Advanced $^{[18}F]FDG$ and $^{[11}C]Flumazenil$ PET analysis for individual outcome prediction after temporal lobe epilepsy surgery for hippocampal sclerosis

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Purpose: We have previously shown that an imaging marker, increased periventricular $^{[11}C]Flumazenil$ ($^{[11}C]FMZ$) binding, is associated with failure to become seizure free (SF) after surgery for temporal lobe epilepsy (TLE) with hippocampal sclerosis (HS). Here, we investigated whether increased preoperative periventricular white matter (WM) signal can be detected on clinical $^{[18}F]FDG$-PET images. We then explored the potential of periventricular $^{[18}F]FDG$ WM increases, as well as whole-brain $^{[11}C]FMZ$ and $^{[18}F]FDG$ images analysed with random forest classifiers, for predicting surgery outcome.

Methods: Sixteen patients with MRI-defined HS had preoperative $^{[18}F]FDG$ and $^{[11}C]FMZ$-PET. Fifty controls had $^{[18}F]FDG$-PET (30), $^{[11}C]FMZ$-PET (41), or both (21). Periventricular WM signal was analysed using Statistical Parametric Mapping (SPM8), and whole-brain image classification was performed using random forests implemented in R (http://www.r-project.org). Surgery outcome was predicted at the group and individual levels.

Results: At the group level, non-seizure free (NSF) versus SF patients had periventricular increases with both tracers. Against controls, NSF patients showed more prominent periventricular $^{[11}C]FMZ$ and $^{[18}F]FDG$ signal increases than SF patients. All differences were more marked for $^{[11}C]FMZ$. For individuals, periventricular WM signal increases were seen at optimized thresholds in 5/8 NSF patients for both tracers. For SF patients, 1/8 showed periventricular signal increases for $^{[11}C]FMZ$, and 4/8 for $^{[18}F]FDG$. Hence, $^{[18}F]FDG$ had relatively poor sensitivity and specificity. Random forest classification accurately identified 7/8 SF and 7/8 NSF patients using $^{[11}C]FMZ$ images, but only 4/8 SF and 6/8 NSF patients with $^{[18}F]FDG$.

Conclusion: This study extends the association between periventricular WM increases and NSF outcome to clinical $^{[18}F]FDG$-PET, but only at the group level. Whole-brain random forest classification increases $^{[11}C]FMZ$-PET’s performance for predicting surgery outcome.

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1. Introduction

Around one-third of patients with epilepsy are refractory to medical treatment, and epilepsy surgery is an efficient therapeutic alternative for some of these patients. Resection surgery is indicated in some focal epileptic syndromes, including mesial temporal lobe epilepsy (mTLE), a common form of adult epilepsy that is often resistant to anti-epileptic drugs (AEDs) and frequently associated with hippocampal sclerosis (HS) (Engel et al., 2003; Spencer and Huh, 2008). Nevertheless, about one-third of patients with HS are not seizure free (NSF) after epilepsy surgery. There is therefore a need of finding relevant biomarkers to anticipate the risk of suboptimal surgery outcome.

The presence of unilateral HS on presurgical magnetic resonance imaging (MRI) ipsilateral to surgery predicts a favourable, i.e. seizure free, outcome (Jeong et al., 2005), as does focal hypometabolism on $^{[18}F]fluorodeoxyglucose$ positron emission tomography (FDG-PET) ipsilateral to affected temporal lobe (Malagren and Thom, 2012). Other potential biomarkers have been investigated for the prediction of TLE-HS surgery outcome and findings are not always consistent (Aull-Watschinger et al., 2008; Janszy et al., 2005). For example, analysis of the ictal semiology as recorded by scalp video-EEG has shown that extra temporal features do not correlate with surgical outcome in
Table 1
Clinical data for 16 patients with refractory mTLE and HS. Upper half: seizure free patients; lower half: non-seizure free patients. Imaging based predictors for seizure status are colour-coded, with green indicating that the particular marker correctly predicted the individual patient’s outcome class according to the method used in that column (see text for details).

| Patient | Age/sex | Onset/duration of epilepsy (years) | HS | Follow-up (months) | Medication at operation: * reduced; ** stopped (medication restarted) | Outcome (Engel class) | FDG hypometabolism on SPM | [11C]FMZ-PET increases in periventricular areas (threshold = 0.01, $z > 3.2, p < 0.046$) | [18F]FDG-PET increases in periventricular areas (threshold = 0.01, $z > 3.2, p < 0.05$) | RF prediction (FMZ) | RF prediction (FDG) | Additional information |
|---------|---------|-----------------------------------|----|--------------------|---------------------------------------------------------------------|----------------------|--------------------------|--------------------------------------------------------------------------------|--------------------------------------------------------------------------------|-----------------|-----------------|-----------------------------------------------|
| #1 24/f | 6/18    | L 31 LEV, LTG                      | IA Yes | None | L ant front lobe $z = 3.89 (-14 28 2), 3.2 cm³ | True False | Temporal-occipital dysplasia |
| #2 22/f | 2/20    | L 58 CEZ, TPM, CLB                | IA Yes | None | None | False False | |
| #3 27/m | 4/23    | L 35 CEZ, VPA, CLB                | IA Yes | None | None | False False | |
| #4 36/f | 3/33    | L 98 CEZ, LTG, TPM                | IA Yes | None | None | True True | |
| #5 46/f | 12/34   | L 52 PR, PRG, LEV                 | IA No | None | R ant front lobe $z = 3.24 (4 22 -4), 2.4 cm³ | True True | L temporal pole signal abnormalities + discrete dysgenesis of left amygdala |
| #6 26/f | 5/21    | R 112 CEZ, CLB, PB                | IA No | None | R lat of ventricle $z = 4.18 (42 -36 8), 2.2 cm³ | True False | L temporal pole signal abnormalities |
| #7 48/f | 6/42    | R 81 CEZ, CLB                      | IA Yes | None | None | True True | |
| #8 30/f | 5/25    | R 24 CEZ, TPM, CLB, LEV           | IA Yes | None | None | True False | |
| #9 46/f | 33/13   | R 23 PH, LEV, PB                  | IB Yes | None | Lateral of posterocentral portion of R lateral ventricle $z = 3.76 (30-34 22), 1.6 cm³ | True False | |
| #10 36/f | 3/33   | R 116 CEZ, PR, CLB                | IVB No | None | None | True True | Seizure onset right temporal and right insular on SEEG |
| #11 29/f | 4/25    | R 44 CEZ, CLB                      | IIA Yes | None | None | True True | Small symmetric hippocampi and temporal pole signal abnormalities on MRI (FDG, SEEG unilateral) |

(continued on next page)
| Patient | Age/sex | Onset/duration of epilepsy (years) | HS | Follow-up (months) | Medication at operation: \(^*\): reduced. \(^\dagger\): stopped (medication restarted) | Outcome (Engel class) | FDG hypometabolism on SPM | \[^{[1]}\]C]FMZ-PET increases in periventricular areas (threshold = 0.01, \(z > 3.2, p < 0.05\)) | \[^{[1]}\]F]DG-PET increases in periventricular areas (threshold = 0.01, \(z > 3.2, p < 0.05\)) | RF prediction (FMZ) | RF prediction (FDG) | Additional information |
|---------|---------|----------------------------------|----|-------------------|-------------------------------------------|----------------------|-------------------------|---------------------------------|---------------------------------|------------------------|------------------------|---------------------|
| #12    | 32/m   | 5/27                             | L  | 119               | PB, CLB \(^\dagger\) (LEV, CLB)           | IIA                  | No                     | Widespread WM increases with the maximum L ant of frontal horn \(z = 6.02 \pm 34 -4\), 15.3 cm\(^3\) but extending bilaterally with other maxima anterior to both frontal horns and postero-lateral of right lateral ventricle. Lateral of posteroentral portion of L lateral ventricle \(z = 4.62 \pm 34 -44, 3.7 \pm 3\) cm\(^3\). Lateral of posterior portion of R lateral ventricle \(z = 4.12 \pm 30 -60, 6.5 \pm 3\) cm\(^3\) | Widespread WM increases with the maximum R lat ventricle \(z = 5.62 \pm 30 -34 20\), 5.02 cm\(^3\) but extending bilaterally with other maxima to the lateral ventricle \(z = 4.92 \pm 36 -42, 7.0 \pm 3\). R ant front lobe \(z = 3.73 \pm 26 32 10\), 2.2 cm\(^3\) | True                   | True                   |                      |
| #13    | 32/f   | 11/21                            | L  | 89                | CLB, CBZ \(^\dagger\) (CBZ, PB)           | IID                   | No                     | Lateral of anteroentral portion of R frontal horn \(z = 3.27 \pm 24 12, 4.0 \pm 3\) cm\(^3\). L anter of frontal horn \(z = 3.82 \pm 18 -36, 5.0 \pm 3\) cm\(^3\) | R anterior of frontal horn \(z = 3.24 \pm 24 -6, 1.9 \pm 3\) | False                  | True                   |                      |
| #14    | 22/m   | 9/13                             | L  | 72                | CBZ, TPM, PB, CLB \(^\dagger\)            | ID                    | Yes                    | Lateral of posteroentral portion of L lateral ventricle \(z = 4.51 \pm 34 -48, 1.7 \pm 3\) cm\(^3\). L anterior of frontal horn \(z = 3.36 \pm 14 38 -2\), 1.7 cm\(^3\) | L ant front lobe \(z = 5.79 \pm 12 -32, 5.5 \pm 3\) cm\(^3\). L sup cella media post horn \(z = 4.7 \pm 32 -50, 3.6 \pm 3\) | True                   | True                   |                      |
| #15    | 39/m   | 1/38                             | L  | 67                | TPM, PB \(^\dagger\)                       | IIA                   | No                     | Lateral and inferior of L ventricle where temporal and occipital horns originate \(z = 4.01 \pm 30 -52, 3.0 \pm 3\) cm\(^3\). Lateral and inferior of R ventricle \(z = 3.29 \pm 10 6, 2.4 \pm 3\) cm\(^3\) | Lateral posteroentral portion of R lateral \(z = 5.11 \pm 24 -38 4, 7.3 \pm 3\) cm\(^3\). L occipital horn \(z = 4.648 \pm 28 -50 6, 4.4 \pm 3\) | True                   | False                  |                      |
| #16    | 41/f   | 3/38                             | R  | 78                | LTG, CBZ, PB \(^\dagger\)                 | IC                    | No                     | None                                    | None                                    | True                   | True                   | Moderated gliosis R temporal pole, L ventricular enlargement |
patients with temporal lobe epilepsy and HS (Borelli et al., 2008). However, other studies have found extratemporal features as relevant prognostic factors for predicting outcome after TLE-HS surgery (Ferrari-Marinho et al., 2012; Najm et al., 2013; Ryvlin and Kahane, 2005). Hence the search of prognostic features is important for identifying reliable predictors of suboptimal outcome as a tool for patient management and counselling, including for avoiding invasive EEG prior to surgery when possible.

PET has long been used for helping to localize the seizure-onset area, especially in TLE (Engel et al., 1990). The most common PET tracer used is [18F]FDG, which measures regional glucose metabolism (Henry et al., 1993). [11C]Flumazenil (FMZ) which binds to the benzodiazepine sites of GABA receptors has been used for predicting surgery outcome in mTLE-HS. Previous studies have shown that increased preoperative periventricular white matter (WM) FMZ binding was associated with NSF outcome after temporal lobe resection for HS. This was shown at the group level using volume-of-distribution images (Hammers et al., 2005) and, with a larger control group, at the individual level using summed images (Yankam Njiwa et al., 2013). The tentative explanation for this association is that increased periventricular binding relates to heterotopic neurons, i.e. a mild form of migration disorder not visible on MRI (Hammers et al., 2003).

Furthermore, in a number of papers, investigation of WM [11C]FMZ volume-of-distribution (VT) by voxel-based analysis has shown a strong correlation between WM [11C]FMZ VT and the number of heterotopic neurons histologically determined in the WM (Hammers et al., 2001; Hammers et al., 2002). PET analysed with SPM detected periventricular [11C]FMZ VT increases in MRI-negative patients with extratemporal epilepsy which were analogous to such increases when analysing MRI visible periventricular heterotopia (Hammers et al., 2003).

In the present study, we first investigated whether increased preoperative WM FDG uptake could also be used for predicting epilepsy surgery outcome. We then assessed if an advanced machine learning technique (random forests; RF; Breiman, 2001) could predict seizure outcome based on voxel-wise signal intensities from preoperative FDG and FMZ whole brain images. Finally, both methods were compared.

2. Material and methods

2.1. Participants

Patients were retrospectively identified for the study and fulfilled the following criteria: 1) suffering from mTLE, 2) HS diagnosed on MRI (confirmed by histopathology for thirteen of the patients), 3) temporal lobe surgery with a postsurgical follow up for at least 12 months, 4) availability of preoperative [18F]FDG PET and [11C]FMZ PET on the Siemens/CT HR+ PET camera installed at the CERMEP imaging centre.

Patient details have been described in a previous study on [11C]FMZ PET alone (Yankam Njiwa et al., 2013). Briefly, sixteen patients (twelve women) fulfilled the criteria. Nine had left-sided HS and six right-sided HS, diagnosed on MRI by an experienced radiologist and confirmed by an experienced neurologist. Outcome was assessed according to Engel’s classification (Engel, 1987) by an epileptologist blinded to the imaging results (PR), for the time point of assessment. Eight patients (50%) were entirely seizure free (SF; Engel class IA) and eight were not (NSF; not Engel class IA either continuously or at some stage during follow-up, at least 4 months after surgery) (Takahashi et al., 2012). Table 1 shows the clinical characteristics of the patients.

Data were available for the following non-imaging potential outcome predictors: age at onset, duration of epilepsy, age at operation.

Fifty control subjects with a mean age of 35 ± 10 years were available for this study. Forty-one had [11C]FMZ PET scans (20 men and 21 women), and 30 had [18F]FDG scans (21 men, 9 women). Twenty patients had both FMZ and FDG-PET scans. All images were acquired on the same camera. All controls had normal MRI and were not treated with any regular medication.

All subjects had signed written informed consent at the time of PET scanning in agreement with the French legislation and after approval by the local Ethics committee.

2.2. PET data acquisition

Three-dimensional PET scans were acquired on an Exact ECAT HR+ scanner (Siemens, Erlangen, Germany) with a full width half maximum (FWHM) spatial resolution of about 5 mm3 (Brix et al., 1997). A 68Ge transmission scan was acquired for attenuation correction.

FMZ (RO 15-1788) was radio labelled with 11C via methylation (Maziere et al., 1984). Unlabelled FMZ (0.01 mg/kg) and [11C]FMZ (2.775 MBq/kg) were injected simultaneously (Delforge et al., 1995), and twelve frames of increasing length were acquired over 55 min (for details, see (Yankam Njiwa et al., 2013)). Summed [11C]FMZ radioactivity images were obtained from the attenuation and scatter corrected dynamic 3D series over the last five frames (20–55 min post injection), as these have been shown to yield particularly reliable measures (Bouvard et al., 2012).

The protocol for FDG-PET imaging consisted of a simultaneous injection of an 18.5 MBq load dose in addition to 1.5 MBq/kg dose of [18F]FDG followed by the acquisition of static images 40–50 min post injection.

2.3. PET data analysis

[11C]FMZ summed radioactivity images and static [18F]FDG images were first analysed using statistical parametric mapping (SPM8; Wellcome Department of Imaging Neuroscience, UCL, London, UK; http://www.fil.ion.ucl.ac.uk/spm) and specific periventricular masks (Hammers et al., 2005; Yankam Njiwa et al., 2013). Voxel-wise PET signal intensities across the whole brain were then used for RF classification.

For group analyses, the images of the seven right HS patients (3/8 SF and 4/8 NSF) were right–left reversed prior to spatial normalization to lateralize the epileptogenic side to the left for all patients.

In-house [11C]FMZ and [18F]FDG templates in the standard Montreal Neurological Institute (MNI) space based on over thirty five healthy subjects scanned on the same scanner were available. Because a proportion of the raw images had been left–right reversed, these templates were symmetrized for group analyses and RF classification as described earlier (Didelot et al., 2010). All images of the study were spatially normalized (Ashburner and Friston, 1999) to the corresponding symmetric template for the appropriate tracer (either [11C]FMZ or [18F]FDG). The resulting spatially normalized images were smoothed using an isotropic Gaussian kernel of 10 mm FWHM.

For individual analyses with SPM8, the original, asymmetric [11C] FMZ and [18F]FDG templates were used, i.e. individual images were not flipped. The preprocessing was otherwise identical.

Global means, i.e. the average radioactivity across the brain determined within a mask consisting of the voxels with intensity above 1/8 of the mean of all voxels in the mask, were compared between SF patients, NSF patients and controls using two-tailed student t-tests.

2.3.1. SPM analyses of periventricular WM increases

For SPM8 analysis, entire WM was extracted through combination of two masks and the assessment of periventricular regions was performed using a mask constructed with basic SPM8 functions, for more details see (Hammers et al., 2005; Yankam Njiwa et al., 2013). A covariance group specific analysis (ANCOVA) model was utilized for global radiactivity correction.

Periventricular increases were assessed within the mask introduced above in the group and individual analyses. We aimed to replicate the earlier finding of periventricular WM increases of [11C]FMZ binding in patients with suboptimal outcome relative to SF patients and controls, using the much more widely used PET tracer [18F]FDG.
At the individual level, each patient was compared against all 30 \( [^{18}F]FDG \) controls in order to investigate whether preoperative periventricular increases can be used to predict postoperative outcome for a given patient. Several combinations of the characteristics of clusters with increased periventricular WM signal were evaluated for the best separation of SF and NSF patients, namely “uncorrected cluster probability” and “z value of peak voxel” (Yankam Njiwa et al., 2013).

A given search space between 0.00 and 0.99, with moving steps of 0.01, for cluster probability and a level of significance of the peak voxel z score at any value above 3.2, step of 0.02 was set. The ability to optimally discriminate SF and NSF was tested for all combinations and after screening the cluster p values.

### 2.3.2. SPM analyses of mesial temporal \( [^{18}F]FDG \) hypometabolism

Presence of focal hypometabolism on presurgical FDG-PET ipsilateral to affected temporal lobe has been shown to predict favourable surgical outcome (Malmgren and Thom, 2012). We have therefore investigated the significance of mesial temporal lobe \( [^{18}F]FDG \) hypometabolism as a predictor of postsurgical outcome in this study. Each patient was compared against all 30 \( [^{18}F]FDG \) controls and clusters with a peak voxel z score \( \geq 3.2 \) and a p value \( \leq 0.5 \) were considered significant.

### 2.3.3. Random forest classification based on whole-brain PET signal intensities

We investigated the capabilities of random forest (RF) classifiers to distinguish patients who became SF and those who remained NSF from each other and from controls based on voxel-wise signal intensities across the full brain volumes for both \( [^{11}C]FMZ \) and \( [^{18}F]FDG \) PET images.

Each subject served as a test dataset for the training set composed of the remaining subjects acquired with each tracer. Feature data from images acquired with each tracer were independently used as inputs to an RF classifier.

RF is an ensemble classifier, consisting of many decision trees, which outputs the class that is the result of combining the prediction by individual trees (Breiman, 2001). Random forests combine bootstrap aggregation (bagging) (Breiman, 1996) and random feature selection (Amit and Geman, 1997; Ho, 1998) to construct a collection of decision trees exhibiting controlled variations. The training set for each individual tree in a random forest is constructed by sampling N examples (subjects) at random with replacement from the N available examples in the dataset. After drawing N bootstrap samples from the original data, an ‘unpruned’ decision tree is grown as follows: at each node, rather than choosing the best split among all predictors, randomly sample \( n_{mp} \) of the predictors and choose the best split from among those variables. New data are classified by aggregating the predictions of the \( N_{tree} \) trees, in the case of classification considering the majority of votes.

Tree nodes are split based on the Gini index (Breiman et al., 1984), which measures the likelihood of an example to be incorrectly labelled if it was randomly classified according to the distribution of labels within the node. A measure of the importance of an individual feature for classification can be computed by summing the decreases in the Gini index occurring at all nodes in the forest, which are partitioned based on that feature.

For the classification results presented in this study, the implementation of random forests in the statistical software package R, a port of Leo Breiman and Adele Cutler’s original Fortran code by Andy Liaw and Matthew Wiener, version 4.6–6 (http://cran.r-project.org/web/packages/randomForest), was used.

At each forest were grown \([1000:step:10,000]\) trees in steps of 1000. The datasets were first tuned, with the specific class implemented in R (rftune), to seek for the optimal number of features (computing the smallest error value) to be selected at each tree node according to the number of available features. The number of features thus computed at each node was 714 for FMZ and 15 for FDG.

### 3. Results

### 3.1. Global values

The global values of the summed radioactivity images differed significantly between the NSF and controls groups before spatial normalization for FDG. After spatial normalization, i.e. on the images used for the present analyses, these values did not differ significantly \( (p_{\text{min}} = 0.11) \). Table 2 illustrates the results.

### 3.2. Group analyses

#### 3.2.1. NSF versus SF patients restricted to WM periventricular signal increases

Higher periventricular ligand uptake was visualized in NSF patients in comparison to SF patients for both areas. Areas of high \( [^{18}F]FDG \) binding were observed around the posterior horn of the ventricles contralateral to the seizure focus \( (z_{\text{max}} = 3.37, 0.4 \text{ cm}^2) \) and ipsilaterally \( (z_{\text{max}} = 3.66, 1.1 \text{ cm}^2) \). \( [^{18}F]FDG \) uptake was increased at very similar locations as \( [^{11}C]FMZ \) uptake (Yankam Njiwa et al., 2013); see Fig. 1a.

There were no significant differences between SF and NSF patients for other potential predictors of postoperative outcome. Results are illustrated in Table 3.

#### 3.2.2. Patient groups against controls restricted to WM periventricular signal increases

As expected, SF patients compared with controls showed a smaller number of periventricular clusters, which were also less significant, than NSF patients for \( [^{18}F]FDG \). \( [^{18}F]FDG \) showed a bigger cluster with a higher z score \( (z = 4.16, 2.1 \text{ cm}^2) \) than \( [^{11}C]FMZ \) had in our previous study (Fig. 1b).

The group of NSF patients compared with controls showed increased FDG uptake in four clusters with higher z scores than SF patients: two ipsilaterally \((8.6 \text{ and } 1.3 \text{ cm}^2, z_{\text{max}} = 4.19)\) and two contralaterally \((5.2 \text{ and } 1.2 \text{ cm}^2, z_{\text{max}} = 4.68)\). Similarly, using \( [^{11}C]FMZ \), more clusters of periventricular signal increases with higher z scores had been found in NSF patients versus controls, compared with SF patients against controls (Yankam Njiwa et al., 2013, Fig. 1c).

#### 3.2.3. Random forest classifier analyses of the entire brain volume

Random forest classifiers were independently applied to the available feature data corresponding to all available voxels \((513,340 \text{ voxels})\) for both radiotracers.

#### 3.2.4. NSF versus SF patient groups

For both FDG-PET and FMZ-PET, RF classification was not able to distinguish between NSF and SF patients. Performance results are

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**Table 2**

|               | NDG | SF      | Controls |
|---------------|-----|---------|----------|
| Before normalization | 2064 ± 946 (p = 0.03) | 2290 ± 986 (p = 0.09) | 2998 ± 652 |
| After normalization | 3268 ± 1807 (p = 0.22) | 3233 ± 1360 (p = 0.11) | 4154 ± 932 |

|               | FMZ | SF      | Controls |
|---------------|-----|---------|----------|
| Before normalization | 1492 ± 300 (p = 0.01) | 1562 ± 207 (p = 0.02) | 1778 ± 256 |
| After normalization | 2183 ± 395 (p = 0.17) | 2202 ± 279 (p = 0.09) | 2409 ± 335 |
illustrated in Table 4. FMZ better performed than FDG for the classification of subjects, except for the NSF/SF comparison in terms of sensitivity.

### 3.2.5. Patient groups against controls

For the comparison between the group of patients and controls, RF classifiers well correctly predicted 7/8 NSF using FMZ images and 6/8 with FDG. FMZ binding was more sensitive in differentiating SF to from controls (7/8 well correctly predicted) than FDG (6/8 well correctly predicted).

With FMZ data, important features for distinguishing NSF patients from controls consisted of more widespread clusters detected principally in the hippocampus, whereas for SF patients those were somewhat more concentrated in the amygdala and in the temporal lobe. Results are shown in Fig. 2.

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**Fig. 1.** Periventricular increases and unmasked statistical results overlaid onto the SPM8 T1-weighted template. FDG results obtained for this study; FMZ results from Yankam Njiwa et al. (2013) and shown for direct comparison. **(a)** Increases in non-seizure free patients (NSF; not Engel IA) compared to seizure-free patients; ipsilateral, left side; \(z_{\text{max}} = 2.69, \text{CE} = 0.56 \text{cm}^3\) and contralateral \(z_{\text{max}} = 4.2, \text{CE} = 0.9 \text{cm}^3\) for FMZ and \(z_{\text{max}} = 3.66, \text{CE} = 1.1 \text{cm}^3\), contralateral \(z_{\text{max}} = 3.37, \text{CE} = 0.4 \text{cm}^3\) for FDG. **(b)** Seizure free (SF; Engel class IA) patients, compared to controls \(z_{\text{max}} = 3.31, \text{CE} = 0.25 \text{cm}^3\), ipsilateral for FMZ and \(z_{\text{max}} = 4.16, \text{CE} = 2.12 \text{cm}^3\), ipsilateral for FDG. The apparent increase near the superior sagittal sinus is a normalization artefact appearing in both modalities. **(c)** NSF patients compared to controls \(z_{\text{max}} = 5.25, \text{CE} = 5.25 \text{cm}^3\), ipsilateral for FMZ and \(z_{\text{max}} = 4.68, \text{CE} = 5.2 \text{cm}^3\), contralateral for FDG. **(d)** Periventricular signal increases in an individual patient (#15), with a suboptimal outcome, for FMZ \(z_{\text{max}} = 4.01, \text{CE} = 3 \text{cm}^3\), ipsilateral, and for FDG \(z_{\text{max}} = 5.11, \text{CE} = 7.3 \text{cm}^3\), contralateral. Significant clusters are pointed with green arrows.
Table 3

| Predictor | Age at onset | Duration of epilepsy | Age at operation |
|-----------|--------------|----------------------|------------------|
| NSF       | 4.5 [6.5]    | 26 [15.3]            | 34 [8.3]         |
| SF        | 5.0 [2.3]    | 24 [12.5]            | 28.5 [13]        |
| MWUT (two tailed) | 32.5 (p = 0.96) | 32.5 (p = 0.96) | 38.5 (p = 0.5) |

4. Discussion

This study shows an association between periventricular WM increases of [18F]FDG binding with NSF status after temporal resection for HS. These results corroborate the observations previously made for [11C]FMZ (Yankam Njiwa et al., 2013). For large numbers of control datasets available, periventricular [11C]FMZ has previously been shown to usefully predict individual postoperative outcome; here we show that this is less the case for [18F]FDG uptake regarding periventricular white matter increases.

Accuracies for distinguishing patients from controls were larger than 0.8 for [11C]FMZ using RF classification. However, this distinction is not clinically meaningful. For the clinically important SF/NSF distinction, [18F]FDG as well as [11C]FMZ performed roughly at chance level.

The previous study (Yankam Njiwa et al., 2013) showed that summed [11C]FMZ radioactivity images, not requiring metabolite corrected arterial plasma input functions, were sufficient for predicting surgery outcome in candidates with TLE based on periventricular WM changes. An important finding of the current study is that the method could in principle, with caution, also be applied to the more clinically widely used FDG.

Increased [11C]FMZ/[18F]FDG signal likely indicates an increased concentration of heterotopic neurons in the sense of a developmental abnormality, i.e. MRI-invisible subependymal heterotopia (Hammers et al., 2003) or a consequence of recurrent seizures. This may imply the existence of a more widespread epileptogenic network beyond the hippocampus and boundaries of the temporal lobes.

As illustrated in the Results section, the presence of ipsilateral mesial temporal lobe hypometabolism to a certain degree predicted SF outcome, and its absence to a degree predicted NSF outcome.

4.1. Clinical outcome

For FMZ, the leave one out test performed with 1000 trees led to five of eight true positive NSF predictions in NSF patients against 35 out of 41 controls correctly predicted. The test also well predicted five SF patients as SF and failed to predict six out of 30 controls.

In contrast, with FDG, the test predicted five out of eight true positive NSF and failed to well predict eleven controls. Fourteen controls were not correctly predicted against four true positive SF patients.

Table 4

| Test output | Clinical outcome |
|-------------|------------------|
| SF          | NSF              |
| FMZ 7/FDG 4 | FMZ 3/FDG 3     |
| NSF         | FMZ 1/FDG 4      |
| FMZ 5/FDG 5 |                  |

These feature importances were highly interesting because voxels were detected outside the periventricular areas, thus opening the door to merge the results of the two methods. Combining RF classification with periventricular masking improved the prediction accuracy for FMZ. If both were used for predicting NSF status and if either test predicted NSF status (data in Table 1), FMZ would correctly predict NSF status in all NSF patients, but assign NSF prognosis incorrectly to 2/8 SF.
patients (sensitivity 100%, specificity 75%). However, combining both predictors for FDG would not be useful, as all patients except two SF patients would be given the NSF prognosis.

Several studies have shown that temporal lobe hypometabolism has postsurgical prognostic significance; patients with at least 15% hypometabolism were likely to become SF (Theodore et al., 1992; Theodore et al., 1994). It has also been illustrated that mean lateral but not mesial temporal asymmetry was significantly higher in patients who did become seizure free than those who did not (Theodore et al., 1992). Here, we investigated relative hypermetabolism in the periventricular white matter. For this feature, NSF patients had more clusters than SF patients (Fig. 2). However, they were seen bilaterally without marked asymmetry. This may reflect a mechanism acting on both sides, either a developmental abnormality of migration (Gleeson and Walsh, 2000) or a response to frequent seizures (Parent et al., 2002). Other attempts to predict postsurgical outcome from presurgical imaging data have also been recently discussed (Chassoux et al., 2010; Feis et al., 2013; Yankam Njiwa et al., 2013). In addition, atypical histological HS patterns, as for example damage restricted to the hilar region, may predict poorer outcome (Blumcke et al., 2007; de Lanerolle et al., 2003; Sagar and Oxbury, 1987). The same held true when HS patterns were analysed quantitatively (Thom et al., 2010). The usefulness of the potential predictors is dependent on the assessment time according to the operation and prognostic factors are also interdependent (Janszky et al., 2005; Najm et al., 2013). Seizure characteristics like high seizure frequency and a history of generalized seizures have also been described as predictors of non-seizure free outcome (Asztely et al., 2007; Janszky et al., 2005). In our study, seizure frequencies had not been prospectively noted at the time of the scan in all patients included, and given the long follow-up, it was not possible to meaningfully reconstruct this information.

Previous studies have sometimes found that “mild” malformations of cortical development, e.g. cortical lamination abnormalities in the temporal lobe associated with HS (focal cortical dysplasia type IIIa according to the 2011 ILAE classification (Blumcke et al., 2011)) may not influence postsurgical outcome in mTLE (Kuba et al., 2012). It is important to note that such findings refer to abnormalities in surgical specimens and hence, by definition, to subtle pathology that has been removed and can therefore no longer contribute to ictogenesis, e.g. via the “short-circuiting” described in animal models associated with heterotopic neurons (Chevassus-au-Louis et al., 1999). This is consistent with our own previous studies, where temporal lobe white matter

| Table 5 | Performance results of the RF classifier for binary classifications. Mean ± SD accuracy, sensitivity and specificity for summed radioactivity images (FMZ) and FDG static images. Performances are averaged over the number of trees for each RF (1000:1000:10,000). NSF = not seizure free, SF = seizure free, C = controls. Performance values of the RF classifier in classifying NSF/SF patients versus controls and in identifying NSF patients versus SF patients. |
|---------|-------------------------------------------------|-------------------------------------------------|-------------------------------------------------|-------------------------------------------------|-------------------------------------------------|-------------------------------------------------|-------------------------------------------------|-------------------------------------------------|-------------------------------------------------|-------------------------------------------------|-------------------------------------------------|-------------------------------------------------|
|         | **FDG**                                         | **FMZ**                                         | **FDG**                                         | **FMZ**                                         | **FDG**                                         | **FMZ**                                         | **FDG**                                         | **FMZ**                                         | **FDG**                                         | **FMZ**                                         | **FDG**                                         | **FMZ**                                         |
|         | NSF/SF                                         | NSF/C                                          | SF/C                                           | NSF/SF                                         | NSF/C                                          | SF/C                                           | NSF/SF                                         | NSF/C                                          | SF/C                                           | NSF/SF                                         | NSF/C                                          | SF/C                                           |
| Accuracy| 0.36 ± 0.03                                    | 0.66 ± 0.001                                   | 0.53 ± 0.02                                    | 0.40 ± 0.06                                    | 0.83 ± 0.01                                    | 0.82 ± 0.01                                    | 0.36 ± 0.03                                    | 0.66 ± 0.001                                   | 0.53 ± 0.02                                    | 0.40 ± 0.06                                    | 0.83 ± 0.01                                    | 0.82 ± 0.01                                    |
| Sensitivity | 0.33 ± 0.06                                    | 0.74 ± 0.04                                    | 0.5 ± 0.00                                     | 0.20 ± 0.06                                    | 0.76 ± 0.04                                    | 0.76 ± 0.04                                    | 0.33 ± 0.06                                    | 0.74 ± 0.04                                    | 0.5 ± 0.00                                     | 0.20 ± 0.06                                    | 0.76 ± 0.04                                    | 0.76 ± 0.04                                    |
| Specificity | 0.40 ± 0.05                                    | 0.63 ± 0.00                                    | 0.53 ± 0.03                                    | 0.54 ± 0.08                                    | 0.85 ± 0.01                                    | 0.83 ± 0.01                                    | 0.40 ± 0.05                                    | 0.63 ± 0.00                                    | 0.53 ± 0.03                                    | 0.54 ± 0.08                                    | 0.85 ± 0.01                                    | 0.83 ± 0.01                                    |

Fig. 2. Feature importances for differentiating between three clinical groups using voxel based FDG-PET (bottom) and FMZ-PET (top). Voxels important for distinguishing patients from controls are overlaid onto an MNI-space FMZ-PET template (top) and FDG-PET template (bottom). Left column, non-seizure free patients; right column, seizure free patients.
increases of flumazenil had been seen bilaterally, but were only associated with suboptimal outcome in the contralateral temporal lobe (Hammers et al., 2001). Similarly, in (Yankam Njiwa et al., 2013) and the present manuscript we describe the same in vivo biomarker in a remote area not removed during surgery.

Both tracers and methods did not always overlap in their prediction of seizure status for individual patients. Periventricular WM signal increases are not always seen at the same locations for both tracers. [18F]FDG uptake assesses metabolism, a more general aspect of brain function than the more specifically neuron-linked [11C]FMZ binding; this could explain the fact that [18F]FDG uptake was less predictive of outcome (see Table 1).

Other studies showed that the complete postoperative absence of seizures widely varies with the considered postoperative time point (Aull-Watschinger et al., 2008; Elsharkawy et al., 2009; Kelemen et al., 2006). For example, in a study of 153 patients with MRI-diagnosed HS, the gender, the seizure frequency before operation, and the unilaterality of interictal EEG discharges were used as variables in a logistic regression equation for predicting outcome after surgery (Aull-Watschinger et al., 2008). Complete seizure freedom (Engel class IA) could not be predicted from conventional preoperative variables; our study provides another element allowing such a prediction even if its accuracy remains limited.

In summary, classification based on RF performed slightly better than the previous method based on periventricular WM signal increases for both used tracers. The accuracy may be sufficient to aid in individual preoperative counselling. However, the classifier constructed will need to be tested on an independent sample.

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