Mesial temporal sclerosis is linked with more widespread white matter changes in temporal lobe epilepsy

Min Liu a, Luis Concha b, Catherine Lebel a, Christian Beaulieu a, Donald W. Gross c,*

a Department of Biomedical Engineering, Faculty of Medicine and Dentistry, University of Alberta, Edmonton, Alberta, Canada
b Instituto de Neurobiologia, Universidad Nacional Autonoma de Mexico, Queretaro, Mexico
c Division of Neurology, Department of Medicine, Faculty of Medicine and Dentistry, University of Alberta, Edmonton, Alberta, Canada

1. Introduction

Temporal lobe epilepsy (TLE) is the most common localization related epilepsy syndrome. Most TLE cases are associated with mesial temporal sclerosis (MTS) which can be detected by abnormal signal intensity and reduced volume of the hippocampus on magnetic resonance imaging (MRI) (Van Paesschen et al., 1997). Along with mesial temporal abnormalities, TLE patients have also demonstrated extensive gray and white matter abnormalities both ipsilateral and contralateral to the seizure onset zone (Keller and Roberts, 2008; Otte et al., 2012). Patients with TLE and unilateral MTS (TLE+uMTS) often demonstrate gray matter atrophy within and beyond the ipsilateral temporal lobe.

Abbreviations: TLE + uMTS, temporal lobe epilepsy with unilateral mesial temporal sclerosis; nl-TLE, temporal lobe epilepsy without mesial temporal sclerosis, non-lesional.

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* Corresponding author at: Division of Neurology, Department of Medicine, 2E3.19 Walter C Mackenzie Health Sciences Centre, Edmonton, AB, Canada T6G 2B7. Tel.: +1 7804073616; fax: +1 7804071325.
E-mail address: donald.gross@ualberta.ca (D.W. Gross).

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abnormalities in TLE+uMTS as compared with nl-TLE in a small number of white matter tracts (Concha et al., 2009). Whether TLE+uMTS and nl-TLE demonstrate differences in other white matter tracts previously reported as abnormal in TLE remains unknown (Otte et al., 2012; Kim et al., 2008; Shon et al., 2010; Focke et al., 2008; Thivard et al., 2005; Kemmotsu et al., 2011; Gross et al., 2006).

The purpose of the current study is to compare the diffusion properties (i.e., fractional anisotropy and mean, parallel and perpendicular diffusivities) of thirteen major white matter tracts in patients with temporal lobe epilepsy and unilateral mesial temporal sclerosis (TLE+uMTS) versus patients with non-lesional TLE (nl-TLE) using diffusion tensor tractography.

2. Materials and methods

Approval of the research protocol was obtained from the local Health Research Ethics Board, and informed consent was obtained from all participants.

2.1. Subjects

TLE with unilateral MTS (TLE+uMTS, n=23, age 40±11 years, range 19–58 years, 14F/9M); Electroencephalogram (EEG) videotelemetry demonstrated unilateral temporal lobe ictal onset in all patients, among which 15 were left (n=15, age 39±13 years, range 19–58 years, 8F/7M) and eight were right (n=8, age 42±5 years, range 36–48 years, 6F/2M). Hippocampal sclerosis was defined based on T2 relaxometry analysis where the ipsilateral hippocampal T2 values were above two standard deviations of the overall mean of the controls (i.e., 120 ms using the methodology below). Patients with contralateral hippocampal T2 greater than two standard deviations of controls were included if the ipsilateral hippocampal T2 was greater than that of the contralateral value and EEG demonstrated ictal EEG onset solely from the ipsilateral temporal region. No other lesions were identified from the clinical imaging for all subjects.

Non-lesional TLE (nl-TLE, n=15, age 35±8 years, range 17–45 years, 6F/9M); All patients had temporal lobe epileptic EEG abnormalities. EEG lateralization was left temporal for four patients, right temporal for seven, and bitemporal for four. No lesions were visually identified on the clinical imaging and hippocampal T2 was within two standard deviations of controls bilaterally.

Healthy controls (n=21, age 37±12 years, range 19–58 years, 8F/13M); All control subjects had no history of any neurological or psychiatric disorders and had no evidence of structural lesions on T1 weighted MRI.

There were no significant age differences among the three groups (analysis of variance (ANOVA), p=0.31). The TLE+uMTS patients had an earlier age of seizure onset than the nl-TLE patients (TLE+uMTS: 14±12 years; nl-TLE: 23±10 years; Mann–Whitney U, p=0.033) and longer disease duration than the nl-TLE patients (TLE+uMTS: 26±14 years; nl-TLE: 12±9 years, Mann–Whitney U, p=0.006).

Note that 17 of the 23 patients in the TLE+uMTS group and 10 of the 15 in the nl-TLE have been previously reported in a more restricted analysis of the thalamus, limbic tracts ( fornix, cingulum), external capsule, and corpus callosum (Concha et al., 2009; Gong et al., 2008).

2.2. Image acquisition

All imaging was performed on a 1.5 T Siemens Sonata scanner (Siemens Medical Systems, Erlangen, Germany). T2 relaxometry with coverage of the hippocampus used a high-resolution, multi-echo sequence with 32 echoes, 10 coronal slices, 3 mm slice thickness with 3 mm inter-slice gap, TR = 4430 ms, TE1 = 9.1 ms, TE spacing = 9.1 ms, NEX = 1, acquisition matrix = 192×176 (interpolated to 384×352), FOV = 230 mm×210 mm, scan time = 8 min 13 s. Both standard DTI and fluid-attenuated inversion recovery (FLAIR) DTI were acquired using the same dual spin-echo, single shot echo planar imaging sequence except FLAIR DTI used an extra inversion pulse and fewer slices. The common parameters include 2 mm slice thickness with no inter-slice gap, TR = 10 s, TE = 88 ms, six diffusion directions with b=1000 s/mm², NEX = 8, acquisition matrix = 128×128 (interpolated to 256×256), FOV = 256 mm×256 mm. Fifty two axial slices with coverage of whole brain were acquired for standard DTI and 26 axial slices with coverage of the fornix plus an inversion time of 2200 ms were used for FLAIR DTI (Concha et al., 2005a). Scan times for standard DTI and FLAIR DTI were 9:30 min and 8:30 min, respectively.

2.3. Quantitative hippocampal T2 relaxometry

The T2 signal decay was fitted to a mono-exponential curve voxel by voxel across the multi-echo coronal images. T2 values for the left and right hippocampi were calculated by averaging within the regions of interest manually drawn on two consecutive slices (Concha et al., 2005b).

2.4. Diffusion tensor tractography

In total, thirteen major white matter tracts were analyzed in this study. Limbic tracts (i.e., fornix and parahippocampal cingulum) and genu, occipital and temporal callosal tracts were included in order to verify our previous findings (Concha et al., 2009) and the other eight tracts were chosen based on previously reported abnormalities from other groups (Otte et al., 2012; Kim et al., 2008; Shon et al., 2010; Focke et al., 2008; Thivard et al., 2005; Kemmotsu et al., 2011; Gross et al., 2006). The tracts were also selected due to their well-established tracking protocols using deterministic tractography (Wakana et al., 2004). Tensor calculation and tractography of the parahippocampal cingulum (pCG), dorsal cingulum (dCG), uncinate fasciculus (UF), inferior/superior longitudinal fasciculus (ILF/SLF), inferior fronto-occipital fasciculus (IFO), anterior limb of the internal capsule (ALIC), corticospinal tracts (CST) and genu and body of the corpus callosum (gCC/bCC) on the whole brain DTI dataset and the occipital and temporal part of the corpus callosum (oCC/tapetum) on the FLAIR DTI dataset (to be consistent with our previous study) (Concha et al., 2009) were identified using a semiautomated deterministic tractography method fully described before (Lebel et al., 2008) (Fig. 1). In brief, a non-diffusion weighted image template was created based on six scans of a 33-year-old male control subject coregistered to each other and averaged together for both whole brain DTI and FLAIR DTI datasets. Regions-of-interest (ROIs) for selecting specific tracts were manually drawn on the template subject based on known anatomical landmarks (Wakana et al., 2004). Each subject’s non-diffusion weighted images were non-linearly normalized to the custom template using SPM8 (Wellcome Department of Cognitive Neurology, London, United Kingdom). The estimated deformation map from normalization step was then inverted and applied to transform the template’s ROIs into each subject’s native space. Fiber tracking was performed in the subject’s native space using the deformed ROIs and a deterministic streamline method provided by ExploreDTI (Leemans et al., 2009). All resultant tracts were visually examined and a limited number of spurious tracts which deviated into other distant anatomical locations were excluded manually. The tractography of the crus of the fornix (Fx) was performed manually (due to the requirement of finer ROIs for this small structure) (Malykkin et al., 2008) on the FLAIR DTI dataset to reduce the signal contamination from the adjacent cerebrospinal fluid (Concha et al., 2005a). Fractional anisotropy threshold was set to 0.25 and the angle threshold was set to 60° for all tracts. In order to reduce the effects of crossing fibers and tract variability near the gray matter boundaries, only the portion between the decusation and the body of the corpus callosum axially and the medial central part before
the lateral corpus callosum fibers turning up to the primary cortices were analyzed in CST and bCC, respectively (Fig. 1). The oCC and tapetum were also analyzed using the same method as our previous study (Concha et al., 2009). Fractional anisotropy (FA), mean diffusivity (MD), and parallel and perpendicular diffusivities were obtained by averaging over all voxels within the tract.

2.5. Statistical analyses

Statistical analyses were performed using SPSS version 18. Paired Student’s t test on FA was performed in control group to evaluate the left and right symmetry of the nine paired tracts (except gCC, bCC, oCC and tapetum). A left-greater-than-right asymmetry was observed for FA in the dCg (left = 0.54±0.02; right = 0.52±0.02; p = 0.001) and ILF (left = 0.48±0.02; right = 0.47±0.02; p = 0.03), while a right-greater-than-left asymmetry was observed in the UF (left = 0.44±0.03; right = 0.46±0.02; p = 0.021). The lateralization of nl-TLE patients was complicated due to the presence of independent bilateral seizure onset in four subjects. The analyses of left and right symmetry in our previous study (Concha et al., 2009) further demonstrated low dependence of seizure lateralization on FA of Fx, pCg and external capsule in nl-TLE patients with different onset sides. As our primary interest in the current study was to look for differences between patients with TLE + uMTS and nl-TLE, and given the fact that the right to left differences for the control group were small (all less than 0.02) and to simplify the analysis from a practical perspective, the paired tracts were collapsed by averaging the value from each hemisphere to yield a single value per tract. The between-group differences of FA were tested by multivariate analysis of covariance (MANCOVA) with age included as a covariate for all thirteen tracts together among TLE + uMTS, nl-TLE and controls. To better understand the basis of any FA differences, the mean, parallel and perpendicular diffusivities were also compared between-groups secondarily. Only if Wilks’ Lambda was significant at p = 0.05, then univariate analysis of covariance (ANCOVA) was performed for each tract followed by post-hoc t-tests for those tracts that survived ANCOVA at p=0.05 between groups. All the post-hoc tests for each diffusion measurement were corrected with False Discovery Rate (FDR) at p = 0.05. Additionally, the FA of the white matter tracts with significant between-group differences were evaluated for linear correlations with disease duration (controlling for age) and age of seizure onset (controlling for disease duration) in each patient group. The correlation between the ipsilateral FA of the white matter tracts and ipsilateral hippocampal T2 were also examined in all TLE + uMTS patients (n=23) and 11 of the 15 nl-TLE patients excluding the four patients with bitemporal seizure onset. Both correlation results with and without FDR correction at p = 0.05 level were reported.

To probe the effect of laterality of MTS, the TLE + uMTS patients were divided into TLE with left MTS (left TLE + uMTS, n=15) and TLE with right MTS (right TLE + uMTS, n=8) and compared with controls (n=21). Statistical tests showed neither age differences among the three groups (left TLE + uMTS: 39±13 years; right TLE + uMTS: 42±5 years; controls: 37±12 years; Student’s t test p = 0.5), nor differences of age of seizure onset and duration of epilepsy between the two TLE + uMTS subgroups (left TLE + uMTS: onset 15±12 years, duration: 24±15 years; right TLE + uMTS: onset 14±13 years, duration: 29±12 years; onset Mann–Whitney U, p = 0.87; duration Mann–Whitney U, p = 0.52). For this analysis, each side of the nine paired tracts (except the four commissural tracts, i.e. gCC, bCC, oCC and tapetum) was considered separately to yield an FA value per side. Z-scores of FA (zFA) were calculated for each side of the nine paired fiber tracts in patients based on the mean FA and standard deviation of the corresponding side of controls. MANCOVA and following ANCOVA and post hoc tests were performed for each side of the tracts to compare the zFA between patients and controls separately for left TLE + uMTS and right TLE + uMTS with age included as a covariate. All post hoc tests were corrected by FDR at p = 0.05. To probe tract asymmetry, the left and right zFA for each tract were tested by paired t tests for
each patient group. For the other four commissural tracts, i.e. gCC, bCC, oCC and tapetum, the FA of the whole tract was compared among left TLE+uMTS, right TLE+uMTS and controls using ANCOVA and post hoc tests with age included as a covariate.

3. Results

The hippocampal T2 values of controls were 112 ± 4 ms for the left side and 111 ± 4 ms for the right side. The overall mean of both sides was 112 ± 4 ms. All TLE + uMTS patients demonstrated ipsilateral hippocampal T2 more than two standard deviations greater than controls (mean: 135 ms, range: 122–160 ms); these differences were highly significant (Student’s t test, p<0.000001). T2 of the contralateral side was also significantly larger than that of controls (mean: 117 ms, range: 107–129 ms, p = 0.0001). Four TLE + uMTS patients had contralateral T2 equal to or higher than 120 ms; however, in each case ipsilateral hippocampal T2 was greater than contralateral T2 (range difference in T2 ipsilateral–contralateral: 6–10 ms). None of the nl-TLE patients demonstrated elevated T2 on either side by definition. The average hippocampal T2 for the nl-TLE group was 114 ms (range: 106–118 ms) for the left and 112 (range: 103–118 ms) for the right side, both of which were not significantly different from the overall mean of controls (left side p = 0.07, right side p = 0.87).

3.1. Primary analysis

The MANCOVA test on FA showed a significant Wilks’ Lambda at p<0.001. ANCOVA and post-hoc tests revealed that ten tracts had significantly lower FA in TLE + uMTS relative to controls (all tracts except gCC, oCC and tapetum) and seven of these (all except the CST, UF and SLF) had lower FA with respect to nl-TLE group (Fig. 2a). The nl-TLE group demonstrated significantly reduced FA of pCg and tapetum only with respect to controls.

3.2. Secondary analysis

The MANCOVA test on MD also demonstrated significant Wilks’ Lambda at p<0.001. The subsequent univariate analyses revealed four tracts (pCg, ILF, IFO and bCC) with significantly higher MD in TLE + uMTS compared to controls, among which the bCC also showed higher MD relative to nl-TLE group (Fig. 2b). There were no significant MD differences between nl-TLE patients and controls. The MANCOVA test on parallel diffusivity showed significant Wilks’ Lambda at p = 0.002. The ANCOVA tests further showed the CST with reduced parallel diffusivity in the TLE + uMTS group versus controls and the bCC with significantly higher parallel diffusivity in TLE + uMTS versus nl-TLE (Fig. 2c). Wilks’ Lambda for perpendicular diffusivity was significant at p<0.001. The elevation of perpendicular diffusivity was observed in ten tracts (all except the CST, gCC and oCC) in TLE + uMTS patients compared to controls, among which the elevation in Fx, pCg, dCg, ILF, IFO and bCC was also significant with respect to nl-TLE patients (Fig. 2d). Only the tapetum showed elevated perpendicular diffusivity in nl-TLE group relative to controls. Uncorrected correlations between disease duration and FA demonstrated no significant findings in 11 of 13 tracts with positive findings being observed in dCg (r = −0.58, p = 0.029) and bCC (r = −0.59, p = 0.027) in nl-TLE patients. However, neither results approached significance following FDR correction (FDR corrected dCg p = 0.19, bCC p = 0.35). For hippocampal T2 versus FA, no significant correlation was found before or after FDR correction.

3.3. Analysis of right and left TLE + uMTS

For the nine paired tracts, right TLE + uMTS had FA reductions in all with seven bilateral and two right only whereas left TLE + uMTS had FA reductions in six tracts with four bilateral and two left only (Fig. 3). Specifically, for patients with left TLE + uMTS (n = 15), four tracts (pCg, dCg, ILF and CST) demonstrated bilateral FA reduction and two tracts (Fx and IFO) showed ipsilateral FA reduction. For patients with right TLE + uMTS (n = 8), significant bilateral FA reduction was shown in...
seven tracts (Fx, pCg, dCg, ILF, SLF, IFO and ALIC) and ipsilateral FA reduction was shown in two tracts (UF and CST). Among the six tracts with significant changes in the left TLE + uMTS group, paired t tests revealed four tracts (dCg, ILF, Fx and IFO) with greater FA reduction on the left side, the UF with greater FA reduction on the right side and two tracts (pCg and CST) without significant asymmetry. Among all the nine paired tracts showing significant change in the right TLE + uMTS group, paired t tests revealed two tracts (pCg and UF) with greater FA reduction on the right side with the other seven tracts not showing any asymmetry. For the four commissural tracts, the only between group difference was shown in bCC (ANOVA, $p < 0.001$) where the FA of all three groups was significantly different from each other with right TLE + uMTS showing the most FA reduction compared to controls (controls: $0.65 \pm 0.02$, left TLE + uMTS: $0.61 \pm 0.03$, right TLE + uMTS: $0.58 \pm 0.05$).

4. Discussion

In the current study, widespread abnormalities of white matter tracts within and beyond the temporal lobe were demonstrated in TLE + uMTS patients, while nl-TLE patients demonstrated fewer white matter abnormalities. These results suggest disruption of brain networks in TLE patients with and without MTS, although the affected network was more extensive in TLE + uMTS. Interestingly, in our cohort of subjects, TLE patients with right MTS showed more extensive bilateral changes than in TLE patients with left MTS, despite similar ages of onsets or disease durations.

These findings on a larger sample (an additional 6 TLE + uMTS and 5 nl-TLE patients) confirm our previous findings of FA reduction in the Fx and pCg in TLE + uMTS patients (Concha et al., 2005b, 2009). Furthermore, the nl-TLE patients here showed abnormalities of the pCg and tapetum, in agreement with our previous report (Concha et al., 2009). However, the gCC and oCC did not show any changes in either the TLE + uMTS or the nl-TLE groups, as was demonstrated before (Gross et al., 2006; Concha et al., 2009). Along with the increased sample size, the different methodologies used for gCC in the previous studies (ROI drawing on a single slice) and here (three-dimensional tractography) may contribute to the discrepancy of results.

The current observation of the abnormal frontal lobe (ALIC) and temporal lobe connected tracts (IFO, ILF, SLF and UF) as well as limbic system involvement (Fx, pCg and dCg) in TLE + uMTS are in agreement with several former DTI reports on cohorts of purely or largely TLE + uMTS patients (Thiward et al., 2005; Ahmadi et al., 2009; McDonald et al., 2008; Rodrigo et al., 2007; Kim et al., 2011). The diffusion abnormalities identified in the central part of the bCC are in accordance with a previous voxel-based study (Knake et al., 2009).

Interestingly, the CST has not been measured in many previous studies in TLE or has been used as a ‘control’ tract (no significant FA or MD changes in 17 unilateral TLE patients, 14 of whom had MTS, in a recent paper) (McDonald et al., 2008); this is in contrast to our study where the CST had reduced anisotropy in adult TLE + uMTS patients. Our results are however consistent with findings of bilaterally decreased FA of CST in 13 children with left TLE (Govindan et al., 2008). Discrepancy between our CST findings and (McDonald et al., 2008) could relate to differences in methodology. For example, our CST measurement was restricted to the central part of the tract (as shown by the blue highlight in Fig. 1) where the fiber orientation is expected to be most uniform while they measured the full extent of the tracts. Our findings are also compatible with a number of earlier MRI-based volumetric and VBM studies in which white matter volume in total (Hermann et al., 2003), ipsilateral temporal lobe and body of the corpus callosum (Bernasco et al., 2004), temporal pole (Coste et al., 2002), and ipsilateral frontal and parietal lobes and cerebellum (Mueller et al., 2006) was reduced in TLE + uMTS.

The reduction of FA observed here was mainly driven by an increase of perpendicular diffusivity, which is consistent with previous reports (Gross et al., 2006; Concha et al., 2009; Afaránakis et al., 2002; Lin et al., 2008). A few histopathological studies based on animal models proposed parallel and perpendicular diffusivities as surrogate indices of axonal membrane and myelination status, respectively (Budde et al., 2009; Song et al., 2002, 2003, 2005; Wu et al., 2007; Sun et al., 2006). Previous human in vivo DTI studies lent support to the notion by demonstrating increased perpendicular diffusivity with unchanged parallel diffusivity at the chronic stage of Wallerian degeneration when the myelin degradation is expected (Pierpaoli et al., 2001; Concha et al., 2006). These observations would suggest that the observed increase in perpendicular diffusion reflects reduced myelin in the chronically degenerated fibers; however, the expected disruption of axonal membranes associated with degeneration could also impact the perpendicular diffusivity (Concha et al., 2010).

The white matter findings in the pCg and tapetum in nl-TLE patients are in line with a previous VBM study demonstrating white matter concentration/volume reduction in the ipsilateral temporal lobe of nl-TLE patients (McMillan et al., 2004). In contrast to our more limited findings in nl-TLE, a recent DTI study detected widespread and bilateral reduction of white matter FA in the temporal lobes, the entire corpus callosum and the thalamus in a group of ten nl-TLE patients using voxel-based statistics (Keller et al., 2011). A possible explanation for the more extensive findings in the Keller et al. study is the fact that qualitative as opposed to quantitative methods were used to classify subjects as non-lesional and therefore, it is possible that their patient group included some TLE + uMTS
patients. One other recent voxel-based DTI study reported significant MD elevation in the ipsilateral posterior fornix and posterior cingulum in ten left nL-TLE patients without changes in seven right nL-TLE patients (Shon et al., 2010). We did not look at left and right nL-TLE patients separately due to a small sample size. This, together with the different patient selection criteria and methodologies used (voxel-based statistics for them and tractography here), may account for the discrepancy among the aforementioned DTI studies.

Overall, we found that the white matter abnormalities were extensive and dramatic in TLE + uMTS and more limited in nL-TLE patients. Given the differences in age of seizure onset and disease duration between our TLE + uMTS and nL-TLE patients, where the MTS patient’s onset was 9 years earlier and had 14 years longer disease duration, a possible explanation for the distinct white matter abnormalities observed is that these changes are secondary to seizure related white matter injury; however, the lack of correlations between FA and age of seizure onset/disease duration would argue against this hypothesis. That said, given the known changes of diffusion parameters in white matter during neurodevelopment (Lebel et al., 2008), it is possible that seizure onset during critical neurodevelopmental periods of childhood and adolescence in the TLE + uMTS patients may have caused aberrant white matter development that is not evident if the seizures occur after the wiring is mostly laid down by adulthood (Kaaden et al., 2011; Herrmann et al., 2002). The distinct white matter differences between TLE + uMTS and nL-TLE agree with previous reports on structural and functional MRI and EEG. A more widespread gray matter volume reduction or white matter diffusivity elevation was seen in TLE + uMTS while nL-TLE patients showed either no change (Mueller et al., 2006) or a less extensive pattern (Riederer et al., 2008; Shon et al., 2010). Evidence from EEG recording analysis (Zaveri et al., 2001), multi-slice proton magnetic resonance spectroscopic imaging (Mueller et al., 2004) and structure-function correlation (Mueller et al., 2012) indicate differences between the two TLE sub-syndromes. Our findings further support the idea that TLE + uMTS and nL-TLE are associated with different epileptic networks. Along with more extensive structural changes, TLE + uMTS have also demonstrated more extensive cognitive dysfunction including measures not only of temporal lobe dysfunction (memory and language) but also executive function (Ahmadi et al., 2009; Mueller et al., 2004, 2012; Diehl et al., 2008; Corcoran and Upton, 1993). Further study is required to assess a possible relationship between white matter integrity and cognitive function in TLE. It has been hypothesized that extratemporal structural abnormalities may predict poor surgical outcome in TLE (Sisodiya et al., 1997). Our observation of more extensive white matter abnormalities in TLE + uMTS (who are expected to have a better surgical outcome Sisodiya et al., 1997) however contradicts this hypothesis.

An increasing number of studies suggest that patients with left TLE have greater impairment of learning and memory (Bell and Davies, 1998) and more distributed gray and white matter changes than patients with right TLE (Riederer et al., 2008; Ahmadi et al., 2009; Bonilha et al., 2007; Coan et al., 2009). One recent DTI study focusing on TLE patients with unilateral MTS reported more widespread diffusivity changes in the ipsilateral temporal lobe and bilateral limbic system in 21 patients with left TLE + uMTS than in 12 patients with right TLE + uMTS, while more prominent contralateral changes were observed in the temporal lobe and the inferior frontal gyrus in the right TLE + uMTS group (Focke et al., 2008). Another DTI tractography study reported overall smaller FA in Fc, pgc, UF, ILF, IFO and arcuate fasciculus in 18 patients with left TLE, among which 14 had left MTS, relative to controls and 18 right TLE patients, among which 11 had right MTS (Kemmonsoti et al., 2011). The exact mechanism of this asymmetry remains unknown. Focke et al. (2008) suggested that the seizure propagation may be more widespread in the language dominant hemisphere, commonly the left side, on the basis of a pre-existing better connectivity (Powell et al., 2007) and Kemmonsoti et al. (2011) speculated that the left hemisphere undergoes a more prolonged maturation process than the right and is therefore more vulnerable to early brain insults (Corballis and Morgan, 1978). Interestingly, despite the comparable age, disease duration, age of seizure onset and ipsilateral T2 relaxation time (left TLE + uMTS: 135 ± 10 ms, right TLE + uMTS: 132 ± 6 ms, p = 0.45) of the two TLE + uMTS subgroups in our study, more extensive white matter abnormalities were observed in patients with right TLE + uMTS in comparison with left TLE + uMTS. The discrepancy between our study and previous reports may be partly attributed to the tighter age range of our right TLE + uMTS patients (mean 42 ± 5 years, range 36–48 years) compared to the other two studies with a wider age range (22–54 years for the right TLE + uMTS group in (Focke et al., 2008) and 38 ± 11 years for right TLE group in (Kemmonsoti et al., 2011)). While many reports suggest more extensive structural changes in left TLE + uMTS, two recent studies using VBM of T1-weighted images or SPECT have demonstrated more extensive abnormalities in right TLE + uMTS (Pail et al., 2010; Tae et al., 2005). More frequent interictal hyperperfusion of the contralateral hippocampus on SPECT was seen in right TLE + uMTS as compared with left TLE + uMTS (Tae et al., 2005) and gray matter volume reduction was significantly more extensive in right TLE + uMTS than left TLE + uMTS (Pail et al., 2010). Given the conflicting observations in the literature more work is needed to better characterize the differences in the extent of structural changes associated with right and left TLE + uMTS.

5. Conclusions

In summary, the current study demonstrated widespread diffusion abnormalities of white matter tracts within and beyond the temporal lobe in patients with TLE and unilateral MTS while, in contrast, patients with non-lesional TLE showed fewer changes. Specifically, patients with TLE and right MTS showed a more extensive bilateral white matter disruption than patients with left MTS. In conjunction with previous findings of white and gray matter abnormalities reported by other independent investigators, there is considerable evidence to suggest different dysfunctional networks in TLE patients with and without MTS.

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