[11C]-Methionine PET: dysembryoplastic neuroepithelial tumours compared with other epileptogenic brain neoplasms

D S Rosenberg, G Demarquay, A Jouvet, D Le Bars, N Streichenberger, M Sindou, N Kopp, F Mauguie`re, P Ryvlin

Background and objectives: Brain tumours responsible for longstanding partial epilepsy are characterised by a high prevalence of dysembryoplastic neuroepithelial tumour (DNT), whose natural evolution is much more benign than that of gliomas. The preoperative diagnosis of DNT, which is not yet feasible on the basis of available clinical and imaging data, would help optimise the therapeutic strategy for this type of tumour. This study tested whether [11C]-methionine positron emission tomography (MET-PET) could help to distinguish DNTs from other epileptogenic brain tumours.

Methods: Prospective study of 27 patients with partial epilepsy of at least six months duration related to a non-rapidly progressing brain tumour on magnetic resonance imaging (MRI). A structured visual analysis, which distinguished between normal, moderately abnormal, or markedly abnormal tumour methionine uptake, as well as various regions of interest and semiquantitative measurements were conducted.

Results: Pathological results showed 11 DNTs (41%), 5 gangliogliomas (18%), and 11 gliomas (41%). MET-PET visual findings significantly differed between the various tumour types (p < 0.0002), regardless of gadolinium enhancement on MRI, and were confirmed by semiquantitative analysis (p < 0.001 for all calculated ratios). All gliomas and gangliogliomas were associated with moderately or markedly increased tumour methionine uptake, whereas 7/11 DNTs had a normal methionine uptake, including all six located in the mesiotemporal structures. No DNT presented with a marked MET-PET abnormality.

Conclusion: Normal MET-PET findings in patient with an epileptogenic and non-rapidly progressing brain tumour are suggestive of DNT, whereas a markedly increased tumour methionine uptake makes this diagnosis unlikely.

In this study of 27 patients with epileptogenic brain tumours, we tested the ability of MET-PET to help distinguish DNTs from gangliogliomas and other gliomas.

PATIENTS AND METHODS

Patients

From 1995 to 2002 we prospectively selected all patients who had been consecutively referred to our epilepsy department for partial epilepsy associated with an MRI abnormality suggesting a non-rapidly progressing brain tumour. All patients gave informed consent to participate in the study, and the local ethic committee approved the study, which conformed to the Declaration of Helsinki. We excluded those patients whose tumour had been symptomatic for less than six months or was associated with the following clinical or MRI signs, indicating a rapid tumour growth:

- a significant progressive neurological deterioration other than changes in seizure frequency and severity
- a clear-cut mass effect with midline displacement on neuroimaging data
- a significant increase in the tumour size on two consecutive MRI.

From the 30 patients who entered this PET study, we selected all those who had been operated on, and who had an
quality of MET-PET and pathological data was considered sufficiently high to allow reliable analysis and correlations. Overall, 27 patients met the above criteria and are included in this report.

MRI
We used a 1.5 T device (Magnetom 63SP; Siemens, Erlangen, Germany) and included an axial or coronal spin echo T1 sequence (TR: 600 ms, TE: 12 ms) with and without gadolinium injection, and an axial spin echo T2 sequence (TR: 3000 ms, TE: 16 ms and 98 ms) perpendicular to the T1 sequence. In 20 patients only, a 3D T1 acquisition sequence (TR: 9.7 ms, TE: 4 ms) was performed.

PET
Data acquisition
We used two different cameras for the MET-PET scanning. The first 11 patients were scanned after receiving an intravenous bolus injection of 440 MBq of $^{11}C$-methionine with an LETI TTV03 (Grenoble, France) camera, providing seven 9 mm thick slices with an in-plane resolution of 7 mm full width at half maximum (FWHM). Data were acquired in two dimensional mode, planes oriented parallel to the orbitomeatal line. The others 16 patients received an intravenous bolus injection of $18.5 + 2.6$ MBq/kg of $^{11}C$-methionine, and they underwent scanning using a high resolution tomograph (HR+ Siemens, Erlangen, Germany). Three dimensional data were acquired, providing 63 slices, 2.4 mm thick, with an isotropic spatial resolution of 5 mm FWHM. Before injection, we carried out transmission scanning for attenuation correction using three $^{68}$Ge rod sources. For both cameras, static emission scanning was performed during a 20 minute period, beginning 35 minutes after the $^{11}C$-methionine injection, as has been done in several other studies. Images were corrected for scatter and attenuation, and reconstructed using a filtered back projection.

PET Data analysis
Visual analysis
Two of the investigators (DR, PR) independently performed a visual analysis of PET data, blinded to all other clinical and MRI data except the anatomical location of the tumour. We first determined whether the degree of methionine uptake within the tumour was significantly increased when compared with the surrounding brain and the contralateral homotopic region. If an increase was observed, we distinguished various degrees of abnormal methionine uptake using the contralateral occipital cortex as a reference. We used the contralateral occipital cortex as a reference because, in our experience, this brain region has consistently proved to display the highest methionine uptake among the entire non-tumoral cortical regions. The result of the visual analysis was classified based on the following definitions:

- Normal tumour methionine uptake—no visually detectable difference between the tumour and the surrounding or contralateral homotopic brain regions
- Moderately increased tumour methionine uptake—the tumour uptake clearly exceeds the uptake in surrounding cortical areas and in the contralateral homotopic region, but remains lower or comparable to that of the contralateral occipital cortex
- Markedly increased tumour methionine uptake—the tumour uptake clearly exceeds that observed in the contralateral occipital cortex

Semi-quantitative analysis
We performed a semi-quantitative analysis, using two sets of regions of interest (ROIs). The placement of the first set of ROIs required coregistered MRI and PET data, which were only available for the 20 patients who benefited from a 3D MRI. We manually traced irregular ROIs along the visually detected tumour borders on each MRI slice displaying the tumour, excluding areas of clear-cut necrosis or cyst. A flipped copy of each of these ROIs was placed over the contralateral homotopic region and readjusted to precisely match the outer border of the cortical ribbon. After coregistration with MRI, we transferred the tumoral (T) as well as the contralateral homotopic ROIs (C) onto the PET images and measured the mean regional count of radio-activity within these ROIs. We then calculated the mean tumour-to-contralateral homotopic ratio (mean T/C = T/C). The second set of ROIs corresponded to radio-circular regions, 5 mm in diameter, which were directly placed onto the PET images. These ROIs were placed over the portion of the tumour displaying the highest $^{11}C$-methionine uptake (Tc = tumour circular ROI), the contralateral homotopic cortical region (Cc = contralateral circular ROI), and the most active area within the contralateral occipital cortex (Oc = occipital circular ROI), on a single slice each. This procedure was undertaken in all patients showing a visually detectable increased tumour methionine uptake, including those who did not benefit from coregistration of MRI and PET images (n = 5). It was also undertaken in the subset of patients with no visually detectable increased tumour methionine uptake, by placing the circular ROI over the most active voxels within the tumour border as delineated by coregistered MRI and PET data (n = 5). Only 2/27 patients (patient 4 and 7), whose MET-PET showed normal findings and who did not benefit from a 3D T1 MRI, were not included in this analysis. In the remaining 25 patients, we calculated two ratios:

- Maximal tumour-to-contralateral homotopic ratio (Max T/C) = T/C/Cc
- Maximal tumour-to-contralateral occipital ratio (OR) = T/C/Oc

Surgical procedure
A total of 24 patients (89%) benefited from a total (n = 18; 67%) or subtotal (n = 6; 22%) removal of their tumour. The remaining three patients (11%) underwent a biopsy, and the size and quality of the specimen enabled a conclusive pathological diagnosis.

Pathological analysis and diagnosis
Sections of formalin-fixed tissues were processed for histological staining using either the haemalum–phloxine safranin or the haematoxylin–eosin technique. Immunohistochemical stains, applied on routinely fixed and paraffin-embedded sections, were prepared for selected cases, using the avidin–biotin complex method and the following antisera: anti-GFAP (glial fibrillary acidic protein), anti-NF (neurofilament protein), anti-NSE (neurone specific enolase), antisynaptophysin, anti-vimentin, anti-S100 protein, and anti-Leu-7. These specimens were analysed at our institution by neuropathologists trained in the evaluation of epileptogenic low grade tumours, and ultimately classified according to the WHO classification. However, where the diagnosis remained uncertain, the specimens were sent to two other pathologists renowned for their experience in epileptogenic brain tumours (C Daumas-Duport, Saint-Anne Hospital, Paris, and B Pasquier, Albert Michallon Hospital, Grenoble, France) for further evaluation, which led to a consensual conclusion in all cases.
Table 1  Individual pathological, clinical, MRI, and PET-MET data

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DNT, dysembryoplastic neuroepithelial tumour; F, female; Fr, frontal; I, insula; L, left; Lt, lateral temporal; M, male; max T/C, maximal tumour-to-contralateral homotopic ratio; mean T/C, mean tumour-to-contralateral homotopic ratio; MET-PET, [123I]-methionine positron emission tomography; mT, mesial temporal; MRI, magnetic resonance imaging; NA, not available; O, occipital; OR, tumour-to-contralateral occipital ratio; P, parietal; R, right.

Statistical analysis

For visual analysis, three types of tumour were considered: DNTs, gangliogliomas, and gliomas (including astrocytomas, oligodendrogliomas, and oligoastrocytomas of various grades). We looked for correlations between the tumour type, its anatomical location, as well as the presence of contrast enhancement on MRI and that of a visually detectable increased tumour methionine uptake on MET-PET images, using the χ² statistic. For the semiquantitative MET-PET analysis we used a one factor analysis of variance statistic. For the semiquantitative MET-PET findings (see table 1)

PET-MET findings (see table 1)

Visual analysis demonstrated an increased methionine uptake within the tumour in 20 patients (74%), which was classified as moderate in 11 (55%), and marked in 9 (45%). No increased uptake was detected in seven patients (26%) (fig 1). There was no correlation between the presence of increased methionine uptake and gadolinium enhancement on MRI, the former being observed in 73% of gadolinium enhancement (table 2).

MET-PET visual findings correlated with the pathological data (p<0.0002) (table 3). In fact, the lack of increased methionine uptake was only observed in DNTs, 7/11 demonstrating that pattern. The difference between DNTs and the other types of tumour was even more pronounced in the subset of mesial temporal lesions, where all DNTs displayed normal findings on MET-PET (n = 6), whereas all the other tumours (n = 4 (3 gangliogliomas, 1 astrocytoma)) demonstrated an increased methionine uptake. Thus, in our study population, normal MET-PET findings were only associated with DNTs, whereas the presence of a marked increase in methionine uptake was not observed in gliuneuronal tumours with the exception of one ganglioglioma. Moderate increase in methionine uptake was associated with all types of tumour except mesiotemporal DNTs.

The semiquantitative analysis also demonstrated significant differences between the four types of tumour for the three calculated ratios (table 4): mean T/C (p<0.001), max T/C (p<0.0001), and OR (p<0.0001). Post hoc tests showed that DNTs were associated with lower ratios than gangliogliomas (mean T/C: p<0.005) as well as low grade (mean T/C: p<0.001; max T/C: p<0.003; OR: p<0.001) and high grade gliomas (max T/C: p<0.001; OR: p<0.0001). Furthermore, gangliogliomas were associated with lower ratios than low grade (OR: p<0.05) and high grade (max T/C: p<0.001).
Figure 1 Examples of T1-weighted magnetic resonance images (left panel), methionine (MET) uptake (middle panel) and coregistered magnetic resonance imaging (MRI) and positron emission tomography (PET) data (right panel) for the three major tumour types. Arrows indicate tumour location on MRI and the corresponding methionine uptake on PET images. (A) Left mesial temporal dysembryoplastic neuroepithelial tumour (DNT), which was not associated with a visually detectable increased methionine uptake on raw MET-PET image, as well as on the coregistered MRI and PET data (patient no 13). The dotted arrow points to the hypophyseal region, which is physiologically associated with a higher methionine uptake than other neighbouring cerebral structures. (B) Left neocortical (parietal) DNT observed on T1-weighted MRI with a moderately increased methionine uptake on raw MET-PET image and coregistered MRI-PET data (patient no 19). (C) Right mesial temporal ganglioglioma associated with a moderately increased methionine uptake (patient no 15). (D) Right frontal glioma (oligodendroglioma grade II) corresponding to a hypodense area without gadolinium enhancement on T1-weighted MRI but associated with a markedly increased methionine uptake (patient no 18).
p<0.001; OR: p<0.003) gliomas. As already reported by others,28–40 we also found an increased methionine uptake in gliomas, more pronounced in high grade than low grade tumours for max T/C and OR.

Results of the semiquantitative data analysis also significantly differed between the three types of visual MET-PET findings (mean T/C: p<0.004; max T/C: p<0.0001; OR: p<0.0001), with the lowest values associated with normal MET-PET findings and the highest with marked abnormalities. Although individual ratios values partly overlapped between the groups (see table 1), there was a cut-off for the max T/C between tumours with normal methionine uptake (<1.3) and those with moderate or marked methionine uptake (≥1.64).

DISCUSSION

In this largest MET-PET series of patients with non-rapidly progressing epileptogenic brain tumours, we have shown that DNT, which is distinct from other low-grade tumours from a prognostic and therapeutic point of view, proved to be associated with a specific MET-PET finding—that is, normal methionine uptake in 7/11 cases, and in all those located in the mesial temporal region. Conversely, all other types of epileptogenic brain tumour displayed an increased methionine uptake in our study population. These results suggest that MET-PET might provide clinically useful and reliable information about the pathology underlying epileptogenic brain tumours.

Some methodological issues must be discussed. Firstly, changes in imaging procedures occurred during the course of this study, resulting in two different PET acquisition protocols, and lack of 3D MRI data in a minority of patients. However, these changes are unlikely to have biased our results, since the proportion of each tumour type was comparable in all the resulting subgroups. Most previous MET-PET studies used circular or irregular ROIs traced around the tumours in comparison with contralateral homotopic or cortical regions, but it is noteworthy that the exact type of ROI has varied from one series to another.22 24–29 32 34 41 We used the different types of ROI and index most commonly reported in other studies to ensure that our results were not dependent on a specific method of PET data analysis. In fact, most of the differences observed between the various types of tumour were demonstrated with all the three parameters employed in this study.

Secondly, visual analysis was performed blinded to all other data except the anatomical location of the tumour to address the issue most relevant to clinical practice—that is, the PET findings within the tumour. Regardless of the limitations and subjectivity of visual analysis, it must be acknowledged that the clinical use of PET imaging partly relies on the possibility of obtaining reliable visual interpretation of the data in daily practice. The criteria proposed in the present study to detect and classify abnormal tumour methionine uptake appear appropriate for such clinical application; they proved easy to use and reliable, allowing for fully consensual reports of two independent investigators and providing results consistent with semiquantitative data.

Lastly, regarding the reliability of our patients’ pathological diagnosis, several authors have emphasised the difficulty in establishing the neuropathological diagnosis of DNT.4 5 The diagnosis is partly based on the presence of the ‘‘glioneuronal element’’, a specific figure which could be missing in non-specific forms of DNT,28 40 or when the tumour has been removed incompletely. In our series, all patients underwent complete removal of their tumour except for three patients, whose biopsy specimens provided sufficient information to ensure a definite diagnosis. In addition, as described in the Methods section our selection criteria included a reliable pathological diagnosis made by pathologists trained to in the evaluation of glioneuronal tumours, including experts from other institutions when necessary.

One of our main findings was that only DNTs, and more specifically mesial temporal DNTs, were associated with a normal tumour methionine uptake, suggesting that this pattern could be specific of this tumour type, at least in the context of a non-rapidly progressing brain mass lesion associated with an epilepsy of more than six months’ duration. Similar findings were recently reported in a small series of seven epileptogenic tumours, including four DNTs, one ganglioglioma, and two low grade gliomas.42 In contrast, several authors have reported the possibility of normal MET-PET findings in low grade astrocytomas.23 24 27 29 34 41 Some of these studies were conducted before DNT was clearly recognised, suggesting the possibility that some of these astrocytomas were DNTs.22 23 29 In other, more recent, studies27 34 pathological examination of up to half of the specimens was based on a biopsy again raised the possibility of misdiagnosed DNT.4 Thus none of the series that have reported normal MET-PET findings in low grade gliomas included DNTs, despite the fact that the latter represents up

<table>
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<th>Visual methionine uptake</th>
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<td>Normal</td>
<td>7 (63)</td>
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<tr>
<td>Moderate increase</td>
<td>4 (37)</td>
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<td>Marked increase</td>
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* p<0.0002.

Table 3 Correlation between qualitative MET-PET data and pathology (n = 27)

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Table 4 Correlation between semiquantitative [11C]-methionine positron emission tomography and pathological data

<table>
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<tr>
<th>Pathological data (n (%))</th>
<th>Mean (SD) T/C (n = 20)</th>
<th>Max (SD) T/C (n = 25)</th>
<th>OR (SD) (n = 25)</th>
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<td>Dysplastic neuroepithelial</td>
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<td>Gangliogliomas</td>
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<td>Low grade gliomas</td>
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<td>2.56 (0.7)</td>
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<td>High grade gliomas</td>
<td>1.48 (0.4)</td>
<td>3.87 (1.3)</td>
<td>2.02 (0.47)</td>
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Max T/C, maximal tumour-to-contralateral homotopic ratio; Mean T/C, Mean tumour-to-contralateral homotopic ratio; OR, tumour-to-contralateral occipital ratio.
to 65% of brain neoplasms responsible for chronic drug resistant epilepsy. Apart from the issue of misdiagnosed DNTs, discrepancies between our results and former series might reflect differences in the rates of chronic drug resistant partial epilepsy in the population studied. In that respect, it is important to stress that the common forms of low grade astrocytomas are rarely encountered in patients with tumour associated chronic epilepsy, and were not observed even in a recent series of 94 epilepsy surgery tumour cases. Finally, some authors have proposed that protoplasmic astrocytomas in cortical locations associated with chronic epilepsy correspond to non-specific forms of DNT. Overall, whether the lack of a visually detectable increased methionine uptake is specific to DNTs, or that it may also occur with other epileptogenic tumours, remains an open issue that needs to be addressed in a larger population.

Semiquantitative analysis of the DNT cases presented in this study resulted in an average max T/C ratio for DNTs within a previously reported range for low grade gliomas. It must be noted, however, that there was a significant difference between DNTs with and without visually detectable increased methionine uptake (average max T/C 1.76 and 1.07, respectively; p<0.05). With regard to the other tumour types, our results are generally in agreement with those of previously published studies. All oligodendrogliomas, as well as all high grade gliomas were associated with an abnormal increased methionine uptake, as has been reported by others. In contrast, gangliogliomas stood apart from gliomas, in that they were only rarely associated with a marked MET-PET abnormality (20%). However, the small number of cases included in this study tempers this finding. We found a single report of a MET-PET study in a patient with a temporal ganglioglioma, which was consistent with our own findings, as well as another report of a patient investigated with single photon emission computed tomography (SPECT) that demonstrated an increased methylyrosine uptake. According to the various ratios that we calculated, gangliogliomas appear to be associated with an average degree of increased methionine uptake intermediate between that of DNTs and gliomas. This is consistent with the risk of malignant transformation of these tumours, estimated around 10%, a low figure compared with that for gliomas in this study, it seems unlikely that the presence of markedly increased methionine uptake makes the diagnosis of either DNT or ganglioglioma unlikely.

ACKNOWLEDGEMENTS

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Authors’ affiliations
D S Rosenberg, G Demarquay, F Mauguire, P Ryvlin, Department of Functional Neurology and Epileptology, Hopital Neurologique, Lyon, France
A Jouvet, N Streichenberger, N Kopp, Department of Neuropathology, Hopital Neurologique, Lyon, France
D Le Bars, P Ryvlin, CERMEP, Hopital Neurologique, Lyon, France
M Sindou, Department of Neurosurgery, Hopital Neurologique, Lyon, France

Competing interests: none declared

This study was approved by the CCPRPB local ethic committee (Comité Consultatif de Protection des Personnes se prêtant à des Recherches Biomédicales—Centre Léon Bérard, Lyon)

REFERENCES