Structural substrates for resting network disruption in temporal lobe epilepsy

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Magnetic resonance imaging methods that measure interregional brain signalling at rest have been advanced as powerful tools to probe organizational properties of functional networks. In drug-resistant temporal lobe epilepsy, resting functional magnetic resonance imaging studies have primarily employed region of interest approaches that preclude a comprehensive evaluation of large-scale functional interactions. In line with the distributed nature of structural damage in this condition, we set out to quantify connectivity across the entire range of resting networks. Furthermore, we assessed whether connectivity is driven by co-localized structural pathology. We obtained resting state, diffusion tensor and anatomical imaging data in 35 patients with temporal lobe epilepsy and 20 healthy subjects on a 3 T scanner. Resting state networks were identified using independent component analysis, which allows an objective whole-brain quantification of functional connectivity. We performed group comparisons before and after correcting for voxel-wise grey matter density. In addition, we identified voxel-wise associations between resting connectivity and white matter coherence indexed by fractional anisotropy. Compared with controls, patients showed altered (typically reduced) functional connectivity between the hippocampus, anterior temporal, precentral cortices and the default mode and sensorimotor networks. Reduced network integration of the hippocampus was explained by variations in grey matter density, while functional connectivity of the parahippocampus, and frontal and temporal neocortices showed atypical associations with white matter coherence within pathways carrying connections of these regions. Our multimodal imaging study suggests that in temporal lobe epilepsy, cortical atrophy and microstructural white matter damage impact functional resting connectivity.

Keywords: functional MRI; diffusion tensor; networks; epilepsy
Abbreviation: TLE = temporal lobe epilepsy

Introduction

Temporal lobe epilepsy (TLE) is the most frequent form of drug-resistant focal epilepsy. Although mesiotemporal sclerosis is the hallmark of this condition, structural changes affecting the neocortex of the fronto-central, temporal and parietal regions (Bernhardt et al., 2009, 2010) as well as the axonal fibre bundles that link them (Focke et al., 2008; Concha et al., 2009), suggest
widespread abnormalities of brain organization. Indeed, using graph theoretical analysis to quantify topological and organizational properties of complex systems, we recently showed evidence for altered large-scale structural networks (Bernhardt et al., 2011).

MRI methods that measure interregional brain signalling have been advanced as powerful tools to probe organizational properties of functional networks (Biswal et al., 1995; McKeown et al., 1998; Beckmann et al., 2005). In particular, studies correlating signal fluctuations while healthy volunteers rest in the scanner have revealed a highly robust set of resting state networks (Greicius et al., 2003; Beckmann et al., 2005) that are believed to reflect synchronous firing of discrete neuronal populations (Fox and Raichle, 2007). In TLE, resting functional MRI studies have mainly focused on quantifying signalling between regions known to be involved in seizure activity, particularly among medial temporal lobe structures (Bettus et al., 2009, 2010; Pereira et al., 2010), and those that form the ‘default mode’ state (Frings et al., 2009; Liao et al., 2010, 2011; Zhang et al., 2010). The majority of these studies used region of interest analytical approaches that preclude a comprehensive evaluation of large-scale functional interactions (Cole et al., 2010).

Notable correspondence between resting functional and structural networks has been shown in healthy subjects (Honey et al., 2007, 2009; Vincent et al., 2007; Skudlarski et al., 2008). Abnormal (typically reduced) resting signalling from the hippocampus in patients with Alzheimer’s disease and schizophrenia (Wang et al., 2006; Zhou et al., 2008), in whom hippocampal atrophy is prevalent (Apostolova et al., 2006; Velakoulis et al., 2006), raises the possibility that abnormal mesiotemporal functional connectivity in TLE may at least partly reflect structural pathology, rather than epileptogenic or functionally compensatory processes to which it has been attributed (Bettus et al., 2009, 2010; Morgan et al., 2010). The potential effect of structural damage on resting signal coupling, however, remains largely unknown.

Our aim was to determine the extent to which resting connectivity in patients with drug-resistant TLE is associated with co-localized structural pathology. In line with the distributed nature of structural damage in this condition, we used independent component analysis that optimizes sensitivity to wide-scale resting signalling, correcting for voxel-wise grey matter density. In addition, we assessed the relationship between resting signalling and white matter microstructure using tract-based spatial statistics.

### Materials and methods

#### Participants

We studied 35 consecutive right-handed patients with drug-resistant TLE being evaluated for epilepsy surgery at the Montreal Neurological Institute (MNI); 16 males, mean age = 34.3 ± 8.9 years, range = 19–53. The diagnosis and lateralization of the seizure focus into left TLE (n = 19) and right TLE (n = 16) were determined by a comprehensive evaluation including detailed history, video-EEG telemetry and neuroimaging in all. None of the patients had a mass lesion (tumour, vascular malformation or malformations of cortical development) or traumatic brain injury. Twenty patients underwent surgery. We determined surgical outcome according to the Engel classification scheme (Engel, 1993) at a mean follow-up time of 18 ± 5.6 months. Fourteen patients had an outcome Class I (seven Class Ia, six Class Ib, one Class IId), three Class II, two Class III and one of Class IV. Following qualitative histopathological analysis, hippocampal sclerosis was detected in all 15 patients in whom specimens were available. In the remaining five, hippocampal specimens were unsuitable for histopathology due to subpial aspiration.

The control group consisted of 20 healthy volunteers (11 males, mean age = 29.2 ± 6.7 years, range = 20–48 years). As controls were slightly younger than patients (left TLE, t = −2.231, P = 0.03; right TLE, t = −1.757, P = 0.09), age was used as a nuisance variable in all analyses. Demographic and clinical data are presented in Table 1.

The Ethics Committee of the Montreal Neurological Institute and Hospital approved this study and written informed consent was obtained from all participants in accordance with the standards of the Declaration of Helsinki.

#### Magnetic resonance imaging acquisition

Magnetic resonance images were acquired on a 3 T Siemens Trio Trim scanner using a 32-channel phased-array head coil. Resting blood oxygen level-dependent data were acquired in each subject using an echo-planar imaging sequence (repetition time = 2020 ms, echo time = 30 ms, flip angle = 90°, field of view = 256 × 256 mm², slice thickness = 4 mm, 34 slices with no gap, voxel size 4 × 4 × 4 mm³, 5 min acquisition and 220 volumes). Participants were instructed to lie still with their eyes closed while remaining awake. To minimize signal loss and distortion affecting orbitofrontal and mesiotemporal regions, slices were angled at an oblique coronal orientation. The diffusion tensor imaging protocol consisted of a twice-refocused echo-planar imaging sequence (63 axial slices, voxel resolution 2 × 2 × 2 mm³, repetition time = 8.4 s, echo time = 90 ms, diffusion-sensitized images in 64 diffusion directions.

### Table 1 Demographic and clinical data

<table>
<thead>
<tr>
<th>Group</th>
<th>Males</th>
<th>Age (years)</th>
<th>Onset</th>
<th>Duration</th>
<th>FC</th>
<th>Surgery</th>
<th>Engel-I</th>
</tr>
</thead>
<tbody>
<tr>
<td>Controls (n = 20)</td>
<td>11</td>
<td>29.2 ± 6.7 (20–48)</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Left TLE (n = 19)</td>
<td>7</td>
<td>34.4 ± 8.0 (19–45)</td>
<td>20.3 ± 10.3 (4–44)</td>
<td>14.6 ± 10 (1–35)</td>
<td>3</td>
<td>8</td>
<td>5</td>
</tr>
<tr>
<td>Right TLE (n = 16)</td>
<td>8</td>
<td>34.1 ± 10.2 (20–53)</td>
<td>21.3 ± 13.6 (2–48)</td>
<td>11.6 ± 10.4 (1–35)</td>
<td>4</td>
<td>12</td>
<td>9</td>
</tr>
</tbody>
</table>

Age, age at seizure onset and duration of epilepsy are presented in mean ± SD (range) years. FC = febrile convulsions.
Grey matter density confounded analysis

To remove potential variations in resting functional MRI signal fluctuations related to grey matter volume, we repeated the resting group comparison using voxel-wise grey matter density as a covariate in the linear model.

Diffusion tract-based spatial statistics analysis

Functional connectivity abnormalities that remained unchanged after correcting for grey matter density were used as covariates of interest in voxel-wise analyses of white matter fractional anisotropy. To minimize potential interactions between grey matter density and white matter microstructure, potentially obscuring the interpretation of resting connectivity, this analysis was performed on the original, non-grey matter corrected clusters.

Following motion/eddy current correction, a diffusion tensor model was fitted at every voxel to derive fractional anisotropy maps, which were non-linearly registered to group-specific templates created from controls and patients. The resulting fractional anisotropy images were temporally concatenated into a single 4D file and averaged to create a mean ‘skeleton’, representing the centre of all white matter tracts for each group-pair (controls and left TLE/controls and right TLE) onto which subject-specific fractional anisotropy values were projected. Finally, using tract-based spatial statistics, we performed group comparisons through a 2D implementation of random permutation testing described above to identify associations between voxel-wise fractional anisotropy and resting connectivity values derived from each cluster showing group differences not accounted for by grey matter volume. In diffusion tensor imaging tractography approaches, white matter tracts are segmented by estimating the directional information of water diffusion of coherently oriented axons. However, fibre bundles may cross, split or merge within individual voxels, affecting fractional anisotropy measurements. To aid interpretation of findings in regions of crossing fibres, TBSS X software (Jbabdi et al., 2010) was used post hoc to visualize directional information of two different fibre populations modelled at the level of single voxels.

Results

Resting state connectivity: relation to grey matter density

Compared with healthy controls, patients with left TLE showed reduced functional connectivity between ipsilateral temporal (lateral and temporopolar) cortices and anterior default mode network (Network 7, \( P = 0.024 \); Fig. 1A), which persisted after correction for grey matter density. Patients with right TLE showed reduced functional connectivity relative to controls between the ipsilateral parahippocampal region and the medial default mode network (Network 1, \( P = 0.043 \); Fig. 1B), between the right hippocampus and left lateral temporal cortex and the posterior default mode network (Network 6, \( P = 0.009 \); Fig. 1C) and between the ipsilateral frontal (premotor and supplementary motor) cortices and the extended sensorimotor network (Network 4, \( P = 0.012 \); Fig. 1D). While removing the variability associated
with grey matter density abolished decreased signalling between the right hippocampus and posterior default mode network, the decreased functional connectivity between this network and the left lateral temporal cortex remained unchanged (Fig. 1C). On the other hand, decreases in functional connectivity between the medial default mode network and the parahippocampal region (Fig. 1B), and between the extended sensorimotor network and the frontal cortices (Fig. 1D) persisted after grey matter correction. Finally, in patients with right TLE, the medial default mode network (Network 1) showed increased functional connectivity with the left temporal pole ($P = 0.024$; Fig. 1E), which also persisted after grey matter correction.

Since permutation tests identify any region where the relationship between the functional MRI signal and the main time-course of a network differs between groups, it is possible that portions of clusters fall outside the boundary of a given network (Fig. 1A and D). We repeated the resting connectivity analyses using the spatial maps for each component to constrain statistical comparisons within network boundaries. This network-specific analysis identified the same clusters found through the whole-brain analysis. Understandably, changes were located within networks, as this approach cannot detect anomalies that fall outside network boundaries. Furthermore, as fewer multiple-comparison corrections are needed to compare with whole-brain analysis, the network-specific approach showed increased sensitivity to within-network group differences in the sensorimotor network in patients with left TLE and in the left fronto-parietal network in patients with right TLE (Supplementary Fig. 2).

**Relationship between altered functional connectivity and fibre pathway coherence**

In left TLE (Fig. 2), the decrease in resting connectivity seen between the left temporal cortices and the anterior default mode network (Network 7) was positively associated with fractional anisotropy in the ipsilateral temporal lobe white matter ($P = 0.02$). Specifically, reduced resting signalling between this region and anterior default mode network in patients correlated...
with reduced fractional anisotropy along the ipsilateral uncinate and thalamic radiation, the inferior and superior longitudinal fasciculi and the corpus callosum ($P = 0.022$). These associations were not seen in controls, resulting in a significant group difference (left TLE controls, $P = 0.018$).

In patients with right TLE (Fig. 2), two of the three resting connectivity findings that were not explained by variations in cortical volume also showed altered associations with white matter coherence compared with controls. While in the latter, functional connectivity between the right parahippocampal gyrus and the medial default mode network (Network 1) was positively associated with fractional anisotropy along the right inferior longitudinal fasciculus and the callosum ($P = 0.05$), this association was not present in patients with right TLE. Similarly, there was a trend for a stronger correlation in controls than in patients with right TLE between functional connectivity in the right precentral gyrus and the sensorimotor network and fractional anisotropy along the callosum ($P = 0.08$). No association was found between fractional anisotropy and increased resting connectivity found between the left temporopolar cortex and the medial default mode network.

Group comparisons of structure–function associations between right TLE and controls co-localized to regions with strong confidence of a single fibre orientation. On the other hand, the left TLE temporal lobe cluster localized to a region with a high probability of crossing fibres (Supplementary Figs 3 and 4).
Discussion

Resting functional MRI studies in TLE have typically shown reduced connectivity involving ipsilateral mesiotemporal structures (Bettus et al., 2009, 2010; Morgan et al., 2010; Pereira et al., 2010) and altered integration with widespread regions of the default mode network (Frings et al., 2009; Zhang et al., 2010, 2011). Given the overlap of these signalling abnormalities with regions implicated electrographically in seizure generation and spread (Chabardes et al., 2005), it is tempting to interpret these findings as indicative of the epileptogenic network. However, previous studies in TLE analysed functional connectivity in predefined regions of interest or single networks, and did not take cortical and white matter damage into account. We opted for an objective whole-brain quantification of resting connectivity through independent component analysis, a technique that decomposes the signal into spatiotemporal components. The benefit of this approach is flexibility in data modelling to identify potentially wide-ranging abnormalities without a priori hypotheses (Cole et al., 2010). This is of particular importance since in TLE pathological interactions involve distributed brain regions (Bernhardt et al., 2011). In addition, abnormal functional connectivity may extend beyond a given network and involve multiple regions, such as the recruitment of the right frontal lobe into the left-lateralized language network (Walters et al., 2006). Specifically, our voxel-wise dual regression approach quantifies the degree to which individual voxels contribute to each resting-state network and compares groups through permutation testing. Thus, clusters may be identified within or outside a given resting network, without bias to either. Moreover, this approach is less affected by subtle variability in the specific regions that cluster into a network across studies, or even during a single resting functional MRI experiment, known as the non-stationarity problem (Cole et al., 2010). For example, the hippocampus is identified in the default mode networks in some but not all studies (Greicius et al., 2004). Finally, the simultaneous analysis of all networks increases sensitivity in cases where individual voxels contribute to multiple networks. On the other hand, the network selection is based on visual inspection of independent spatial components. We have addressed this limitation by performing a cross-correlation with template resting state networks (Beckmann et al., 2005) and selecting only the 10 that were strongly correlated with the Beckmann et al. (2005) set.

We found resting anomalies in patients with left TLE restricted to the ipsilateral temporal lobe, whereas in right TLE, we observed additional changes in the ipsilateral sensorimotor cortices and contralateral temporal lobe. These regions overlap notably with neural systems supporting memory, language and motor functions. Behavioural explanations for resting findings can only be speculative at the moment, as definitive studies relating resting functional MRI, task functional MRI and neuropsychological performance in the same patients remain lacking. Impaired tactile performance in patients with TLE [see Grant (2005) for review] and may account for our findings of reduced functional integration of precentral regions in the sensorimotor network. Activity within and correlation between midline posterior cingulate and hippocampal structures that form core parts of the default mode network can be specifically manipulated through episodic memory tasks (Vincent et al., 2006) on which patients with TLE are often impaired (Bell et al., 2011). The spatial correspondence between brain regions forming these neurocognitive systems and those showing altered resting functional connectivity, therefore, offer the tentative hypothesis that reduced functional connectivity in these networks may reflect patient-specific behavioural organization.

Observed differences in the extent of resting connectivity abnormalities between left and right TLE’s are in agreement with a previous resting default mode network connectivity study using independent component analysis (Zhang et al., 2010), but are in contrast with others describing bilateral temporal changes in left TLE (Bettus et al., 2009; Pereira et al., 2010). Aside from possible differences related to patient selection criteria, divergences may also stem from the analytical approach (region of interest-based versus whole-brain), making a direct comparison between studies difficult. Nevertheless, our observation of greater functional disconnectivity in right TLE supports the possibility of hemisphere-specific vulnerability to injury potentially related to asymmetric brain development (Sun and Walsh, 2006). In this scenario, abnormalities would occur during a time window of left hemisphere susceptibility when right hemisphere homologues, due to a more advanced stage of development, are relatively spared. Asymmetric resting functional connectivity differences notably parallel our previous structural observations in a separate group of patients with TLE. When assessing morphometric markers of brain development, we found bilaterally increased sulco-gyral complexity in right TLE, whereas such changes were unilateral in left TLE (Voets et al., 2011).

In TLE, the most parsimonious explanation for widespread structural abnormalities is that, at the scale of neuroimaging, this condition is represented by an altered configuration of grey matter regions and their interconnecting white matter tracts, which is somehow reflected in abnormal functional connectivity. Our multimodal MRI study indeed revealed that structural variations influence resting connectivity in TLE. Notably, reduced functional connectivity between the epileptic hippocampus and the posterior default mode network in our patients with right TLE was explained by variability in grey matter volume, suggesting altered signal coupling may be an indirect marker of hippocampal damage. Outside of the mesiotemporal lobe, while grey matter variations seemed not to alter resting connectivity, complex interactions were found with white matter microstructure. However, we cannot confidently disentangle grey from white matter contributions, as fractional anisotropy may co-vary with grey matter volume (Douaud et al., 2007). While there was a direct relation between reduced functional connectivity and fractional anisotropy in left TLE, in right TLE associations normally seen in controls were lost. These findings parallel dichotomies in cortical thickness correlations were reported previously (Bernhardt et al., 2008). Reduced coherence of white matter bundles secondary to axonal degeneration could explain the loss of normally present structural-functional associations. On the other hand, abnormal myelination (Thom et al., 2000) or maturation (Thom et al., 2001) may alter...
the efficiency of functional communication, resulting in a pathological association not seen in controls.

In left TLE, the decrease in resting connectivity seen between the left temporal neocortex and the default mode network was positively associated with fractional anisotropy in the ipsilateral temporal lobe white matter. On the other hand, in our patients with right TLE, the left temporal neocortex showed both reduced and increased functional connectivity with default mode network findings that were not accounted for by our structural measures and possibly indicate co-existing pathological processes. Notably, our previous morphometric analyses in right TLE showed extensive thinning (Bernhardt et al., 2010), as well as bilaterally increased gyral complexity (Voets et al., 2011) in this region. While atrophy may be associated with decreased functional communication, it is conceivable that increased cortical complexity is reflective of aberrant, potentially increased fibre connectivity, which could in turn enhance functional interactions.

Importantly, even though it is commonly assumed that functional connectivity reflects structural connectivity, the exact relationship between structure and function might not be straightforward (Damoiseaux and Greicius, 2009). While diffusion tensor imaging supports dependence between resting functional and structural connectivity in specific regions in both the healthy and diseased brain (van den Heuvel et al., 2008; Greicius et al., 2009; Mars et al., 2011), functional connectivity has also been observed in regions where there is little or no structural connectivity (Lowe et al., 2008; Uddin et al., 2008). This probably indicates signal correlations mediated by indirect structural connections (i.e. via a third region) or polysynaptic pathways that do not have a structural correlate (Honey et al., 2009; Lu et al., 2011). Finally, intensity abnormalities in the temporo-polar region may blur the grey–white matter transition (Ryvlin et al., 2002; Sankar et al., 2008), making it difficult to segment confidently tissue compartments and obscuring the interpretation of the underlying pathophysiological substrate.

Overall our results suggest that morphological impacts function in TLE, although a contribution of the epileptic activity cannot be entirely excluded. Structural damage may explain previously reported discrepancy between increased interictal EEG synchronicity and decreased mesiotemporal functional MRI signalling (Bettus et al., 2009).

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Supplementary material

Supplementary material is available at Brain online.

References


Fox MD, Raichle ME. Spontaneous fluctuations in brain activity observed with functional magnetic resonance imaging. Nat Rev Neurosci 2007; 8: 700–11.
Functional networks in TLE

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