

Detection of residual metastatic tumor in the brain following Gamma Knife radiosurgery using a single or a series of magnetic resonance imaging scans: An autopsy study

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Abstract. The aim of the present study was to investigate the usefulness of magnetic resonance image (MRI) for the detection of residual tumors following Gamma Knife radiosurgery (GKR) for brain metastases based on autopsy cases. The study investigated two hypotheses: i) Whether a single MRI may detect the existence of a tumor; and ii) whether a series of MRIs may detect the existence of a tumor. The study is a retrospective case series in a single institution. A total of 11 brain metastases in 6 patients were treated with GKR between 2002 and 2011. Histopathological specimens from autopsy were compared with reconstructed follow-up MRIs. The maximum diameters of the lesions on MRI series were measured, and the size changes classified. The primary sites in the patients were the kidneys (n=2), lung (n=1), breast (n=1) and colon (n=1), as well as 1 adenocarcinoma of unknown origin. The median prescribed dose for radiosurgery was 20 Gy (range, 18-20 Gy),

and median time interval between GKR and autopsy was 10 months (range, 1.6-20 months). The pathological outcomes included 7 remissions and 4 failures. Enhanced areas on gadolinium-enhanced MRI contained various components: Viable tumor cells, tumor necrosis, hemorrhage, inflammation and vessels. Regarding the first hypothesis, it was impossible to distinguish pathological failure from remission with a single MRI scan due to the presence of various components. Conversely, in treatment response (remission or failure), on time-volume curves of MRI scans were in agreement with pathological findings, with the exception of progressive disease in the acute phase (0-3 months). Thus, regarding the second hypothesis, time-volume curves were useful for predicting treatment responses. In conclusion, it was difficult to predict treatment response using a single MRI, and a series of MRI scans were required to detect the existence of a tumor.

Introduction

The incidence of brain metastases may be increasing, due to both improved detection of small metastases by magnetic resonance imaging (MRI) and improved control of extracerebral disease as a result of improved systematic therapy (1,2). Whole brain radiotherapy (WBRT) and Gamma Knife radiosurgery (GKR) are standard modalities of treatment for brain metastases. WBRT has been a classical treatment for almost all cases of brain metastases. GKR has been performed for a limited number and small sizes of brain metastases (≤ 4 and < 3 cm in diameter) (3,4), but recent multiple randomized trials support the use of GKR in the initial management of patients with < 5 brain metastases (5). GKR provides a high control rate due to spatially accurate and high-conformal isodose curve to the target. However, the recurrence of brain metastases has been identified to occur in 10-16% of lesions following treatment with GKR (5-9). Tumor recurrence usually presents with a progressive increase in lesion size on gadolinium-enhanced MRI (Gd-MRI) (10-13). Certain enlargements are caused by

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Abbreviations: GKR, Gamma Knife radiosurgery; Gd-MRI, gadolinium-enhanced magnetic resonance imaging; WBRT, whole brain radiotherapy; T1WI, T1-weighted imaging; T2WI, T2-weighted imaging; FLAIR, fluid-attenuated inversion recovery; DWI, diffusion-weighted imaging; FDG-PET, fluoro-deoxyglucose-positron emission tomography; PR, partial response; PD, progressive disease; SD, stable disease; BBB, blood-brain barrier; NSCLC, non-small-cell lung cancer; BED, biological effective dose

Key words: brain metastases, Gamma Knife radiosurgery, follow-up magnetic resonance imaging, pathology, autopsy

tumor progression, whereas others are caused by radiation effects without tumor progression. It is often difficult to distinguish radiation effects from tumor progression. Pathological examination, including surgical resection or biopsy, is performed to predict the cause of enlarged lesion size on MRI scans. Although there are numerous studies (14-17) comparing the pathological findings from surgical resection or biopsy with MRI findings, this method may not always be accurate because of the uncertain location information. Investigations comparing pathological findings from autopsy with MRI findings is required to acquire accurate interpretations of MRI.

The aim of the present study was to investigate the usefulness of MRI for the detection of residual tumors following GKR for brain metastases. Two hypotheses were investigated: i) Whether a single MRI may detect tumor existence; and ii) whether a series of MRIs may detect tumor existence. Follow-up MRI and pathological results in brain metastases were compared using autopsy cases to answer these questions.

Materials and methods

Study design. This study was a retrospective case series performed in a single institution; The University of Tokyo Hospital (Tokyo, Japan).

Ethics statement. The study protocol was approved by the ethical review board of the University of Tokyo Hospital (authorization no. 10857). Written informed consent was acquired from patients in advance, or from surrogates following mortality.

Patients. Patients with metastatic tumors in the brain, who were treated with GKR and received autopsy with craniotomy between August 2003 and April 2011 at The University of Tokyo Hospital were included. Patients with brain metastases without follow-up MRIs were excluded. The characteristics of the included patients are summarized in Table I.

Radiation techniques. All lesions were treated with GKR. A Leksell frame (Elekta Instruments AB, Stockholm, Sweden) was attached to secure the head of the patient in place. Subsequently, contrast-enhanced stereotactic MRI was performed to obtain precise data of the 3D coordinates of the tumors. The Leksell Gamma Plan (Elekta Instruments AB) was used to plan the treatment for tumors by stereotactic Gd-MRI and thin-slice diagnostic Gd-MRI. Treatments were delivered using the Leksell Gamma Knife B or 4C (Elekta Instruments AB). A dose of 20 Gy was prescribed for patients in whom GKR was the initial radiotherapy, and 18 Gy for patients who had undergone WBRT prior to GKR. The appropriate isodose curves were calculated considering the shape of the target tumor. Target volumes were defined based on the area of contrast enhancement plus a margin of 1 mm. For each patient, the recommendation for WBRT was based on the attending physician's discretion, using a linear accelerator with a median prescribed dose of 30 Gy in 3-Gy fractions or 40 Gy in 2-Gy fractions.

Follow-up MRI and other modalities. Patients received follow-up MRIs every 1-3 months following GKR to detect

recurrence and/or newly occurring tumors. Routine MRI sequences included axial T1-weighted imaging (T1WI), axial T2-weighted imaging (T2WI), coronal fluid-attenuated inversion recovery (FLAIR), coronal and axial Gd-MRI, axial diffusion-weighted imaging (DWI), and axial apparent diffusion coefficient. Fluorodeoxyglucose-positron emission tomography (FDG-PET) was performed to distinguish tumor recurrence from radiation effects in an enlarged lesion, or to perform systemic investigation for the patient with the adenocarcinoma of unknown origin.

Autopsy and histological examination. Autopsy was performed within 24 h after mortality. Brain sections were fixed in 20% neutral-buffered formaldehyde for 2-4 weeks. The cerebrum was cut into coronal slices 7-10-mm thick. The specimens were cut into small quadrangular pieces and embedded in paraffin, then cut again to produce 4-6- μ m-thick paraffin-embedded tissue sections. The specimens were mounted on glass microscope slides coated with 0.01% poly-L-lysine, and maintained at 37°C in an incubator overnight. Subsequent to dehydration with an alcohol and xylene series, hematoxylin staining (2 min 30 sec x3) and eosin staining (2 min x1) at room temperature was performed for primary examination. Sections from formalin-fixed paraffin-embedded tissue blocks were subjected immunohistochemical analysis using a Ventana BenchMark XT automated immunostainer (Roche Diagnostics, Basel, Switzerland). Staining conditions were as follows; samples were incubated with primary antibody incubation for 30 min at room temperature, following addition of CC1 standard (CC1-buffer, 60 min, 95°C). The applied primary antibody was Ki-67 (cat. no. M7240; Clone MIB-1; dilution, 1:200; Dako; Agilent Technologies, Inc., Santa Clara, CA, USA). Immunohistochemical results of Ki-67 were evaluated for identifying tumor proliferation. Pathological failure was defined as the detection of Ki-67-positive tumor cells by histological examination in treated brain metastases of autopsy specimens.

Analysis. First, the most recent MRI scans and the pathological results from autopsy were compared. The MRI scans were reconstructed using a free 3D imaging program, OsiriX (Pixmeo, Geneva, Switzerland), to adapt the slices of the scans to the cut slices of brain at autopsy. The last single MRI was compared with pathological specimen and pathological results in the gadolinium-enhanced area were obtained. Subsequently, the maximum diameters of the lesions were measured on a series of Gd-MRI scans, the percentage of the tumor diameter was calculated, referring to the diameter of the lesion at the time of GKR as baseline, and a time-volume curve was produced. The treatment responses of the tumors were classified as follows: Complete response (CR), disappearance of target lesion; partial response (PR), $\geq 30\%$ decrease in the diameter of the target lesion, taking as reference the baseline diameter; progressive disease (PD), $\geq 20\%$ increase in the diameter of the target lesion, taking as reference the baseline diameter; stable disease (SD), neither sufficient shrinkage to qualify for PR nor sufficient increase to qualify for PD, taking as reference the baseline diameter. The follow-up period following GKR was divided into three phases: Acute (0-3 months); sub-acute (3-6 months); and late (>6 months).

Table I. Summary of clinical data and pathological outcomes of 11 lesions.

Lesion no.	Patient age/sex	Primary site	ExCr lesion ^a	BT location	Size (mm)	Volume (cc)	Margin dose (Gy)	Isodose curve (%)	WBRT	Total BED (3Gy) ^b (Gy)	CTx ^c	Interval ^d (months)	Acute MRI ^e	Late MRI ^f	Pathology
1a	71/M	Colon (AC)	Bone	R. frontal	30	9.3	20	50	-	153	-	10	SD	SD	Failure
2a	64/M	Unknown (AC)	Unknown	L. thalamus	30	8.8	18	50	30 Gy/10 fr (pre-GKR)	186	-	1.6	SD	-	Failure
3a	77/M	Kidney (ccRCC)	Kidney	R. parietal	20	3.01	20	60	-	153	-	19	PD	SD	Remission
4a	35/F	Breast (AC)	Bone	L. frontal	15	1.88	18	40	40 Gy/20 fr (pre-GKR)	193	+	19	SD	PD	Failure
4b ^g			Bone	R. temporal	1st 18; 2nd 32	1st 1.76; 2nd 9.85	1st 18; 2nd 18	1st 40; 2nd 40	40 Gy/20 fr (pre-GKR)	319	+	1st 19; 2nd 7	1st SD	1st PD	Failure
5a	63/M	Lung (NSCLC)	Lung (primary)	L. parietal	29	9.01	20	40	40 Gy/20 fr (post-GKR)	220	+	7	SD	SD	Remission
5b			Lung (primary)	R. temporal	13	0.934	20	40	40 Gy/20 fr (post-GKR)	220		7	PR	CR	Remission
6a	59/M	Kidney (ccRCC)	-	R. frontal	25	6.6	20	40	-	153	+	20	SD	PR	Remission
6b			-	R. frontal	8	0.27	20	60	-	153	+	18	PD	CR	Remission
6c			-	R. frontal	8	0.27	20	50	-	153	+	10	PR	CR	Remission
6d			-	R. occipital	11	0.63	20	75	-	153	+	2	PD	-	Remission
Median (range)	-	-	-	-	18 (8-30)	-	20 (18-20)	50 (40-75)	-	-	-	10 (1.6-20)	-	-	-

^aExCr metastases at GKR; ^bTotal BED (3 Gy) was defined as sum of BED (3 Gy) of GKR and WBRT; ^cCTx following GKR; ^dtime between GKR and autopsy; ^eMRI classification in acute phase; ^fMRI classification in late phase; ^gLesion no. 4b was treated by GKR twice. 2nd GKR was performed 11 months after 1st GKR because of clinical recurrence. ExCr, extracranial; BT, brain tumor; MRI, magnetic resonance imaging; WBRT, whole brain radiotherapy; fr, fractions; BED, biological effective dose; CTx, chemotherapy; M, male; F, female; AC, adenocarcinoma; ccRCC, clear cell renal cell carcinoma; NSCLC, non-small cell lung cancer; R., right; L., left; GKR, Gamma Knife radiosurgery; SD, stable disease; PD, progressive disease; PR, partial response; CR, complete response.

Treatment responses were categorized by the size changes (CR, PR, SD or PD) and the follow-up period (acute or late phases). In the present study, 'temporary enlargement' was defined as an initial growth of >20%, and reduction to less than the baseline size within 3 months. To assess the total radiation effect of GKR and WBRT, the total biological effective dose (BED; 3 Gy) of the tumor margins was calculated. The present study presumed that BED (3 Gy) reflected late biological effect from radiation therapy.

Results

Patient characteristics and treatments. A total of 9 patients with 14 metastatic lesions in the brain were treated with GKR and received autopsy with craniotomy between June 1995 and June 2013 at The University of Tokyo Hospital. Brain metastases without follow-up MRIs were excluded. As a result, 6 patients with a total of 11 brain metastases were eligible for the present study. These lesions were treated with GKR between October 2002 and February 2011.

The median age at diagnosis was 63.5 years (range, 35-77 years). The male:female ratio was 5:1. Sites of primary tumors were the kidneys in 2 patients, lung in 1 patient, breast in 1 patient, colon in 1 patient and an adenocarcinoma of unknown origin in 1 patient. The numbers of brain metastases at diagnosis were 1 tumor in 3 patients, 2 tumors in 2 patients, and 4 tumors in 1 patient, respectively. WBRT was performed in 3 patients prior or subsequent to GKR. Surgery was performed in 1 patient prior to GKR. The median follow-up time was 15 months (range, 1.6-20 months). None of the patients succumbed with any clear evidence of neurological symptoms. Table I summarizes the clinical data and pathological outcomes. A total of 11 brain metastases in 6 patients were treated with GKR. The median prescribed dose was 20 Gy (range, 18-20 Gy) at the tumor margin, with a median maximal dose of 40 Gy (range, 27-50 Gy). GKR was performed twice in the same lesion in 1 brain metastasis from breast cancer (lesion no. 4b).

The pathological outcomes were 7 remissions and 4 failures. The pathological outcomes of the all lesions were the same in every patient: Either all remissions or all failures. Although 1 lesion (no. 4b) received the highest total BED (3 Gy) of 319 Gy due to repeated GKR, it did not exhibit radiation necrosis.

Comparison of the last MRI results with pathological results. Table II summarizes the associations between the details of the last MRI and pathological results. A case of pathological failure is presented in Fig. 1 (lesion no. 1a). The contrast-enhanced areas on Gd-MRI contained various pathological components, including viable tumor cells, tumor necrosis, hemorrhage, inflammation and vessels. The degree of contrast on Gd-MRI was similar in all components. All viable tumor tissue and hemorrhages demonstrated contrast enhancement. Some areas of tumor necrosis, inflammation and vessels exhibited contrast enhancement, while others did not.

Time-volume curves from a series of MRI scans. Time-volume curves determined by Gd-MRI are shown in Fig. 2, a summary of which is listed in Table III. PD indicated pathological failures in the late phase. Treatment response, remission or failure and

Table II. Comparison of the last MRI results with pathological results.

Lesion no.	Primary site	Size of lesion (mm)	Pathology	Final MRI classification	Final MRI results	Pathological results
1a	Colon	30	Failure	SD	CE+ (heterogeneity)	Degenerated tumor (center)+viable tumor (periphery)+TN+ICs
2a	Unknown	30	Failure	SD	CE+ (solid)	Viable tumor (periphery)+TN+ICs
3a	Kidney	20	Remission	SD	CE+/hetero-nodule/CE-	TN/hemorrhage/fibrosis+TN+ICs
4a	Breast	15	Failure	PD	CE+/CE-	Viable tumor (periphery)+TN+ICs/fibrosis
4b ^a		1st 18	Failure	1st PD	CE+/CE-	Viable tumor (periphery)+TN+ICs/fibrosis
5a	Lung	29	Remission	SD	CE+/CE-	ICs/lack of tissue
5b		13	Remission	CR	CE-	TN+ICs
6a	Kidney	25	Remission	PR	CE+	Scar+ICs
6b		8	Remission	CR	CE-	Fibrosis+ICs+hemosiderin-phagocytes
6c		8	Remission	CR	CE-	Fibrosis+TN
6d		11	Remission	PD	CE+ (heterogeneity)	Hemorrhage+TN

^aLesion no. 4b was treated by GKR twice due to clinical recurrence. MRI, magnetic resonance imaging; SD, stable disease; PD, progression disease; CR, complete response; PR, partial response; CE, contrast enhancement; TN, tumor necrosis; ICs, inflammatory cells.

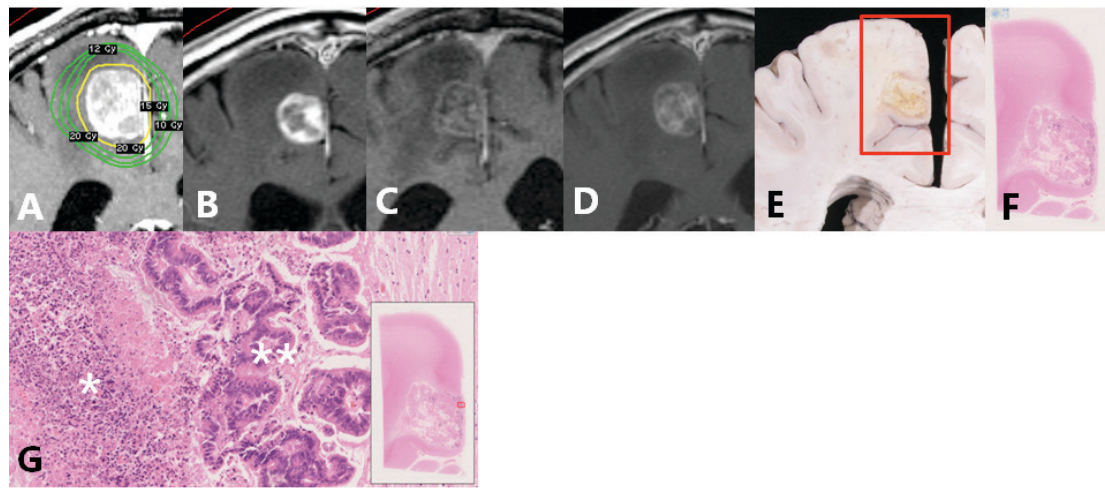


Figure 1. Images and pathological findings of one case (lesion no. 1a). (A) Dose distribution at GKR. (B) Gd-MRI 1 month following GKR. (C) Gd-MRI 3 months after GKR, classified as SD in acute phase. (D) final Gd-MRI 7 months after GKR, classified as SD in late phase. (E) Macroscopic view at autopsy 10 months after GKR. (F) Histological examination. (G) *Tumor necrosis and **viable tumor cells were observed at the periphery and center of the lesion, respectively (magnification, 20x10). GKR, Gamma Knife radiosurgery; Gd-MRI, gadolinium-enhanced magnetic resonance imaging; SD, stable disease.

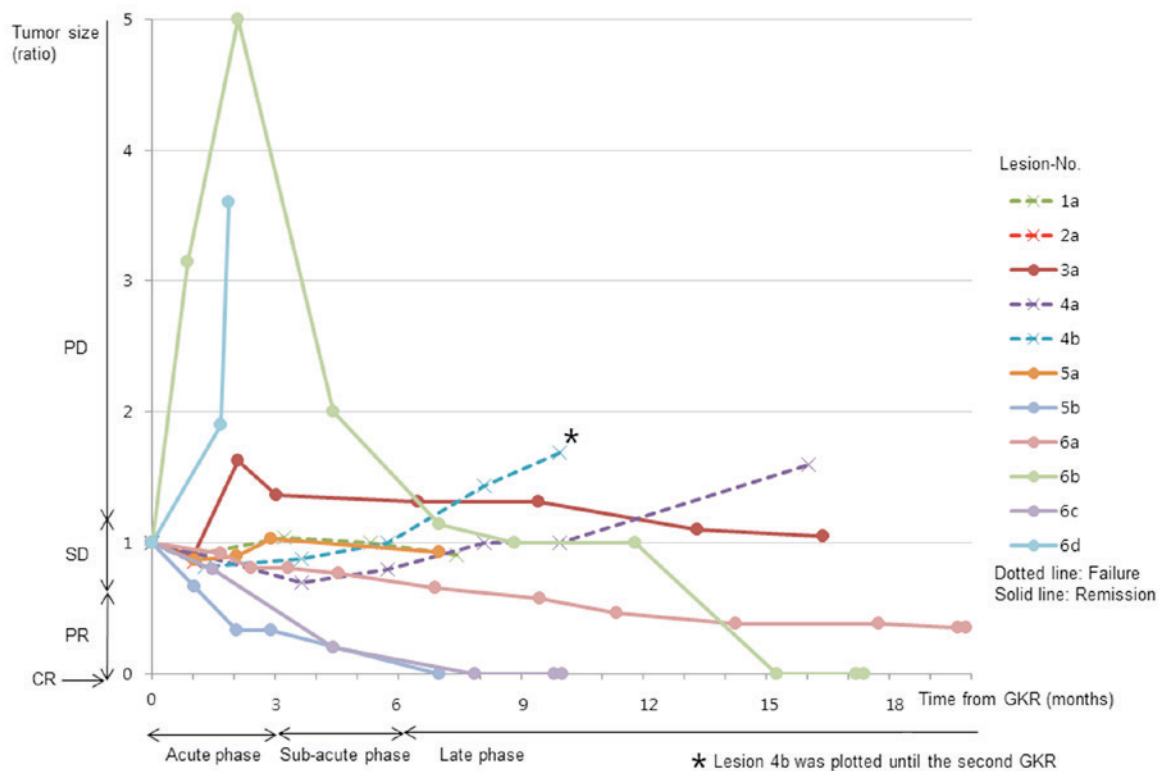


Figure 2. Time-volume curves on Gd-MRI for 11 lesions. The maximum diameters of the lesions were measured on a series of Gd-MRI scans, the percentage of the tumor diameter was calculated, referring to the diameter of the lesion at the time of GKR as baseline, and a time-volume curve was produced. Pathological failures are shown by dotted lines and pathological remissions are shown by solid lines. GKR, Gamma Knife radiosurgery; Gd-MRI, gadolinium-enhanced magnetic resonance imaging; PD, progressive disease; SD, stable disease; PR, partial response; CR, complete response.

on time-volume curves of MRI scans were in agreement with pathological findings, excluding acute PD lesions (Table II; lesion nos. 3a, 6b and 6d). A total of 2 acute PD lesions (nos. 6b and 6d) demonstrated high intensity on non-enhanced T1WI, indicating hemorrhage. Histopathological examination revealed hemosiderin-phagocytes (lesion no. 6b) and hemorrhages (lesion no. 6d) without viable tumor tissue. A partially enlarged, high-intensity area on Gd-MRI was observed in 1

acute PD lesion (no. 3a). The enlarged area on Gd-MRI was iso-intense on T2WI, indicating an expansion of contrast enhancement, and not hemorrhage. As the enlarged area on Gd-MRI reduced in size within 3 months, enlargement was judged to be temporary. Histopathological examination indicated only edema in the temporarily enlarged area. Certain SD lesions (nos. 1a and 2a) demonstrated remnant viable tumor, whereas others (nos. 3a and 5a) did not in either the acute

Table III. Association between MRI classification and pathological outcomes.

MRI classification	Pathological remission	Pathological failure	Total
Acute phase			
CR	0	0	0
PR	2	0	2
SD	2	4	6
PD	3	0	3
Late phase			
CR	3	0	3
PR	1	0	1
SD	2	1	3
PD	0	2	2
No follow-up until late phase	1	1	2
Total	7	4	11

MRI, magnetic resonance imaging; CR, complete response; PR, partial response; SD, stable disease; PD, progression disease.

Table IV. Results of DWI, FLAIR, and FDG-PET scans.

Scan findings	Pathological remission	Pathological failure	Total
DWI			
Normal intensity	7	3	10
Abnormal intensity	0	1	1
FLAIR			
Stable	7	2	9
Enlarged peritumoral high intensity	0	2	2
FDG-PET			
Normal uptake	0	1	1
Increased uptake	0	2	2
No image	7	1	8

Values indicate number of lesions. DWI, diffusion-weighted imaging; FLAIR, fluid-attenuated inversion recovery; FDG-PET, fluorodeoxyglucose-positron emission tomography.

or the late phase. Late PD lesions (nos. 4a and 4b) exhibited tumor recurrences (no. 4b).

Other sequences and modalities. Regarding DWI, FLAIR and FDG-PET scans, the examination outcomes are summarized in Table IV. From the results of the DWI scans, only 1 lesion (no. 2a) exhibited abnormal intensity among the 4 pathological failures (nos. 1a, 2a, 4a and 4b). As for FLAIR, enlarged peritumoral high intensity was observed in 2 lesions (nos. 4a and 4b) among the 4 pathological failures. Concerning

FDG-PET, increased uptake was observed in 2 lesions (nos. 2a and 4b) but not in 1 lesion (no. 4a) among the 4 pathological failures.

Discussion

To the best of our knowledge, the present study is the first to report a precise correlation between MRI and pathological results in autopsy cases of brain metastases treated with GKR. In previous studies, correlations between MRI and pathology using surgical or biopsy specimens were studied while relying on insufficient orientation; the locations of the MRI features were not adapted to the location of pathological features directly (15,18). To resolve this problem, the slices of the reconstructed MRIs were adapted to the macroscopically viewed slices of the brain in autopsy cases. This enabled the correct study of the correlations between MRI and pathological findings. The treatment method of GKR was standardized, making it easy to compare its effects. At various timings from acute to late phase, the histopathological outcomes were evaluated.

Although pathological failure was evident in 4/11 lesions (36%), no patient succumbed to brain metastases during the course of the present study. It is often hypothesized that pathological confirmation of remnant viable tumors by surgical specimen may be an overdiagnosis in view of the clinical course (15,19).

To investigate the usefulness of MRI for the detection of residual tumor following GKR for brain metastases, two hypotheses were investigated: i) Whether a single MRI scan may detect tumor existence; and ii) whether a series of MRI scans may detect tumor existence.

According to certain studies (20,21), contrast enhancement in the central nervous system is a combination of two primary processes: Vascular (intravascular) enhancement and interstitial (extravascular) enhancement. New blood vessels (angiogenesis), active inflammation (infectious and noninfectious), cerebral ischemia and pressure overload (eclampsia and hypertension) are known as extravascular enhancement. All of these changes are associated with alterations in permeability of the blood-brain barrier (BBB). In the present study, five pathological components were observed: Viable tumor cells, tumor necrosis, hemorrhage, inflammation and vessels in the area of contrast enhancement on Gd-MRI. There were abundant tumor vessels adjacent to viable tumor tissue and tumor necrosis. BBB of tumor vessels were vulnerable. Hemorrhage indicated injury of the vessels, and inflammation occasionally indicated active inflammation. As all of these components were associated with alterations in BBB permeability, contrast enhancement itself was not useful for distinguishing viable tumor tissues from the other components. Other MRI sequences, such as DWI and FLAIR, were not always useful. Thus, regarding the first hypothesis, a single MRI was not able to assess the existence of viable tumor tissue.

Of 3 acute PD lesions, 2 (lesion nos. 6b and 6d) were confirmed as hemorrhage by histopathology. These two lesions were metastases from renal cell carcinoma. Brain metastases, particularly from renal cell carcinoma, tend to exhibit bleeding as part of the natural course of the disease (22,23), and bleeding was identified in 9% of patients following GKR

in a previous study (24). Bleeding at the tumor site is not a rare event following stereotactic irradiation in renal cell carcinoma.

The third acute PD lesion (no. 3a) exhibited temporary high-intensity enlargement on Gd-MRI, which demonstrated iso-intensity on T1WI/T2WI. The histopathology indicated that the temporarily enlarged area consisted only of edema. Several previous studies (25,26) demonstrated temporary enlargement of brain metastases following stereotactic radiosurgery: One study reported a rate of 12% (11/87 cases) and a median duration of 3 months (range, 2-6 months) (25); while the other study reported temporary enlargement in 7% (5/73 cases) of all lesions following GKR (26). However, the detailed mechanism of temporary enlargement is not clear. It was hypothesized that extravascular enhancement by vessel changes and consequential increased BBB permeability caused temporary, enhancing enlargement on MRI. Park *et al* (27) suggested that endothelial cells were injured and the permeability of the vessels rose to a peak within 24 h after irradiation when normal vessels were irradiated. This increased permeability then remained for ~1 month. The magnitude of these changes depended on the radiation dose. An increase in the permeability of vessels led to invasion of inflammatory cells and fibroblasts, facilitating fibrosis (22). Similar changes may have occurred in 1 patient of the present study (no. 3a). As chronic changes of normal vessels, such as increased wall thickness, were observed in the area of temporary enlargement, these normal vessels were assumed to have presented increased permeability as a result of radiation injury in the acute phase. Following these acute changes, permeability of these normal vessels would decrease with increasing wall thickness.

In the present study, 2 acute PR lesions (nos. 5b and 6c) continued to shrink, becoming CR on the last MRI in the late phase, and these lesions were confirmed as pathological remission. Da Silva *et al* (28) identified that tumors with a histology associated with high radiosensitivity, such as breast cancer and non-small-cell lung cancer (NSCLC), tended to shrink in the acute phase. In the present study, acute shrinkage was observed in the metastases that arose from NSCLC (lesion no. 5b) and renal cell carcinoma (lesion no. 6c). Lesions from breast cancer were reduced by a small amount, but were diagnosed as SD. Tumor necrosis, inflammation and hemorrhages occurred in the acute or late phases, and exhibited contrast enhancement in the absence of viable tumor tissues.

The time-volume curves demonstrated a good association with pathological outcomes, excluding acute PD. The determination of acute PD was due to hemorrhage or temporary enlargement on Gd-MRI. Acute PD lesions decreased in size during the late phase, indicating that they should be observed for an extended period. In addition, tumor necrosis should be monitored in the late phase. There was no case of radiation necrosis in the present study, but generally it is difficult to distinguish radiation necrosis from tumor recurrence by MRI results in late PD lesions (15). Thus, as regards the second hypothesis, a series of MRI scans may detect the presence of residual tumor. For acute PD, subsequent time-volume curves were useful to distinguish pathological remission from failure.

High intensity on DWI, enlarged size of high intensity on FLAIR, and increased uptake on FDG-PET were occasionally useful for detecting tumor recurrence, but the absence of these signs did not always rule out tumor existence.

The present study was subject to certain limitations. First, it comprised a small number of patients, and more patients are required to confirm the findings. In addition, there was a selection bias due to the use of only autopsy cases. Furthermore, there was a delay between the last MRI and autopsy, and the MRI results may change within that interval. Additionally, there was no patient with radiation necrosis included in the present study.

It may be concluded that time-volume curves from a series of MRI scans are useful for predicting pathological tumor responses. Particularly for acute PD lesions exhibiting hemorrhage or temporary enlargement, subsequent time-volume curves were useful to distinguish pathological remission from failure. It is impossible to distinguish pathological failure from remission with a single MRI. This is because contrast enhancement on Gd-MRI contained viable tumor tissues, tumor necrosis, hemorrhage, inflammation and vessels.

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