Remote Effects of Hippocampal Sclerosis on Effective Connectivity during Working Memory Encoding: A Case of Connectional Diaschisis?

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Accumulating evidence suggests a role for the medial temporal lobe (MTL) in working memory (WM). However, little is known concerning its functional interactions with other cortical regions in the distributed neural network subserving WM. To reveal these, we availed of subjects with MTL damage and characterized changes in effective connectivity while subjects engaged in WM task. Specifically, we compared dynamic causal models, extracted from magnetoencephalographic recordings during verbal WM encoding, in temporal lobe epilepsy patients (with left hippocampal sclerosis) and controls. Bayesian model comparison indicated that the best model (across subjects) evidenced bilateral, forward, and backward connections, coupling inferior temporal cortex (ITC), inferior frontal cortex (IFC), and MTL. MTL damage weakened backward connections from left MTL to left ITC, a decrease accompanied by strengthening of (bidirectional) connections between IFC and MTL in the contralateral hemisphere. These findings provide novel evidence concerning functional interactions between nodes of this fundamental cognitive network and sheds light on how these interactions are modified as a result of focal damage to MTL. The findings highlight that a reduced (top-down) influence of the MTL on ipsilateral language regions is accompanied by enhanced reciprocal coupling in the undamaged hemisphere providing a first demonstration of "connectional diaschisis."

Keywords: dynamic causal modeling, effective connectivity, magnetoencephalography, temporal lobe epilepsy, working memory

Introduction

Extensive evidence indicates that medial temporal lobe (MTL) is not exclusively involved in long-term memory (LTM). Human neuroimaging studies have reported activation of MTL during working memory (WM) tasks that engage informational encoding (Campo et al. 2005; Karlgodt et al. 2005; Mainy et al. 2007), maintenance of information (Ranganath and D’Esposito 2001; Axmacher et al. 2007), and retrieval (Cabeza et al. 2002; Schon et al. 2009). Also supporting this view, neuropsychological and neuroimaging studies have revealed impaired performance and abnormalities in MTL activity during WM tasks in patients with MTL damage with various causes (Owen et al. 1996; Krauss et al. 1997; Abrahams et al. 1999; Grady et al. 2001; Lancelot et al. 2003; Lee et al. 2006; Olson et al. 2006; Piekmala et al. 2007; Ezzyat and Olson 2008; Wagner et al. 2009). However, based on the assumption that cognitive processes engage distributed neural networks, if we want to gain a clearer understanding of the functional role of MTL in WM it cannot be considered as an independent processor. It is, therefore, necessary to characterize that role from the perspective of the functional systems (Bullmore and Sporns 2009). Accordingly, the goal of the current study was to investigate the conjoint function of MTL and other functionally related brain regions involved in verbal WM as a large-scale network (Bressler and Menon 2010). We obtained whole-head magnetoencephalographic (MEG) recordings during a verbal WM task, which was designed to ensure that participants encoded words semantically (Campo et al. 2005, 2009), as prior neuroimaging investigations have demonstrated that depth processing modulates MTL activity (Kapur et al. 1994; Lepage et al. 2000).

Although few previous studies have used connectivity analyses to investigate the interactions between MTL and other key structures in the neural network involved in visual and verbal WM (Petersson et al. 2006; Nee and Jonides 2008; Rissman et al. 2008), our study diverges from those in 2 main aspects. First, we studied temporal lobe epilepsy (TLE) patients with left hippocampal sclerosis (HS) (Trenerry et al. 1993; Thom et al. 2005) in order to evaluate the impact of unilateral MTL pathology on functional organization and connectivity among brain regions engaged in verbal WM encoding. This is considered as a useful approach that allows the characterization of changes in the functional organization of interconnected brain regions following focal brain damage (Guye et al. 2008). Second, we used dynamic causal modeling (DCM) (Friston et al. 2003; David et al. 2006; Daunizeau et al. forthcoming) to characterize the effective connectivity in the WM network, in subjects with and without MTL damage (Seghier et al. 2010). Effective connectivity denotes "directed or causal relationships between elements" (Bullmore and Sporns 2009) and in the present context refers to the change that the activity in one brain region causes in the activity of another, and how this is modulated by experimental factors (Stephan and Friston 2007). Effective connectivity can be estimated with Bayesian model inversion by perturbing the system and measuring its response (Friston and Price 2001; Garrido, Kilner, Kiebel, and Friston 2007)—this is DCM. DCM represents a fundamental variation from alternative methods to estimate connectivity because it employs a generative model of measured brain responses that takes into account their nonlinear and dynamic nature. As opposed to functional connectivity measures that explore nondirectional statistical dependencies between brain regions,
DCM explicitly estimates the causal influence of one area over another.

On the basis of previous findings showing changes in functional connectivity (correlations) during declarative memory in TLE patients (Addis et al. 2007; Wagner et al. 2007; Bettus et al. 2009; Frings et al. 2009; Voets et al. 2009), we hypothesized that TLE patients with left HS would show decreased connectivity between left MTL and ipsilateral brain regions (prefrontal and temporal cortices) and an increased connectivity in contralateral homologous structures (Bettus et al. 2009). More specifically, we expected changes in the connectivity of MTL with inferior temporal language cortex and IPC/ventrolateral prefrontal cortex (VLPFC) (Fiebach et al. 2006; Nee and Jonides 2008; Rissman et al. 2008; Ojemann et al. 2009; Saling 2009; Hashimoto et al. 2010). Neurons in ITC respond selectively to task relevant features of stimuli in visual WM (Fuster 1990), are active during verbal WM when semantic processing is required (Fiebach et al. 2006, 2007), and have been shown to be affected in patients with semantic WM deficits (Hoffman et al. 2009). Interestingly, an interaction of ITC with rhinal cortex is considered to be part of a semantic associative memory subsystem (Salig et al. 1993; Salig 2009). Furthermore, previous studies have highlighted the relevance of hippocampus--ITC connectivity in strengthening mnemonic traces during visual WM (Axmacher et al. 2008; Rissman et al. 2008), while VLPFC-MTL interactions have been associated with semantic memory processing during WM in a prior study (Nee and Jonides 2008) and have been proposed to be fundamental for memory formation (Ranganath et al. 2005).

Materials and Methods

Participants

Eleven patients (6 males) with refractory MTLE epilepsy were consecutively recruited following presurgical evaluation at the "Hospital Ruber Internacional" and participated in the study. They ranged in age from 24 to 43 years (mean = 32.91; standard deviation [SD] = 6.89). Diagnosis was established according to clinical EEG and magnetic resonance imaging (MRI) data. All patients underwent neurological examination, continuous video-EEG monitoring, and high-resolution 1.5-T brain MRI. Patients were included in the study when clinical data and MRI and EEG findings were suggestive of unilateral mesial TLE related to left HS. All patients had: 1) seizures with typical temporal lobe semiology that were not controlled with antiepileptic drugs (AEDs) and 2) moderate to severe decreased volume (and abnormally increased T2 and FLAIR signal) of the left hippocampus on brain MRI. No lesions were observed in other structures beyond left MTL. Bedside video-EEG monitoring showed interictal epileptiform activity ipsilateral to the side of HS and in 5 cases complex partial seizures with an ictal onset in left anterior temporal electrodes. No seizure occurred within 24 h prior to the experiment. At the time of study, patients were on monotherapy or multitherapy. There was no significant difference between groups in terms of age (t20 = 0.82, P > 0.20).

Demographic and clinical information about patients and controls is provided in Table 1.

Table 1

<table>
<thead>
<tr>
<th></th>
<th>TLE (n = 11)</th>
<th>Controls (n = 11)</th>
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<tbody>
<tr>
<td>Age</td>
<td>32.91 (6.89)</td>
<td>31.09 (2.63)</td>
</tr>
<tr>
<td>Years of education</td>
<td>15.18 (2.40)</td>
<td>16.91 (1.04)</td>
</tr>
<tr>
<td>Duration of epilepsy (years)</td>
<td>18.23 (11.70)</td>
<td>14.88 (10.76)</td>
</tr>
<tr>
<td>Age at epilepsy onset (years)</td>
<td>14.88 (10.76)</td>
<td>2.54 (0.93)</td>
</tr>
<tr>
<td>Seizure frequency (per month)</td>
<td>1.73 (0.47)</td>
<td>1.73 (0.47)</td>
</tr>
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procedures of the study in accordance with the Declaration of Helsinki (1991).

Stimuli and Tasks

We used the same verbal WM task as in our previous studies (Campo et al. 2005, 2009). In each trial, subjects first saw a stimulus array comprising 4 words, located centrally in the display. The to-be-remembered array remained on the screen for 3000 ms. After a 2500 ms delay interval, participants were presented with 3 consecutively presented comprising a semantic category name for 500 ms. They were required to make a push-button response to indicate whether any of the words belonged to the semantic category represented by one of probe words. Thus, correct performance required subjects to maintain the target words in memory and make a deep categorization; ensuring a deep processing of probe words. There was an interval between probes of 500–700 ms. Match and no-match trials occurred with equal probability.

Concrete words were used, 4–7 letters in length (5.62 ± 1.57) and of moderate frequency (Algarabel 1996). A total of 120 trials were presented. The stimuli were projected through a LCD video projector (SONY VPL-X600E), situated outside the shielded room, onto a series of in-room mirrors, the last of which was suspended approximately 50 cm above the subject’s face and subtended a visual angle of 1–3° horizontally and 0.5° vertically.

Data Acquisition and Analysis

All MEG recordings were obtained using a whole-head neuromagnetometer comprising an array of 148 magnetometers (4-D 2500, San Diego) housed in a magnetically shielded room. Neuromagnetic signals were digitized continuously at 678 Hz and were band-pass filtered between 0.1 and 100 Hz. MEG data were submitted to an interactive noise reduction procedure to reduce environmental noise (4-D 2500, San Diego). Data were analyzed using SPM8 (Wellcome Trust Centre for Neuroimaging, London; http://www.fil.ion.ucl.ac.uk/spm/). The continuous time series for each participant was subjected to a Butterworth band-pass filter at 3–30 Hz. We analyzed epoched encoding period activity for each trial, for each participant. Trials including eye blinks or other myogenic or mechanical artifacts were removed using the thresholding artifact rejection algorithm implemented in SPM8 (trials containing signal strength exceeding 3000 FT were excluded). After artifact rejection, epochs were baseline corrected from −100 to 0 ms and then averaged.

Source Localization

Multiple sparse priors (as implemented in SPM8) were used to estimate the cortical origin of the neuronal response during the encoding period (Friston et al. 2008). This model specifies 512 sparse patches of activation and then iteratively reduces them until an optimal number and location of active patches are found using a (variational) Bayesian scheme. The hyperparameters of these multiple sparse priors are optimized using a greedy search. A tessellated cortical mesh template surface in canonical (Montreal Neurological Institute [MNI]) served as a brain model to estimate the current source distribution (Mattout et al. 2007). This dipole mesh was used to calculate the forward solution using a spherical head model. The inverse solution was calculated over a time window from 0 to 1000 ms during the encoding epoch. These reconstructions were analyzed using a general linear model (Kilner and Friston 2010), as described in Furl et al. (2010).
Effective Connectivity Analysis: DCM
Determining effective connectivity requires a causal model of the interactions among the constituents of the neural network subject to study (Stephan and Friston 2007). DCM considers the brain as "a deterministic nonlinear dynamical system that is subject to inputs and produces outputs" (David 2007).

DCM is a hypothesis-driven method that relies on the specification of a plausible biophysical and physiological model of interacting brain regions (Stephan and Friston 2007). The model is specified by its regions connections and by whether these connections are unidirectional (forward or backward) or bidirectional (both forward and backward). Forward and backward connections are defined according to the connectivity rules outlined in Felleman and Van Essen (1991) and specified in DCM to convey bottom-up and top-down effects, respectively. This model is then supplemented with a forward model of how neuronal or synaptic activity is transformed into a measured response (Kiebel et al. 2006). This enables the parameters of the neuronal model (i.e., effective connectivity) and spatial model (i.e., dipole orientations) to be estimated from observed data using a Bayesian scheme. Estimating the parameters of a DCM model relies on estimating the hidden states and parameters of the modeled system, which corresponds to the sources that comprise the model (David et al. 2006). DCM for MEG uses a neural mass model to explain source activity (David and Friston 2003) and has been described in detail elsewhere (David et al. 2006).

DCM Specification: Hypotheses Tested
Network architecture was specified on the basis of the inverse solutions (source localizations; see Fig. 1) for single subjects using multiple sparse priors (Friston et al. 2008) and was constrained by recent studies of functional connectivity on verbal WM (Fiebach et al. 2006; Nee and Jonides 2008). Accordingly, we considered for our models 6 regions that corresponded to ITC, MTL, and VLPFC/IFC bilaterally. These sources were modeled as equivalent current dipoles, whose prior mean locations coordinates (x; y; z) are: bilateral ITC: -13, -54, -15 (left); 13, -54, -15 (right); bilateral MTL: -27, -15, -20 (left); 27, -15, -20 (right); and bilateral IFC/VLPFC: -54, 35, 6 (left); 54, 35, 6 (right). Twelve models were specified and inverted separately for each subject (Fig. 2b). In all models, left and right ITC were chosen as visual input nodes for semantic processing of words (Bitan et al. 2005; Heim et al. 2009). The models were specified starting with simple architectures and adding hierarchical levels (i.e., sources and extrinsic connections). The simplest models only included the ITC and IFC/VLPFC sources, while more complex models included MTL sources. The sources were left unilateral, right unilateral, or bilateral. Models also differed in terms of their connections; forward only or both forward and backward. Accordingly, model 1F+ included left unilateral and forward connections, while model bFF+ included bilateral sources with forward and backward connections. Models with MTL sources were created by simply adding MTL sources; that is, model bFF included left unilateral and forward connections and the MTL. See Figure 2b for details. The MTL models allowed an evaluation of the involvement of MTL in verbal WM and the functional relevance of the connections of this region within the network.

Model Comparison
One of the advantages of DCM is that it can be used to compare competing hypotheses about functional architectures (David 2007; Garrido, Kilner, Kiebel, Stephan, et al. 2007; Friston 2009; Garrido, Kilner, Kiebel, Stephan, et al. 2009). This is accomplished by specifying a model (hypothesis), in terms of anatomical connections between brain regions. Using Bayesian model selection, DCM tests a group of competing models and provides evidence in favor of one model, relative to others (Penny et al. 2004). The model log-evidence or the marginal log-likelihood of each model is compared against the remaining models. The model with the highest evidence (i.e., the model with the best balance of accuracy and complexity) is then considered the best or optimal model. A difference of 3 or more in favor of one model as compared with others is required (Penny et al. 2004). We performed a fixed-effect analysis for comparing model log-evidence at the group level (i.e., patient group and control group), which is accomplished by summing the log-evidence of each participant for each model, finding the highest valued model and comparing it with the summed log-evidence of the next highest model (Garrido, Kilner, Kiebel, and Friston 2007; Garrido, Kilner, Kiebel, Stephan, et al. 2009; Stephan and Friston 2007; Heim et al. 2009).

Figure 1. Source localization for a representative subject using multiple sparse priors (upper panel). Sources of activity, modeled as dipoles (estimated posterior moments and locations) superimposed in an MRI of a standard brain in MNI space (lower panel).
Reyt et al. forthcoming). We also performed a random-effect analysis for comparing model evidence, an approach that admits different models for different subjects and that is relevant when investigating "cognitive tasks that can be performed with different strategies" (Stephan et al. 2009; Penny et al. 2010; Reyt et al. forthcoming). The 12 models were also compared at a single subject level (Penny et al. 2004; Garrido, Kilner, Kiebel, and Friston 2009). After selecting the optimal model, its subject-specific parameters (restricted to posterior probabilities of 90% or more) were analyzed using paired *t*-tests, to test for group differences in the usual way (Noppeney et al. 2006; Werner and Noppeney 2010). Following previous studies (Mechelli et al. 2007; Benetti et al. 2009), we controlled for Type-I error derived from multiple comparisons using a statistical threshold of *P* < 0.025.

### Results

#### Behavioral Performance

We assessed performance in the verbal WM task was in terms of correct hits for each stimulus set. We observed a mean accuracy level of 75.55% (SD = 8.79) in the control group and mean accuracy of 58.93% (SD = 12.53) in the patient group. Control subjects performed significantly better than patients in terms of accuracy (*t*20 = 3.60; *P* < 0.001). No significant differences were found for reaction time (RT) measure between groups (*t*20 = 0.76, *P* > 0.20). Average RTs were

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**Figure 2.** (A) Significant differences between groups in left MTL derived from conventional SPM8 analysis rendered on an averaged normalized brain. (B) Outline of the 12 DCM models for the effective connectivity analysis shown on axial brain schematics (see text for coordinates of all regions). The brain regions comprising the network architecture for each model are represented by circles. Arrows between the regions indicate the directionality of the connections (i.e., forward or forward and backward). IFG, inferior frontal gyrus; ITC, inferior temporal cortex; MTL, medial temporal lobe.
776.73 for patients (SD = 83.37) and 753.36 for controls (SD = 57.86).

**Group Differences in Spatiotemporal Activation**

Differences between groups during verbal WM encoding were associated with greater left MTL activation in the control group between 200 and 800 ms ($t_{20} = 3.17$, $P < 0.003$, uncorrected). Further analyses were conducted on more specific time windows of 200 ms duration. These analyses revealed a greater activation in left MTL for the control group between 200 and 400 ms ($t_{20} = 3.59; P < 0.001$) (Fig. 2a), between 400 and 600 ms ($t_{20} = 3.28; P < 0.002$, uncorrected), and between 600 and 800 ms ($t_{20} = 2.89; P < 0.005$, uncorrected). We detected no differential activity for the reverse contrast (patients > controls).

**Bayesian Model Selection**

To determine changes that left MTL damage can produce in the functional organization of interconnected brain regions during WM encoding, we evaluated the model evidence for 12 (DCM) models described in the Materials and Methods section (see Fig. 2b). This established the best functional architecture over all subjects, which we then used to test for group differences in connection strengths. A fixed-effects analysis (Garrido, Kilner, Kiebel, Stephan, et al. 2007; Garrido, Kilner, Kiebel, Stephan, et al. 2009) revealed that model bFB+ (i.e., bilateral forward and backward connections) supervened (Bayes factor relative to the second best model [model lFB+] = 452.07). These results constitute “very strong” evidence in favor of model bFB+ (Penny et al. 2004). A random-effect analysis (allowing for random effects on models) yielded similar results (exceedance probability for model bFB+ = 0.965). A representation of the fixed effect and random effect approaches is shown in Figure 3a. Model comparison was also performed for each participant individually. This confirmed that, for the majority of the patients (8 of 11) and controls (6 of 11), model bFB+ was superior to all other models (see Table 2).

Once the best model had been determined, group differences in effective connectivity were assessed using subject-specific (maximum a posteriori) parameter estimates (Fig. 3c). A two-sample $t$-test revealed that the extrinsic backward connection from left MTL to left ITC was stronger in controls (mean = 1.40; SD = 0.64) than patients (mean = 0.74; SD = 0.33) ($t_{20} = 2.99; P < 0.01$). In the right hemisphere, contralateral to the lesion, backward connection between VLPFC/IFC and MTL was greater for patients (mean = 0.97; SD = 0.34) as compared with controls (mean = 0.51; SD = 0.17) ($t_{20} = 3.98, P < 0.001$). A significant greater forward connection from right MTL to right VLPFC/IFC in favor of patients (mean = 1.31; SD = 0.44) (for controls, mean = 0.72; SD = 0.37) was also observed ($t_{20} = 3.36, P < 0.005$). We failed to detect differences in the remaining connections ($P > 0.15$).

To assess the functional significance of group differences in effective connectivity, connectivity strengths were correlated with task performance. A linear regression analysis showed that the backward connections from right VLPFC/IFC to right MTL were inversely related to task performance ($R^2 = 0.649, P < 0.002$). We also observed a trend for a positive correlation between backward connections from left IFG to left MTL and task performance in the control group ($R^2 = 0.519, P = 0.10$). When an outlier (a control subject with very high performance but low connectivity strength) was eliminated from this analysis, this correlation reached significance ($R^2 = 0.641, P < 0.05$).

A factor that may have influenced these results is the difference in task performance between epilepsy patients and controls. We used a median-split approach to identify a group of patients and controls that were matched on performance because neural activity from patients cannot be unambiguously interpreted unless the 2 groups are matched on this variable (Brown and Eyler 2006). Consequently, a subgroup of patients ($n = 7$) and a subgroup of controls ($n = 7$) with similar task performance were selected. These groups did not significantly differ in terms of age ($t_{12} = 1.12; P > 0.20$), level of education ($t_{12} = 0.32; P > 0.50$), nor in task performance ($t_{12} = 1.04; P > 0.30$). Bayesian model comparison showed that model bFB+ (i.e., bilateral forward and backward connections) supervened (fixed effects Bayes factor relative to the second best model [model lFB+] = 391.33). These results constitute very strong evidence in favor of model bFB+ (Penny et al. 2004). Furthermore, we found the same differences in connectivity. That is, extrinsic backward connections from left MTL to left ITC were reduced in patients (mean = 0.81; SD = 0.38) as compared with controls (mean = 1.50; SD = 0.65) ($t_{12} = 2.37, P < 0.02$). In the nonlesional hemisphere, a significantly greater forward connection from right MTL to right VLPFC/IFC was seen in patients (mean = 1.34; SD = 0.42) (vs. controls mean = 0.71; SD = 0.39; $t_{12} = 2.85, P < 0.01$). Backward connections between VLPFC/IFC and MTL were greater in patients (mean = 0.80; SD = 0.28) than in controls (mean = 0.57; SD = 0.13; $t_{12} = 2.05, P = 0.031$).

**Discussion**

The main focus of the current study was the functional organization expressed in terms of effective connectivity among MTL and other functionally related brain regions subserving verbal WM encoding. This is the first study of effective connectivity in relation to the impact of MTL damage on mnemonic function and we note that all previous work has examined functional connectivity that may or may not be mediated by directed neuronal connections (i.e., effective connectivity). This is because functional connectivity simply establishes a statically dependency between sources and does not address how these dependencies are mediated. Studying effective connectivity allows us to understand the effect of MTL damage precisely since we can examine the causal influences in the network and ask what connections to or from the MTL are affected. The findings of this study corroborate a framework that the MTL is a part of an extended neural network engaged in verbal WM. Moreover, the data provide new evidence about its functional interactions during a WM task and sheds light on how these interactions are modified as a result of localized damage to MTL. Interestingly, we found a bilateral network model with forward and backward connections including MTL, ITC, and IFC/VLPFC (see Materials and Methods section) was the best model across participants. Note that because the prior source locations were based on source reconstructions of the channel data, they are optimized for the particular subjects we studied. The bilateral nature of the model is in agreement with functional imaging studies showing that left and right MTL contribute to verbal memory processes, especially with semantic encoding (Leppage et al. 2000; Davachi and Wagner 2002). However, a left MTL
Figure 3. (A) Group level Bayesian selection of the 12 tested models. Left: fixed effect analysis (FFX) showing log-evidence and model posterior probability. Right: random fixed effects (RFX) showing model expected probability and model exceedance probability. Results indicate the best model is one with bilateral forward and backward connections comprising IFG, ITC, and MTL. (Bayes factor relative to the second best model [model lFB+] = 452.07; exceedance probability for model bFB+ = 0.965). 1. LF/C0; 2. RF/C0; 3. B/F/C0; 4. LFB/C0; 5. RFB/C0; 6. BFB/C0; 7. LF+; 8. RF+; 9. BF+; 10. LFB+; 11. RFB+; 12. BFB+. L, left; R, right; B, bilateral; F, forward; FB, forward and backward; + model architecture not including MTL; + model architecture including MTL. (B) Predicted (blue) and observed (red) responses in measurement space for the best model. (C) Group differences in effective connectivity assessed using subject-specific (maximum a posteriori) parameter estimates.
In fact, important reductions of functional connectivity of the function and use-dependent changes in gray and/or white interaction (Squire 1991; Saling 2009). Hence, reduced reduced in TLE, which has been proposed to regulate this damage to rhinal or parahippocampal cortex, commonly (Yogarajah et al. 2008; Voets et al. 2009). Alternatively, white matter connections due to MTL pathology in TLE between these regions could be mediated by alterations in subjects (Powell et al. 2004). Hence, changes in coupling demonstrated in vivo using diffusion tensor imaging in healthy network regions, we found both an attenuation in effective connectivity between MTL regions and ITC has been demon-

### Table 2

<table>
<thead>
<tr>
<th>Model</th>
<th>Bayes Factor</th>
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<td>bFB^-rFB^+</td>
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Note: P, patient; C, control; l, left; r, right; b, bilateral; F, forward; FB, forward and backward; − model architecture not including MTL; + model architecture including MTL.

The IFC/VLPFC has been shown to manifest increased its connectivity with MTL and ITC during verbal and visual WM tasks (Grady et al. 2001; Simons and Spiers 2003; Fiebach et al. 2007; Nee and Jonides 2008). It has also been shown that the pattern of fronto--limbic interactions in the hemisphere ipsilateral to the lesioned MTL are impaired during memory tasks in patients with TLE (Addis et al. 2007; Voets et al. 2009). Surprisingly, our connectivity analyses failed to identify any differences in the pattern of connectivity of left IFC/VLPFC, either with left ITC or left MTL between groups. Despite this lack of significant differences, we observed a trend to a significant positive correlation between backward connec-
tions from left IFC/VLPFC to left MTL in the control group. A significant correlation emerged when we one control with very high performance but low connectivity value was excluded from the analysis. Increased connectivity between VLPFC and MTL during encoding of words has been previously found to correlate with better performance in healthy young subjects (Grady et al. 2003). It is possible that this interaction constitutes a common mechanism supporting encoding process both in WM and LTM (Fuster 1995), as has been recently suggested for retrieval processes (Öztekín et al. 2010).

In contrast, compared with controls, the coupling between IFC/VLPFC and the MTL in the contralesional hemisphere was enhanced in patients. This strengthening of connections was bidirectional. This change in the pattern of connectivity between these regions is interesting, considering that brain activation analyses did not reveal evidence for substantial differences in activation in the contralesional MTL between groups and reflect a dissociation between regional activation and connectivity measures (Grady et al. 2001; Ranganath et al. 2006, 2007; Nee and Jonides 2008). It has also been shown that the pattern of fronto--limbic interactions in the hemisphere ipsilateral to the lesioned MTL are impaired during memory tasks in patients with TLE (Addis et al. 2007; Voets et al. 2009). Surprisingly, our connectivity analyses failed to identify any differences in the pattern of connectivity of left IFC/VLPFC, either with left ITC or left MTL between groups. Despite this lack of significant differences, we observed a trend to a significant positive correlation between backward connec-
tions from left IFC/VLPFC to left MTL in the control group. A significant correlation emerged when we one control with very high performance but low connectivity value was excluded from the analysis. Increased connectivity between VLPFC and MTL during encoding of words has been previously found to correlate with better performance in healthy young subjects (Grady et al. 2003). It is possible that this interaction constitutes a common mechanism supporting encoding process both in WM and LTM (Fuster 1995), as has been recently suggested for retrieval processes (Öztekín et al. 2010).
In this sense, our findings provide evidence for a "connectional diaschisis." Diaschisis (from Greek, meaning "shocked throughout") usually refers to loss of neuronal activity, due to lost afferents from a lesion area. It has been generalized to cover "dynamic diaschisis" (Price et al. 2001), which refers to a selective changes in neuronal responses, due to lost afferents. We suggest that the phenomenon we report here reflects a connectional diaschisis—a selective change in coupling due to lost afferents; in this case, from the contralateral (lesioned) nodes of the network.

Enhanced recruitment of right prefrontal cortex has been previously reported in left TLE patients while processing verbal material (Maccotta et al. 2007) as well as increased basal functional connectivity involving contralesional MTL in patients with intractable epilepsy of MTL origin (Bettus et al. 2009). Interpretation of these effects as reflecting compensatory mechanisms is controversial (Maccotta et al. 2007; Vlooswijk et al. 2008; Saling 2009). Increased levels of activity or changes in connectivity dynamics in response to a pathological state can be interpreted in different ways, taking into account their relation with task execution (Maccotta et al. 2007). A linear regression analysis showed that the backward connections from right VLPFC/IFC to right MTL was inversely related to task performance (see Fig. 4). Crucially, the relationship between coupling and behavior is seen over both patients and controls, suggesting that the remote effects of lesions on connectivity are functionally (behaviorally) relevant and may reflect compensatory or adaptive changes that are similar to differences among normal subjects. More generally, this correlation lends the coupling estimates predictive validity, in relation to function or task performance. Interestingly, when we controlled for behavioral differences between patients and normal subjects (by examining a subset of subjects), there was still evidence for significant differences in coupling. This suggests that both compensatory and pathophysiological mechanisms underlie the increased connection strength in patients. In this sense, enhanced contralesional fronto–limbic coupling can be regarded "as a marker of network disruption in the presence of mesial temporal pathology" (Saling 2009), which is in agreement with previous work (Dupont et al. 2000; Maccotta et al. 2007; Powell et al. 2007; Vlooswijk et al. 2008). The pathological nature of this pattern of connectivity is reinforced by recent findings showing an increased functional connectivity between hippocampus and diffuse areas of prefrontal cortex which was negatively correlated with performance on a memory task in amnestic mild cognitive impairment patients showing hippocampal atrophy (Bai et al. 2009).

As both groups differed in task performance, correct inferences about connectivity measures require comparisons between a subgroup of patients and controls that were matched on performance. We found that the differences in connectivity measures observed at the group level were maintained at the subgroup level. In view of these results, differences in connectivity between MTLE patients and controls cannot be attributed completely to performance differences. This could suggest that group differences are mediated by the disruption of the network supporting WM encoding due to MTL damage (Mueller et al. 2011). That is, MTL lesion not only affects local neural processing but also interactions with other brain regions that constitute a network supporting a specific cognitive process. Nonetheless, further studies including patients with epileptogenic lesions in other locations (i.e., right MTLE, extratemporal epilepsy) will be needed to evaluate this interpretation.

The potential impact of AEDs on cognitive functioning, brain activation, and connectivity measures cannot be discounted as contributing to the differences reported here. AEDs have been reported to have both positive and negative effects on cognition, in patients, and in healthy controls (Prevost et al. 1996; Thompson et al. 2000; Aldenkamp et al. 2002; Meador et al. 2007; Seo et al. 2007; Park and Kwon 2008) and vary in the type and degree of their associated side effects, depending upon several factors such as the type and dosage of AED used (Meador 2006; Schilbach et al. 2007; Baxendale et al. 2010; Canevini et al. 2010; Hermann et al. 2010). Additionally, it is difficult to dissociate AEDs effects in epileptic patients from the effect of epilepsy itself and associated psychosocial variables (Gualtieri and Johnson 2006; Bocquillon et al. 2009). Although, we can discount an effect of AEDs that is mediated through performance differences (see above), the role of AEDs on brain activation differences cannot be excluded. The effects of AEDs in brain activation are difficult to disentangle since previous studies have shown a decrease in electrophysiological measures of amplitude (Tuunainen et al. 1995) and power (Zaveri et al. 2010) or hemodynamic signals (Chen et al. 2009) either during

![Figure 4](http://cercor.oxfordjournals.org/)

**Figure 4.** Correlation of task performance and effective connectivity measures between right VLPFC/IFC and right MTL in patients and controls.
AED discontinuation or associated with AEDs (Sun et al. 2007). However, it is important to emphasize that activation differences between patients and controls in the current study were restricted to the lesional temporal lobe, a finding that matches with a previous study showing a decrease in power in signals recorded from epileptogenic mesial temporal lobe structures as compared with nonepileptogenic regions (Bettus et al. 2008). Finally, the impact of AEDs on connectivity measures must be also considered. Studies addressing the effects of AEDs on functional connectivity in epilepsy patients are scarce. Chen et al. (2009) have shown an attenuation of frontal-hippocampal connections after AED withdrawal. Similarly, Fingelkurts et al. (2004) observed a widespread increase in functional connectivity after administering Lorazepam to a group of healthy volunteers. Contrary to these findings, van Dellen et al. (2009) found a significant lower phase-lag index in epilepsy patients on multiple AED therapy, compared with those on monotherapy, although no effect of AEDs were found on network configuration measures. It is important to highlight that these results were restricted to the lesional temporal lobe and contradict those from a study showing an increase functional connectivity within the MTL when comparing epilepsy patients and healthy controls at rest (Liao et al. 2010). In relation to our findings, it is unlikely that AED effects could account for both an increase and a decrease in specific connections observed in the current study, especially those observed in the nonlesional hemisphere.

The current study has some limitations. One is the relatively small size of the patient group. Therefore, the findings should be considered preliminary and need to be replicated in further patient cohorts. The fact that we obtained significant results with such a small sample size suggests that the sizes of the effects reported above are large; however, our homogenous patient group was selected carefully, and our findings may or may not generalize to other groups. As we have mentioned before, another limitation is that only patients with left MTLE were included. Studies of patients with right MTLE, patients with temporal neocortical lesions and extratemporal epilepsy, should afford a more reliable test of our hypothesis.

In summary, our findings revealed that, 1) MTL is part of a network of functionally related regions subserving verbal WM encoding, 2) this network is best defined as a bilateral corticocortical network encompassing IFG/vLIFG, MTL, and ITC, with forward and backward connections, 3) changes caused by damage within left MTL in the aforementioned network are characterized by weakened connections from left MTL to left ITC and by the strengthening of forward/backward connections between IFG/vLIFG and MTL in the contralesional hemisphere, and 4) the pattern of connectivity identified in the patient group may not be an effective compensation for MTL damage but reflect a greater engagement of the remaining components of a damaged network subserving verbal WM (i.e., constituting a connectional diaschisis) and could be considered an indication that the network supporting a specific process (i.e., encoding) has been perturbed by pathophysiology (Mueller et al. 2011).

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Notes
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