodes $w$.

To determine which changes are statistically significant a random permutation test has been applied: subjects were randomly assigned an age (12–23 years) while ensuring that each age group had the same size as for the original data. Subsequently, each group’s mean $w$ and $k$ were compared to those of the original data to compute the $p$-values (using a total of 500,000 permutations). Results, shown directly in Figure 4, represent how far each group is to the average over age. Note that except for the lowest network resolution, the large spike at age 21 does not lead to significant increase in $w$ and $k$. This is because our data only include one subject aged 21 and further data would be needed to confirm that the deviations at this age are indeed significant across all scales of resolution.

A pattern that is very similar to that of the outlier number $w$ is the diversity $k$ of expressed motifs. The number $k$ is high and low for nearly the same ages as was the number of outliers $w$ before (Figure 4, dashed orange lines), which is also reflected by the strong correlation between $w$ and $k$ (Pearson correlation 0.94, 0.88, and 0.87 for 414-, 813-, and 1615-node networks, respectively). Peaks at low (13–14 years) and high (21 years) ages can be seen with a dip in between. This pattern fits networks with 414 nodes best, whereas networks with the higher resolution (813 and 1615 nodes) show an increasingly flat profile except for the later peak (21 years). Note that absolute motif-diversity $k$ is relatively similar for all network resolutions, which was not the case for the actual number of outliers $w$.

Both, number of outliers $w$ and motif-diversity $k$ are interesting on their own, but their strong correlation provides additional information. Although the absolute values differ, similarity of the curves’ profiles is independent of network resolution. We thus find the very robust effect that motif-diversity is coupled to the number of outlier nodes. In other words, whenever more outliers occur, they also become more diverse.

### 3.2 Motif Changes with Network Resolution

The previous section has already shown that network resolution affects outlier quantity, $w$, but not the relative distribution across the age range (Figure 4, solid blue lines). Different from that, motif-diversity, i.e., how many different motifs are expressed, $k$, is nearly invariant on an absolute scale, in contrast to its relative distribution (Figure 4, dashed orange lines). We tested whether the number $w$ and diversity $k$ of outliers, at different spatial scales, are correlated with cognitive skills of our subjects. Unfortunately, measures of intelligence yielded by four sub-scores of the Wechsler intelligence test were only available for 44 out of our 53 subjects. For all comparisons, the Pearson correlation was $r \leq 0.5$. Therefore, we cannot confirm a link between intelligence and $w$ or $k$. However, intelligence might still be related to a specific combination of motifs; a possibility which could only be tested given a larger cohort of subjects. Next, we analyze the actual frequencies with which motifs are expressed by different age groups (Table 1).

Using the same raw DTI-data, networks have been constructed with different resolutions, i.e., differently sized ROIs, for each subject. More fine-grained ROIs parcelate the brain into more elements, each of which is represented by a network node. We were interested in the effect that different network resolutions have on the occurring node motifs (Table 3; Figure 5) and therefore, networks with 414, 813, and 1615 nodes were analyzed and compared to each other.
that the spatial resolution is critical for assessing the frequency of motifs and, presumably, also for its underlying spatial and topological features.

Motif counts also depend on age. For 414-node networks, many motifs only occur for one or two out of the four age groups. Whereas this might be influenced by the low number of nodes in the network, we also see age-based motif frequency changes for higher spatial resolutions. Looking at the resolution of 1615 nodes, the results show that motif-expression strongly depends on network resolution (Figure 6). Whereas motif 4 is most frequent for the 414-node resolution, it is nearly absent in the highest resolution of 1615 nodes. Other motifs also show large fluctuations between spatial resolutions with either increasing or decreasing frequency. Motifs can also be characteristic for a single scale. For example, motifs 8, 9, and 10 are most prevalent across age groups at an intermediate resolution of 813 nodes. This shows that the spatial resolution is critical for assessing the frequency of motifs and, presumably, also for its underlying spatial and topological features.

Motif counts also depend on age. For 414-node networks, many motifs only occur for one or two out of the four age groups. Whereas this might be influenced by the low number of nodes in the network, we also see age-based motif frequency changes for higher spatial resolutions. Looking at the resolution of 1615 nodes, the

Figure 4 | Motif-expression changing with age: both the number of outlier nodes in a network w (solid blue line) and the diversity of motifs expressed k (dashed orange line) vary with subject age. The time-dependent patterns of w and k are shown for different network resolutions (rows). Age groups indicated by dashed vertical lines. Significantly de- or increased values for w and k are indicated by symbols < and >, respectively (*90, **95, and ***99% significance). Note that our data only include one subject aged 21 and further data would be needed to confirm significance of deviations at this age.
Symbols ..., ↑, and ↓ indicate normal, elevated, and decreased values of \( K \), normalized node degree; \( r \), normalized average degree; \( cv \), coefficient of variation of neighbors’ degrees; \( loc \), locality index; \( cc \), clustering coefficient; \( cc_2 \), hierarchical clustering coefficient of level two; \( acl \), average connection length; \( mcl \), maximum connection length; \( air \), average indirect reach, respectively.

| Motif | \( K \) | \( r \) | \( cv \) | \( loc \) | \( cc \) | \( cc_2 \) | \( acl \) | \( mcl \) | \( air \) |
|-------|--------|--------|--------|--------|--------|--------|--------|--------|--------|
| 3     | ↓      | ↓      | ↓      | ↑      | ↑      | ↓      | ...    | ↓      | ↓      |
| 4     | ...    | ↑      | ...    | ↑      | ↓      | ...    | ...    | ↓      | ↓      |
| 5     | ↓      | ↓      | ↓      | ↑      | ↑      | ↑      | ↑      | ↑      | ↑      |
| 8     | ↓      | ...    | ↓      | ↑      | ↑      | ↑      | ↑      | ↑      | ↑      |
| 9     | ↓      | ↓      | ↑      | ↑      | ↑      | ↑      | ↑      | ↑      | ↑      |
| 10    | ↓      | ...    | ↓      | ↑      | ↑      | ↑      | ↑      | ↑      | ↑      |

Table 3 | Properties of most frequent motifs.

Figure 5 | Stylized illustration of most frequent motifs (Table 3): motif 3 Provincial node with connections in the direct (high \( cc \)) but not the larger neighborhood (low \( cc \), \( air \), and \( mcl \)). Motif 4 Provincial hub with more connections than its neighbors (high \( r \)) that are less connected between themselves (low \( cd \)). Motif 5 Global bottleneck with few (low \( K \)) but long-range connections (high \( acl \), \( mcl \), \( air \)) that reach beyond the local neighborhood (low \( cd \)). Motif 8 Global uniform bottleneck sharing properties of motif 5 but connected to nodes with similar degrees (low \( cv \)). Motif 9 Global local bottleneck sharing properties of motif 5 but also having well-connected neighbors (high \( cd \)) thus better informing local circuits. Motif 10 Provincial bottleneck with only short-range connections (low \( mcl \)) and few connections between its neighbors (low \( cd \)) leading to a large influence on the local circuit.

3.3 Fiber Length Distribution for Different Network Sizes

A possible explanation for the observed dependence on network resolution could be fiber length based network measures (namely: average connection length, maximum connection length, and average indirect reach). These measures take into account the determined fiber trajectory between ROIs, i.e., their fiber-tract length. It is thus to be expected that the number of ROIs impacts on the corresponding fiber lengths. We tested this by observing the fiber length distribution for different network resolutions (Figure 7). Across all resolutions, short fibers are more frequent than long fibers. This exponential tail, which can be fitted by a Gamma distribution (Kaiser et al., 2009), has been reported before not only at the level of fiber tracts (Kaiser and Hilgetag, 2004b), but also at the level of connections between neurons within cortical areas (Hellwig, 2000). One notable difference between spatial resolutions is the increase in the number of short fibers (length < 10 mm) increasing by more than 50% from the lowest to the highest spatial resolution. This can be due to the smaller size of network nodes where surface regions that before belonged to the same node are now separate nodes on the surface. Therefore, a short fiber tract between these nodes becomes feasible. Looking at the maximum range of fibers, spatial resolution does not seem to influence maximum fiber lengths. However, differences in average fiber length, in particular due to an increased number of short fiber connections, might occur for particular nodes. The pattern for average indirect reach is more complex and can be influenced by changes in both short and long fibers. Overall, given a three-fold increase in spatial resolution, the fiber length distributions remain remarkably similar.

3.4 Motif Consistency Across Spatial Scales

How consistent are motifs across different resolutions? If a region splits into several regions for higher resolutions, do the daughter regions show the same motif as the parent region? Interestingly, this does not seem to be the case. Figure 8 shows a typical example of motif locations for one subject. For 414 nodes, one location shows motif 3. One part of this location then shows motif 8 for the resolution of 813 nodes. For 1615 nodes, part of the original region shows motif 3 again, whereas another part shows motif 9. This is an interesting point when talking about the connectivity between brain regions: a pattern shown at the low-resolution level (e.g., cortical regions or Brodmann areas) may consist of several diverse patterns when observing connectivity at a higher resolution.

4 Discussion

We observed how the distribution (fingerprint) of characteristic motifs of single network nodes changed for human structural connectivity (DTI) during brain development. We studied the role of both temporal scales (age groups from 12 to 23 years) and spatial scales (414, 813, and 1615 nodes) and found that, first, the number and diversity of motifs in a network are strongly correlated. Second, comparing different scales, the number and diversity of motifs varied across the temporal (subject age) and spatial (network resolution).
Therefore, both the type and localization of motifs differ for different spatial resolutions. This indicates that spatial resolution is crucial when comparing characteristic node fingerprints given by topological and spatial network features. This result is in line with previous studies that observed the role of sampling on brain network properties (Antiqueira et al., 2010) and it additionally shows the influence of the temporal scale.

Networks can be characterized at different levels. Aggregate measures that characterize a network as a whole can be used to distinguish different network types, e.g., small-world (Watts and Strogatz, 1998; Hilgetag et al., 2000; Sporns et al., 2000), scale-free (Eguiluz et al., 2005; Kaiser et al., 2007b), modular (Hilgetag et al., 2000), or hierarchical (Kaiser et al., 2010). Networks can also be characterized at the level of individual components. Looking at the topological features of individual nodes, a first study by Passingham et al. (2002) has linked the network features of macaque cortical areas with the function of each area. Whereas this approach considered all nodes of the network, there are also ways to search for “special” nodes of a system. Following the ideas of scale-free networks, brain areas with a large number of connections – so-called network hubs – might be crucial for integrating or distributing information (Kaiser et al., 2007b). Sporns et al. (2007) were able to detect and classify different kinds of hubs of cortical structural connectivity. However, hubs are just one type of outliers where one or more node features (here: the number of connections of a node) differ from the average value of nodes in the network. In two previous manuscripts (Costa et al., 2009; Echtermeyer et al., 2011), we have developed a tool that systematically searches for and classifies nodes which differ from the majority of nodes in a network. This approach includes the joint analysis of multiple features, which include topological as well as spatial properties (Costa et al., 2007a).

Our measures are based on outliers, which do not show the overall trend as shown in previous studies (spatial resolution dependent changes: Zalesky et al. (2010), Bassett et al. (2010), Fornito et al. 2011).
per age group was not identical (Table 1). Therefore, findings in age groups with fewer members might have been influenced by relatively few outliers. Whereas an identical number of subjects per group is desirable, for our study there was also a trade-off with the age range: using the same subject numbers would have led to groups where the age ranges differ, e.g., covering 3 years for one group and 5 years for another. We decided for a different number of subjects per group in order to perform the comparison across the temporal scale. Second, single node motifs become more robust for larger network sizes. As node motifs are outliers, only a small percentage of all nodes will be characterized as node motifs. Therefore, for low-resolution networks, node motifs might only occur for few subjects. In addition, their number might be a poor estimate of the underlying frequency in human connectivity networks. For this reason, we have only reported networks with at least 414 nodes, leaving out networks with a parcelation into 110 cortical and subcortical regions. Third, node features but not necessarily motifs depend on the edge density of the network. In our developmental networks, we observed significant edge density changes for different spatial resolutions ranging from more than 7% for 414 nodes to 2% for 1615 nodes. However, for each network resolution, edge densities remained comparable for different temporal scales (age groups; Figure 1). Fourth, deterministic tracking cannot capture crossing fibers which can be done in probabilistic tracking (Behrens et al., 2007), in Diffusion spectrum imaging (Wedeen et al., 2008; Bassett et al., 2010), or in high-angular resolution diffusion imaging (2010), Hayasaka and Laurienti (2010), age-dependent changes: Fan et al. (2010), Hagmann et al. (2010), Fair et al. (2009); Uhlhaas et al. (2009), Table A1 in Appendix). Thus, it is difficult to directly relate our results to their findings. However, our report that edge density decreased with increased spatial resolution is consistent with previous studies of structural connectivity (Bassett et al., 2010; Zalesky et al., 2010). For the temporal resolution, most previous studies showed no overlap with the age range of our study. Hagmann et al. (2010) performed a study with overlapping temporal scales, but only observed ages up to 18 years and thus only about half of the temporal scales considered in our study. Studies with larger overlap focused on functional rather than structural connectivity (Uhlhaas et al., 2009; Fan et al., 2010).

Here we noted that surface areas of used ROIs varied around 33–43% compared to their mean (shown only for the cortical ROIs because subcortical ROIs were not subdivided, Table 2). This variation stems from the variation of anatomical regions’ surface areas (ratio of its SD to its mean is 73.5%). The ratio of interquartile range to median showed comparatively small ROI variation (Fornito et al., 2010). To keep the topological consistency, this might be the best we can do. At least, this is better than using just anatomical regions whose variation is much higher (73.5% variation).

This node fingerprint study is a proof of principle for the current technique showing the influence of spatial and temporal scales on network comparison. However, the current results could also be influenced by the following points. First, the number of subjects per age group was not identical (Table 1). Therefore, findings in age groups with fewer members might have been influenced by relatively few outliers. Whereas an identical number of subjects per group is desirable, for our study there was also a trade-off with the age range: using the same subject numbers would have led to groups where the age ranges differ, e.g., covering 3 years for one group and 5 years for another. We decided for a different number of subjects per group in order to perform the comparison across the temporal scale. Second, single node motifs become more robust for larger network sizes. As node motifs are outliers, only a small percentage of all nodes will be characterized as node motifs. Therefore, for low-resolution networks, node motifs might only occur for few subjects. In addition, their number might be a poor estimate of the underlying frequency in human connectivity networks. For this reason, we have only reported networks with at least 414 nodes, leaving out networks with a parcelation into 110 cortical and subcortical regions. Third, node features but not necessarily motifs depend on the edge density of the network. In our developmental networks, we observed significant edge density changes for different spatial resolutions ranging from more than 7% for 414 nodes to 2% for 1615 nodes. However, for each network resolution, edge densities remained comparable for different temporal scales (age groups; Figure 1). Fourth, deterministic tracking cannot capture crossing fibers which can be done in probabilistic tracking (Behrens et al., 2007), in Diffusion spectrum imaging (Wedeen et al., 2008; Bassett et al., 2010), or in high-angular resolution diffusion imaging.
(HARDI; Tuch, 2004; Zalesky et al., 2010). However, fiber length distributions, influencing our three spatial features, were similar across resolutions (Figure 7).

Network Science has led to a wide range of tools for analyzing neural systems (Sporns et al., 2004; Costa et al., 2007b; Bullmore and Sporns, 2009; Rubinov and Sporns, 2010; Kaiser, 2011). Whereas the characterization of individual networks, e.g., as small-world or scale-free, is now possible, comparing different networks is still a challenge. Our study shows how node properties, given by characteristic single node motif fingerprints, can be compared between different networks. Studying single node properties is only one way to characterize networks with alternatives being (a) the study of global properties including local and global efficiency (Latora and Marchiori, 2001; Achard and Bullmore, 2007) as well as modularity (Newman, 2006), (b) the comparison of the cluster organization of different networks, and (c) direct comparison of network matrices (Crofts and Higham, 2009). However, it is problematic for these measures that network features are influenced by the number of nodes and edges of a network; feature changes between networks might simply be due to different edge densities in the compared systems (van Wijk et al., 2010). For our networks, edge densities did not significantly differ across age but they differed across spatial resolutions. Whereas the underlying topological and, to some extent, spatial features might have changed in our networks, our motif fingerprints do not depend on the absolute values for network measures but on their distribution: an overall increase or decrease in the average value of a measure will have less of an influence on the number of node motifs as these depend on statistical outliers and not on absolute values of a measure.

Nonetheless, nodes that present characteristic node motifs, as well as hubs, are rare within networks. These fingerprints can therefore only be applied for high-resolution networks (≥2414 nodes) and large groups of subjects. Fortunately, large subject cohorts are currently being recruited in several initiatives including the Human Connectome Project for structural connectivity and the 1000 Functional Connectome Project (Biswal et al., 2010). In addition, fingerprints will also be useful for high-resolution networks at the scale of the micro-connectome observing connections between individual neurons (DeFelipe, 2010). Another potential application lies in the analysis of multi-electrode array recordings, which nowadays can record from more than 4000 channels (Sernagor et al., 2010). We therefore made this tool available within the CARMEN initiative3 for developing electrophysiology analysis tools (Smith et al., 2007). The tool is also available on our website4.

5 CONCLUSION

In this study we found that (a) node motifs change over time and can therefore be used to find characteristic changes in the developing brain and (b) motifs change over spatial scales, therefore comparisons between studies are only meaningful when the same (or a comparable) spatial resolution has been used. Our results also indicate that spatial resolution has a higher effect on topological measures whereas spatial measures, based on fiber lengths, remain more comparable between resolutions. As node motifs are based on topological and spatial properties of brain connectivity networks, these conclusions are also relevant to other studies using network analysis. Another important aspect is the analysis of differences between healthy controls and subjects with brain disorders that can arise during brain network development, such as schizophrenia and epilepsy; we hope that our node fingerprint approach will be useful for detection and specification in these cases.

ACKNOWLEDGMENTS

Marcus Kaiser and Christoph Echtermeyer were funded by EPSRC (EP/G093950X/1) and the CARMEN e-science Neuroinformatics project (http://www.carmen.org.uk) funded by EPSRC (EP/E002331/1). Marcus Kaiser and Cheol Han also acknowledge support by the WCU program through the National Research Foundation of Korea funded by the Ministry of Education, Science and Technology (R32-10142).

REFERENCES

Achard, S., and Bullmore, E. (2007). Efficiency and cost of economical brain functional networks. PLoS Comput. Biol. 3, e17. doi: 10.1371/journal.pcbi.0030017

Albert, R., and Barabási, A.-L. (2002). Statistical mechanics of complex networks. Rev. Mod. Phys. 74, 47–97.

Antiqueira, L., Rodrigues, F. A., van Wijk, B. C. M., Costa, L. D. F., and Daffertshofer, A. (2010). Estimating complex cortical networks via surface recordings – a critical note. Neuroimage 53, 439–449.

Bassett, D. S., Brown, J. A., Deshpande, V., Carlson, J. M., and Grafton, S. T. (2010). Conserved and variable architecture of human white matter connectivity. Neuroimage 54, 1262–1279.

Behrens, T. E., Berg, H. J., Jbabdi, S., Rushworth, M. F., and Woolrich, M. W. (2007). Probabilistic diffusion tractography with multiple fibre orientations: what can we gain? Neuroimage 34, 144–155.

Biswal, B. B., Mennes, M., Zuo, X. N., Gohel, S., Kelly, C., Smith, S. M., Beckmann, C. F., Adelstein, J. S., Buckner, R. L., Colcombe, S., Dogonowski, A. M., Ernst, M., Fair, D., Hampson, M., Hoptman, M. J., Hyde, J. S., Kiviniemi, V. J., Kotter, R., Li, S. J., Lin, C. P., Lowe, M. I., Mackay, C., Madden, D. J., Madsen, K. H., Margulies, D. S., Mayberg, H. S., McMahan, K., Monk, C. S., Mostofsky, S. H., Nagel, B. J., Pekar, J. I., Peltier, S. J., Petersen, S. E., Riedl, V., Rombouts, S. A., Rypma, B., Schlaggar, B. L., Schmidt, S., Seidler, R. D., Siegle, G. J., Sorg, C., Teng, G. J., Veijola, J., Villringer, A., Walter, M., Wang, L., Weng, X. C., Whitfield-Gabrieli, S., Williamson, P., Windischberger, C., Zang, Y. F., Zhang, H. Y., Castellanos, F. X., and Milham, M. P. (2010). Toward discovery science of human brain function. Proc. Natl. Acad. Sci. U.S.A. 107, 4734–4739.

Bullmore, E., and Sporns, O. (2009). Complex brain networks: graph theoretical analysis of structural and functional systems. Nat. Rev. Neurosci. 10, 186–198.

Bullmore, E. T., and Bassett, D. S. (2011). Brain graphs: graphical models of the human brain connectome. Annu. Rev. Clin. Psychol. 7, 113–140.

Butz, M., Wörgötter, E., and Van Ooyen, A. (2009). Activity-dependent structural plasticity. Brain Res. 60, 287–305.

Buzsaki, G. (2006). Rhythms of the Brain. Oxford: Oxford University Press.

Costa, L. D. F., Kaiser, M., and Hilgetag, C. C. (2007a). Predicting the connectivity of primate cortical networks from topological and spatial node properties. BMC Syst. Biol. 1, 16. doi: 10.1186/1752-0509-1-16

Costa, L. D. F., Rodrigues, F. A., Travieso, G., and Boas, P. R. V. (2007b). Characterization of complex networks: a survey of measurements. Adv. Phys. 56, 167–242.

Costa, L. D. F., Rodrigues, F. A., Hilgetag, C. C., and Kaiser, M. (2009). Beyond the average: detecting global singular nodes from local features in complex networks. Europhys. Lett. 87, 18008.

Costa, L. D. F., and Silva, F. N. (2006). Hierarchical characterization of complex networks. J. Stat. Phys. 125, 841–876.

Crofts, J. J., and Higham, D. J. (2009). A weighted communicability measure applied to complex brain networks. J. R. Soc. Interface 6, 411–414.

3http://www.carmen.org.uk

4http://www.biological-networks.org/
Desikan, R., Segonne, F., Fischl, B., Quinn, B., Dickerson, D., Blacker, D., Buckner, R., Dale, A., Maguire, R., Hyman, B., Albert, M., and Killiany, R. (2006). An automated labeling system for subdividing the human cerebral cortex on mri scans into gyral based regions of interest. Neuroimage 31, 968–980.

Duda, R. O., Hart, P. E., and Stork, D. G. (2001). Pattern Classification. 2nd Edn. New York, NY: Wiley Interscience.

Echtermeyer, C., da Fontoura Costa, L., Rodrigues, F. A., and Kaiser, M. (2011). Automatic network fingerprinting through single-node motifs. PLoS ONE 6, e15765. doi: 10.1371/journal.pone.0015765

Eguiluz, V. M., Chialvo, D. R., Cecchi, G. A., Baliki, M., and Apkarian, A. V. (2005). Scale-free brain functional networks. Proc. Natl. Acad. Sci. U.S.A. 102, 11773–11778.

Fair, A. D., Cohen, A. L., D’Esposito, M., Dosenbach, N. U. F., Church, J. A., Miezin, F. M., Schlaggar, B. L., and Petersen, S. E. (2009). Functional brain networks develop from a ‘local to distributed’ organization. PLoS Comput. Biol. 5, e1000381. doi: 10.1371/journal.pcbi.1000381

Fan, Y., Shi, F., Smith, J. K., Lin, W., Gilmore, J. H., and Shen, D. (2010). A tutorial in connectivity analysis: from the connectome to the synaptome: an epic love story. Front. Syst. Neurosci. 4:8. doi: 10.3389/fnnsys.2010.00022

Fischl, B., Cammoun, L., Nguyen, N., Jezzard, P., Di Martino, A., Busa, E., Maquet, P., Vidal, M., Concha, L., Della Riuta, D., and Busatto, G. F. (2010). Mapping human whole-brain structural networks with diffusion mri. PLoS ONE 2, e951. doi: 10.1371/journal.pone.0000957

Fischl, B., Sorg, S., Dale, A. M., Sereno, M. I., and Toga, A. W. (2001a). Whole brain segmentation: automated labeling of neuroanatomical structures in the human brain. Neuron 33, 315–327.

Fischl, B., Salat, D., Busa, E., Albert, M., Dieterich, M., Haselgrove, C., van der Kouwe, A., Killiany, R., Kennedy, D., Klaveness, S., Montillo, A., Makris, N., Rosen, B., and Dale, A. (2002). Whole brain segmentation: automated labeling of neuroanatomical structures in the human brain. Neuroimage 13, 515–529.

Fischl, B., Salat, D., van der Kouwe, A., Makris, N., Ségonne, F., Quinn, B., and Dalea, A. (2004a). Segmentation-independent representation of magnetic resonance images. Neuroimage 23(Suppl. 1), S69–S84.

Fischl, B., Van der Kouwe, A., Destrieux, C., Halgren, E., Ségonne, F., Salat, D., Busa, E., Seidman, L. D., Goldstein, J., Kennedy, D., Caviness, V. S., Makris, N., Rosen, B., and Dale, A. (2004b). Automatically parcellating the human cerebral cortex. Cereb. Cortex 14, 11–22.

Fornito, A., Zalesky, A., and Bullmore, E. T. (2010). Network scaling effects in graph analytic studies of human resting-state fMRI data. Front. Syst. Neurosci. 4:22. doi: 10.3389/fnssy.2010.00022

Hagmann, P., Cammoun, L., Gigandet, X., Meuli, R., Honey, C. J., Wedeen, V. I., and Sporns, O. (2008). Mapping the structural core of human cerebral cortex. PLoS Biol. 6, e159. doi: 10.1371/journal.pbio.0060159

Hagmann, P., Kurant, M., Gigandet, X., Thiran, P., Wedeen, V., Meuli, R., and Thiran, J. (2007). Mapping human whole-brain structural networks with diffusion mri. PLoS ONE 2, e951. doi: 10.1371/journal.pone.0000957

Hagmann, P., Sporns, O., N. M. F., Cammoun, L., R. P., Thiran, P., Wedeen, V., Meuli, R., and Thiran, J., and Grant, P. (2010). White matter maturation reshapes structural connectivity in the alte developing human brain. Proc. Natl. Acad. Sci. U.S.A. 107, 19067–19072.

Hayasaki, S., and Lauriacci, P. I. (2010). Comparison of characteristics between region- and voxel-based network analyses in resting-state fMRI data. Neuroimage 50, 499–508.

Hellyer, B. (2000). A quantitative analysis of the local connectivity between pyramidal neurons in layers 2/3 of the rat visual cortex. Biol. Cybern. 82, 111–121.

Hilgetag, C. C., and Barbas, H. (2006). Role of mechanical factors in the morphology of the primate cerebral cortex. PLoS Comput. Biol. 2, e22. doi: 10.1371/journal.pcbi.0020022

Hilgetag, C. C., Burns, G. A. P. C., O’Neill, M. A., Semaille, J. L., and Young, M. P. (2000). Anatomical connectivity defines the organization of clusters of cortical areas in the macaque monkey and the cat. Philos. Trans. R. Soc. Lond. B Biol. Sci. 355, 91–110.

Horton, J. C., and Adams, D. L. (2005). The cortical column: a structure without a function. Philos. Trans. R. Soc. B Biol. Sci. 360, 837–862.

Hubel, D., Wiesel, T., and LeVay, S. (1977). Plasticity of ocular dominance columns in monkey striate cortex. Philos. Trans. R. Soc. Lond. B Biol. Sci. 278, 377–409.

Johnson, R. A., and Wichern, D. W. (2007). Applied Multivariate Statistical Analysis, 6th Edn. Englewood Cliffs, NJ: Prentice Hall.

Kaiser, M. (2011). A tutorial in connectome analysis: topological and spatial features of brain networks. Neuroimage 57, 892–907.

Kaiser, M., Goerner, M., and Hilgetag, C. C. (2004a). Criticality of spreading dynamics in hierarchical cluster networks without inhibition. New J. Phys. 9, 110.

Kaiser, M., Martin, R., Andras, P., and Young, M. P. (2007b). Simulation of abnormal neural oscillations and cognitive outburst. Brain 130, 8577–8582.

Kaiser, M. E. J. (2003). The structure and function of complex networks. SIAM Rev. 45, 174–205.

Kaiser, M. E. J. (2006). Modularity and community structure in networked dynamical systems. Proc. Natl. Acad. Sci. U.S.A. 103, 8577–8582.

Kaiser, M. E. J., Barabasi, A. L., and Watts, D. J. (2006). The Structure and Dynamics of Networks. Princeton, NJ: Princeton University Press.

Kaiser, M. (2012). On estimation of a probability density function and mode. Ann. Math. Stat. 33, 1065–1076.

Passingham, R. E., Stephan, K. E., and Kotter, R. (2002). The anatomical basis of functional localization in the cortex. Nat. Rev. Neurosci. 3, 606–616.

Roozendaal, B., Rabin, M. A., Crevilla, M., De Jong, P., and Daffertshofer, A. (2010). Comparing brain networks of different size and connectivity density using graph
theory. PLoS ONE 5, e13701. doi: 10.1371/journal.pone.0013701
Wang, R., Benner, T., Sorensen, A., and Wedeen, V. (2007). Diffusion
toolkit: a software package for diffusion imaging data processing
and tractography. Int. Soc. Magn. Reson. Med. 15, 3720.
Watts, D. J., and Strogatz, S. H. (1998). Collective dynamics of
‘small-world’ networks. Nature 393, 440–442.
Wedeen, V. J., Wang, R. P., Schmahmann, J. D., Benner, T., Tseng, W. Y. I., Dai,
G., Pandya, D. N., Hagmann, P., D’Arceul, H., and de Crespigny, A. J.
(2008). Diffusion spectrum magnetic resonance imaging (DSI)
tractography of crossing fibers. Neuroimage 41,
1267–1277.
Zalesky, A., Fornito, A., Harding, I. H., Cocchi, L., Yücel, M., Pantelis, C.,
and Bullmore, E. T. (2010). Whole-brain anatomical networks: does
the choice of nodes matter? Neuroimage 50, 970–983.

Conflict of Interest Statement: The authors declare that the research was
conducted in the absence of any commercial or financial relationships that could be
construed as a potential conflict of interest.
Received: 15 April 2011; accepted: 02 July 2011; published online: 22 July 2011.
Citation: Echtermeyer C, Han CE, Rotarska-Jagiela A, Mohr H, Uhlhaas PJ
and Kaiser M (2011) Integrating temporal and spatial scales: human structural network motifs across age and region of
interest size. Front. Neuroinform. 5,10.
doi: 10.3389/fninf.2011.00010
Copyright © 2011 Echtermeyer, Han, Rotarska-Jagiela, Mohr, Uhlhaas and
Kaiser. This is an open-access article subject to a non-exclusive license between the
authors and Frontiers Media SA, which permits use, distribution and reproduction in
other forums, provided the original authors and source are credited and other Frontiers
conditions are complied with.
### APPENDIX

Table A1 | Comparison of our findings with previous studies.

| Study                  | Findings                                                                 |
|------------------------|---------------------------------------------------------------------------|
| **Spatial scale**      |                                                                           |
| Ours                   | As the number of nodes increases                                          |
| Zalesky et al. (2010)  | In SC (414, 813, and 1615 nodes), ↓ ED, ↑ w (see Figure 4)                |
| Bassett et al. (2010)  | In SC (6 different scales between 82 and 4000 nodes), ↓ ED, ↑ small world, ↓ γ, ↑ global efficiency, changes in nodal rank degree, and betweenness centrality |
| Hagmann et al. (2010)  | In SC (12 different scales between 54 and 880), ↓ ED, ↓ γ, ↑ λ, conserved hierarchy, ↑ Rentian scaling |
| Fornito et al. (2010)  | In rsFC (7 different scales between 84 and 4320 nodes), ↓ average correlation, ↑ size of largest component, ↓ path length, ↑ CC (for low ED), ↓ CC (for high ED) ↑ small worldness, changes in nodal rank degree |
| Hayasaka and Laurienti (2010) | In rsFC (voxel-based nodes with three different voxel sizes and region-based nodes), changes in node degree distribution ↑ CC, ↑ path length (while increasing voxel sizes, but it decreased with region-based node) |
| **Temporal scale**     |                                                                           |
| Ours                   | As the age increases                                                      |
| Hagmann et al. (2010)  | In SC (18 m/o to 18 y/o), ↓ mean ADC, ↑ mean FA, ↓ ED, ↓ CC, ↑ Efficiency, ↑ Node strength, ↑ SC–FC correlation, no changes in modularity and no major changes in module composition after 2 y/o, no significant changes in betweenness centrality |
| Fan et al. (2010)      | In SC (1 y/o, 1 y/o, 2 y/o, and adult), ↑ global efficiency, ↑ cost efficiency, a peak at 2 y/o in local efficiency and modularity, ↑ size of largest component in modules, changes in module assignment, and participation coefficient |
| Fair et al. (2009)     | In rsFC (7–31 y/o), no significant changes in optimized modularity Q, CC and CPL. Changed module assignment and the number of long-distance correlations ↑ |
| Uhlhaas et al. (2009)  | In EEG (6–21 y/o), strong correlation between neural synchrony and cognitive performance. As ages increase, the neural synchrony was increasing in early adolescence, then decreasing in late adolescence, and finally increasing again in adult |

SC, structural connectivity; rsFC, resting state functional connectivity; m/o, months-old; y/o, years-old; ED, edge density; CC, clustering coefficient; CPL, characteristic path length; ADC, apparent diffusion coefficient; FA, fractional anisotropy; γ, CC/CCrand; λ, CPL/CPLrand.