Comparison of $^{11}$C-Methionine, $^{11}$C-Choline, and $^{18}$F-Fluorodeoxyglucose-Positron Emission Tomography for Distinguishing Glioma Recurrence from Radiation Necrosis

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Abstract
The aim of this study is to assess the different metabolic activities characteristic of glioma recurrence and radiation necrosis (RN) and to evaluate the diagnostic accuracy for differentiation of the two conditions using $^{11}$C-methionine (MET), $^{11}$C-choline (CHO), and $^{18}$F-fluorodeoxyglucose (FDG)-positron emission tomography (PET). Fifty patients with lesions suggestive of recurrent glioma by magnetic resonance imaging (MRI) underwent MET, CHO, and FDG-PET. All patients who had previously been treated with radiotherapy for malignant glioma were subjected to open surgery and pathological diagnosis (17 recurrent grade 3 gliomas (Gr.3s) comprising 7 anaplastic astrocytomas (AAs) and 10 anaplastic oligodendrogliomas (AOs), 17 recurrent glioblastomas (Gr.4s), and 16 RNs). We measured the PET/Gd volume ratio, the PET/Gd overlap ratio, and the lesion/normal brain uptake ratio (L/N ratio) and determined the optimal index of each PET scan. The PET/Gd volume ratio and the PET/Gd overlap ratio for RN were significantly lower than those of glioma recurrence only with MET-PET ($P < 0.05$). The L/N ratio of RN was significantly lower than that of Gr.4 with all PET imaging ($P < 0.001$) and was significantly lower than that of Gr.3, especially for AO, only with MET-PET images ($P < 0.005$). Receiver operating characteristic (ROC) analysis showed that the area under the curve of MET, CHO, and FDG was 92.5, 81.4, and 77.4, respectively. MET L/N ratio of greater than 2.51 provided the best sensitivity and specificity for establishing glioma recurrence (91.2% and 87.5%, respectively). These results demonstrated that MET-PET was superior to both CHO and FDG-PET for diagnostic accuracy in distinguishing glioma recurrence from RN.

Key words: $^{11}$C-methionine, positron emission tomography, radiation necrosis, glioma

Introduction
Radiation necrosis (RN) is a serious clinical complication in the diagnosis and treatment of patients with malignant gliomas. Because the imaging features of most RN appear similar to those of malignant gliomas by computed tomography (CT) or magnetic resonance imaging (MRI), it is difficult to distinguish glioma recurrence from RN. Since therapeutic strategies for these pathological entities are fundamentally different, their differential diagnosis is crucial. Recently, several clinical studies using diffusion MRI,1–4 perfusion MRI,3 MR spectroscopy,3–5 and $^{203}$thallium single photon emission computed tomography (SPECT)6 have been undertaken in attempts to distinguish between the two conditions. These modalities have made it possible to easily diagnose some cases compared to protocols from the previous era in which only conventional CT or MRI was used. Furthermore, $^{11}$C-methionine (MET) and $^{18}$F-fluorodeoxyglucose (FDG)-positron emission tomography (PET) have been reported to be more useful for differential diagnosis between glioma recurrence and RN.6–13 These PET methods

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Diagnostic Accuracy of MET-PET for Radiation Necrosis

In this retrospective study from 2002 to 2008, we examined PET scans from 50 consecutive patients with supratentorial space-occupying lesions following radiotherapy for malignant gliomas at the Chubu Medical Center for Prolonged Traumatic Brain Dysfunction, Kizawa Memorial Hospital. All supratentorial space-occupying lesions were Gd-enhanced, and interpretation of the lesions as glioma recurrence or RN was unclear. Presurgical radiologic evaluation was performed using the World Health Organisation (WHO) classification system. Of the 50 patients, 17 had recurrent grade 3 glioma (Gr.3), 17 had recurrent glioblastoma (Gr.4), and 16 had RN. The 17 Gr.3s were further classified as 7 anaplastic astrocytomas (AAs) and 10 anaplastic oligodendrogliomas (AOs). RN was pathologically diagnosed in the limited cases in which the surgical specimen showed typical necrotic tissues including thickness and fibrinoid necrosis of the vascular walls, multiple microcysts, coagulation necrosis, endothelial proliferation, and inflammatory cells interspersed with or without scattered tumor cells. The clinical features of the patients are summarized in Table 1. All patients gave written informed consent, and the study protocol was approved by the research committee of the Kizawa Memorial Hospital Foundation.

The PET study was carried out according to standardized procedures recommended by the Japan Radioisotope Association. The PET scanner was an ADVANCE NXi Imaging System (General Electric Yokokawa Medical System, Hino, Tokyo), which provided 35 transaxial images at 4.25 mm intervals covering a 25.6 cm in-plane field of view (FOV). The in-plane spatial resolution (full width at half maximum) was 4.8 mm, and the scan mode was the standard 2D mode. Before the emission scan was performed, a 3 minute transmission scan was performed to correct photon attenuation with a ring source containing $^{68}$Ge. Patients had fasted for at least 4 hours before PET studies. A venous cannula was inserted into the forearm for injection of radiopharmaceuticals. From this cannula, blood samples could also be collected if necessary. A dose of 7.0 MBq/kg of MET, 7.0 MBq/kg of CHO, or 5.0 MBq/kg of FDG was injected intravenously, depending on the particular examination. Emission scans were acquired as follows: (1) for 30 minutes, beginning 5 minutes after MET injection, (2) for 7 minutes, beginning 2 minutes after CHO injection, and (3) for 7 minutes, beginning 35 minutes after FDG injection. During PET data acquisition, head motion was continuously monitored using laser beams projected onto ink marks drawn on the forehead and was corrected manually, as necessary. Scan images were reconstructed using the ordered-subsets expectation maximization algorithm (2 iterations, 14 subsets). Images were reconstructed into a 128 × 128 matrix with a pixel size of 2 × 2 mm.

MR imaging was performed with a 1.5 T system (Signa; GE Medical Systems, Milwaukee, Wisconsin, USA). Axial T₁-weighted images (TR/TE/NEX = 350/9/2), T₂-weighted images (2300/100/2), and fluid attenuated inversion recovery (FLAIR) images (800/110/1, inversion time = 2400 ms) (FOV 24 × 24 cm, matrix size 512 × 256) were acquired. The slice thickness was 6 mm, with a 3-mm slice gap. For co-registration of metabolic and anatomic data, 3D spoiled gradient-echo images were also acquired after administration of 0.2 ml/kg of gadopentate dimeglumine (Gd-DTPA) (Magnevist; Nihon Shering, Osaka) using the following parameters: no gap, 1.0 mm thickness, TR/TE = 20.0/1.6 ms, flip angle = 15°, NEX = 1, and axial views.

Tracer accumulation in the regions of interest (ROIs) was analyzed as the standardized uptake value (SUV), which is the activity concentration in the ROI at a fixed time point divided by the injected dose normalized to the patient’s measured weight. MET, CHO, and FDG lesion/normal brain uptake ratios (L/N ratios) were calculated...
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by dividing the maximum SUV for the enhanced lesion on the MR image by the mean SUV of the contralateral normal frontal cortex. The lesion SUVs were selected at the highest accumulation, and reference ROIs on each of the three axial planes were drawn with a diameter of 10 mm. Co-registration of PET and MR imaging was accomplished with an analysis software package (AJS, Tokyo), using the method described by Kapouleas et al.20 We used the L/N ratio instead of the absolute SUV because of the high, unexplained intersubject variability of the SUV.21 We used the lesion maximum SUV instead of lesion mean SUV to minimize the effect of lesion heterogeneity. For each PET tracer, we defined regions with L/N ratios greater than 1.5 as PET abnormal high uptake regions and measured the volumes of these regions in each PET image and also the volumes of the Gd-enhanced area in the MRI using an analysis software package (AJS, Tokyo). The volume of the PET abnormal high uptake region overlap with the Gd-enhanced area was measured by the same method for each case. The volume ratio of the PET abnormal high uptake area to Gd-enhanced MR area (PET/Gd volume ratio) was calculated as follows: PET/Gd volume ratio (%) = [PET abnormal high uptake area (volume) ÷ Gd-enhanced area (volume)] × 100.

The ratio of the PET abnormal high uptake area overlapping the Gd-enhanced MR area (PET/Gd overlap ratio) was calculated as follows: PET/Gd overlap ratio (%) = [PET abnormal high uptake area overlapping Gd-enhanced area (volume) ÷ Gd-enhanced area (volume)] × 100.

Data are presented as means ± standard deviations (SDs). To compare the L/N ratios of the three PET modalities at the best distinction between glioma recurrence and RN, statistical analysis was performed using analysis of variance and Tukey’s test for multiple comparisons. Receiver operating characteristic (ROC) curves were calculated to determine the cut off values for differential diagnosis of glioma recurrence and RN. P values less than 0.05 were considered statistically significant.

**Results**

I. Volume comparison between MRI and PET studies

The MET-PET/Gd volume ratios of RN, AA, AO, and Gr.4 were 21.7% ± 20.9%, 164.3% ± 158.5%, 185.5% ± 162.6%, and 123.6% ± 66.4%, respectively (Fig. 1A). The MET-PET/Gd overlap ratios of RN, AA, AO, and Gr.4 were 20.7% ± 21.4%, 63.5% ± 40.3%, 74.8% ± 34.0%, and 64.6% ± 29.4%, respectively (Fig. 1B). Both the MET-PET/Gd volume ratio and the MET-PET/Gd overlap ratio of RN were significantly lower than those of AA, AO, and Gr.4, respectively (P < 0.05).

The CHO-PET/Gd volume ratios of RN, AA, AO, and Gr.4 were 100.5% ± 20.5%, 110.2% ± 17.3%, 99.9% ± 15.9%, and 104.1% ± 13.7%, respectively (Fig. 1C). The CHO-PET/Gd overlap ratios of RN, AA, AO, and Gr.4 were 83.6% ± 15.1%, 97.4% ± 3.9%, 92.5% ± 10.3%, and 96.1% ± 7.1%, respectively (Fig. 1D). There were no significant differences of the CHO-PET/Gd volume ratios and the CHO-PET/Gd overlap ratios among RN, AA, AO, and Gr.4.

The FDG-PET/Gd volume ratios of RN, AA, AO, and Gr.4 were 0.4% ± 1.5%, 0.5% ± 1.3%, 0.0% ± 0.0%, and 12.1% ± 20.6%, respectively (Fig. 1E). The FDG-PET/Gd overlap ratios of RN, AA, AO, and Gr.4 were 0.4% ± 1.5%, 0.5% ± 1.3%, 0.0% ± 0.0%, and 11.7% ± 19.4%, respectively (Fig. 1F). Both the FDG-PET/Gd volume ratio and the FDG-PET/Gd overlap ratio of Gr.4 were significantly higher than those of RN (P < 0.05).

II. Semiquantitative analysis of PET studies

The mean SUVs of MET, CHO, and FDG from the contralateral normal frontal cortex were 1.30 ± 0.25, 0.26 ± 0.94, and 6.31 ± 1.71, respectively. MET L/N ratios of RN, Gr.3, and Gr.4 were 1.95 ± 0.60, 3.40 ± 1.04, and 4.29 ± 1.45, respectively. There was a significant difference between the MET L/N ratios of RN and Gr.3 (P < 0.005) and of RN and Gr.4 (P < 0.001). However, there was no significant difference between the MET L/N ratios of Gr.3 and Gr.4 (Fig. 2A). MET L/N ratios of AA and AO were 2.79 ± 0.68, and 3.83 ± 1.06, respectively. There was a significant difference between the MET L/N ratios of AA and AO (P < 0.001) and of RN and AA (P < 0.05), but not of RN and AO (Fig. 2B).

CHO L/N ratios of RN, Gr.3, and Gr.4 were 6.90 ± 4.30, 11.18 ± 6.75, and 18.09 ± 10.82, respectively. There was a significant difference only between the CHO L/N ratios of RN and Gr.3 (P < 0.001) and of Gr.3 and Gr.4 (P < 0.05) (Fig. 2C). CHO L/N ratios of AA and AO were 9.21 ± 4.19, and 12.56 ± 8.01, respectively. There was no significant difference between CHO L/N ratios of RN and any of the Gr.3 histological types (Fig. 2D).

FDG L/N ratios of RN, Gr.3, and Gr.4 were 1.15 ± 0.50, 1.26 ± 0.23, and 1.97 ± 0.64, respectively. There was a significant difference only between the FDG L/N ratios of RN and Gr.4 (P < 0.001) and of Gr.3 and Gr.4 (P < 0.001) (Fig. 2E). FDG L/N ratios of AA and AO were 1.24 ± 0.33, and 1.27 ± 0.16, respectively. There was no significant difference between FDG L/N ratios of RN and any of the Gr.3 histological types (Fig. 2F).

Representative PET and MRI images from RN, AA,
Fig. 1 Graphs showing $^{11}$C-methionine (MET)-PET/Gd (A), $^{11}$C-choline (CHO)-PET/Gd (C), and $^{18}$F-fluorodeoxyglucose (FDG)-PET/Gd (E) volume ratios, and MET-PET/Gd (B), CHO-PET/Gd (D), and FDG-PET/Gd (F) overlap ratios of radiation necrosis (RN), anaplastic astrocytoma (AA), anaplastic oligodendroglioma (AO), and recurrent glioblastoma (Gr.4). The significant low values ($P < 0.05$) of both the PET/Gd volume ratio and the PET/Gd overlap ratio of RN compared with glioma recurrence were shown to be characteristic only for MET-PET. *$P < 0.05$, **$P < 0.005$, ***$P < 0.001$.

AO, and Gr.4 cases are shown in Fig. 3.

III. ROC analysis of PET studies

Fig. 2G shows the ROC curves of the 3 PET modalities. The area under the curve of MET, CHO, and FDG-PETs were 0.925, 0.814, and 0.774, respectively. Table 2 shows the best cutoff values, diagnostic sensitivities, and specificities of the 3 PET modalities for recurrent gliomas. The best MET L/N ratio cutoff value was 2.51, which provided a sensitivity of 91.2% and a specificity of 87.5% for diagnosis of glioma recurrence. These results indicate that MET-PET is the most informative method for differentiating tumor recurrence from RN.

Discussion

Radiotherapy has been used for the past four decades as a standard treatment following surgical mass reduction in malignant gliomas. More recently, conventional external radiotherapy has been expanded to include stereotactic radiotherapy, intensity modulated radiotherapy, boron neutron captured therapy, and radiotherapy using heavy ions. The usefulness of radiotherapy for malignant gliomas is not in doubt as it has been verified by improved patient survival and local control. However, identifying RN, which deteriorates the clinical condition of patients, is still a critical problem. Normally, 60 Gy of whole brain external irradiation induces necrosis in about 50% of patients up to 5 years after irradiation. Although the therapeutic strategy for RN is different from that for glioma recurrence in most cases of malignant gliomas, it has been difficult to distinguish these pathological entities from each other even using conventional neuroradiological modalities.
With advancements in metabolic neuroimaging, $^{201}$thallium-SPECT and FDG-PET have been anticipated to be useful for differential diagnosis between glioma recurrence and non-neoplastic lesions. Gómez-Río et al. prospectively evaluated $^{201}$thallium-SPECT and FDG-PET in 76 patients with suspicion of glioma recurrence after surgical excision and radiotherapy. Their results showed that although FDG-PET yielded a slightly higher specificity for diagnosis of glioma recurrence, the sensitivity was considerably lower than that of $^{201}$thallium-SPECT. This means that FDG-PET does not clearly improve upon the diagnostic accuracy of $^{201}$thallium-SPECT.

CHO is another PET tracer recently used for neuroradiological evaluation of gliomas, and it was reported to be a diagnostic agent which was able to differentiate between low-grade gliomas and high-grade gliomas in PET studies, but had not been used for studies of RN. Apart from RN, a high uptake of CHO is also reported in non-neoplastic lesions including brain abscess, inflammatory granulomas, tuberculosis, and some demyelinating diseases which present Gd-enhancement by MRI. A study by Ohtani et al. showed that CHO-PET did not differentiate, in particular, between low-grade gliomas and non-neoplastic lesions. Utriainen et al. described that an association between CHO uptake measured with PET and the concentration of choline containing components measured by 1H-MR spectroscopy was not statistically significant. This data suggests that CHO uptake is scarcely related to intracellular metabolite pools of phosphocholine and glycerophosphocholine. In this study, both the CHO-PET/Gd volume ratio and the CHO-PET/Gd overlap ratio of RN, AA, AO, and Gr.4 were all at levels near 100%. This suggests that there is a regional correspondence between areas of high CHO uptake on PET images and areas with Gd-enhancement on the MRI. These results imply that CHO uptake is mostly dependent on the enhancement effect, which is related to the passive diffusion of materials in regions with BBB disruption, rather than tissue biological activity, which is related to the active transport of materials.

One of the most promising modern neuroimaging protocols in this regard is MET-PET, a popular amino acid imaging modality in oncology indications. MET-PET has been a useful and reliable diagnostic tool for the differentiation of glioma recurrence from non-neoplastic lesions.
Fig. 3 Representative PET and MRI images of radiation necrosis (RN), anaplastic astrocytoma (AA), anaplastic oligodendroglioma (AO), and glioblastoma (GBM) are shown. RN: A 45-year-old man. $^{11}$C-methionine (MET)-PET/Gd volume ratio = 57.0%, MET-PET/Gd overlap ratio = 57.0%, $^{11}$C-choline (CHO)-PET/Gd volume ratio = 81.5%, CHO-PET/Gd overlap ratio = 81.5%, $^{18}$F-fluorodeoxyglucose (FDG)-PET/Gd volume ratio = 0%, FDG-PET/Gd overlap ratio = 0%, MET lesion/normal brain uptake ratio (L/N ratio) = 3.34, CHO L/N ratio = 2.03, and FDG L/N ratio = 1.57. AA: A 67-year-old man. MET-PET/Gd volume ratio = 189.4%, MET-PET/Gd overlap ratio = 100%, CHO-PET/Gd volume ratio = 121.3%, CHO-PET/Gd overlap ratio = 100%, FDG-PET/Gd volume ratio = 0%, FDG-PET/Gd overlap ratio = 0%, MET L/N ratio = 3.39, CHO L/N ratio = 7.7, and FDG L/N ratio = 1.65. AO: A 51-year-old man. MET-PET/Gd volume ratio = 172.7%, MET-PET/Gd overlap ratio = 100%, CHO-PET/Gd volume ratio = 107.8%, CHO-PET/Gd overlap ratio = 95.3%, FDG-PET/Gd volume ratio = 0%, FDG-PET/Gd overlap ratio = 0%, MET L/N ratio = 5.03, CHO L/N ratio = 14.41, and FDG L/N ratio = 1.31. GBM: A 35-year-old man. MET-PET/Gd volume ratio = 164.9%, MET-PET/Gd overlap ratio = 100%, CHO-PET/Gd volume ratio = 109.2%, CHO-PET/Gd overlap ratio = 98.7%, FDG-PET/Gd volume ratio = 74.3%, FDG-PET/Gd overlap ratio = 67.5%, MET L/N ratio = 5.21, CHO L/N ratio = 17.94, and FDG L/N ratio = 2.33. MRI: magnetic resonance imaging, PET: positron emission tomography.
Overlap ratio of RN compared with glioma recurrence showed a significant low value with previous reports. The significant low values of both the PET/Gd volume ratio and the PET/Gd overlap ratio of RN compared with glioma recurrence were characteristic only with the MET-PET and provide additional evidence for distinguishing glioma recurrence from RN.

The main mechanism for MET accumulation in RN; BBB disruption-related passive diffusion, is presumed to differ from that in tumor recurrence which is active transport affected by cell proliferation. The different mechanisms of MET accumulation for the two pathological processes are the means of potentially distinguishing glioma recurrence from RN by MET-PET. However, because of the substantial tissue biological activity in RN due to cells related to immunological and inflammation reactions and reactive glia cells with a high proliferation potential, some degree of active transport for MET may increase the MET uptake in RN. Additionally, there should be mixed tissues with both RN and residual/recurrent tumor cells around the irradiated region, because it is not feasible to completely kill the malignant glioma cells by clinical irradiation doses. These factors contribute to the continuing difficulty of distinguishing glioma recurrence from RN even using MET-PET in some cases, and further studies for a resolution of this problem are needed.

Recently, 3,4-dihydroxy-6-18F-fluoro-L-phenylalanine (FDOPA) has been utilized as another promising amino acid PET tracer for distinguishing tumor recurrence from RN. Chen et al. reported 98% sensitivity and 86% specificity for the detection of glioma recurrence using FDOPA-PET. 3'-Deoxy-3'-18F-fluorothymidine (FLT) is another recently developed PET tracer for imaging tumor cell proliferation that correlated with Ki-67 values. These tracers appear to be powerful predictors of tumor progression and survival, and comparative studies to evaluate which of the tracers, MET, FDOPA, and FLT, is the most accurate for distinguishing glioma recurrence from RN is needed.

In this study, three PET scans were taken on a single day. This introduced an increase of radiation exposure to patients compared with a single PET scan. “Cross-talk” between PET tracers during subsequent imaging was considered to be minimal, because 11C-labeled tracers such as MET and CHO have short half-lives and sufficient time was allowed between PET scans. However, from this minimal “cross-talk”, the order of PET scans (MET, CHO, FDG) could have slightly contributed to our observed result.

MET-PET appears to be superior to both CHO and FDG-PET in diagnostic accuracy for distinguishing glioma recurrence from RN on the basis of intensity as well as extent of tracer uptake volume, and it could play an important role in monitoring newly appearing Gd-enhanced lesions on MRI following neuroimaging modality for diagnosis of gliomas because of the correlation of MET-uptake with malignancy and proliferative activity in gliomas and its accumulation during glioma cell invasion. Normally, MET uptake is reported to be lower in RN than in glioma recurrence. Tsuyuguchi et al. reported that the mean L/N ratios for RN and glioma recurrence were 1.31 and 1.87. In a comparative study, Sonoda et al. showed that MET-PET was superior to 201tallium-SPECT for the differentiation of tumor recurrence from RN. In a comparative study of FDG and MET-PET, van Laere et al. reported that MET was superior to FDG as a diagnostic agent for the evaluation of glioma recurrence because of its higher sensitivity for differentiation from RN.

This study showed the superiority of MET-PET for distinguishing glioma recurrence from RN based on evaluation of intensity of tracer uptake in agreement with previous reports. The significant low values of both the PET/Gd volume ratio and the PET/Gd overlap ratio of RN compared with glioma recurrence were characteristic only with the MET-PET and provide additional evidence for distinguishing glioma recurrence from RN.

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Table 2. The best cutoff values and diagnostic accuracy for distinguishing glioma recurrence from RN

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radiotherapy in patients with malignant gliomas.

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Conflicts of Interest Disclosure

The authors have no personal, financial, or institutional interest in any of the drugs, materials, or devices in the article. All authors who are members of The Japan Neurosurgical Society (JNS) have registered online Self-reported COI Disclosure Statement Forms through the website for JNS members.

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with Positron Nuclides Approved as Established Techniques for Medical Use and Recommendations on Practices of Their Clinical Use [Supplement of 1999 Revision]. Radioisotopes 50: 38–41, 1999


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