Spatiotemporal Dependency of Age-Related Changes in Brain Signal Variability

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Recent theoretical and empirical work has focused on the variability of network dynamics in maturation. Such variability seems to reflect the spontaneous formation and dissolution of different functional networks. We sought to extend these observations into healthy aging. Two different data sets, one EEG (total n = 48, ages 18–72) and one magnetoencephalography (n = 31, ages 20–75) were analyzed for such spatiotemporal dependency using multiscale entropy (MSE) from regional brain sources. In both data sets, the changes in MSE were timescale dependent, with higher entropy at fine scales and lower at more coarse scales with greater age. The signals were parsed further into local entropy, related to information processed within a regional source, and distributed entropy (information shared between two sources, i.e., functional connectivity). Local entropy increased for most regions, whereas the dominant change in distributed entropy was age-related reductions across hemispheres. These data further the understanding of changes in brain signal variability across the lifespan, suggesting an inverted U-shaped curve, but with an important qualifier. Unlike earlier in maturation, where the changes are more widespread, changes in adulthood show strong spatiotemporal dependence.

Keywords: electroencephalography, functional connectivity, magnetoencephalography, multiscale entropy, nonlinear dynamics

Introduction

It is becoming clear that brain signal variability (i.e., transient temporal fluctuations in brain signal) conveys important information about network dynamics (Deco et al. 2011). In neural network modeling, information integration across widespread neural networks is achieved through the emergence and disappearance of correlated activity between brain regions over time and across multiple timescales (Jirsa and Kelso 2000; Honey et al. 2007). Such transient changes cause fluctuations in the temporal dynamics of the corresponding brain signal; networks with more potential configurations elicit a more variable response. In turn, signal variability may represent the information processing capacity of the system, where higher variability reflects greater information integration across the network.

Our empirical studies have noted developmental increases of brain signal variability that could be related to the idea that variability reflects the capacity for information processing. The first study (McIntosh et al. 2008) examined EEG signal variability using two measures; (1) principal components analysis (PCA) of single trial data, where a more variable brain would produce a greater number of principal components, and (2) multiscale entropy (MSE), a measure that is sensitive to linear and nonlinear variability and is able to differentiate variability of a complex system such as the brain, from a purely random system (Costa et al. 2002; Costa et al. 2005). Both PCA and MSE analyses indicated that EEG signal variance increased from ages 8 to 15 and was even higher in young adults. Furthermore, behavioral stability (measured by accuracy and intra-subject variability of reaction time) was greater with a more variable brain signal, suggesting that a more variable brain produces more stable behavior. Misic et al. (2010) replicated this finding across a slightly wider age range (6–16 years) and using magnetoencephalography (MEG). Finally, Lippe et al. (Lippe et al. 2009) extended these observations to infants, showing that EEG signal complexity increased from 1 month to 5 years in response to auditory and visual stimuli. Interestingly, the trajectories of the change were not the same for the auditory and visual responses, in that auditory responses were more complex than visual in infants and complexity in both modalities gradually merged to be indistinguishable in adults.

During maturation from childhood to adulthood, the brain becomes more specialized in that a larger repertoire of individualized physiological states for separate brain regions develops. Integration between distributed neuronal populations also increases. Vakorin et al. (Vakorin et al. 2011) introduced a method to decompose the total variability of signals into local entropy that characterizes the dynamics within brain areas and distributed entropy that characterizes the signal variability attributed to information exchange between areas. Using these measures, the present study explored the interplay between specialization and integration and how these factors contribute to changes in brain complexity in development. The results suggested that increased integration was the key factor contributing to the developmental increase in complexity of brain signals. Specifically, developmental changes were characterized by a decrease in the amount of information processed locally, accompanied by an increase in distributed entropy.

The structural and functional changes during adult aging are less global than those during early maturation. Extant literature has characterized more heterogeneous changes in structure, with areas showing different rates of change with age (Guttmann et al. 1998; Courchesne et al. 2000; Raz and Rodrigue 2006). Similarly, functional networks seem to show different rates of change that seem to reflect changes in the cognitive functions they support (Grady 1998; Greenwood 2000; Grady et al. 2003; Andrews-Hanna et al. 2007; Damoiseaux et al. 2007; Park and Reuter-Lorenz 2009). If we consider such alterations in the context of nonlinear systems, one may predict that brain signal variability should also show changes, but perhaps less globally than was observed for children. Indeed some of our work with functional MRI suggests both regional increases and decreases of variability in normal aging (Garrett et al. 2011; Garrett et al. 2012).
Our measures of brain signal variability can be most readily appreciated from the perspective of complex systems. Compared with deterministic or random systems, complex systems such as the brain have a greater capacity for information processing (Tononi et al. 1994; Sporns et al. 2000a; Sporns et al. 2000b). Entropy measures are sensitive to the information content in signals, and thus the measures of MSE, distributed and local entropy, are estimates of the information and so too, complexity. Thus, we can reframe the focus of the study as follows: development brings with it the evolution of a brain that has greater capacity for information processing. Whereas this evolution likely continues into early adulthood, it is less clear how the trend changes in senescence. A handful of studies that have measured complexity of EEG in aging have produced conflicting results (e.g., Anokhin et al. 1996; Pierce et al. 2000; Gaal et al. 2010). Thus, a key aspect of our current work is to characterize the different temporal scales over which information integration may take place in the brain. Although we found that information processing capacity increases across most scales in development, given that changes in early adulthood are not as global as they are early in life, it is likely that the characterization of changes in information processing capacity will depend on spatiotemporal scale.

We sought to characterize the changes in brain signal variability across two different adult samples from two studies. One study tested adults from 19 to 72 years and measured EEG in a range of visual perception tasks. The second study tested adults from 20 to 75, with no participants in the middle-age range and examined MEG response in a multisenсорy task (Diaconescu et al. 2012). For both data sets, analysis was done based on cortical source solutions rather than in original sensor space to increase the spatial location precision, given the spatial differences reported for age-related changes and to minimize the artifact of volume conduction for analyses of functional connectivity (Srinivasan et al. 1998). The purpose of the present work was to get general characterizations of age-related changes in variability to see how they complement those we observed early in development.

Methods

The study was based on the analysis of two different data sets. Both focused on the age-related changes in evoked activity to sensory stimuli. Both studies were approved by the joint Baycrest Centre–University of Toronto Research Ethics Committee, and the rights and privacy of the participants were observed. All participants gave formal informed consent before the experiment and received monetary compensation.

Study 1—EEG

Sixteen young adults (6 males, mean age 22 ± 3 years), 16 middle-age adults (7 males, mean age 45 ± 6 years), and 16 older adults (5 males, mean age 66 ± 6 years) participated in the EEG study. All participants had healthy neurological histories, and normal or corrected-to-normal vision.

Apparatus and Task

EEG recordings from 76 electrodes were collected using BioSemi Active Two system with a bandwidth 99.84 (0.16–100) Hz and sampling rate of 512 Hz. Data were recorded reference free, but were converted to an average reference at Cz during the pre-processing.

The EEG data were part of a larger study that contained six different conditions (similar to the configuration of tasks used in a fMRI study by Grady et al. 2010). For the purposes of the present work, we used two tasks, visual perceptual matching (PM) and delayed-match-to-sample (DMS). Presentation (version 10.3, Neurobehandral Systems, Inc.) and Matlab (version 7, Mathworks) were used to control visual stimulus delivery and to record participants’ response latency and accuracy. Stimuli were one-dimensional Gaussian white noise fields with a two-octave frequency filter and were presented simultaneously in a triangular array. In the PM task, subjects indicated which of the three bottom stimuli matched the one on the top by pressing one of three buttons. The task instructions for DMS were the same as for PM, except that in DMS, the top row stimulus appeared first for 1.5–2 s and then disappeared before the bottom row stimuli were shown. There was a 4-s delay between the onset of top (encoding phase) and bottom stimuli (recognition phase). For the present study, only the data from PM and the encoding phase of DMS were used.

We used a psychophysical thresholding procedure to ensure that subjects were matched in terms of accuracy by adjusting stimulus discriminability so that each person was 80% accurate (Protzman and McIntosh 2007). Stimulus discriminability was manipulated by modifying the center frequency ratio. For example, at a ratio of 2, and a base frequency of 2 c/deg, the center frequencies were 2 c/deg, 4c/deg, and 8c/deg. To decrease stimulus discriminability, a ratio of 1.5 would produce center frequencies of 2 c/deg, 3 c/deg, and 4.5 c/deg.

Data Pre-processing

Continuous EEG recordings were bandpass filtered from 0.5 to 55 Hz. Only trials with correct responses were analyzed. Data were epoched and base-lined into 500-2000 ms epochs with a 500-ms pre-stimulus baseline. Artifact removal was performed using independent component analysis in EEGLAB (Delorme and Makeig 2004). There was an average of 182 trials for the PM task and 171 for the DMS task after data processing was completed for all subjects. There was no significant difference in the number of useful trials between conditions or groups.

To further localize the dynamics of source activity at specific locations, we identified 72 ROIs in Talairach space (Diaconescu et al. 2011) and performed source estimation at these locations using sLORETA (Pascual-Marqui 2002), as implemented in Brainstorm (Tadel et al. 2011), which is documented and freely available for download under the GNU general public license (http://neuroimage.csc.edu/brainstorm). Source reconstruction was constrained to the cortical mantle of a brain template MNI/Colin27 defined by the Montreal Neurological Institute. Current density for three source orientations (X, Y, and Z components) was mapped at 72 brain regions of interest adapting the regional map coarse parcellation scheme, as developed in Kotter and Wanke (2005) (see Table 1). For each subject, MSE measures of source waveforms were calculated for the PM task and the encoding part of the DMS task.

Study 2—MEG Data

Fifteen young adults (7 males, mean age 23 ± 3 years) and 16 older adults (8 males, mean age 70 ± 5 years) with an average of 16.5 years of education participated. All participants were right handed with healthy neurological histories, and normal to corrected-to-normal vision. All participants were audiometrically screened to determine hearing thresholds for each ear separately; adults whose hearing thresholds exceeded 15dB hearing level were excluded from participation, as that was considered below normal levels. The young adults who participated in the study had average hearing thresholds of 2 dB (range 0–8 dB), and older adults had average hearing thresholds of 10 dB (range 5–15 dB).

Apparatus and Task

Auditory and visual stimuli were used in this study. Black and white line drawings selected from a database (Smogdass and Vandewart 1980) were used for visual presentations. All visual stimuli were matched according to size (in pixels), brightness, and contrast. Auditory stimuli were selected from a local database of non-speech, complex sounds (e.g., animal calls, car horns). Complex sounds were matched according to amplitude. Complex sounds were delivered...
MEG data were co-registered to each participant’s individual structural MRI to constrain the sources of activation to each participant’s head shape and structural anatomy. MRI scans were acquired for each participant using a 3.0T Siemens TIM MAGNETOM Trio MRI scanner (Software level Syngo MR, Siemens Medical, Germany) with 12-channel head coil.

To obtain spatial precision without integrating power over long temporal windows, we used an event-related version of the synthetic aperture magnetometry analysis technique introduced by Cheyne et al. (2006) to identify evoked brain responses from unaveraged, single trial data. The spatial filter included the same 72 brain regions of interest in Talairach coordinate space as used in the EEG data set (Table 1). The individual functional maps were overlaid on the individual participant’s MRI based on co-registration with the indicator coils placed on the nasion and bilateral pre-auricular points. The functional data were then transformed to the standard Talairach–Tournoux space using the same transformation applied to the structural MRI [AFNI software, Cox (1996)].

### MultiScale Entropy

Full details of MSE and its relevance for the analysis of signal complexity are given in Costa et al. (2002) and Costa et al. (2005). The utility of MSE for characterizing complexity of brain signals has been confirmed by numerous studies (Bhattacharya et al. 2005; McIntosh et al. 2008; Lippe et al. 2009; Takahashi et al. 2009; Misco et al. 2010; Protzner et al. 2010; Catarino et al. 2011).

To calculate MSE, we used the algorithm available at [www.physionet.org/physiotools/mse/](http://www.physionet.org/physiotools/mse/) that computes MSE in two steps. First, the algorithm progressively down-samples the EEG post-stimulus time series per trial and per condition (i.e., for timescale \( t \)), the coarse-grained time series is constructed by averaging data points within non-overlapping windows of length \( \ell \). Second, the algorithm calculates the sample entropy for each coarse-grained time series. Sample entropy quantifies the variability of a time series by estimating the predictability of amplitude patterns across a time series. The pattern length was set to \( m = 2 \); that is, two consecutive data points were used for pattern matching. The similarity criterion was set to \( r = 0.5; \) data points were considered to have indistinguishable amplitude values (i.e., to ‘match’) if the absolute amplitude difference between them was \( \leq 5\% \) of the time series standard deviation. For each subject, a source specific MSE estimate was obtained as a mean across single trial entropy measures for each timescale.

### Local and Distributed Entropy

The specific derivation of local versus distributed entropy has been described in our previous publication (Vakorin et al. 2011). In information theory (Shannon 1949), entropy \( H(X) \) of a single random variable \( X \) can be defined as a measure of uncertainty associated with \( X \). Conditional entropy \( H(X|Y) \) of \( X \) given another random variable \( Y \) is the entropy of \( X \), provided that the uncertainty about \( Y \) is excluded. The reduction in uncertainty due to another variable is called mutual information. Specifically, the mutual information between two random variables \( X \) and \( Y \) is defined as

\[
I(X;Y) = H(X) - H(X|Y).
\]

The mutual information is a measure of affiliation between two variables, similar to a correlation coefficient.

The joint entropy \( H(X,Y) \), which represents the uncertainty of a pair of random variables \( (X,Y) \), can be partitioned into the conditional entropies of the variables \( X \) and \( Y \) and the mutual information between them. Specifically,

\[
H(X,Y) = H(X|Y) + H(Y|X) + I(X;Y).
\]

Suppose that a network of \( M \) interacting neural sources is identified, and source dynamics are described by \( M \) variables \( X_i \), where \( i = 1, \ldots, M \). For a given pair of sources, \( i \) and \( j \), the information contained in the dynamics of source \( i \) can be partitioned into the local entropy associated only with source \( i \) and the distributed entropy that is shared between sources \( i \) and \( j \). The local entropy corresponds to the conditional entropy \( H(X_i|X_j) \). For a given source \( i \), the local entropy, \( E_{local}(i) \), can be computed by averaging conditional entropy \( H(X_i|X_j) \)

| Region Short name | x | y | z |  
|-------------------|---|---|---|---|
| Frontal eye fields | 36 | 8 | 56 |  
| Anterior insula | 36 | 16 | −8 |  
| Lateral prefrontal cortex | 36 | 16 | −4 |  
| Superior parietal cortex | 24 | −24 | 56 |  
| Angular gyrus | 44 | −8 | 28 |  
| Precuneus | 8 | −64 | 54 |  
| Dorsolateral prefrontal cortex | 48 | 36 | 32 |  
| Dorsomedial prefrontal cortex | 8 | 36 | 40 |  
| Medial prefrontal cortex | 8 | 48 | 20 |  
| Orbitalis | 24 | 44 | −20 |  
| Frontal polar | 24 | 64 | −4 |  
| Ventral prefrontal cortex | 48 | 32 | −8 |  
| Ventral pallidum | 28 | −16 | −16 |  
| Dorsal prefrontal motor cortex | 48 | 0 | 60 |  
| Medial prefrontal cortex | 4 | 0 | 60 |  
| Ventral prefrontal motor cortex | 16 | 4 | 24 |  
| Pulvinar | 40 | −28 | 64 |  
| S2 | 56 | −16 | 16 |  
| Middle temporal cortex | 64 | −24 | −12 |  
| Inferior temporal cortex | 64 | −24 | −24 |  
| Superior temporal cortex | 52 | −12 | 28 |  
| Ventral temporal cortex | 32 | −28 | −28 |  
| Thalamus | 8 | −8 | 4 |  
| V1 | 4 | −84 | −4 |  
| V2 | 4 | −96 | 8 |  
| Cuneus | 20 | −88 | 20 |  
| Fusiform gyrus | 20 | −84 | −12 |  

Table 1: Regional map coordinates with reference to the Talairach–Tournoux Atlas

Binaurally at an intensity level of 60 dB HL based on the audiometric mean across both ears. There were 40 trials for each condition.

Presentation software (version 10.3, Neurobehavioural Systems, Inc.) was used to control visual and auditory stimulus delivery and to record participants’ response latency and accuracy. The time interval between the end of the stimulus presentation and the beginning of the next trial was between 2 and 4 s (equiprobable). Participants were instructed to respond to any trial type, auditory or visual, as quickly as possible with their left index finger response.

The MEG recordings were acquired in a magnetically shielded room at the Rotman Research Institute, Baycrest Centre using a 151-channel whole head neuro-magnetometer (OMEGA, VSM Medtech Inc., Vancouver, Canada). Participants sat in upright position and viewed the visual stimuli on a back projection screen that subtended approximately 30° of visual angle when seated 70 cm from the screen. With regard to the visual presentations, the MEG collection was synchronized to the onset of each stimulus by recording the luminance changes of the screen with a photodiode. Binaural auditory stimuli were presented at 60 dB HL via OB 822 Clinical Audiometer through ER30 transducers (Etymotic Research, Elk Grove, USA) and connected with 1.5 m of length matched plastic tubing and foam earplugs to the participants’ ears. With respect to the auditory stimuli, the MEG data collection was synchronized to the onset of the auditory sound envelope.

Neuromagnetic activity was sampled at a rate of 1250 Hz. Third gradient noise correction was applied to the continuous MEG data. Afterward, the MEG data were parsed into epochs including a 200 ms pre- and 1000 ms post-stimulus activity window, and DC offsets were removed from the entire epoch. The data were bandpass filtered between 0.1 and 55 Hz. A principal component analysis was performed on each epoch, and components larger than 2.0 picoTesla (pT) at any time point were subtracted from the data to remove large artifacts caused by eye blinks.

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of time series at various scales that is similar to the approach used for MSE. Downsampling alleviates linear stochastic effects, such as autocorrelation of the brain signals, that might lead to a bias in estimation of the entropy and mutual information (Kaffashi et al. 2008).

**PLS Analysis.** Partial least squares analysis (PLS McIntosh et al. 1996; McIntosh and Lobaugh 2004; Krishnan et al. 2011) was used to assess age-related changes in spatiotemporal distributions of the entropy measures. Similar to multivariate techniques like canonical correlation analysis, PLS operates on the entire data structure at once, extracting the patterns of maximal covariance between two data sets.
matrices, in the present case group membership (age) and the entropy measures.

Statistical assessment in PLS is done across two levels. First, the overall significance of each pattern, or latent variable (LV), that relates the two data sets is assessed with permutation testing (Good 2000). An LV was considered significant if the observed pattern was present less than 1% of the time in random permutations (e.g., $P < 0.01$). The second level uses bootstrap resampling to estimate confidence intervals around the individual weights in each LV, allowing for an assessment of relative contribution of particular locations and timescales, in the case of the brain side of the equation, and the stability of the relation with age group in the case of the other side of the equation (Efron and Tibshirani 1986; Efron and Tibshirani 1993). For the brain data, we plot bootstrap ratios (ratio of the individual weights over the estimated standard error) as a proxy for the confidence interval. Confidence intervals are plotted for group effects. A minimum threshold of a stable 95% confidence interval was used for all analysis.

Results

Behavior Analysis

EEG Experiment: all groups maintained an accuracy level at or greater than 80% for all tasks, and this level did not differ

Figure 2. PLS result from the MEG data set for the comparison of age-related changes in MSE. (A) The bar graph depicts the data-driven contrast between age groups that was significantly expressed across the entire data set (source $\times$ timescales) as determined by permutation tests. (B) The statistical image plot represents the regional sources and timescale at which the contrast was most stable as determined by bootstrapping. Values represent the ratio of the parameter estimate for the source divided by the bootstrap-derived standard error (roughly $z$ scores). Positive values indicate timescales and sources showing decreased MSE with age and negative values denote age-related increases. Text labels in bold font indicate regions where the bootstrap ratio is greater than 4.0. (C) Average MSE ($\pm SE$) curves for each group for the regional source in right dorsolateral prefrontal cortex for auditory (top) and visual (bottom). Circle above the curves indicate the timepoints with reliable confidence intervals.
between groups. Mean reaction time was different between groups, showing the typical slowing effect with age (PM: $F(2,45) = 4.14$, $P < 0.05$; DMS: $F(2,45) = 8.92$, $P < 0.01$). There was a slight group difference in the PM task only, with the youngest age group showing more variation than the middle age or oldest group ($F[2,45] = 4.3$, $P < 0.05$).

MEG Experiment: accuracy was above 90% for both groups in the two detection tasks, with the older subjects showing slightly lower accuracy in the visual condition (91 vs. 98%). Mean reaction time was slower in general for older subjects ($F[1,29] = 5.88$, $P < 0.05$), and the coefficient of variation of reaction time was higher in the older group ($F[1,29] = 12.36$, $P < 0.01$).

Aging and MSE
In both data sets, we observed significant changes in MSE distribution with age. There were no reliable interactions of age with task in both data sets. Although we treated age as a categorical grouping factor, the results do not change if age is treated as a continuous measure. Unlike what we have observed in children, this age effect depended on temporal scale.

In the EEG data set, we observed a progressive increase in sample entropy at fine temporal scales with age (Fig. 1A), roughly from 2 to 15 ms and then a cross-over at coarse scales with young subjects showing greater sample entropy at approximately 20 ms (Fig. 1, B, C). These differences were not homogeneous across source space, with the strongest effects in medial cortical sources such as precuneus and superior parietal lobe (Fig. 1C).

For the MEG data set, a similar pattern was observed, although the age differences were much stronger (Fig. 2A). At fine scales, the older adults showed greater sample entropy from 2 to 15 ms. The cross-over of the curves took place around 16 ms at which point sample entropy was higher in young adults (Fig. 2B, C), similar to the EEG data. The effects of fine scale were quite robust across most sources, whereas the difference at coarse scales showed more spatial variation concentrating mainly on temporal and occipital cortical sources (Fig. 2C).

Local Versus Distributed Entropy

Local Entropy
For local entropy, PLS analyses revealed the existence of one significant LV in both MEG and EEG data. Specifically, Figure 3 shows the age-related patterns of changes in local entropy for MEG (left column) and EEG (right column). The upper panels in Figure 3 illustrate the corresponding age-related trends of changes in the variability of brain signals associated with the information processed locally.

The overall distributions of the bootstrap values for all sources and timescales are shown in Figure 3B and E, for MEG and EEG, respectively. Figure 3C and F shows the same bootstrap ratio values as the function of timescales for all brain sources, each associated with a curve. As can be seen,
the overall distributions of the bootstrap values are significantly skewed toward negative numbers. Considered together with the effect in Figure 3A and D, this would imply that the dominant effect in aging is an increase of amount of information processed locally.

**Distributed Entropy**

One significant LV was found both for the EEG and MEG data in the analysis of distributed entropy. The upper panels A and E in Figure 4 illustrate the corresponding effects (P-value < 0.001 in both cases) underlying the group differences. The corresponding overall distributions of the bootstrap ratio values associated with the pairwise connections between the brain regions and timescales are given in Figure 4B and G. As can be seen, there are relatively large negative and positive bootstrap ratio values, although the bulk of the connections show positive ratios. In conjunction with the effect in Figure 4A and E, this would indicate that there are more decreases than increases of distributed entropy in relation to aging. This difference is strongest in the MEG data set.

The difference between decreased and increased distributed entropy is broken down further in Figure 4, which shows how the strength of effects attributed to age-related changes in distributed entropy varies across timescales. Specifically, we identified the 2.5%-tails, cut off by 0.025- and 0.975-quantiles, of the overall distributions of the bootstrap ratio values in Figure 4B and F. At each timescale, the number of connections with the bootstrap ratio values larger than the 0.975-quantile was computed. These numbers are plotted in Figure 4C and G. In a similar way, the number of connections with the bootstrap ratio values smaller than the 0.025-quantile is plotted in Figure 4C and G, as a function of timescale.

Figures 5 and 6 show the spatial patterns supporting the corresponding age-related changes in connectivity between the brain regions. We plotted the bootstrap ratio values at scales around 7 ms for decreased entropy, and 27 ms for increased entropy, for MEG, and around 20 ms (decreased) and 8 ms (increased), for EEG. These time points correspond to the peak distribution of bootstrap by time shown in Figure 4, hence the difference in the peaks between the data sets. The spatial distribution of these changes did not change appreciably across other time points.

Pairwise connections that show decreased entropy with age are depicted in Figure 5. A threshold of >1 was used for the figure to emphasize the spatial distribution. The corresponding lower panels of Figure 5 show the distribution of the bootstrap ratio values associated with local entropy. Similar to Figure 5, Figure 6 shows the distribution of connections showing increased entropy with age.

An obvious spatial pattern is evident in Figures 5 and 6. The networks can be differentiated from the perspective of inter- versus intra-hemispheric connections. As can be seen in

![Figure 4](image-url). Age-related changes in distributed entropy: MEG (left) and EEG (right), from the PLS analysis. Panels (A) and (E) illustrate the data-driven contrasts. Expression of these trends is shown in (B) and (G) as the distributions of all the bootstrap ratio values for the distributed entropy between each pair of regional sources. The strengths of the effects related to a decrease in distributed entropy are plotted as the number of connections whose bootstrap ratio values belong to the corresponding 2.5% tails.
Figure 5. Matrix showing bootstrap ratios for pairwise age-related decreases in distributed entropy based on (A) MEG data; (B) EEG data. Plotted in the matrices, from the 7 ms timescale for MEG and 20 ms for EEG, are the connections associated with the bootstrap ratio values that are larger than one. The corresponding distribution of the bootstrap ratio values for local entropy across brain regions is shown below the matrix.
Figure 6. Matrix showing bootstrap ratios for pairwise age-related increases in distributed entropy based on (A) MEG data; (B) EEG data. Plotted in the matrices, from the 27 ms timescale for MEG and 8 ms for EEG, are the connections associated with the bootstrap ratio values that are larger than one. The corresponding distribution of the bootstrap ratio values for local entropy across brain regions is shown below the matrix.
Figure 5, the dominant effect in the pattern of age-related decreases in connectivity is represented by the communication between the hemispheres (right/left and left/right quadrants). In contrast to Figure 5, Figure 6 reveals that the increase in connectivity is supported mostly by the connections linking the brain areas that belong to the same hemispheres (left/left and right/right quadrants). It is worth noting that there is an overall agreement between MEG and EEG modalities with respect to the inter- and intra-hemispheric connections. It is noteworthy that in the case of EEG, however, the effects related to increased connectivity as a result of normal aging are expressed most strongly in the right hemisphere. We emphasize again that for both figures, we purposely chose a liberal statistical threshold for the plots to indicate that the pattern is not an artifact of an arbitrary threshold.

Discussion

Across two independent data sets, one using EEG and one MEG and each using different source reconstruction algorithms, we observed similar changes in brain signal variability, measured using MSE, with normal aging. As was hypothesized, these differences were dependent on the temporal scale of investigation. Finer timescales showed an increase with normal aging, whereas more coarse timescales showed a decrease in normal aging. The MSE results were then broken down further using the estimation of local and distributed entropy. The significance of the MSE findings can be appreciated in two different perspectives. First, there has been an emphasis in the neurophysiology literature on long-range correlations within time series as a reflection of the formation of associations and networks (Nunez 1989). Although it is likely the case that finer timescales reflect local dynamics, in this case reflecting neural populations, the more coarse timescales would reflect longer-range interactions with other populations. The observations in this paper suggest that with age, comes a shift from long-range connections (captured by coarse scales in MSE and distributed entropy estimates) to more local processing (captured at fine scales in MSE and local entropy estimates). This stands in a nice juxtaposition to the observations early in life that seemed to reflect an increase in long-range interactions and a concomitant decrease in local entropy (Vakorin et al. 2011). We found that in aging, this situation is reversed, suggesting that the interplay between integration and specialization can be described as a U-shaped function of age over the lifespan. We also observed a widespread increase in local entropy. This may suggest more functional independence for different brain areas in normal aging. There is correspondence between this idea and observations from graph theory metrics applied to functional connectivity in fMRI across young and old adults (Meunier et al. 2009). The essential finding from that study was that older adults on average had more modules than younger adults, with few that integrated spatially distant nodes. It is reasonable to link this finding with the current ones by suggesting that an increase in modularity would correspond to the observed increase in local entropy and potentially the concomitant decrease of distributed entropy.

An important observation from our study is that the age-related changes are temporally dependent. This is quite evident from the MSE analysis, where fine scales showed higher entropy and coarse scales showed lower entropy with age. In some respects, such temporal dependence could have been predicted from the work on spectral power changes in normal aging, in which low frequencies, in general, decrease in power whereas higher frequencies tend to increase (Dustman et al. 1993; Dustman et al. 1999), which was also observed in our data sets (see Supplementary Figures). Our MSE results mirror this. Moreover, the analysis of local and distributed entropy extend the interpretation by showing that local entropy, which is most highly correlated with fine scales in MSE, increases with age, and is mostly reflected in high frequency dynamics.

Although tempting to make a general statement that distributed entropy shows a complementary decrease in aging, the picture is not so simple. We do note that there are more decreases than increases in distributed entropy, but the decreases predominantly involve cross-hemispheric interactions. Studies of EEG coherence have also noted the reduction in interhemispheric functional connections with age (Duffy et al. 1996; Kikuchi et al. 2000). The observed decrease in distributed entropy seems to be consistent with previous studies, indicating that both the axonal and myelin integrity of the white matter is compromised in aging (Bartzokis et al. 2004; Head et al. 2004; Persson et al. 2006; Makris et al. 2007; Kennedy and Raz 2009; Seidler et al. 2010). In particular, a number of studies reported age-related deterioration of white-matter microstructure of corpus callosum (Doraismamy et al. 1991; Sloane et al. 1999; Abe et al. 2002; Sullivan et al. 2010) that is in accordance with our finding indicating age-related decrease in distributed entropy between the hemispheres.

Conclusions

The intersection of studies in human neuroimaging from EEG and fMRI have emphasized that the brain operates at many different spatial and temporal scales, whereas theoretical expositions underscore the importance of the space–time structure as key to understanding processing capacity of brain networks (Jirsa and Kelso 2000; Deco et al. 2011). The essential points of this study revolve around spatiotemporal dependency as captured by measures of brain signal variability. In particular, we complete the picture of maturational changes in signal variability showing a general inverted-U trend from childhood to old age. There is an important caveat here—the nature of the latter part of this trend is critically dependent on temporal scale.

Supplementary Material

Supplementary material can be found at: http://www.cercor.oxfordjournals.org/.

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Notes

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