Disrupted functional connectivity affects resting state based language lateralization

Alex Teghipco\textsuperscript{a}, Ali Hussain\textsuperscript{b}, Madalina E. Tivarus\textsuperscript{a,b,}\textsuperscript{*}

\textsuperscript{a}Rochester Center for Brain Imaging, University of Rochester, USA
\textsuperscript{b}Department of Imaging Sciences, University of Rochester, USA

**Abstract**

Pre-operative assessment of language localization and lateralization is critical to preserving brain function after lesion or epileptogenic tissue resection. Task fMRI (t-fMRI) has been extensively and reliably used to this end, but resting state fMRI (rs-fMRI) is emerging as an alternative pre-operative brain mapping method that is independent of a patient’s ability to comply with a task. We sought to evaluate if language lateralization obtained from rs-fMRI can replace standard assessment using t-fMRI. In a group of 43 patients scheduled for pre-operative fMRI brain mapping and 17 healthy controls, we found that existing methods of determining rs-fMRI lateralization by considering interhemispheric and intrahemispheric functional connectivity are inadequate compared to t-fMRI when applied to the language network. We determined that this was attributable to widespread but nuanced disturbances in the functional connectivity of the language network in patients. We found changes in interhemispheric and intrahemispheric functional connectivity that were dependent on lesion location, and particularly impacted patients with lesions in the left temporal lobe. We then tested whether a simpler measure of functional connectivity to the language network has a better relation to t-fMRI based language lateralization. Remarkably, we found that functional connectivity between the language network and the frontal pole, and superior frontal gyrus, as well as the supramarginal gyrus, significantly correlated to task based language lateralization indices in both patients and healthy controls. These findings are consistent with prior work with epilepsy patients, and provide a framework for evaluating language lateralization at rest.

© 2016 The Authors. Published by Elsevier Inc. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/).

1. Introduction

1.1. Resting state fMRI for pre-surgical brain mapping

Variability in the functional anatomy of the individual makes it essential to map the borders of the eloquent cortex prior to resection if functional tissue is to be preserved. Invasive subdural methods remain the gold standard of functional localization (De Witte and Mariën, 2013; Daffau et al., 2005; Miller et al., 2007), however neuroimaging presents a useful and crucially non-invasive approach that can be leveraged prior to surgery to help plan resection, shorten the surgery, and predict its outcome (e.g., De Witte and Mariën, 2013; Petrella et al., 2006; Rabin et al., 2004; Rutten et al., 2002a,c; Tharin and Golby, 2007). Successful use of functional magnetic resonance imaging (fMRI) for pre-surgical functional mapping is established (Buxhoeveden, 2004; Brodmann, 1909; Lehericy et al., 1999; Schiebinger et al., 2002; Voyvodic et al., 2009), but contingent on patient participation in potentially difficult tasks that they may not be able to perform, or that may distract from avoiding movement and thus compromise the quality of the data being collected. As a result, fMRI is often used in conjunction with intraoperative mapping procedures although they also rely on task participation (Buxhoeveden, 2004). Recent advances in the field of neuroimaging may circumvent this issue by isolating complex neural networks in the brain at rest.

With resting-state functional magnetic resonance imaging (rs-fMRI), low-frequency fluctuations in BOLD signal across the brain can be measured (although note resting state MEG and EEG methods exist as well; e.g., Stam et al., 2007), and used to isolate a multitude of neural networks from a single scan thereby providing surgeons with greater information at a lower resource cost (Shimony et al., 2009). Indeed, robust methods such as Independent Component Analysis (ICA), seed-based, and voxel-level functional connectivity have been used to extract sensorimotor, language, auditory, and visual networks (Baldassano et al., 2012; Biswal et al., 1995; Smith et al., 2009; Tomasi and Volkow, 2012a; Yeo et al., 2011), as well as default mode, executive, salience, and attentional networks (Fox and Raichle, 2006; Power et al., 2011; Raichle et al., 2001; Seeley et al., 2007). Particularly, recent studies have demonstrated the feasibility of extracting the language network from rs-fMRI in healthy controls using ICA, and have suggested that...
The degree of lateralization in the language network is a useful metric by which decisions may be made regarding the selection of patients for subdural localization procedures, especially for patients with epilepsy, a group that presents higher instances of atypical language laterality (Rasmussen and Milner, 1977). The Wada test has been largely used for this purpose, however t-fMRI has been gaining traction as an alternative and non-invasive method by which to determine lateralization. For example, t-fMRI can be used to predict language lateralization as determined by the Wada test (Adcock et al., 2003; Binder et al., 1996; Desmond et al., 1995; Lehericy et al., 2000; Ruten et al., 2002b; Woermann et al., 2003), but critically this is contingent on the soundness of the task design and methodology employed during the t-fMRI (e.g., Binder et al., 1999). Rs-fMRI has also been used for localization and lateralization in patients. On the basis of functional connectivity to the basal ganglia, the epileptogenic zone in patients with mesial temporal lobe epilepsy (MTLE) can be lateralized (Bettus et al., 2009). Rs-fMRI has also been successfully used to lateralize and localize sensorimotor areas pre-surgically (Kokkonen et al., 2009; Zhang et al., 2009), but remarkably little work has been done on the use of rs-fMRI for pre-surgical localization or lateralization of the language network despite the advantages of this method over t-fMRI. Recently, some research has indicated that unique functional connectivity patterns in temporal lobe epilepsy patients may be able to predict language network lateralization derived from task (Doucet et al., 2015). These authors found that resting functional connectivity between regions of the inferior frontal gyrus and the mesial temporal lobe correlated with the degree of task language lateralization in epilepsy patients, but not in healthy controls, for whom they found it was functional connectivity to the bilateral frontal cortex that correlated with task based language lateralization. This presents the exciting possibility that intrinsic functional connectivity may be used in place of t-fMRI for presurgical lateralization of the language network.

1.3. Altered functional connectivity in the language network of patients with brain disease

Brain damage has been linked to disorganized topology of a variety of networks. Patients with left and right mesial temporal lobe epilepsy have weaker functional connectivity from the temporal lobe to the medial frontal cortex that corresponds to memory impairment (Doucet et al., 2013a). Patients with gliomas have also shown a default mode network (DMN) deficit in hippocampal and prefrontal regions, as a function of lesion grade (Laufs et al., 2007). However, few studies have addressed language network topology as a function of lesion location using rs-fMRI, although there is mounting evidence that lesions may induce functional reorganization at rest (Bosma et al., 2008; Carpenti et al., 2001; Diessen et al., 2013; Douw et al., 2008; Doucet et al., 2013a; Duffau, 2005; Duffau et al., 2005; Goldmann and Golby, 2005; Haneef et al., 2014; Maccotta et al., 2013; Negishi et al., 2011; Partovi et al., 2012; Pittau et al., 2012; Voets et al., 2012; Waites et al., 2006; Woodward et al., 2014). Those that have addressed this reorganization using fMRI have supported the general pattern of results reported from MEG studies (Bartolomei et al., 2006a,b; Guggisberg et al., 2008). In one study Briganti et al. (2012) showed more diffuse disruptions in functional connectivity, and more widespread reorganization of functional connectivity associated with the language networks of patients with left hemisphere lesions. In another fMRI study, Pravata et al. (2011) evaluated functional connectivity in patients with intractable epilepsy, and found reorganization in the language network, and reduced functional connectivity in brain regions spanning the left hemisphere, regardless of epileptogenic focus side. Doucet et al. (2013a) also reported that left hemisphere damage in mesial temporal lobe epilepsy was more disruptive of functional connectivity than right hemisphere damage. More recently, Doucet et al. (2015) have reported that in temporal lobe epilepsy, strong left hemisphere language lateralization corresponds to high functional connectivity between the left inferior frontal cortex, and left frontal lobe areas. Kinno et al. (2015) have also looked at the effect of frontal gliomas on functional connectivity, primarily finding changes in connectivity within several syntax-related networks following the ventral and dorsal streams (see also Kinno et al., 2014). These findings from MEG and fMRI showing global effects on functional connectivity from relatively isolated damaged brain tissue are consistent with a rich literature of clinical findings showing transient deficits in regions unrelated and proximally distant from the area of damage, also known as diaschisis (Nomura et al., 2010; Von Monakov, 1969), and provide evidence that suboptimal neural network architecture affects complex brain function (Heimans and Reijneveld, 2012 for review).

2. Methods

2.1. Participants

Twenty healthy controls were recruited and gave informed consent to be scanned at the Rochester Center for Brain Imaging following the guidelines of the IRB at the University of Rochester. One participant was excluded for not being a native English speaker, one participant withdrew from the study due to anxiety, and one participant was excluded for taking psychoactive drugs. All remaining 17 healthy subjects (11 female; age range 18–33, mean age 21.2 years old, SD = 3) were right handed as determined by the Edinburgh Handedness Inventory (Oldfield, 1971). All healthy controls performed resting state and language t-fMRI.
Forty-four native English-speaking patients referred for pre-surgical mapping of language or motor function were enrolled in the study and gave written informed consent following the guidelines of the IRB at the University of Rochester. Motor mapping was not analyzed here as it does not relate to the topic of language lateralization, but was performed during the same session as part of these patients’ clinical care. One patient was excluded due to excessive head motion, leaving 43 patients (23 female; age range 18–74, mean age 51 years old, SD = 17.5) with diverse diagnoses in the group (6 mesial temporal sclerosis-MTS; 13 vascular lesions, including 11 cavernomas and 2 arteriovenous malformations; 18 gliomas including 9 low grade oligodendroglioma, oligoastrocytoma, astrocytoma, and 9 high grade oligodendroglioma, oligoastrocytoma, astrocytoma, glioblastoma multiforme; 1 meningioma; 1 arachnoid cyst; 1 lung cancer metastasis; 3 no apparent lesion). Thirty-six patients were right handed, 5 were left-handed and 2 were ambidextrous as determined by the Edinburgh Handedness Inventory (Oldfield, 1971). This reflects the heterogeneity of the typical patient population encountered in clinical practice. The patient group was scanned as part of their clinical workup, and as a result all patients participated in a resting state scan, but 31 patients underwent language mapping while the others participated in motor mapping (not analyzed here). Correspondingly, the analysis we performed to compare resting state and task language lateralization indices were done over this smaller subset of patients who have had both resting state and language mapping, while group comparisons of intrinsic functional connectivity involved the entire dataset of patients independent of task fMRI. Previous research has suggested that lesions significantly reorganize the topology of functional connectivity within the brain, including within the language network (Doucet et al., 2015; Briganti et al., 2012; Pravata et al., 2011). Thus, all group comparisons of functional connectivity were done both across the entire patient group (n = 43), and also several smaller patient groups based on lesion location. Twenty-nine patients had damaged tissue in the left hemisphere, 13 patients had damaged tissue in the right hemisphere, 17 patients had damaged tissue in the left frontal lobe, and 9 patients had damaged tissue in the left frontal lobe. Due to the low number of patients with right hemisphere frontal (n = 5) or temporal (n = 6) lesions, these smaller groups were not analyzed independently (see Table 1 for more details and Fig. 1 for visualization of lesion overlap across all 43 patients).

### 2.2. Language paradigms

The language tasks administered to both patients and healthy controls were presented visually and, included Definition Naming (DN), Word Generation in a Category (WGC), Word Generation from a Letter (WGL) and Verb Generation (VG). Healthy controls performed each of these tasks, but some patients could not complete all of the tasks. Of these tasks, but some patients could not complete all of the tasks. Of those patients undergoing language mapping, 29 performed WGC, 28 performed WGL and 31 performed both VG, and DN.

For the DN task, participants viewed short descriptions of an object (e.g., ‘jewelry you wear on your finger’) presented for 3 s and were instructed to covertly name the object (‘ring’). Blocks of 34 s sentence presentation were alternated with blocks of 20 of fixation cross for a total scan time of 4 min 8 s. During the VG task, participants viewed one concrete noun every 3 s and had to silently generate a semantically related verb (e.g., ‘car – drive’). Blocks of 30 s noun presentation alternated with control blocks (a string of the symbol ‘#’, matched to the approximate length of the nouns), for a total scan time of 3 min 30 s. During WGC task, participants viewed one word indicating a category and had to silently generate as many items in that category as they could in 30 s (e.g., ‘animals - dog, cat, cow, pig, lion, zebra, etc.’). Blocks of 30 s category presentation alternated with control blocks (a string of the symbol ‘#’, matched to the approximate length of the words), for a total scan time of 3 min 30 s. During WGL task, participants viewed one letter and had to silently generate as many words beginning with that letter as they could in 30 s (e.g., ‘A - animal, arch, arrive, army, alphabet, etc.’). Blocks of 30 s letter presentation alternated with 30 s fixation cross blocks for a total scan time of 3 min 30 s. Finally, during the 5 min long resting state scan, the patients were asked to lie in the scanner with their eyes closed.

### 2.3. MRI data parameters

Patient imaging data was collected on a 3T GE Discovery MR750 MRI scanner (General Electric, Milwaukee WI, USA) equipped with an 8-channel head coil. Due to better availability, volunteer imaging data was acquired on a 3T Siemens MAGNETOM Trio MRI scanner (Siemens Healthcare, Erlangen, Germany) equipped with a 32-channel head coil. Imaging parameters were similar on both scanners. Since the two groups were evaluated separately, the use of different scanners was not considered critical. High resolution structural T1 contrast images were acquired using either MPRAGE (Siemens) or Bravo FSPGR (GE) sequences (FOV = 256 mm, resolution 1 × 1 × 1 mm³). Functional images were acquired during each of the task paradigms described above using BOLD echo-planar imaging pulse sequence (TR = 2000/3000 ms, TE = 30 ms, resolution 4 × 4 × 4 mm³). The first four imaging volumes were acquired to allow stabilization of longitudinal magnetization, and were discarded before data analysis.

### 2.4. fMRI analysis

Functional language and anatomical images were analyzed with FSL package (Jenkinson et al., 2012; Smith et al., 2004; Woolrich et al., 2009). Data preprocessing consisted of motion correction (Jenkinson et al., 2002), slice-timing correction, non-brain signal removal (Smith, 2002), Gaussian spatial smoothing (FWHM 5 mm), intensity normalization, registration to the high resolution de-skulled anatomy and MNI space, and high-pass temporal filtering. The general linear model (GLM) was used to fit beta estimates to events of interest, and each voxel’s time series was whitened nonparametrically using FILM. Motion was included as a regressor of no interest. Lesion masks were drawn manually and checked for accuracy and extent by CAQ certified neuroradiologist and co-author, AH. Edemas were excluded from the lesion masks and either T1 or T2 images (with and without contrast) were

| Group                  | Number of patients | Female | Average age (± std dev) | Range of age | Left hemisphere pathology | Right hemisphere pathology | Number of right handed patients | Lesion volume in mm (± std dev) |
|------------------------|--------------------|--------|-------------------------|--------------|---------------------------|---------------------------|-------------------------------|---------------------------------|
| All patients           | 43                 | 23     | 51 (17.5)               | 18–74        | 29                        | 13                        | 36                            | 33,120 (37,659)                 |
| Temporal lobe lesions  | 23                 | 12     | 34 (12.1)               | 21–61        | 17                        | 6                         | 16                            | 26,402 (27,804)                 |
| Frontal lobe lesions   | 14                 | 9      | 40 (15.3)               | 20–74        | 9                         | 5                         | 14                            | 42,548 (49,075)                 |
| Parietal lesions       | 5                  | 2      | 39 (19.1)               | 18–71        | 3                         | 2                         | 2                            | 11,140 (12,229)                 |
| Other lesions          | 1                  | 0      | 61                      | –            | 1                         | 1                         | 1                            | 16,256 (–)                     |
| MTS or epilepsy        | 6                  | 2      | 42 (13.9)               | 25–63        | 5                         | 1                         | 6                            | (–)                             |
| Vascular lesions       | 6                  | 2      | 30 (8.1)                | 21–44        | 4                         | 2                         | 3                            | 10,453 (15,051)                 |
used to guide the drawings on a case-by-case basis, which was determined by the sequence used by clinicians in preparation for a particular surgery. The same sequence was used for outlining the borders of the lesion. These lesion masks were registered in standard space and added for patients as pre-thresholding masks during statistical analysis. First level Z statistic images were thresholded using clusters determined by Z > 2.3 and a corrected cluster significance threshold of p = 0.05.

Functional connectivity (FC) at rest was generated with the CONN toolbox (Whitfield-Gabrieli and Nieto-Castanon, 2012), relying on SPM (Penny et al., 2011) and MATLAB (MATLAB and Statistics Toolbox Release, 2014). Preprocessing of resting state volumes included realignment and normalization to MNI space. A component-based noise correction method was employed for noise reduction. This allowed us to avoid global signal regression, and thereby also avoid spurious negative correlations and spurious group level region to region interactions (e.g., Murphy et al., 2009; Saad et al., 2012). Segmentation of T1 images was performed to derive white matter, and cerebrospinal fluid masks. These masks were eroded to minimize partial volume effects, and regressed in the general linear model along with estimated subject movement (across 6 rotation/translation parameters and another 6 parameters representing their first-order temporal derivatives). After regression, a band-pass filter from 0.008 to 0.09 Hz was applied to the data. FC was calculated using a general linear model weighing the HRF and applying bivariate correlations to the BOLD signal at every voxel.

**2.5. Language laterality index analysis**

Comparison of language lateralization indices using both rs-fMRI and t-fMRI was done by correlating the indices generated by each method across a group of 31 patients and within smaller patient subgroups based on lesion location. From the subset of patients who underwent language mapping, 26 had left hemisphere lesions and 17 had lesions in the left temporal lobe. Because only 8 patients had lesions in the left frontal lobe, and 3 had lesions in the right hemisphere, these groups were not included in this analysis.

To calculate t-fMRI based language lateralization indices (LLI) we employed the most common threshold dependent method. A total of 6 large ROIs (Fig. 2A) were created from the probabilistic Harvard-Oxford structural atlas available in FSL to account for possible atypical activation in the patient group and included areas known to be involved in language processing: all gray matter of the frontal lobe, posterior half of the inferior, middle and superior temporal lobe, and temporoparietal regions, angular gyrus and supramarginal gyrus (Backes et al., 2005; Binder et al., 2008; Sabsevitz et al., 2003; Szafarski et al., 2002). LLI were calculated for each participant and each task, using (L − R) / (L + R), where L and R are the number of voxels significantly activated (Z > 2.3, p = 0.05) in the ROIs in each hemisphere. As in previous studies, higher positive numbers correspond to left lateralized language (Binder et al., 1996; Backes et al., 2005; Szafarski et al., 2002).

For rs-fMRI, we computed lateralization index measures as developed by Liu et al. (2009b). To determine this lateralization index, intrahemispheric FC between language network ROIs and their right hemisphere homologues were used to calculate lateralization using:

\[
\frac{(LL-RL)-(RR-LR)}{LL+|RL|+|RR|+|LR|}
\]

where \(LL = \text{LSTG-to-LIFG FC; } RR = \text{RSTG-to-RIFG FC; } LR = \text{LSTG-to-RIFG FC; and } RL = \text{RIFG-to-LSTG (see Supplemental Fig. 1S, panel A).}

Higher positive indices indicate stronger left lateralization just as for task lateralization. One crucial difference between the methods we employed here and those of Liu et al. (2009b) is our lack of global signal regression in the process of deriving FC. Global signal regression has been shown to affect functional connectivity patterns across the brain (Weissenbacher et al., 2009), including lateralization of functional connectivity (McAvoy et al., 2015), likely as a result of ignoring the locus of neural signal (Scholvinck et al., 2010). We then correlated resting state

---

**Fig. 1.** Lesion overlap in patient group. Lesion masks in MNI space corresponding to all 43 patients. Brighter red colors correspond to more lesion overlap across the patient group. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

**Fig. 2.** Regions of interest. A: Large ROIs were created from the probabilistic Harvard-Oxford structural atlas available in FSL, to account for possible atypical activation in the patient group. B: Smaller language ROIs (left superior temporal gyrus, and left inferior frontal gyrus) were also atlas derived and used for FC calculations.
LLls to task LLls in our group of patients and healthy controls to evaluate if resting state LLls can replace standard task LLls.

2.6. Group comparisons of functional connectivity

The pairwise FCs used to calculate resting lateralization indices were compared between patients and healthy controls to determine how FC within the language network is disrupted in patients with brain damage, and to evaluate the resting language lateralization index in a more nuanced manner by decomposing it into individual FCs. In particular, we characterized patterns of intrahemispheric and interhemispheric FC within the language network and its homologues as function of lesion location using two-tailed t-tests. We compared the magnitude of difference between each patient group (i.e., patients with lesions in the left hemisphere, right hemisphere, left temporal lobe, and left frontal lobe) and healthy controls to determine whether certain lesion groups showed more impaired FC.

In a subsequent analysis, differences in brain connectivity to the language network were explored using the whole brain FC maps created using the same language network seed regions and their homologues. We compared whole brain FC maps by applying two-tailed t-tests to FC at every voxel in the brain between pairs of patient groups. To test how FC differed as a result of damage to a hemisphere, we compared FC between patients with lesions in the left and right hemispheres. To test how location within hemisphere impacted FC, we then tested patients with lesions in the left temporal and frontal lobes. This resulted in whole brain t-statistic maps showing significant differences in FC between the language network and all other voxels of the brain. These maps were thresholded at $p < 0.01$ (cluster threshold $p < 0.05$). In order to address patient group differences that may attributable to lesion volume, functional connectivity in these regions was also correlated across subjects to their lesion volume as defined by the mm volume of their lesion mask, and reported in the table associated with each figure ($p$-values only reported for significant relationships).

2.7. Functional connectivity as an index of language network lateralization

Finally, in light of recent research suggesting FC predicts language lateralization in epilepsy patients (Doucet et al., 2015) we also correlated whole brain FC computed from the language network and its homologues, to task based LLls. FC from a seed to every voxel of the brain was extracted for every patient and healthy control, and correlations were performed across subjects (including separately in each patient group), between this measure and task based LLls. The resulting whole brain correlation maps were thresholded at $p < 0.05$ (cluster threshold $p < 0.05$), and again the influence of lesion volume was evaluated in the resulting effects by correlating LLls directly to lesion volume (see Supplemental section Lesion Volume Analysis on page 55). Doucet et al. (2015) report different regions for patients and healthy controls that exhibit functional connectivity that correlates with that groups’ corresponding language lateralization index. In our analysis we were also interested in determining whether some regions of the brain show functional connectivity to the language network that could replace language lateralization across both healthy controls and patients. As a result, we identified voxels with FC to the language network that correlated significantly to language lateralization both in patients and controls. The effect that lesions have on task activation, and therefore t-fMRI based language lateralization, may correspond to altered functional connectivity within that region of the brain, allowing for a spatial overlap between the healthy and diseased brain that exhibits functional connectivity related to task based language lateralization. Determining brain regions that show this relationship consistently across diverse populations would help facilitate employment of similar approaches in the clinic as this would not require specialized treatment of patient groups.

3. Results

3.1. Language lateralization indices in patients and healthy controls

Consistent with previous studies, when using task fMRI we found typical left-sided dominance for language in most participants (across all tasks: 2 of 31 patients had LLI < 0.1). LLls were consistent across tasks but the VG task elicited the strongest left lateralization both in patients and healthy controls (Fig. 3A). We also generated combined task LLls, as prior research has suggested that language lateralization analysis over several tasks may produce more reliable results (Ramsey et al., 2001). Two-tail t-tests did not reveal any between-group differences in t-fMRI based LLls. Comparison of rs-fMRI LLls between patients ($n = 31$) and healthy controls ($n = 17$) revealed a difference trending towards significance ($t(58) = 1.97, p < 0.06$). It is worth noting that in comparison to t-fMRI LLls, the rs-fMRI LLls were generally low, indicating more bilateral language representation (Fig. 3A).

We found significant correlations between resting state and task LLls only for WGC (Fig. 3B), which elicited more bilateral language activity (see Fig. 3A). Although there was no effect across the entire patient group (Fig. 3B), we tested correlations between t-fMRI and rs-fMRI based LLls within our two largest patient groups: patients with lesions in the left hemisphere and patients with lesions in the left temporal lobe. This analysis yielded a significant correlation between rs-fMRI based LLls and language lateralization during WGC in patients with lesions in the left hemisphere (r(24) = 0.43; $p < 0.05$), and in patients with lesions in the left temporal lobe (r(17) = 0.66; $p < 0.004$, Fig. 3BII). Patients with lesions in the left temporal lobe also showed a significant relationship between rs-fMRI LLls and language lateralization indices generated from activations across all 4 t-fMRI scans (r(17) = 0.51; $p < 0.04$), but again this LLI was more bilateral relative to individual tasks. In healthy controls the relationship between rs-fMRI LLls and LLls from WGC approached significance (r(17) = 0.42; $p = 0.1$), and the stronger correlation in patients is consistent with the finding that patients showed a general trend for weaker left lateralized language representation during task than healthy controls (see Fig. 3A). Tasks that elicited more left lateralized language activity, such as VG, produced LLls that correlated weakly with rs-fMRI. Thus, a clear picture emerges showing that rs-fMRI LLls may correlate to t-fMRI LLls for tasks that exhibit more bilateral language network activation.

3.2. Group comparisons of functional connectivity in the language network

3.2.1. Patients vs. healthy controls

Comparisons between patient groups and healthy controls were then investigated to determine whether altered intrahemispheric or interhemispheric FC in the language network (as used for the rs-fMRI LLI measure) may be affected by lesion location and thus help explain the correspondence between bilateral LLls from t-fMRI, and LLls from rs-fMRI for certain patient lesion groups. Intrahemispheric FC between the right homologues of the language network was lower in the patient group relative to controls (Fig. 4A; decrease of 0.12 in FC from RIFG to RSTG: t(41) = 2.9; $p < 0.006$ and decrease of 0.1 in FC between RSTG and RIFG: t(41) = 2.7; $p < 0.01$), but interhemispheric FC between the left language network ROIs and their right hemisphere homologues was significantly higher (Fig. 4A; increase of 0.11 in FC from RIFG to RSTG: t(41) = 3.7; $p < 0.0007$ and increase of 0.14 in FC from LSTG to RIFG: t(41) = 3.1; $p < 0.004$). We also found that correlations between the timecourse of a whole ROI in the language network, or its homologues, and voxels within that ROI itself (i.e., intraregional functional connectivity) were significantly lower in patients, describing generally depressed FC in the patient group (Fig. 4A; LIFG: decrease of 0.11 in FC t(41) = 4; $p < 0.0003$; LSTG decrease of 0.11: t(41) = 3.4; $p < 0.002$; RIFG decrease of 0.14: t(41) = 3.8; $p < 0.0006$; RSTG decrease of 0.16: t(41) = 4.7; $p < 0.00004$).
**3.2.2. Patient lesion groups vs. healthy controls**

Patients with lesions in the right hemisphere had more disrupted FC in the right hemisphere language network homologues than patients with left hemisphere lesions when we compared each of these groups to healthy controls (Fig. 4B; **patients with lesions in the right hemisphere**: decrease of 0.15 in FC between RSTG and RIFG: t(11) = 2.7; p < 0.02 and decrease of 0.14 in FC between RIFG and RSTG: t(11) = 2.8; p < 0.02; **patients with lesions in the left hemisphere**: decrease of 0.08 in FC between RSTG and RIFG: t(27) = 2.1; p < 0.05 and decrease of 0.1 in FC between RIFG and RSTG: t(27) = 2.3; p < 0.04). Patients with lesions in the left hemisphere had more disrupted intrahemispheric connectivity between the language network and its homologues than patients with right hemisphere lesions, when compared to healthy controls (**patients with lesions in the right hemisphere**: increase of 0.11 in FC from LSTG to RIFG: t(11) = 2.4; p < 0.04; **patients with lesions in the left hemisphere**: increase of 0.14 in FC between LSTG and RIFG: t(27) = 3.6; p < 0.002; increase of 0.13 in FC between RIFG and LSTG: t(27) = 3.9; p < 0.0006).

Intrahemispheric FC changes in patients with lesions in the left hemisphere were stronger for patients with lesions in the temporal lobe, who showed larger changes in FC when compared to controls than patients with lesions in the frontal lobe did (**patients with lesions in the left temporal lobe**: decrease of 0.1 in FC between LSTG and LIFG: t(15) = 3.3; p < 0.005; increase of 0.14 in FC between LSTG and RIFG: t(15) = 3.3; p < 0.005).

**3.2.3. Patients with lesions in the left versus right hemispheres**

In a whole brain contrast analysis, we observed differences in FC generated from the language network and its homologues between patients with lesions in the right and left hemispheres. When seeding from the LIFG, we found significantly higher connectivity primarily to bilateral postcentral gyri (PostCG; though the peak coordinate in the right hemisphere is in the central opercular cortex), bilateral frontal poles (FP), and bilateral occipital poles (OP) in patients with lesions in the
left hemisphere relative to patients with lesions in the right hemisphere (Table 2, Fig. 5). In order to address whether any of the effects within these regions are related to lesion volume, a correlation was performed across subjects between functional connectivity within each cluster, and lesion volume. This analysis suggests that the greater connectivity from the LIFG to the LFP for patients with lesions in the left hemisphere is related to lesion volume such that patients with smaller lesions have higher FC ($r = -0.52; p < 0.02$). In contrast, patients with lesions in the right hemisphere showed higher connectivity from the LIFG to the right inferior temporal gyrus (RITG) (Table 2, Fig. 5). When whole brain FC was computed seeding from the RIFG, we found higher FC to the periaqueductal cortex bilaterally, including subcallosal cortices (SC) and parahippocampal gyrus (PHG), but also to the right posterior cingulate gyrus (RPCG) in patients with left hemisphere lesions. Greater connectivity from the RIFG to the right supplemental motor area (RSMA), right temporal occipital fusiform cortex (RTOF), RITG, and right precuneus (RP) was observed in patients with right hemisphere lesions (Table 2, Fig. 5).

When seeding from the LSTG we found that the FC to the ITG bilaterally, as well as to the right parietal lobe, including right supramarginal
Table 2
Peak differences in FC between patients with lesions in the left hemisphere and patients with lesions in the right hemisphere.

### Seeding from the LIFG

| Region                                               | Peak t-value | Correlation to lesion volume t-value (p-value) | x, y, z-coordinate (MNI) |
|------------------------------------------------------|--------------|-----------------------------------------------|-------------------------|
| Patients with lesions in left hemisphere > patients with lesions in the right hemisphere |              |                                               |                         |
| R Frontal orbital cortex                             | 3.63         | 0.12                                          | 14, 20, −18             |
| R Occipital pole                                     | 4.18         | 0.32                                          | 22, −95, −24            |
| L Postcentral gyrus                                  | 4.03         | −0.08                                         | −68, −2, 18             |
| R Frontal pole                                       | 3.27         | 0.1                                           | 24, 40, 0               |
| R Intracalcarine cortex                              | 3.77         | −0.21                                         | 8, −82, 2               |
| R Central opercular cortex                           | 3.20         | −0.05                                         | 56, −4, 6               |
| L Lateral occipital cortex                           | 3.75         | −0.18                                         | −50, −70, −20           |
| L Occipital pole                                     | 3.34         | 0.16                                          | −10, −98, 2             |
| L Frontal orbital cortex                             | 3.16         | 0.2                                           | −18, 18, −16            |
| R Paracingulate gyrus                                | 3.01         | −0.06                                         | 0.38, −8                |
| L Frontal medial cortex                              | 3.2          | 0.32                                          | −8, 34, −18             |
| L Lingual gyrus                                      | 3.44         | −0.14                                         | −4, −84, −10            |
| L Frontal pole                                       | 3.62         | −0.52 (p < 0.02)                              | −18, 70, 14             |
| L Insular cortex                                     | 3.12         | 0.08                                          | −32, 12, −12            |
| R Middle frontal gyrus                               | 3.75         | −0.32                                         | 38, 2, 08               |
| L Intracalcarine cortex                              | 3.08         | −0.27                                         | −10, −72, 10            |
| T-L Temporal pole                                    | 2.94         | −0.33                                         | −58, 8, 0               |
| R Frontal orbital cortex                             | 3.41         | 0.03                                          | 38, 28, −8              |
| L Frontal medial cortex                              | 2.99         | 0.45 (p < 0.04)                               | −12, 42, −14            |
| Patients with lesions in right hemisphere > patients with lesions in the left hemisphere |              |                                               |                         |
| R Precuneous cortex                                  | 3.32         | −0.31                                         | 6, −58, 48              |
| L Parahippocampal gyrus, anterior division           | 3.61         | 0.17                                          | −32, −4, −26            |
| R Postcentral gyrus                                  | 4.16         | −0.28                                         | 30, −36, 46             |
| L Lingual gyrus, posterior                           | 3.67         | −0.92 (p < 0.0005)                            | 8, −42, 50              |
| R Middle frontal gyrus                               | 3.44         | −0.14                                         | −8, −42, −10            |
| R Occipital gyrus                                    | 4.27         | 0.06                                          | 22, −14, 44             |
| L Frontal pole                                       | 3.7          | −0.1                                          | 14, 60, 16              |
| T-L Inferior temporal gyrus, anterior division       | 4.24         | −0.02                                         | 56, 4, −42              |
| T-L Temporal fusiform cortex                         | 3.34         | 0.26                                          | 30, 10, −46             |
| R Precuneous cortex                                  | 3.28         | −0.43                                         | 16, −60, 24             |
| R Temporal occipital fusiform cortex                 | 3.11         | −0.07                                         | 42, −62, −10            |
| L Occipital fusiform gyrus                           | 3.22         | −0.1                                          | −30, −70, −8            |
| T-L Temporal fusiform gyrus                          | 3.02         | −0.28                                         | −34, 8, −42             |

### Seeding from the RIFG

| Region                                               | Peak t-value | Correlation to lesion volume t-value (p-value) | x, y, z-coordinate (MNI) |
|------------------------------------------------------|--------------|-----------------------------------------------|-------------------------|
| Patients with lesions in left hemisphere > patients with lesions in the right hemisphere |              |                                               |                         |
| R Frontal pole                                       | 4.09         | −0.26                                         | 30, 42, 26              |
| R Supplementary motor area                           | 3.6          | 0.09                                          | 10, −8, 56              |
| R Superior frontal gyrus                             | 3.71         | −0.65                                         | 6, −2, 72               |
| L Temporal fusiform cortex                            | 4.21         | −0.25                                         | −30, 8, −48             |
| R Precuneous cortex                                  | 4.95         | −0.26                                         | 14, −58, 46             |
| R Parahippocampal gyrus, anterior division            | 3.9          | 0.41                                          | 38, −8, −18             |
| L Middle temporal gyrus                              | 3.96         | −0.21                                         | −70, −42, −4            |
| L Frontal pole                                       | 3.3          | −0.2                                          | −20, 54, 12             |
| R Postcentral gyrus                                  | 4.52         | −0.4                                          | 8, −46, 66              |
| R Cingulate gyrus, anterior division                 | 3.88         | −0.14                                         | 18, 30, 20              |
| R Temporal occipital fusiform cortex                 | 3.86         | 0.24                                          | 40, −48, −10            |
| L Cingulate gyrus, anterior division                 | 3.36         | 0.19                                          | −10, 20, 24             |
| L Inferior temporal gyrus                            | 3.14         | 0.39                                          | −48, −6, −40            |
| R Supramarginal gyrus                                | 3.45         | −0.06                                         | 26, −32, 36             |
| R Occipital fusiform gyrus                           | 3.22         | 0.31                                          | 18, −80, −24            |
Seeding from the LSTG

| Region | Peak t-value | Correlation to lesion volume r-value (p-value) | x, y, z-coordinate (MNI) |
|--------|--------------|-----------------------------------------------|------------------------|
| Patients with lesions in left hemisphere > patients with lesions in the right hemisphere | | | |
| L Cingulate gyrus, anterior division | 3.95 | −0.22 | −14, −16, 30 |
| L Frontal medial cortex | 4.79 | 0.06 | −2, −42, −24 |
| L Frontal orbital cortex | 4.08 | 0.33 | −20, 18, −2 |
| L Parahippocampal gyrus, dposterior division | 5.11 | 0.13 | −12, −24, −10 |
| L Frontal orbital cortex | 3.83 | −0.02 | −30, −18 |
| L Paracingulate gyrus | 3.64 | 0.42 (p < 0.05) | −48, −14, −9.6 |
| L Subcallosal cortex | 3.7 | 0.16 | −6, 20, −12 |
| L Superior frontal gyrus | 4.01 | 0.02 | −2, 66, 28 |
| L Frontal pole | 3.7 | −0.1 | −32, 66, −6 |
| L Precenral gyrus | 3.18 | 0.21 | −2, −26, 76 |
| R Middle temporal gyrus, posterior division | 3.65 | −0.31 | 6, −18, −12 |
| R Parahippocampal gyrus, anterior division | 3.38 | 0.36 | −44, −24, −4 |
| R Temporal fusiform cortex | 4.07 | −0.11 | 36, −26, −40 |
| R Frontal pole | 3.28 | −0.27 | 12, −12, −50 |
| L Occipital pole | 3.06 | 0.01 | −10, −102, −22 |
| L Lateral occipital cortex | 3.1 | −0.03 | −62, −58, 18 |

Patients with lesions in right hemisphere > patients with lesions in the left hemisphere

| Region | Peak t-value | Correlation to lesion volume r-value (p-value) | x, y, z-coordinate (MNI) |
|--------|--------------|-----------------------------------------------|------------------------|
| R Supramarginal gyrus | 3.76 | −0.05 | 44, −34, 28 |
| R Temporal fusiform cortex | 3.89 | −0.17 | 24, −42, −18 |
| R Occipital fusiform gyrus | 3.61 | 0.4 | 26, −66, −24 |
| L Superior frontal gyrus | 5.04 | 0.23 | −18, −8, 52 |
| L Inferior temporal gyrus | 3.57 | −0.22 | −48, −50, −20 |
| L Precuneous cortex | 3.52 | −0.5 | −22, −56, 22 |
| R Precuneous cortex | 3.4 | −0.46 | 10, −46, 46 |
| R Superior parietal lobe | 3.57 | 0.03 | 48, −42, 64 |
| R Middle frontal gyrus | 3.48 | 0.11 | 38, 24, 56 |
| L Precenral gyrus | 3.33 | −0.74 (p < 0.03) | −30, −24, 38 |
| L Precuneous cortex | 3.26 | −0.23 | −6, −66, 58 |
| R Superior frontal gyrus | 3.25 | 0.18 | 28, 4, 72 |

Seeding from the RSTG

| Region | Peak t-value | Correlation to lesion volume r-value (p-value) | x, y, z-coordinate (MNI) |
|--------|--------------|-----------------------------------------------|------------------------|
| Patients with lesions in left hemisphere > patients with lesions in the right hemisphere | | | |
| L Precentral gyrus | 4.32 | 0.07 | −8, −18, 76 |
| R Precentral gyrus | 3.71 | 0.3 | 58, −2, 22 |
| R Postcentral gyrus | 4.26 | 0.22 | 64, −14, 44 |
| L Planum temporale | 3.47 | 0.06 | −52, −12, 12 |
| R Cingulate gyrus, anterior division | 4.2 | −0.17 | 0, 30, 6 |
| R Parahippocampal gyrus, posterior division | 4 | −0.2 | 16, −26, −8 |
| R Subcallosal cortex | 3.4 | 0.24 | 14, 20, −6 |
| L Postcentral gyrus | 4.14 | −0.31 | −60, −10, 54 |
| L Parahippocampal gyrus, posterior division | 3.66 | 0.07 | −18, −28, −18 |
| L Superior temporal gyrus | 3.94 | 0.14 | 42, −26, −4 |
| L Temporal fusiform cortex | 5.01 | −0.38 | −34, −28, −38 |
| L Insular cortex | 3.2 | 0.25 | −32, −20, 6 |
| L Cingulate gyrus, posterior division | 4.25 | −0.47 (p < 0.03) | −8, −24, 10 |
| L Paracingulate gyrus | 3.79 | 0.04 | −12, 36, −10 |
| L Parahippocampal gyrus, anterior division | 3.96 | 0.11 | −22, −16, −40 |
| L Postcentral gyrus | 2.95 | 0.16 | −28, −30, 68 |

Patients with lesions in right hemisphere > patients with lesions in the left hemisphere

| Region | Peak t-value | Correlation to lesion volume r-value (p-value) | x, y, z-coordinate (MNI) |
|--------|--------------|-----------------------------------------------|------------------------|
| L Lateral occipital cortex | 3.93 | −0.18 | −42, −86, −18 |
| R Inferior temporal gyrus | 4.54 | −0.03 | 48, −18, −42 |
| R Middle frontal gyrus | 3.94 | −0.02 | 18, 4, 52 |
| L Inferior temporal gyrus | 4.34 | −0.1 | −58, −44, −32 |
| R Occipital fusiform gyrus | 3.76 | −0.05 | 44, −66, −20 |
| R Superior frontal gyrus | 3.31 | −0.2 | 20, 4, 66 |
| R Frontal pole | 3.39 | 0.03 | 42, 60, 6 |
| R Lateral occipital cortex | 3.31 | 0.16 | 44, −80, −20 |
| L Frontal pole | 3.09 | 0.32 | −26, 58, −14 |
| R Middle temporal gyrus, temporooccipital part | 3.78 | −0.18 | 56, −44, −8 |
| L Occipital fusiform gyrus | 3.3 | 0.04 | −24, −66, −20 |
| L Middle frontal gyrus | 3.12 | 0.1 | −30, 16, 40 |
| L Supramarginal gyrus, posterior division | 3.26 | −0.17 | −48, −44, 52 |
| R Inferior temporal gyrus | 3.07 | 0.07 | 58, −44, −18 |
| L Lateral occipital cortex | 3.45 | −0.15 | 28, −82, −42 |

gyrus (RSG), as well as the right superior parietal lobe (RSPL), was higher in patients with lesions in the right hemisphere (Table 2, Fig. 5). Mainly medial regions of the surface of the brain showed higher FC in patients with lesions in the left hemisphere, including bilateral anterior cingulate gyrus (ACG), frontal medial cortex (FMC), and frontal orbital cortex (OFC; but note that the peaks for these regions are in the left hemisphere). Higher FC in left hemisphere lesion patients was observed in the LPHG, and right middle temporal gyrus (RMTG) also (Table 2, Fig. 5). Meanwhile, seeding from the RSTG generated higher FC to precentral gyrus (PreCG), PostCG, PHG, and cingulate gyrus (CG)
bilateral (although note that connectivity to the left cingulate gyrus was correlated with lesion volume; \( r = -0.47, \ p < 0.03 \)), as well as to other portions along RSTG in patients with lesions in the left hemisphere. Patients with lesions in the right hemisphere showed higher connectivity from the RSTG primarily to ITG and lateral occipital cortex (LOC) bilaterally, but also to regions in the right frontal lobe including MFG, and superior frontal gyrus (SFG) (Table 2, Fig. 5).

3.2.4. Patients with lesions in the left temporal versus frontal lobes

The comparison between FC in the group of patients with left frontal lobe lesions, and the group of patients with left hemisphere temporal lobe lesions is shown in Table 3, Fig. 6. Seeding from the LIFG, we found widespread patterns of higher connectivity in patients with left hemisphere temporal lobe lesions. These patients had higher connectivity from the LIFG to the left and right PreCG, LPostCG, LMTG, LSTG, right frontal orbital cortex (ROFC), RMFG, LACG, left supplementary motor area (LSMA) and RIGF. In contrast, patients with lesions in the left frontal lobe had higher functional connectivity in the left inferior anterior temporal regions, and visual areas including LOC and ITG (Table 3, Fig. 6). Connectivity from the RIFG was higher to the LIFG, LPostCG, bilateral PreCG, ROFC, and RSGF in patients with lesions in the left temporal lobe, and higher connectivity to the right postcentral gyrus, RSPIL, right supramarginal gyrus (RSG), RPHG, and ITG bilaterally in patients with lesions in the left frontal lobe. In patients with lesions in the left temporal lobe FC from the LSTG was higher to bilateral PostCG, the LPreCG, bilateral STG, and areas in the parietal and occipital lobes. Patients with lesions in the left frontal lobe meanwhile showed higher FC from the LSTG to regions in the frontal lobe particularly frontal poles (FP), as well as ITG, bilaterally. When seeding from the RSTG we found a similar pattern of FC. Patients with lesions in the left temporal lobe showed higher FC to the right parietal lobe (right angular gyrus), right frontal lobe (RFP), occipital lobe bilaterally (LOC), and PostCG bilaterally. Patients with lesions in the left frontal lobe showed fewer areas of increased FC from the RSTG on the cortex, but included the right frontal lobe (RFP).

3.3. Functional connectivity to the language network that is related to task lateralization both in patients and healthy controls

We evaluated what brain regions show FC to the language network or its homologue that correlates to task based lateralization indices in both patients, and controls (Table 4). This analysis was performed over all tasks (i.e., LLIs generated by collapsing activation across all tasks patients performed) and VG as this particular index was most strongly left lateralized in Fig. 3. Clusters of overlap were generally consistent across these two measures, and particularly implicated FC from the language network to parietal and frontal lobes.

Intrahemispheric FC from the language network to the frontal lobe significantly correlated with LLIs derived from the combined BOLD-contrast of all four language tasks we employed. FC from LIFG, LSTG, RIFG and RSTG to the frontal lobe significantly correlated with language lateralization derived from all tasks (FC from LIFG to LFP in patients: \( r = 0.47, \ p < 0.008 \) and healthy controls: \( r = 0.68, \ p < 0.003 \); FC from LSTG to LFP in patients: \( r = 0.46, \ p < 0.01 \) and healthy controls: \( r = 0.52, \ p < 0.04 \); FC from RIFG to RMFG in patients: \( r = 0.47, \ p < 0.008 \) and healthy controls: \( r = 0.58, \ p < 0.02 \); FC from LSTG to RFP in patients: \( r = 0.51, \ p < 0.004 \) and healthy controls: \( r = 0.62, \ p < 0.008 \)). VG based LLIs also correlated with FC from the language network to regions of the temporal lobe tended not to correlate with LLIs, however interestingly we did find that intra-regional FC in the LSTG (i.e. between the timecourse of the entire LSTG ROI, and individual voxels of the LSTG ROI) correlated significantly with LLIs derived from VG (FC from LSTG to LSTG in patients: \( r = 0.63, \ p < 0.0002 \) and healthy controls: \( r = 0.49, \ p < 0.05 \)). VG based LLIs also correlated with FC between LSTG and the RITG (FC from LSTG to RITG in patients: \( r = 0.37, \ p < 0.05 \) and healthy controls: \( r = 0.75, \ p < 0.0006 \)).

Connectivity between the language network’s right hemisphere homologues and the left parietal lobe, specifically LSG, also significantly correlated to VG based LLIs (FC from RIFG to LSG in patients: \( r = 0.39, \ p < 0.03 \) and healthy controls: \( r = 0.68, \ p < 0.02 \); (FC from RSTG to LSG in patients: \( r = 0.44, \ p < 0.02 \) and healthy controls: \( r = 0.53, \ p < 0.03 \)). FC to the parietal lobe also correlated to LLIs derived from all tasks patients and healthy controls performed, but the relationship was only present when seeding from the LSTG, and implicated the parietal lobe bilaterally (FC from LSTG to LSG in patients: \( r = 0.42, \ p < 0.02 \) and healthy controls: \( r = 0.62, \ p < 0.008 \); FC from LSTG to RSG in patients: \( r = 0.44, \ p < 0.02 \) and healthy controls: \( r = 0.6, \ p < 0.01 \)). In the occipital cortex, the LOC was identified to show FC to the language network that correlated with LLIs derived from all the language tasks we employed. FC from the LIFG to bilateral LOC correlated with LLIs across all tasks (FC from LIFG to LLOC in patients: \( r = 0.42, \ p < 0.02 \) and healthy controls: \( r = 0.49, \ p < 0.05 \); FC from LIFG to RLOC in patients: \( r = 0.48, \ p < 0.007 \) and healthy controls: \( r = 0.49, \ p < 0.05 \)), as well as FC from the LSTG to the RLOC (FC from LSTG to RLOC in patients: \( r = 0.43, \ p < 0.02 \) and healthy controls: \( r = 0.59, \ p < 0.02 \)).

4. Discussion

In this study we evaluated whether rs-fMRI can be used to generate language lateralization indices that are comparable to the current standard of t-fMRI. Resting state is an attractive alternative to t-fMRI for numerous reasons such as shorter scan times that reduce participant movement, and the ability to administer the task to patients that may otherwise perform poorly on task fMRI due to inability to comply with complex task demands. Here we found that a previously proposed method of determining lateralization (Liu et al., 2009b) produces indices suggesting bilateral language network, and as such is inadequate for determining the degree of lateralization in this network. While we did observe some relationship between this resting state language lateralization index and task based language lateralization indices in patients with left hemisphere lesions, this was only the case for tasks that elicited bilateral language activation in the scanner, and thus would be inappropriate for clinical consideration. In contrast, we found no correspondence between this resting state language lateralization method and task based language lateralization in healthy controls. We traced this effect to the underlying FC of the language network, where we found that relative to healthy controls, patients had increased intrahemispheric connectivity to the LSTG, and decreased intrahemispheric connectivity. However, FC between regions in the language network may still be used in place of t-fMRI. In line with previous research on epilepsy, we evaluated FC to the language network that correlated with t-fMRI language lateralization indices, finding that intrahemispheric connectivity involving the frontal cortex, and intrahemispheric connectivity involving the left parietal cortex produces potential candidates for language lateralization measures across patients with lesions, and healthy controls.

4.1. Increased functional connectivity to the left superior temporal gyrus in lesion patients

One surprisingly consistent pattern we found was increased intrahemispheric connectivity between the LSTG and the RIFG across patient lesion groups, which suggests that this may be a general compensatory mechanism induced by lesion pathology within the language network.
Intrahemispheric hyperconnectivity has been found ipsilateral to epileptogenic sites in temporal lobe epilepsy (Maccotta et al., 2013), and between subregions of the hippocampus in patients with left medial temporal lobe epilepsy (Bettus et al., 2009). It has also been previously found in childhood survivors of brain lesions, but only in frontal functional networks using ICA (Chen et al., 2016), and in the motor cortex of juvenile myoclonic epilepsy patients (Vollmar et al., 2011). Patterns of hyperconnectivity have also been observed in psychiatric illness, for example patients with schizophrenia show hyperconnectivity in the default mode network correlated to subpar

**Fig. 5.** Brain regions with higher FC in patients with lesions in the left hemisphere than patients with right hemisphere lesions. A two-sample, two-tailed t-test was performed across subjects on every voxel of the brain, between FC from a seed region in the language network to that voxel (and its homologues) in patients with lesions in the left hemisphere, and patients with lesions in the right hemisphere. Blue colors correspond to regions where FC in patients with lesions in the right hemisphere was significantly greater than FC in patients with lesions in the left hemisphere. Red colors show the reverse, where FC is significantly higher in patients with lesions in the left hemisphere. This analysis is performed across FC maps seeding from all regions of the language network and their right hemisphere homologues. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
Table 3
Peak differences in FC between patients with lesions in the left temporal lobe and patients with lesions in the left frontal lobe.

| Seeding from the LIFG Region | Peak t-value | Correlation to lesion volume r-value (p-value) | x, y, z-coordinate (MNI) |
|------------------------------|--------------|-----------------------------------------------|-------------------------|
| Patients with lesions in left temporal lobe > patients with lesions in the left frontal lobe | | | |
| L Postcentral gyrus | 5.17 | −0.09 | −38, −34, 58 |
| L Superior temporal gyrus | 5.18 | −0.02 | −70, −18, 2 |
| R Inferior frontal gyrus, pars opercularis | 5.07 | 0.05 | 34, 24, 16 |
| R Frontal orbital cortex | 4.05 | 0.2 | 42, 28, −10 |
| R Precentral gyrus | 4.03 | 0.29 | 50, 6, 32 |
| L Precentral gyrus | 3.9 | 0.42 | −48, 0, 30 |
| L Lingual gyrus, anterior division | 3.75 | 0.26 | −14, 4, 26 |
| L Temporal pole | 4.28 | −0.46 | −60, 10, −6 |
| R Frontal pole | 3.3 | −0.14 | −48, 54, 0 |
| R Postcentral gyrus | 3.57 | 0.31 | 46, −16, 46 |
| L Supplementary motor area | 3.76 | 0.35 | −6, −6, 50 |
| R Inferior frontal gyrus, pars triangularis | 3.88 | −0.25 | −38, 24, 12 |
| R Paracingulate gyrus | 3.56 | −0.7 (p < 0.02) | 6, 30, 36 |
| R Insular cortex | 4.43 | 0.76 (p < 0.007) | 42, −2, −2 |
| R Temporal occipital fusiform cortex | 4.08 | 0.11 | 22, −32, −26 |
| R Frontal pole | 3.45 | −0.32 | 42, 54, 0 |
| L Middle frontal gyrus | 3.61 | −0.13 | −34, 28, 28 |
| R Frontal pole | 3.13 | −0.17 | 24, 40, 24 |
| R Supramarginal gyrus | 3.5 | 0.12 | 46, −38, 40 |
| R Superior frontal gyrus | 3.54 | 0.56 | −30, 39, 32 |
| R Middle frontal gyrus | 3.48 | −0.21 | 18, 16, 52 |
| R Frontal pole | 3.08 | −0.14 | 16 |

Patients with lesions in left frontal lobe > patients with lesions in the left temporal lobe

| Seeding from the RIFG Region | Peak t-value | Correlation to lesion volume r-value (p-value) | x, y, z-coordinate (MNI) |
|------------------------------|--------------|-----------------------------------------------|-------------------------|
| Patients with lesions in left temporal lobe > patients with lesions in the left frontal lobe | | | |
| R Inferior temporal gyrus | 4.52 | 0.03 | 42, −12, −40 |
| R Occipital pole | 4.13 | −0.37 | 12, −96, 8 |
| L Lingual gyrus | 4.21 | −0.71 (p < 0.04) | −6, −80, −8 |
| L Temporal occipital fusiform cortex | 4.22 | 0.1 | −26, 52, −12 |
| R Precuneous cortex | 6.47 | −0.44 | 30, −50, 10 |
| R Temporal pole | 3.58 | 0.22 | 46, 10, −42 |
| L Temporal pole | 3.76 | 0.01 | −34, 20, −42 |
| L Lingual gyrus, posterior division | 4.09 | 0.46 | −16, −46, 8 |
| R Lateral occipital cortex, inferior division | 5.04 | −0.28 | 30, −72, 6 |
| R Middle frontal gyrus | 3.65 | −0.25 | 30, 0, 68 |
| L Middle temporal gyrus, posterior division | 4.48 | −0.22 | −62, −6, −36 |
| R Cuneal cortex | 3.42 | −0.29 | 0, −78, 30 |
| R Middle frontal gyrus | 3.65 | −0.25 | 30, 0, 68 |
| R Middle temporal gyrus, posterior division | 4.48 | −0.22 | −62, −6, −36 |
| R Cuneal cortex | 3.42 | −0.29 | 0, −78, 30 |
| L Frontal pole | 3.3 | 0.16 | −30, 48, 38 |
| L Superior frontal gyrus | 3.2 | 0.48 | −10, 30, 62 |
| R Temporal occipital fusiform cortex | 3.47 | −0.05 | 40, −48, −2 |
| R Lingual gyrus | 3.59 | 0.22 | 32, −38, −4 |
| L Occipital fusiform cortex | 3.28 | −0.46 | −20, −72, −6 |
| R Occipital fusiform gyrus | 3.18 | 0.2 | 32, −64, −14 |
| L Lateral occipital cortex, superior division | 3.15 | 0.01 | −54, −68, 40 |
| L Precuneous cortex | 3.23 | 0.03 | −16, −50, 22 |

(continued on next page)
### Table 3 (continued)

#### Seeding from the RIFG

| Region                             | Peak t-value | Correlation to lesion volume r-value (p-value) | x, y, z-coordinate (MNI) |
|-----------------------------------|-------------|-----------------------------------------------|-------------------------|
| R Occipital fusiform gyrus        | 4.43        | 0.19                                          | 20, –74, –20            |
| L Temporal pole                   | 3.76        | –0.6                                          | –46, 12, –30            |
| R Precentral cortex               | 3.54        | –0.22                                         | 8, –56, 60              |
| L Parahippocampal gyrus, anterior division | 3.71 | –0.57                                         | –24, 6, –40             |
| L Precuneus cortex                | 3.66        | –0.59                                         | –14, –58, 44            |
| R Lateral occipital cortex        | 3.17        | –0.06                                         | 50, –66, –16            |

#### Seeding from the LSTG

| Region                             | Peak t-value | Correlation to lesion volume r-value (p-value) | x, y, z-coordinate (MNI) |
|-----------------------------------|-------------|-----------------------------------------------|-------------------------|
| Patients with lesions in left temporal lobe > patients with lesions in the left frontal lobe |
| R Postcentral gyrus               | 5.42        | –0.19                                         | 64, –10, 32             |
| L Postcentral gyrus               | 3.94        | –0.38                                         | –54, –12, 24            |
| R Superior temporal gyrus         | 4.33        | 0.01                                          | 58, –10, 10             |
| L Precentral gyrus                | 3.83        | –0.12                                         | –62, –2, 18             |
| R Supramarginal gyrus             | 4.05        | 0.24                                          | –40, –38, 24            |
| R Frontal pole                    | 3.71        | 0.09                                          | 42, 40, 44              |
| L Superior parietal lobule        | 3.7         | –0.51                                         | –36, –40, 58            |
| L Superior temporal gyrus         | 3.8         | 0.05                                          | –70, –16, 0             |
| R Frontal pole                    | 3.76        | 0.05                                          | 10, 68, 32              |
| L Central opercular cortex        | 3.33        | –0.16                                         | –42, –12, 10            |
| L Cingulate gyrus, anterior division | 3.44   | 0.13                                          | –8, 14, 22              |
| L Lateral occipital cortex        | 3.25        | 0.48                                          | –24, –68, 54            |
| R Central opercular cortex        | 3.14        | –0.34                                         | 62, –4, 10              |

#### Seeding from the RSTG

| Region                             | Peak t-value | Correlation to lesion volume r-value (p-value) | x, y, z-coordinate (MNI) |
|-----------------------------------|-------------|-----------------------------------------------|-------------------------|
| Patients with lesions in left temporal lobe > patients with lesions in the left frontal lobe |
| R Angular gyrus                   | 8.3         | 0.12                                          | 64, –46, 12             |
| L Postcentral gyrus               | 5.82        | 0.01                                          | –22, –18, 80            |
| R Cingulate gyrus, posterior division | 4.81   | 0.19                                          | 6, –50, 16              |
| R Lateral occipital cortex        | 5.62        | 0.24                                          | 54, –72, 2              |
| R Precentral gyrus                | 4.43        | 0.08                                          | 8, –28, 80              |
| L Lingual gyrus                   | 4.26        | –0.13                                         | –16, –56, –10           |
| R Frontal pole                    | 4.64        | 0.18                                          | 22, 46, 50              |
| L Lateral occipital cortex        | 3.92        | –0.06                                         | –46, –72, 2             |
| R Postcentral gyrus               | 3.67        | 0.19                                          | 50, –16, 54             |
| L Intracalcarine cortex           | 3.63        | 0.09                                          | –22, –68, 12            |
| L Central opercular cortex        | 3.63        | 0.64 (p < 0.04)                               | –50, –12, 18            |
| R Cuneal cortex                   | 2.99        | –0.07                                         | 14, –82, 26             |

| Region                             | Peak t-value | Correlation to lesion volume r-value (p-value) | x, y, z-coordinate (MNI) |
|-----------------------------------|-------------|-----------------------------------------------|-------------------------|
| Patients with lesions in left temporal lobe > patients with lesions in the left frontal lobe |
| R Frontal opercular cortex        | 5.16        | –0.2                                          | 48, 22, –4              |
| L Superior frontal gyrus          | 6.57        | –0.57                                         | –8, 24, 52              |
| R Precentral gyrus                | 7.58        | 0.26                                          | –76, 46, 36             |
| L Supramarginal gyrus             | 5.57        | –0.19                                         | 32, –40, 36             |
| R Frontal orbital cortex          | 4.95        | –0.29                                         | –36, 20, –8             |
| L Supramarginal gyrus             | 4.32        | 0.14                                          | –48, –38, 44            |
| R Frontal pole                    | 3.85        | 0.13                                          | 38, 42, –12             |
| L Occipital fusiform gyrus        | 4.1         | –0.39                                         | –16, –76, –20           |
| R Angular gyrus                   | 3.56        | –0.05                                         | –44, –54, 50            |
| L Precuneus gyrus                 | 3.97        | –0.19                                         | –54, 6, 4               |
| R Temporal fusiform cortex        | 3.35        | –0.2                                          | 22, –2, –44             |
| L Supplementary motor cortex      | 3.26        | –0.4                                          | –6, –8, 62              |
| R Parahippocampal gyrus, anterior division | 4.33   | –0.41                                         | 20, –16, –38            |
| L Temporal pole                   | 3.3         | –0.09                                         | –28, 8, –48             |
| L Precuneus cortex                | 3.24        | –0.27                                         | –10, –74, 40            |
| L Precuneus gyrus                 | 3.34        | –0.74 (p < 0.03)                              | –28, –4, 44             |
| R Cingulate gyrus, anterior division | 3.15   | 0.08                                          | 6, 40, 10               |
working memory performance (Whitfield-Gabrieli et al., 2009). Patients with mild TBI also show hyperconnectivity in fronto-parietal task-related networks (Mayer et al., 2011). To the extent of our knowledge this is the first study showing hyperconnectivity in the language network of lesion patients. The pattern of interhemispheric connectivity we observed suggests that right hemisphere brain regions may be coming online to support left hemisphere language function lost due to pathology. This is supported by the finding that generally, lesioned patients showed decreased intrahemispheric FC, thus interhemispheric hyperconnectivity may reflect an internal compensatory mechanism to this decreased connectivity in the language network.

Fig. 6. Brain regions with higher FC in patients with left hemisphere lesions in the temporal lobes than frontal lobes. A two-sample, two-tailed t-test was performed across subjects on every voxel of the brain, between FC from a seed region in the language network to that voxel (and its homologues) in patients with lesions in the left temporal lobe, and patients with lesions in the left frontal lobe. Blue colors correspond to regions where FC in patients with lesions in the left frontal lobe was significantly greater than FC in patients with lesions in the left temporal lobe. Red colors show the reverse, where FC is significantly higher in patients with lesions in the left temporal lobe. This analysis is performed across FC maps seeding from all regions of the language network and their right hemisphere homologues. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
Table 4

Clusters of voxels with functional connectivity to the language network that correlates with task language lateralization indices in both patients and healthy controls.

| Region                        | Peak voxel patients r-value (p-value) | Peak voxel healthy controls r-value (p-value) | x, y, z-coordinate (MNI) |
|-------------------------------|--------------------------------------|-----------------------------------------------|--------------------------|
| FC seed in LIFG               | L Frontal pole 0.47 (p < 0.008)      | 0.68 (p < 0.003)                              | −26, 48, 48              |
| L Lateral occipital cortex    | 0.42 (p < 0.02)                      | 0.49 (p < 0.05)                               | −42, −70, 36             |
| R Lateral occipital cortex    | 0.48 (p < 0.007)                     | 0.49 (p < 0.05)                               | 44, −70, 36              |
| R Superior temporal gyrus     | 0.49 (p < 0.006)                     | 0.51 (p < 0.04)                               | 66, −6, −6               |
| FC seed in LSTG               | L Supramarginal gyrus 0.42 (p < 0.02) | 0.62 (p < 0.008)                              | −60, −44, 20             |
| R Lateral occipital cortex    | 0.43 (p < 0.02)                      | 0.59 (p < 0.02)                               | 46, −74, 40              |
| L Frontal pole                | 0.46 (p < 0.01)                      | 0.52 (p < 0.04)                               | −28, 52, 44              |
| L Angular gyrus               | 0.37 (p < 0.05)                      | 0.57 (p < 0.02)                               | −46, −52, 48             |
| R Supramarginal gyrus         | 0.44 (p < 0.002)                     | 0.6 (p < 0.01)                                | 60, −46, 40              |
| FC seed in RIFG               | R Middle frontal gyrus 0.47 (p < 0.008) | 0.58 (p < 0.02)                              | 30, 30, 20               |
| L Frontal pole                | 0.55 (p < 0.002)                     | 0.6 (p < 0.01)                                | −26, 52, 44              |
| L Superior temporal gyrus     | 0.51 (p < 0.004)                     | 0.58 (p < 0.02)                               | −54, 4, −10              |
| L Putamen                     | 0.46 (p < 0.01)                      | 0.52 (p < 0.04)                               | −26, 10, −4              |
| FC seed in RSTG               | R Frontal pole 0.51 (p < 0.004)      | 0.62 (p < 0.008)                              | 32, 38, −6               |

Task lateralization indices generated from Verb Generation

| Region                        | Peak voxel patients r-value (p-value) | Peak voxel healthy controls r-value (p-value) | x, y, z-coordinate (MNI) |
|-------------------------------|--------------------------------------|-----------------------------------------------|--------------------------|
| FC seed in LIFG               | L Frontal pole 0.44 (p < 0.02)      | 0.59 (p < 0.009)                              | −24, 48, 50              |
| L Superior frontal gyrus      | 0.52 (p < 0.003)                     | 0.55 (p < 0.02)                               | −8, −2, 78               |
| L Supramarginal gyrus         | 0.39 (p < 0.03)                      | 0.68 (p < 0.02)                               | −58, −44, 18             |
| FC in RIFG                    | L Superior temporal gyrus 0.63 (p < 0.0002) | 0.49 (p < 0.05)                              | −52, 6, −12              |
| R Inferior temporal gyrus     | 0.37 (p < 0.05)                      | 0.75 (p < 0.0005)                             | 48, −38, −32             |
| FC seed in RSTG               | L Supramarginal gyrus 0.44 (p < 0.02) | 0.53 (p < 0.03)                               | −50, −46, 22             |

However, it should be noted that these patterns of interhemispheric and intrahemispheric connectivity in our patient group were contingent on lesion location. For example, lesions in the right hemisphere elicited a weaker interhemispheric effect within the language network, and lesions in the left hemisphere frontal lobe showed more widespread changes in interhemispheric language network connectivity. That right hemisphere lesion patients showed this compensatory hyperconnectivity at rest may be explained by the fact that their lesions still induced decreased left hemispheric FC in the language network. This phenomenon is consistent with the general model of connexional diaschisis (see Carrera and Tononi, 2014 for review), with lesions showing distal effects on functional connectivity that may be independent of task activation in the same regions (e.g., Campo et al., 2012). Interhemispheric hyperconnectivity and decreased left hemisphere connectivity in the language network may be related, and occurred even in patients with lesions in the right hemisphere. These findings parallel recent research showing increased FC in multiple sclerosis correlates with impairment (Hawellek et al., 2011), and increased FC in stroke patients may underlie motor recovery (Sharma et al., 2009).

4.2. Altered functional connectivity depends on lesion location

Prior research in patients with drug-resistant epilepsy has found that left hemisphere lesions show widespread changes in FC in the language network using pairwise FC (Briganti et al., 2012) and whole brain FC to the language network (Pravata et al., 2011). Our research confirms these findings in a larger group of patients, and demonstrates that patients with lesions in the right hemisphere showed decreased FC in the left hemisphere (Pravata et al., 2011). By comparing FC within the language network and its homologues between healthy controls, and subgroups of patients categorized on the basis of their lesion location, we determined that patients with lesions in the left hemisphere generally showed lower FC, and patients with temporal lobe lesions within the left hemisphere in particular showed more widely disrupted connectivity patterns than those with left frontal lobe, or right hemisphere lesions. This pairwise FC analysis corroborates previous research using the same method to indicate that patients with posterior gliomas show more differences in the FC of the language network than patients with anterior gliomas (Briganti et al., 2012). Whole brain FC from the language network supported these findings, and implicated regions within the left lateral temporal cortex, including anterior and inferior temporal regions as showing significant patient lesion group differences. Patterns of connectivity from the language network to the rest of the brain also emphasized the subtlety of these effects, as left hemisphere and left hemisphere temporal lesion patient groups showed a considerable network of regions with higher FC outside the language network, particularly the precentral and postcentral gyri.

4.3. Lateralization indices for resting state

We found that the method of calculating lateralization proposed by Liu et al. (2009b) is not sensitive to language lateralization as determined by t-fMRI. Although this method was more favorable in our group of patients with lesions in the left hemisphere, this was only the case for lateralization indices in tasks that generally produced more bilateral language activation. The correspondence in patients is also likely affected by brain pathology, which we determined induced widespread and nuanced changes to FC depending on the lesion location. Thus, this method which takes into account many FC measures, including interhemispheric and intrahemispheric FC, is problematic as our findings
indicate subtle patterns of both increased and decreased interhemispheric FC from the language network, as well as decreased intrahemispheric FC within the language network of patients. It is important to note that one critical difference that may affect the correspondence of the task based and resting state lateralization measures may be the application of global signal regression in the preprocessing of resting state fMRI data, which we did not perform due to mounting evidence that it may induce spurious correlations and negatively affect connectivity patterns (e.g., Murphy et al., 2009; Saad et al., 2012).

Substantial future work will be needed to determine a reliable resting state based language lateralization measure. In line with previous research in epilepsy (Doucet et al., 2015), we propose that using simple FC to the language network may be a good candidate for such a measure. We performed whole brain correlations between task based language lateralization indices and FC from areas of the language network and its homologues to the rest of the brain. Critically, this analysis highlighted that FC from the language network to areas in the frontal and parietal lobes consistently correlated with language lateralization across many tasks. The supramarginal gyrus, superior frontal gyrus, and frontal pole emerged as particularly consistent brain regions that showed the effect, suggesting that this connectivity could be used in lieu of task language lateralization indices in both patients, regardless of lesion location, and healthy controls. Interestingly, the effect can be achieved with homologues of the language network but only using regions within the contralateral hemisphere. These results identify FC to the language network that warrants additional investigation as a potential alternative to language lateralization as assessed using t-fMRI, and are consistent with previous work indicating intrahemispheric FC from the left inferior frontal cortex to regions in the frontal lobe predicts task language lateralization in temporal lobe epilepsy (Doucet et al., 2015).

4.4. Limitations

There are several limitations to the current study. First and foremost, we perform an analysis of functional connectivity between patients and healthy controls (Fig. 4) with the caveat that our controls are not matched for age. As a result, we have no way to elucidate whether the differences we are seeing are attributable to the effect of age on functional connectivity in the language network. That being said, we observed differences in functional connectivity that are consistent with those reported elsewhere with similar patient groups, and using age-matched controls (Briganti et al., 2012; Doucet et al., 2015; Pravata et al., 2011). With regards to interhemispheric hyperconnectivity in the language network of patients that we describe, we are not aware of any reports detailing a similar effect in the language network as a result of aging and as such consider this finding at least mostly attributable to brain damage. Nonetheless, the results we present from our analysis comparing patients to healthy controls (Fig. 4) must be considered carefully as we cannot rule out the influence of age. This limitation does not affect our analysis of rs-fMRI and t-fMRI based language lateralization as we focused on within group effects (Fig. 3), and detailed overlapping effects between the patient and control group. We would expect to see a greater overlap between these groups if they were more homogeneous. It must also be considered that lesions may affect the hemodynamic response of neighboring and distal gray and white matter and as such affect FC measures. We have taken that into account as best as we could through the use of lesion masks, thus eliminating analysis of matter in the lesion, however there is no way for us to tell how other brain matter may have been affected by lesion dynamics using fMRI. Additionally, we were unable to compare FC in patients with lesions in the right temporal and frontal lobes due to sample size, and one question that remains to be answered is whether these patients will show similar effects to patients with lesions in the left hemisphere. Finally, it may be the case that FC between different brain regions in the diseased and healthy brain may better correspond to task based language lateralization. However, identifying regions with FC that show consistent correspondence with task based language lateralization across diverse patient populations would help facilitate the adoption of intrinsic FC for use in the clinic. We show evidence that there may be some regions that can be utilized to this end as their FC correlates with task based language lateralization in both healthy controls and patients with diverse lesions.

5. Conclusions

We conclude that existing methods of determining language lateralization in resting state are inadequate compared to t-fMRI, especially in lesion patients that show systemic differences in FC. Further research is needed to determine how best to identify lateralization within the resting state language network in specific patient populations. To this end, future research may be directed towards obtaining FC measures between regions of the language network, but should consider the relationship between the language network and parietal and frontal regions, especially the supramarginal gyrus, frontal pole, and superior frontal gyrus, which show a consistent relationship to language lateralization determined using a variety of tasks in the scanner.

Acknowledgements

This work was supported by funding from the Harry W. Fischer Research Fund, Department of Imaging Sciences at the University of Rochester Medical Center.

Appendix A. Supplementary data

Supplementary data to this article can be found online at doi:10.1016/j.nicl.2016.10.015.

References

Adcock, J.E., Wise, R.G., Oxbury, J.M., Oxbury, S.M., Matthews, P.M., 2003. Quantitative fMRI assessment of the differences in lateralization of language-related brain activation in patients with temporal lobe epilepsy. NeuroImage 18 (2), 423–438.

Backes, W.H., Deblare, K., Vonck, K., Kessels, A.G., Boon, P., Hofman, P., et al., 2005. Language activation distributions revealed by fMRI in post-operative epilepsy patients: differences between left- and right-sided resections. Epilepsia Res. 66 (1–3), 1–12.

Baladassane, C., Jordan, M.C., Beck, D.M., Fei-Fei, L., 2012. Voxel-level functional connectivity using spatial regularization. NeuroImage 63 (3), 1099–1106.

Bartolomei, F., Bosma, I., Klein, M., Baayen, J.C., Reijnolfs, J.C., Postma, T.J., ... Cover, K.S., 2006a. Disturbed functional connectivity in brain tumour patients: evaluation by graph analysis of synchronization matrices. Clin. Neurophysiol. 117 (9), 2039–2046.

Bartolomei, F., Bosma, I., Klein, M., Baayen, J.C., Reijnolfs, J.C., Postma, T.J., ... Cover, K.S., 2006b. How do brain lesions alter functional connectivity? A magnetoencephalography study. Ann. Neurol. 59 (1), 128–138.

Bettus, G., Bartolomei, F., Confort-Gouny, S., Soulier, E., Laguitton, V., ... Guye, M., 2009. Decreased basal fMRI functional connectivity in epileptogenic networks and contralateral compensatory mechanisms. Hum. Brain Mapp. 30 (5), 1580–1591.

Biswal, B., Zerrin Yetkin, F., Haughton, V.M., Hyde, J.S., 1995. Functional connectivity in the motor cortex of resting human brain using echo-planar MRI. Magn. Reson. Med. 34 (4), 537–541.

Binder, J.R., Swanson, S.J., Hammeke, T.A., Morris, G.L., Mueller, W.F., Fisher, M., ... Haughton, V.M., 1996. Determination of language dominance using functional MRI. A comparison with the Wada test. Neurology 46 (4), 978–984.

Binder, J.R., Frost, J.A., Hammeke, T.A., Bellgowan, P.S.F., Rao, S.M., Cox, R.W., 1999. Conceptual processing during the conscious resting state: a functional MRI study. J. Cogn. Neurosci. 11 (1), 80–93.

Binder, J.R., Sabsevitz, D.S., Swanson, S.J., Hammeke, T.A., Raghavan, M., Mueller, W.M., 2008. Use of preoperative functional MRI to predict verbal memory decline after temporal lobe epilepsy surgery. Epilepsia 49 (8), 1377–1394.

Bizzi, A., Blasi, V., Faioli, A., Ferroli, P., Cardini, M., Danesi, U., ... Broggi, G., 2008. Presurgical functional MR imaging of language and motor functions: validation with intraoperative electrocorticographic mapping. 1 Radiology 248 (2), 579–589.

Bookheimer, S., 2007. Pre-surgical language mapping with functional magnetic resonance imaging. Neurophysiol. Rev. 17 (2), 145–155.

Bosma, I., Bouw, L., Bartolomei, F., Heijmans, J.J., Van Dijk, B.W., Postma, T.J., ... Klein, M., 2008. Synchronized brain activity and neurocognitive function in patients with low-grade glioma: a magnetoencephalography study. Neuro-Oncology 10 (5), 734–744.
