RESEARCH ARTICLE

Combined Ex Vivo 9.4T MRI and Quantitative Histopathological Study in Normal and Pathological Neocortical Resections in Focal Epilepsy

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Keywords
9.4T MRI, dysplasia, focal epilepsy, quantitative neuropathology.

Abstract
High-resolution magnetic resonance imaging (MRI) may improve the preoperative diagnosis of focal cortical dysplasia (FCD) in epilepsy. Quantitative 9.4T MRI was carried out on 13 cortical resections, representing pathologically confirmed FCD (five cases) and normal cortex. Quantitative immunohistochemistry for myelination (myelin basic protein/SMI94), neuronal populations [microtubule-associated protein 2 (MAP2), neurofilament (SMI31, SMI32), synaptophysin, NeuN, calbindin], reactive glia (GFAP), microglia (CD68) and blood–brain barrier permeability (albumin) was carried out in 43 regions of interest (ROI) from normal and abnormal white matter and cortex. MRI was spatially aligned and quantitative analysis carried out on corresponding ROI. Line profile analysis (LPA) of intensity gradients through the cortex was carried out on MRI and immunostained sections. An inverse correlation was noted between myelin/SMI94 and T1, T2 (P < 0.05) and T2* (P < 0.05; Spearman’s correlation) and a positive correlation between neuronal MAP2 and T1 (P < 0.005) and T2* (P < 0.05) over all ROI. Similar pathology–MRI correlations were observed for histologically unremarkable white matter ROI only. LPA showed altered gradient contours in regions of FCD, reflecting abnormal cortical lamination and myelo-architecture, including a preoperatively undetected FCD case. This study demonstrates the ability of quantitative 9.4T MRI to detect subtle differences in neuronal numbers and myelination in histologically normal appearing white matter and LPA in the evaluation of cortical dyslamination. These methods may be translatable to the in vivo detection of mild cortical malformations.

INTRODUCTION
A proportion of localized structural brain abnormalities and malformations causing epilepsy remain “occult” preoperatively, with current magnetic resonance imaging (MRI) technologies. These include cases of pathologically confirmed focal cortical dysplasia (FCD) type II (33) as well as milder malformations of cortical development [mild malformations of cortical development (MCD)] (3, 22). Advanced MRI techniques have the potential to improve the identification and characterization of these lesions. This improvement, however, is dependent on a thorough understanding of the cellular basis of these techniques, best achieved by correlating the lesion’s magnetic resonance (MR) and its pathology directly with each other (22). High-resolution MRI (7T and 9.4T) of tissue samples followed by histology has been utilized to study post-mortem (PM) cortical alterations in multiple sclerosis (17, 27), Alzheimer’s disease (14, 29), schizophrenia (18) as well variations in normal brain myelo-architecture and cyto-architecture (10, 16, 34) including epilepsy surgical samples (11, 12). Many of these previous studies have used traditional Nissl and myelin stains for pathological qualitative or quantitative analysis (10, 12, 16, 34). With the application of immunohistochemistry to specific cellular and structural components of cortical laminae and white matter, there is further potential to dissect out the cellular basis of MRI signal alterations in “cryptogenic” focal epilepsies. Furthermore, varying reactive pathological changes are frequently noted in surgical epilepsy specimens, including gliosis, inflammation and blood–brain barrier breakdown (1, 32), which could potentially also influence MRI signals and need to be accounted for.

Previous quantitative neuroimaging and pathology correlative studies have generally relied on the analysis of small regions of interest (ROIs) in tissue samples (9, 21), which is dependent on accuracy of co-registration between sample and MRI (8) and may not be representative of the entire pathology, particularly in more ill-defined lesions as mild MCD. Whole slide quantitative analysis...
methods (WSA) maximize the tissue areas analyzed, and we have recently shown this to be a more reliable quantitative application in the study of mild MCD (19). Confirmed pathology–imaging correlations based on ex vivo high-field MRI of tissue specimens have the potential to be translated to the in vivo characteristics of these lesions and ultimately to improve detection, and potentially delineate the boundaries of these lesions, prior to surgery (22).

We present a series of cases where high-field 9.4T MRI and quantitative analysis has been carried out on surgical samples from patients with focal epilepsy including histologically normal and dysplasia cases. We employed quantitative immunohistochemistry for a broad range of neuronal, glial, microglial, myelin and vascular markers, including the application of WSA analysis, in order to further explore the relationship between pathological features and MR signal change in epilepsy cortical resections. Our aims were to identify quantitative MRI (qMRI) measures that could be of predictive value in the preoperative evaluation of focal epilepsy, in particular FCD types and mild MCD.

METHODS

Case selection

The specimens were selected prospectively from adult patients undergoing surgical treatment for epilepsy at the National Hospital for Neurology and Neurosurgery, Queen Square, during the period 2011–2013. Twelve cases were studied after obtaining preoperative consent for use of surgical tissue, and the study has approval through the ethics committee of the UCL, Epilepsy Society Brain and Tissue Bank. To correlate these samples with a reference, normal control tissue, a further sample from a PM of a patient without cortical neuropathology or history of epilepsy was included. The clinical details including epilepsy history and preoperative MRI diagnosis are presented in Table 1. The preoperative suspected diagnosis included FCD subtypes (five cases), hippocampal sclerosis (1) and dysembryoplastic neuroepithelial tumor (1). The resections were from the frontal lobe (eight cases), temporal lobe (three cases), parietal lobe (one case) and occipital lobe (one case); the control PM tissue was from the temporal lobe.

Sample preparation

In each of the 13 cases, a section of cortex and underlying white matter of 5-mm thickness was sliced using a cradle as previously described (8) and fixed for a minimum period of 5 days in formalin. Either one or two such samples from each case were selected for 9.4T imaging, aiming to include macroscopically pathological and normal cortex as well as white matter in all cases. The sample was fixed rigidly in a tissue-tek® (Sakura, Tatcham, UK) cassette, which was held tightly within a plastic tube to minimize movement artifacts and filled with fomblin® perfluoro polyether (LC 08, Solvay Solexis, Milan, Italy) to avoid susceptibility artifacts. In a previous pilot study carried out on PM tissues, there were no adverse effects of the fomblin medium on tissue processing and immunostaining protocols.

MRI protocol

The samples were scanned on a 9.4T MR scanner (Agilent Technologies, Santa Clara, CA, USA) using a volume coil with a diameter of 33 mm (Rapid Biomedical GmbH, Rimpar, Germany). The applied overnight MRI protocol consisted of T2-weighted images and T1, T2, T2* and magnetization transfer ratio (MTR) maps. Multi-slice spin echo sequences were used for the acquisition of these data except for the T2* and MTR maps where multi-slice gradient echo sequences were applied. After manual global shimming and acquiring fast localization images, 12 slices were acquired using a field of view of 35 × 35 mm², an in-plane spatial resolution of 1.36 × 1.36 μm² and a slice thickness of 500 μm. Depending on the contrast, the echo time (TE) and the repetition time (TR) were:

(i) T2-weighted images: TE = 55 ms, TR = 4500 ms and the acquisition time (TA) was 96 minutes.
(ii) T1 maps: TE = 10 ms, TR = 0.3/0.5/0.8/1.3/2/4 s and TA = 145 minutes.
(iii) T2 maps: TR = 2 s, TE = 12.6/20/35/60 ms and TA = 102 minutes.
(iv) T2* maps: TR = 290 ms, TE = 5.2/7/9/11/13/20 ms, flip angle (FA) = 40° and TA = 75 minutes.
(v) MTR maps: TR = 260 ms, TE = 5 ms and FA = 40°. Two frequency offsets of the saturation RF pulse were used: 6 kHz (MT weighted) and 100 kHz (control). TA = 65 minutes.

The total scanning time for the entire protocol was approximately 8 h. T1, T2, T2* and MTR maps were obtained by processing the data using both an in-house developed script running under Matlab (Mathworks, Massachusetts, USA) and ImageJ (National Institutes of Health, Bethesda, MD, USA; http://imagej.nih.gov/ij/index.html).

Immunohistochemistry and quantitation on ROIs

The tissue samples were then routinely processed, sections cut at 7 μm and an immunohistochemistry panel applied as detailed in Table 2. All sections were assessed qualitatively for the pathology present on the scanned section as detailed in Table 3. Between three and four ROIs were outlined on each of the pathology sections to include normal, as well as pathological, white matter and cortex as detailed in Table 3. The histological sections were manually spatially aligned with the MRI-scanned images, matching anatomical landmarks in the section, corresponding ROIs identified and qMRI values for T1, T2, T2* and MTR calculated for each area. Quantitative evaluation of immunohistochemistry staining was carried out on neuropilform (SMI32), myelin basic protein (MBP/SMI94), microtubule-associated protein 2 (MAP2), synaptophysin, microglial marker CD68, albumin (for loss of integrity of blood–brain barrier) and GFAP (for astrogliosis) stained sections with Image Pro Plus software (Media Cybernetics, Buckinghamshire, UK). Using RGB thresholding, the immunostaining reaction was detected in all fields within the ROI, captured sequentially at ×40 objective (Nikon Eclipse microscope and camera). An average of 71 high power fields (SD 25) were captured for white matter ROI and 69 fields (SD 27) for gray matter. A mean labeling index (LI) was then calculated for each ROI. Repeat measurements were carried out on MAP2, SMI94 and GFAP section (between 12 and 30 ROIs), which showed reproducibility; intraclass correlation coefficients of 0.81, 0.67 and 0.88, respectively. NeuN was not included in quantitative analysis due to the noted variability in the intensity of staining with longer fixation.
Table 1. Clinical data of patients and fixation times.

Abbreviations: DNT = dysembryoplastic neuroepithelial tumor; EEG = electroencephalogram; FCD = focal cortical dysplasia; FS = febrile seizures; GS = generalized seizures; HS = hippocampal sclerosis; icEEG = intracranial EEG from depth electrodes; ILAE = International League Against Epilepsy; LPA = line profile analysis; MRI = magnetic resonance imaging; NA = not applicable; PM = post-mortem; qMRI = quantitative MRI; SE = status epilepticus; Sz = comprises focal seizures with and without loss of consciousness.

<table>
<thead>
<tr>
<th>Case</th>
<th>Age onset epilepsy (years)</th>
<th>Age at surgery (years)</th>
<th>Duration of epilepsy (years)</th>
<th>Seizure types</th>
<th>Gender</th>
<th>Preop MRI diagnosis</th>
<th>Localization of resection</th>
<th>Postoperative outcome</th>
<th>Fixation time* (days)</th>
<th>Used in study</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>8</td>
<td>31</td>
<td>23</td>
<td>Focal Sz, GS</td>
<td>F</td>
<td>Extensive malformation with subcortical changes</td>
<td>Right frontal</td>
<td>Reduced frequency of seizures</td>
<td>452 (T2, T2*)</td>
<td>qMRI</td>
</tr>
<tr>
<td>2</td>
<td>30 (FS—2 years)</td>
<td>54</td>
<td>24</td>
<td>Focal Sz</td>
<td>F</td>
<td>Left HS: left hemisphere smaller</td>
<td>Left temporal lobectomy</td>
<td>Not seizure free: new right HS</td>
<td>464 (T1)</td>
<td>qMRI</td>
</tr>
<tr>
<td>3</td>
<td>15</td>
<td>32</td>
<td>17</td>
<td>Focal Sz, GS</td>
<td>M</td>
<td>FCD in calcarine tissue</td>
<td>Left temporal lobectomy (icEEG guided)</td>
<td>Not seizure free</td>
<td>142 (T2, T2*)</td>
<td>qMRI</td>
</tr>
<tr>
<td>4</td>
<td>3</td>
<td>27</td>
<td>24</td>
<td>Focal S</td>
<td>F</td>
<td>Minor ventricular prominence</td>
<td>Left frontal resections (icEEG guided)</td>
<td>Psychiatric illness</td>
<td>152 (T1)</td>
<td>qMRI</td>
</tr>
<tr>
<td>5</td>
<td>25</td>
<td>58</td>
<td>33</td>
<td>Focal Sz</td>
<td>F</td>
<td>FCD in left middle frontal gyrus, extending to ventricular surface</td>
<td>Left frontal resection (icEEG guided)</td>
<td>Seizure free</td>
<td>378</td>
<td>qMRI</td>
</tr>
<tr>
<td>6</td>
<td>14</td>
<td>49</td>
<td>34</td>
<td>Focal Sz</td>
<td>M</td>
<td>Previous surgery for DNT residual tumor</td>
<td>Left frontal lobe</td>
<td>Seizure free</td>
<td>21</td>
<td>qMRI</td>
</tr>
<tr>
<td>7</td>
<td>11</td>
<td>23</td>
<td>12</td>
<td>Focal S, GS, SE</td>
<td>M</td>
<td>No FCD</td>
<td>Left frontal resection (icEEG guided)</td>
<td>Seizure free</td>
<td>10</td>
<td>qMRI</td>
</tr>
<tr>
<td>8</td>
<td>8</td>
<td>24</td>
<td>16</td>
<td>Focal Sz, GS</td>
<td>F</td>
<td>No specific abnormality</td>
<td>Left frontal resection (icEEG guided)</td>
<td>ILAE 1b (some minor seizures)</td>
<td>23</td>
<td>qMRI, LPA</td>
</tr>
<tr>
<td>9</td>
<td>16</td>
<td>32</td>
<td>16</td>
<td>Focal S</td>
<td>M</td>
<td>Blurring of the right frontal lobe</td>
<td>Right frontal, basal resection (icEEG guided)</td>
<td>Seizure free</td>
<td>29 (sample 1)</td>
<td>qMRI, LPA</td>
</tr>
<tr>
<td>10</td>
<td>7</td>
<td>28</td>
<td>21</td>
<td>Focal Sz</td>
<td>M</td>
<td>FCD in right supra-marginal gyrus</td>
<td>Right parietal (icEEG guided)</td>
<td>One seizure since operation</td>
<td>36 (sample 2)</td>
<td>qMRI, LPA</td>
</tr>
<tr>
<td>11</td>
<td>NA—no epilepsy</td>
<td>62 (PM control)</td>
<td>NA</td>
<td>NA</td>
<td>F</td>
<td>No cortical lesion, salivary gland tumor</td>
<td>NA</td>
<td>568</td>
<td>qMRI, LPA</td>
<td></td>
</tr>
<tr>
<td>12</td>
<td>18</td>
<td>30</td>
<td>12</td>
<td>Focal Sz, GS</td>
<td>M</td>
<td>Normal</td>
<td>Right frontal (icEEG guided)</td>
<td>ILAE 2b</td>
<td>122 (T2)</td>
<td>qMRI</td>
</tr>
<tr>
<td>13</td>
<td>11</td>
<td>23</td>
<td>12</td>
<td>Focal Sz, GS</td>
<td>M</td>
<td>Perinatal infarct/FCD IId</td>
<td>Right occipital lobe</td>
<td>Two seizures following operation—now seizure free for 8 months</td>
<td>132 (T2)</td>
<td>LPA</td>
</tr>
</tbody>
</table>

Seizure outcome is taken at the time point for evaluation in this study.*The fixation time is recorded as time from resection to MRI scan acquisition.
MAP2 and 40 mm² (range 13.5–84 mm²) for SMI94/MBP. The calculated for the entire outlined white matter.

A selected area then the entire image was processed and the LI intensity threshold for detection of immunostaining was preset in Neurofilament: phosphorylated Neurofilament: non-phosphorylated NeuN Neuronal marker Chemicon; 1/2000 LPA Myelin basic protein (SMI94) Myelinated axons Sternberger monoclonals; 1/5000 ROI, WSA, LPA NeuN Neuronal marker Chemicon; 1/2000 LPA Neurofilament: non-phosphorylated (SMI32) Pyramidal cell bodies (layer III, V; normal cortex), axons; in FCD type II, DN Neurofilament: phosphorylated (SMI31) Axons; in FCD type II, DN Synaptophysin Synaptic density Dako; 1/100 ROI, LPA MAP2 Neuronal cell body and dendrites Sigma; 1/5000 ROI, LPA Albumin Regions of blood–brain barrier breakdown Dako; 1/80 000 ROI, LPA GFAP Reactive cellular and chronic fibrillary gliosis Dako; 1/2500 ROI, LPA Nestin Acute gliosis/BC Chemicon; 1/8000 LPA Calbindin Interneurons (cortical layers II–III mainly) Abcam; 1/40 LPA CD68 Microglia Dako; 1/100 ROI, LPA CD34 Vasculature: tumor cells in LEAT, BC in FCD Leica/Novocastra; 1/50 Qualitative

*Indicates which markers were used in different studies.

times (20); CD34- and nestin-labeled sections were also not evaluated quantitatively due to the overall paucity of positively labeled cells.

Whole slide scanning

Based on initial findings from the ROI analysis, SMI94/MBP- and MAP2-stained sections from each case were selected for whole slide scanning analysis. Slides were scanned with a Leica SCN400F digital slide scanner (Leica Microsystems, Wetzlar, Germany); images were processed into a Pyramid TIFF file, stored on a fileserver and viewed and managed on the SlidePath Digital Image Hub (Leica). Analysis was carried out with Definiens Tissue Studio 3.6 (Definiens AG, Munich, Germany) as follows. The entire area of white matter was outlined on each image using a stylus pen on a touch-sensitive computer screen, drawn at a set distance of 0.5 mm from the cortical margin, to exclude layer VI neurons. The mean area was 29.4 mm² (range 7–63 mm²) for MAP2 and 40 mm² (range 13.5–84 mm²) for SMI94/MBP. The intensity threshold for detection of immunostaining was preset in a selected area then the entire image was processed and the LI calculated for the entire outlined white matter.

Line profile analysis (LPA) in the assessment of laminar cortex. Similar LPA of matched cortical regions was carried out on all MRI sequences.

Statistical analysis was carried out with SPSS (version 21 for Windows, IBM Corporation, Armonk, NY, USA) using nonparametric correlation tests for the comparison of MRI and pathology measures (Spearman’s correlation and Mann–Whitney test).

RESULTS

The pathological diagnosis on the sections selected for 9.4T was FCD IIa in one case, FCD IIb in two cases, FCD IIIId in one case and “pathology negative” (no focal lesional pathology) in nine cases; in three cases intracranial electrode (ICE) injury sites were also present (Table 3).

Quantitative pathology and MRI measurements

There were 43 ROI analyzed in all; 24 in white matter and 19 in cortex with 31 in histologically normal appearing areas and 12 in pathological areas. There were significant differences in quantitative pathology measures between normal cortical ROI and normal white matter ROI for GFAP (P < 0.05), SMI94/MBP (P < 0.001), MAP2 (P < 0.0001), CD68 (P < 0.05) and synaptophysin (P < 0.001) but not for neurofilament or albumin (Figure 1A). When these measures were compared between normal white matter ROI and FCD white matter ROI, a significant reduction was observed for SMI94/MBP (P < 0.05) (Figure 1B); although the number of FCD cases is small, this finding is in keeping with previous observations of reduced myelin in dysplastic region (28). There were no differences for normal cortical ROI and FCD cortical ROI. For qMRI measures, significant differences were observed for T1 (P < 0.01), T2 (P < 0.05), T2* (P < 0.0001) and MTR (P < 0.0001) between cortex and white matter ROI (Figure 1C), between normal white matter and FCD white matter.
Table 3. Neuropathological diagnosis represented in resections, the whole sections subjected to 9.4T MRI and regions of interest (ROI) for analysis.

<table>
<thead>
<tr>
<th>Case number</th>
<th>Underlying pathology diagnosis</th>
<th>Pathology represented</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Extensive FCD type IIB</td>
<td>FCD IIB, WM–FCD II, WM–normal, Cortex–FCD II</td>
</tr>
<tr>
<td>2</td>
<td>HS (ILAE type 1): no cortical lesion</td>
<td>Path negative, WM–normal, WM–normal, Cortex–normal</td>
</tr>
<tr>
<td>3</td>
<td>No lesion</td>
<td>Path negative, WM–normal, WM–normal, Cortex–normal</td>
</tr>
<tr>
<td>4</td>
<td>FCD IIA with PAS-positive dysmorphic neurons</td>
<td>FCD IIA, WM–normal, Cortex–normal, WM–FCD</td>
</tr>
<tr>
<td>5</td>
<td>FCD IIB</td>
<td>FCD IIB, WM–normal, Cortex–normal, Cortex–FCD (Few BC)</td>
</tr>
<tr>
<td>6</td>
<td>Dysembryoplastic neuroepithelial tumor</td>
<td>No tumor, WM–normal, Cortex–normal, WM–ICE injury and increased normal neurons</td>
</tr>
<tr>
<td>7</td>
<td>FCD IIA with PAS-positive dysmorphic neurons</td>
<td>Path negative, WM–normal, WM–normal, Cortex–normal</td>
</tr>
<tr>
<td>8</td>
<td>No lesion</td>
<td>Path negative, WM–normal, WM–normal, Cortex–normal</td>
</tr>
<tr>
<td>9</td>
<td>Focal FCD IIA</td>
<td>Path negative, WM–normal, WM–normal, Cortex–normal</td>
</tr>
<tr>
<td>10</td>
<td>FCD IIB</td>
<td>ICE injury, WM–normal, WM–normal, Cortex–ICE injury</td>
</tr>
<tr>
<td>11</td>
<td>Post-mortem control case. No FCD/pathology</td>
<td>Path negative, WM–normal, WM–normal, Cortex–normal</td>
</tr>
<tr>
<td>12</td>
<td>Focal FCD IIA</td>
<td>Path negative, WM–normal, WM–normal, Cortex–normal</td>
</tr>
</tbody>
</table>

Abbreviations: BC = balloon cells; DN = dysmorphic neurons; FCD = focal cortical dysplasia; HS = hippocampal sclerosis; ICE injury = intracranial electrode injury site; ILAE = International League against Epilepsy; MRI = magnetic resonance imaging; PAS = periodic acid schiff stain; WM = white matter.

*The age of the electrode injury site is recorded as the time between electrode implantation and resection of tissue.
for T1 \((P < 0.05)\) and T2* \((P < 0.01)\) (Figure 1D) and between normal cortex and FCD cortex for MTR \((P < 0.05)\).

**Quantitative pathology and MRI correlations**

**Myelin**

In SMI94/MBP-stained sections there was a significant inverse correlation between the LI and qMRI measures in the 43 ROI for T1 \((P < 0.005)\) (Figure 2A,C,D), T2 \((P < 0.005;\) Spearman’s correlation) (Figure 2B,E) and T2* \((P < 0.05)\); there was a positive correlation between SMI94/MBP and MTR \((P < 0.05)\). When considering normal white matter ROI alone, a significant inverse correlation between SMI94/MBP and both T1 and T2 was still observed \((P < 0.05)\). With WSA, there was also a significant inverse correlation between SMI94/MBP and T2 \((P < 0.05)\).

**Neuronal density**

In MAP2-stained sections, there was a significant positive correlation between the LI and qMRI measures over all 43 ROI for T1 \((P < 0.005)\) (Figure 3A,D,E) and T2* \((P < 0.05)\). When considering normal white matter ROI alone, there was also a positive correlation between MAP2 LI and T1 \((P < 0.05)\) (Figure 3B). With WSA for pathology-negative cases, there was a positive correlation between MAP2 LI and T2*, T2 \((P < 0.05)\) (Figure 3F) and an inverse correlation with MTR \((P < 0.01)\). There were no significant correlations noted for neurofilament LI and MRI measures.

When the MAP2 and SMI94/MBP LI were expressed as a ratio, over the 43 ROI there was a more significant positive correlation with T1 \((P < 0.0001)\), T2 \((P < 0.001)\), T2* \((P < 0.01)\) and negative correlation with MTR \((P < 0.01)\); with normal white matter ROI alone there was a positive correlation between the MAP2/SMI94 ratio with both T1 and T2 values \((P < 0.01)\) (Figure 3C).

**Reactive pathological changes**

There was a positive correlation between T2 measures in the normal cortical ROIs and GFAP LI \((P < 0.05)\). There was also a positive correlation between albumin labeling in normal appearing cortex and T1 measures \((P < 0.05)\). Albumin was qualitatively observed to stain the neuropil adjacent to vessels and some reactive astrocytes, as previously described (31). Correlations were not seen for GFAP or albumin in other ROI classes, and no significant correlations were noted for microglial marker CD68.
In three cases, localized ICE injuries were identified in the sections submitted for 9.4T; the age of the injuries, taken as the time from electrode insertion to tissue resection, were 3 days (case 6), 14 days (case 10) and 22 days (case 1). The high frequency of ICE injuries in this cohort is accounted for by the high proportion of cases that had undergone intracranial monitoring for “MRI-negative” epilepsy. The injury site was clearly visible as a small empty hypo-intense “cavity” in the cortex on MRI, and in the marginal zones on T2, there was impression of increased signal from 3- to 14-day-old injuries (Figure 4). Quantitative evaluation of CD68 showed a trend for increased CD68 \((P = 0.064\) Mann–Whitney) in these lesions compared with normal cortex, a decline in MAP2 \((P = 0.07\), Mann–Whitney) but no significant alteration in GFAP with days post-injury in this small group (Figure 4). There was a significant reduction in T2* values in injured compared with normal cortex \((P = 0.02)\) (Figure 4).

**Clinical correlations**

There were no significant correlations between MRI and pathology quantitative parameters in relation to age of epilepsy onset, duration of epilepsy or localization of the resection. There was also no effect of fixation times on MR measurements; there was an inverse correlation between fixation time and MAP2 and SMI32 staining \((P < 0.032\) Spearman’s) but not for the other markers, including SM194/MBP.

**LPA**

**Normal laminar cortex**

LPA of normal cortex and white matter with neuronal- and layer-specific markers are illustrated in case 8 (Figure 5); NeuN showed a peak in layer II corresponding to the high packing density of small neurons (Figure 5A,C) whereas neurofilament confirmed increased labeling in lower compared with superficial cortical layers (Figure 5B,C) with a small fluctuation in mid-cortical layers, reflecting the region between the laminae-like bands of labeling of the dendrites of the pyramidal neurons in layers III, V and VI with their high neurofilament content. Synaptophysin by contrast showed a flat labeling intensity profile through the cortex reflecting the more homogenous staining with a relatively sharp drop at the cortical–white matter boundary (Figure 5C). LPA for SM194/MBP showed increased intensity in deeper cortical layers corresponding to the radial fibers with a small wave in mid-layers (layer IV) likely reflecting horizontal myelinated fibers.
GFAP line profiles were highest in layer I and white matter corresponding to the Chaslin’s gliosis and white matter gliosis, respectively (Figure 5A,C). Calbindin showed lower intensities overall, reflecting low density of labeled neurons, but higher in outer cortical layers (I–III) compared with deeper cortex in keeping with cell distribution (Figure 5C). When ratios of microtubule-associated protein 2 : SMI94/MBP were calculated for, there was a significant positive correlation overall ROI for all MRI sequences (T1, T2, T2* and MTR; *P < 0.01 to *P < 0.0001 see text) and as shown here for normal white matter ROI alone with T2 (*P < 0.01) (C). (D) MAP2 stained section of case 2 (pathology negative; inset shows the single white matter neurons and processes within the normal white matter. (E) Case 2 with T1- and (F) T2-weighted MRI sequences at 9.4T. Bar = 5 mm in D (~300 μm).

Dysplastic cortex in FCD

LPA of dysplastic cortex with dyslamination is illustrated by case 4, 10 and 13 (Figure 6). In case 4, preoperative in vivo MRI was normal, resected frontal lobe was macroscopically unremarkable (Figure S4A) but FCD IIA confirmed histologically in the depth of one sulcus, with numerous dysmorphic neurons scattered within the cortex but absent balloon cells (Figure S4A). LPA carried out on the FCD region compared with adjacent normally laminated cortex showed lower overall intensity values for NeuN (Figure S4A) and altered line profile for MAP2 (Figure S4B,C) in FCD, correlating with the changes seen in LPA in MRI sequences confirmed the qualitative impression of increased signal intensity in outer cortical layers in T2 as previously described (12) (Figure 5D,G) and to a lesser extent in T1 (Figure 5F,H). In addition, waves in the LPA intensity were observed in mid-cortex (Figure 5H) and likely correspond to the profiles observed in neurofilament and myelin-stained sections in layer IV region. The MTR showed a more uniform, flat line profile through the cortex, changing at the white matter. LPA in further cases of normal cortex (Figures S1–S4b) supported these findings, with corresponding fluctuation in the profile intensity of T2 and T2* sequences in mid-cortex.

Dys laminar cortex in FCD

LPA of dysplastic cortex with dyslamination is illustrated by case 4, 10 and 13 (Figure 6). In case 4, preoperative in vivo MRI was normal, resected frontal lobe was macroscopically unremarkable (Figure S4A) but FCD IIA confirmed histologically in the depth of one sulcus, with numerous dysmorphic neurons scattered within the cortex but absent balloon cells (Figure S4A). LPA carried out on the FCD region compared with adjacent normally laminated cortex showed lower overall intensity values for NeuN (Figure S4A) and altered line profile for MAP2 (Figure S4B,C) in FCD, correlating with the changes seen in LPA in MRI sequences confirmed the qualitative impression of increased signal intensity in outer cortical layers in T2 as previously described (12) (Figure 5D,G) and to a lesser extent in T1 (Figure 5F,H). In addition, waves in the LPA intensity were observed in mid-cortex (Figure 5H) and likely correspond to the profiles observed in neurofilament and myelin-stained sections in layer IV region. The MTR showed a more uniform, flat line profile through the cortex, changing at the white matter. LPA in further cases of normal cortex (Figures S1–S4b) supported these findings, with corresponding fluctuation in the profile intensity of T2 and T2* sequences in mid-cortex.

Figure 3. Quantitative pathology and magnetic resonance imaging (MRI) correlations with microtubule-associated protein 2 (MAP2). There was a significant positive correlation with T1 in all 43 regions of interest (ROI) (P < 0.005, Spearman’s correlation) (A). There was a significant positive correlation with T1 in normal white matter regions of interest (P < 0.05, Spearman’s correlation) (B). When ratios of microtubule-associated protein 2 : SMI94/MBP were calculated for, there was a significant positive correlation overall ROI for all MRI sequences (T1, T2, T2* and MTR; *P < 0.01 to *P < 0.0001 see text) and as shown here for normal white matter ROI alone with T2 (*P < 0.01) (C). (D) MAP2 stained section of case 2 (pathology negative; inset shows the single white matter neurons and processes within the normal white matter. (E) Case 2 with T1- and (F) T2-weighted MRI sequences at 9.4T. Bar = 5 mm in D (~300 μm).
with reduced neuronal labeling and reflecting abnormal neuronal density and distribution, in keeping with previous stereological quantitative studies in FCD type II (30). Similarly, LPA of T2 and T2* (Figure S4A,B,C) also showed differences in intensities between the normal and FCD regions, but there were no obvious differences for T1 and MTR (Figure S4B,C). These observations on LPA were also reflected in the qMRI values for this region of FCD IIA that showed lower T2 and T2* values compared with both the adjacent cortex as well as mean values for normal cortex in the whole series. In case 13 with a perinatal infarct and FCD type IIId (Figure 6A,B), with severe abnormalities of the cortical myeloarchitecture and cyto-architecture (Figure 6E,G), LPA of T2-weighted images (Figure 6C) showed distinct lamination corresponding to the abnormal neuronal and myelin cortical layering.

LPA in a third case with dysplasia (FCD IIB) (case 10; Figure S5) also showed abnormal line profiles, with flatter contours observed on NeuN-, myelin- and neurofilament-stained sections as well as T2*, T2 and T1, reflecting abnormal cortical lamination and myelo-architecture.

**DISCUSSION**

Further technical advances in MRI, including increased field strength, improvements in image resolution and post-processing quantitative analysis, are likely to improve future detection of subtle FCD and mild malformations in epilepsy (22). Such advances, however, need to be validated with comparative neuropathology, still regarded as the gold standard in the diagnosis and

*Figure 4. Intracranial electrode (ICE) injuries.*
In three cases ICE injury sites were represented within the tissue blocks scanned at 9.4T magnetic resonance imaging. The ages of these ICE injuries (time from electrode insertion to the time of tissue resection) were 3, 14 and 22 days, representing acute to subacute stages of cellular organization. The position of the injury is arrowed in T2 image with evidence of intensity at the margin of the injury cavity noted. The ICE injury site is shown on the GFAP stain (with the cavity indicated by a black arrow), indicating the reactive cellular margin of astrocytes (shown in high magnification in the corresponding column on the right side). Graphical representation of values for the region of interest corresponding to the ICE injury is shown in the bar chart with an increase in GFAP and CD68 labeling index (LI), decline in microtubule-associated protein 2 LI and T2* alterations with age of the ICE injuries. (T2 and T2* values are shown at $10^1$). Bar equivalent to ~1000 μm in middle column GFAP and ~50 μm in right hand column GFAP).
classification of lesion-based epilepsies (3, 22). In this study we have aimed to address this need through correlating *ex vivo* 9.4T qMRI measures with quantitative immunohistochemistry. We have shown significant correlations between MRI measures and myelin and neuronal density measures, even in histologically “normal” appearing white matter. Such correlations show the potential of high-resolution MRI in the detection of FCD and mild MCD with subtle white matter pathological changes.
Figure 5. Line profile analysis (LPA) in normal cortex (case B). (A) Merged and superimposed images, recolored for illustrative purposes, of NeuN (blue), myelin (green) and GFAP (red); same area is shown in (B) labeled with neurofilament/SMI32 in gray scale. Gray scale monochrome images were used for LPA along vectors drawn at an axis perpendicular to the pial surface through all cortical layers (indicated with the red lines SMI32/neurofilament stain in B). (C) Graphical representation of LPA of mean intensities of labeling between the two vectors in this case; the x-axis represents the distance through the cortex scaled as 0 = pial boundary to 100 = cortical white matter boundary (shown in red on panel B and the approximate position of cortical layers, as measured with image pro plus on NeuN-stained sections, is shown at the bottom of graph in C in red text). Intensity of gray-scale signal is shown in the y-axis (note: the graphs have been inverted for illustration so higher y-value corresponds to lower gray scale/greater immuno-positive reaction). Synaptophysin shows a relatively flat intensity line profile through the cortex until the white matter junction; neurofilament (SMI32) shows greater immunostaining intensity in deeper compared with superficial cortex with irregularities in the line profile in the mid-layers (double arrows in B and C) reflecting the zones between the pyramidal cell dendrites in layers III and IV and V. The profile for MBP/SMI94 also shows increased intensity of immunostaining on LPA in deeper cortical layers corresponding to the radial fibers, with an impression of a small peak in mid-layers (arrow in A and C), which is likely detecting the horizontal myelinated fibers. NeuN LPA is low in cell-free layer I and shows peak intensity in LPA in layer II, corresponding to the greater neuronal packing density of small neurons, whereas the neurons in deeper cortical layers are more widely spaced and the line profile flattens off until the white matter. GFAP LPA intensities peaked in layer I (corresponding to the Cushing’s gliosis) and in the white matter. Calbindin showed lower overall intensities, due to the overall low density of immunopositive cells, but with highest intensities in outer cortical layers (I–III). MRI sequences are T2 (D); bar indicates region of LPA and shown in higher magnification in (G)], MTR (E) and T1 (F). (H) LPA of MR sequences in T2 and T2* and to some extent T1 show higher intensities of LPA in superficial cortical layers with a small peak in mid-layers on T2 (arrow) approximately corresponding to the region of altered intensity in the neurofilament and myelin sections in the region of layer IV.

FCD

Established in vivo MRI hallmarks of FCD type II include thickening of the cortex, a less distinct gray–white matter interface and increased signal intensity in T2/FLAIR, with reduced or hypointense T1 signals in the white matter (5, 22, 33). It is recognized that not all confirmed FCD II show these alterations and there is evidence for improved MRI detection of FCD with the application of quantitative methods (2, 33). The pathological correlate of white matter T2 and FLAIR signal abnormalities is considered to reflect reduced myelin content (3, 6). Hypomyelination of the white matter in the region of FCD II is a common, but not essential, histological feature (23), confirmed through quantitative immunohistochemistry in more subtle cases (23, 28). The etiology of white matter hypomyelination is less well understood; it may represent a developmental deficit in local oligodendrogial maturation and myelination or reduced density of normally myelinated axons in the region of FCD (23, 28). In the current study in non-FCD epilepsy cases, we demonstrated an inverse correlation between myelin immunolabeling and T2 over all ROI (including cortex and white matter) as well as a significant correlation when including only ROI from histologically unremarkable appearing white matter. This demonstrates the potential of 9.4T MRI to detect subtle differences in white matter myelination that could be implemented in the localization of occult FCD in patients with cryptogenic focal epilepsy as well as in the detection of other white matter myelination abnormalities recognized to occur in temporal lobe epilepsy (4, 11, 26). The present series also included a 3T “MRI occult” FCD II case, in which we confirmed a reduction in white matter myelin labeling and significant increase in T1 and T2* with 9.4T MRI in the dysplasia region with quantitative analysis compared with normal values.

Mild MCD

Mild MCD encompass subtle (presumed developmental) cortical and white matter abnormalities and remain one of the more controversial entities described in focal epilepsies, both in view of their preoperative detection, histological criteria, biology and significance (3). Nevertheless they may represent a “biomarker” for focal epilepsy. In this study we confirmed a significant correlation between MAP2 neuronal labeling in the white matter and quantitative T1 measures, supporting the ability of 9.4T MRI to detect pathologically elevated white matter neuronal numbers. Mild MCD type II is currently poorly defined as an “excess” of single neurons in the white matter (3). In a recent quantitative analysis of 142 temporal lobe epilepsy cases utilizing WSS methods (19) we confirmed higher white matter neuronal densities in epilepsy patients (2298 cells/mm³) compared with controls; however, the clinical significance of this finding, in terms of epilepsy control following surgery, remained unclear. In the current study we also implemented WSS, which we have shown a more efficient and reliable method for the quantitative analysis of larger ROI such as the white matter (19). We observed a correlation between white matter MAP2 neuronal labeling, T2*, T2 and MTR. This supports continued implementation and development of WSS technology for the rapid analysis of epilepsy lobectomy specimens. Indeed, limitations of the size of white matter ROI, which were necessary in our previous studies of in vivo 3T MRI/pathology correlations implementing stereology, may be one explanation for the lack of correlation between white matter neuronal number and qMRI then observed (9). Furthermore, in this current study we included MAP2 for the assessment of neuronal density as an alternative to previously employed NeuN marker. Threshold-based analysis of the area of MAP2 immunostaining, which labels both neuronal cell body and dendrites, as an alternative to more traditional neuronal density measurements based on nuclear NeuN labeling (9, 19), may prove a more sensitive marker in the evaluation of altered white matter neuronal populations. Furthermore, Mild MCD II-like pathologies are also recognized in other neurological conditions as schizophrenia, autism spectrum disorders (24, 35) as well as epilepsy and although its relevance, in terms of a prognostic marker, remains uncertain its in vivo detection may be important as a biomarker of abnormal neurodevelopment in these conditions.
Dyslamination (abnormal neuronal layering) is a hallmark of FCD and be utilized as a potential diagnostic technique for currently no neurological conditions. Schmierer et al (27), in a study of PM tissues from patients with multiple sclerosis using 9.4T MRI, showed an inverse correlation between MBP and T2 in the cortex and an inverse correlation between MBP and T1 in the white matter. Spencer et al (29), in a mouse model of AD, showed that T1 correlated better with myelin density than cell density in the cortex. In a study of PM AD brains, Gouw et al also noted that increased white matter myelin loss was associated with increased T2 signal (14). In previous studies of 7T MRI of temporal lobe dysplasia samples, areas of in-homogenous signal intensity in the white matter on T2 corresponded to fewer myelinated and unmyelinated axons on ultrastructural analysis (11). It is also plausible that the observed white matter microstructural alterations of reduced and abnormal white matter myelination (11, 23, 28) and increased neuronal density operate synergistically to influence MRI signal alterations. Indeed, in the current study, when we expressed MAP2 and myelin as a ratio, there was an augmented statistical association with qMRI measures.

LPA and dyslamination

We utilized LPA to compare intensity gradients in normal laminar cortex and regions with histologically proven FCD types. Dyslamination (abnormal neuronal layering) is a hallmark of FCD type II and the only pathological abnormality in FCD type I (3). It can be evident on cresyl violet (Nissl) stained sections, although it can be subtle with improved detection shown through application of cortical layer-specific markers (15, 25), some of which were applied in the current study. Although less recognized, the cortical myelo-architecture is also frequently altered in FCD, supported by previous quantitative studies (28) and is more dramatic in FCD type IIIa as exemplified by the case in this series. Clarification of dyslamination in in vivo MRI could facilitate detection of subtle FCD and be utilized as a potential diagnostic technique for currently “occult” patients with electro-clinically focal seizures. Previous MRI studies of human brain tissue samples have employed similar LPA methods. Fatterpekar et al. (10) imaged PM samples of the calcarine cortex at 9.4T and correlated this with Nissl and Luxol fast blue stains for myelin; they visualized bands of low signal in mid-layers on MRI images corresponding to the myelinated fibers in the external band of Baillarger. WALTERS et al also confirmed the ability of high-resolution MRI to detect cortical lamination in PM samples in regions outside the calcarine cortex (34) and noted that this correlated overall better with cortical myelo-architecture than cyto-architecture, as also in a study by Eickhoff et al (7). We were also able to visualize cortical lamination with MRI in a PM specimen taken from the temporal cortex, where the intracortical horizontal myelin plexus is less pronounced than in the visual striate cortex. Using similar LPA method, Garbelli et al (12) carried out an ex vivo 7T MRI of surgical epilepsy resections of FCD type IIIa to (a temporal cortical dysplasia adjacent to hippocampal sclerosis).

In normal cortex, they noted intensity differences on T2 between the upper and lower cortex separated by a dip of lower signal intensity correlating with cortical myelination patterns and the neuronal density of layer IV on NeuN sections; in FCD IIIa cases, however, this dip in mid-layer intensity was lost and the T2 line profile through the cortex was flatter without a clear difference between supragranular and infragranular cortex (12). They concluded that altered LPA in dysplasia is strongly correlated with abnormal cortical myelination patterns.

With LPA we confirmed similar intensity profiles for T1 and T2 in normal cortex from varied cortical locations, with NeuN, neurofilament and MBP markers more consistently revealing contours and linear gradients across cortical layers and correlating with MRI line profiles. Furthermore, we observed differences in LPA on 9.4T MRI in histologically confirmed FCD II and FCD IIId reflecting the abnormal lamination, altered neuronal populations and abnormal myelo-architecture. These studies support the potential application of LPA techniques in the identification and discrimination of FCD types.

Reactive and secondary cortical changes

Gliosis is a ubiquitous finding in surgical resections from patients with epilepsy but of unknown cause. Furthermore, inflammatory changes and blood–brain barrier breakdown are reported that could influence MRI appearances (1). In our quantitative analysis of normal cortex, we noted a positive correlation between T2 and gliosis (GFAP) and albumin leakage and T1, suggesting that 9.4T MRI signals may be influenced by subtle reactive cellular changes.

There is a great incentive to understand processes of gliosis and brain repair responses, both the pathological mechanisms and MRI correlates, in particular the time course of early cellular changes, reparative neuro-glial interactions as well as identifying any maladaptive, detrimental effects. We have previously used ICE “injuries” in patients with epilepsy as a model to study the time course of cellular responses and gliogenesis (13). In the current series we had three injuries of varying ages (acute to subacute) and although a small number, quantitative histological analysis confirmed the expected microglial responses accompanied by progressive loss of...
neurons and dendrites as highlighted with a reduction in MAP2 and a marginal astroglial reaction. Despite the gliotic margin and the qualitative impression of a rim of increased signal intensity around the injury cavity we noted a significant fall in T2* with days post-injury compared with normal cortex. Although we have only three samples, it is likely that the T2* measure in these cases reflects the overall loss of neurons rather than the cellular glial reaction. Nevertheless, these cases highlight the potential of 9.4T MRI in the evaluation of this “model” of cortical injury; by including more ROI, representing zones of injury from the necrotic center to the penumbra as well as greater number and age of injuries, the sequence of cellular alterations in brain repair and imaging alterations could be teased out.

In conclusion, this study demonstrates the feasibility of quantitative 9.4T MRI to detect subtle differences in cortical and white matter neuronal numbers and myelination in histologically normal appearing cortex and white matter and LPA as a useful tool in the detection of subtle cortical dyslamination. These methods have the potential to be translated to the in vivo detection of mild MCD and occult FCD cases in epilepsy.

ACKNOWLEDGMENTS

We are grateful to Derek Marsdon and Andrew Beckett in the Department of Neuropathology for their invaluable expertise in the processing and preparation of the epilepsy tissue samples for this study. This study was supported by the Medical Research Council, UK (Grant no. MR/JO127OX/1). This work was undertaken at the University College London and University College London Hospital, who receive part of their funding from the Department of Health Biomedical Centre scheme.

CONFLICT OF INTEREST

All authors have no conflicts of interest to declare.

REFERENCES


SUPPORTING INFORMATION

Additional Supporting Information may be found in the online version of this article at the publisher’s web-site:

Figure S1. Line profile analysis (LPA) for case 6 (no cortical dyslamination).
Figure S2. Line profile analysis (LPA) for case 9 (no cortical dyslamination).
Figure S3. Line profile analysis (LPA) for case 11 (post-mortem sample with no dyslaminatation).
Figure S4. (A) Line profile analysis (LPA) in FCD IIA (case 4).
Figure S5. Line profile analysis (LPA) for case 10 in FCD IIB.