Radiation-indicated cavernous hemangioma (RICH) refers to a localized vascular, tumor-like lesion that develops after cerebral radiation [1-3]. The lesion occurs mainly in children and young people, with variable latency periods after radiation treatment and diverse original lesions, including arteriovenous malformation (AVM), glioma, and metastatic tumor [4,5]. Similar to de novo cavernous hemangioma (CH), RICH appears as an enhancing lesion with popcorn-like appearance and partial hemosiderin rim on magnetic resonance imaging (MRI) and is observed histologically as a vascular-rich hemorrhagic lesion [6-8]. Thus, RICH has been regarded as a sporadic form of de novo CH that appears as a late complication of cerebral radiation [9,10].

Recently, in several studies in which RICH was compared with de novo CH, RICH was shown to have differences in clinicopathological features compared with previously known CHs [6,11-13]. Previous research has reported that RICH following stereotactic radiosurgery shows some distinguishing features on MRI such as an unilocular cystic area with some solid component and prominent perilesional edema, and de novo CH appears to have a complete hemosiderin rim and less prominent perilesional edema [6]. Furthermore, RICH is more likely to be an inactive organizing hematoma rather than a vascular malformation. Therefore, the term “radiation-induced organizing hematoma (RIOH)” was proposed to describe the lesions more appropriately and replace the term RICH, which might be a misnomer for these lesions [6]. All the authors of the current study agreed to this concept, and the term RIOH, instead of RICH, was used throughout the manuscript.
Detailed studies on RIOH are limited. In particular, RIOH lesions have been reported irrespective of radiation dose, type of malignancy, or radiation type, such as gamma knife surgery (GKS) or conventional radiation therapy (RT) [11]. However, studies to improve the understanding of RIOH lesions have not been conducted. In the present study, RIOH lesions were better defined, and their clinicopathological characteristics, including original pathology and type of radiation treatment, were compared.

MATERIALS AND METHODS

Patient selection and clinicopathological examination

Between January 2009 and May 2020, a total of 37 cases of pathologically confirmed RIOH was selected from Severance Hospital. For each case, the size of both the original tumor and RIOH was obtained. The size of original tumor was measured from preoperative imaging, and the size of RIOH were obtained through microscopic examination. The difference between the maximal diameter of primary tumor and RIOH was measured. All the pathologic slides were reviewed, and the tumor wall thickness of RIOH was measured at the thickest point of the submitted tissue under light microscope.

All medical records were reviewed, and clinical data including age at the time of RIOH detection, sex, radiologic findings, locations of the original tumor and RIOH, multiplicity, and latency period were compared between the original tumor and RIOH. A radiologist (M.P.) reviewed all MRI scans of patients, focusing on differences in radiologic findings including perilesional edema and hemosiderin rim depending on type of RT. Furthermore, information on previous treatments was collected. Cases were classified into two groups based on previous type of radiation (GKS, n = 24 vs. conventional RT, n = 13) and into four groups based on preceding pathology of the original tumor (AVM, n = 14; glioma, n = 12; metastasis, n = 4; other tumors, n = 7). The clinicopathological parameters were analyzed. For patients who had been treated multiple times or who underwent both GKS and RT, the groups and latency were divided and calculated based on timing and method of the last treatment.

Statistical analysis

Statistical analyses were performed using SPSS ver. 21.0 (IBM Corp., Armonk, NY, USA). Continuous and categorical variables were analyzed using the non-parametric Mann-Whitney U test and the chi-square test, respectively, and p < .05 was considered statistically significant.

RESULTS

Patient characteristics

A total of 37 samples was included in the study. Baseline patient characteristics are summarized in Table 1. RIOH samples were from 14 males (37.8%) and 23 females (62.2%). Twenty-four patients (64.9%) underwent GKS, and 13 (35.1%) were treated with conventional RT. Original tumor pathologies were as follows: AVM (n = 14, 37.8%), brain tumors (11 gliomas and 1 ependymoma; n = 12, 32.4%), metastasis (n = 4, 10.8%), and other tumors (3 schwannomas, two nasopharyngeal cancers, one pituitary tumor, and one craniopharyngioma; n = 7, 18.9%). The cases of nasopharyngeal cancer were subcategorized into ‘other tumors’ in this study because postoperative RT often is used for nasopharyngeal cancer, and the central nervous system area usually is included in the radiation field.

Clinicopathological differences between RIOH lesions

The overall results of the clinical and pathological comparisons are summarized in Tables 2 and 3. RIOH cases were divided into two groups based on previous type of radiation (GKS or conventional RT). Histologically, RIOH shows a hematoma-like area with a reduced number of hyalinized thin-walled vessels with fibrin and infiltrating foamy macrophages in the vessel walls. Conversely, de novo CH shows a thick, hyalinized wall without prominent macrophage infiltration (Fig. 1). The results

Table 1. Baseline characteristics of the patients (n=37)

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>No. (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age at the time of RIOH detection (yr), mean ± SD</td>
<td>46.57 ± 13.79</td>
</tr>
<tr>
<td>Sex</td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>14 (37.8)</td>
</tr>
<tr>
<td>Female</td>
<td>23 (62.2)</td>
</tr>
<tr>
<td>Original pathology</td>
<td></td>
</tr>
<tr>
<td>AVM</td>
<td>14 (37.8)</td>
</tr>
<tr>
<td>Brain tumor</td>
<td>12 (32.4)</td>
</tr>
<tr>
<td>Metastasis</td>
<td>4 (10.8)</td>
</tr>
<tr>
<td>Other tumors</td>
<td>7 (19.0)</td>
</tr>
<tr>
<td>Type of treatment</td>
<td></td>
</tr>
<tr>
<td>GKS</td>
<td>24 (64.9)</td>
</tr>
<tr>
<td>RTx</td>
<td>13 (35.1)</td>
</tr>
<tr>
<td>Tumor location</td>
<td></td>
</tr>
<tr>
<td>Frontal lobe</td>
<td>10 (27.0)</td>
</tr>
<tr>
<td>Parietal lobe</td>
<td>5 (13.5)</td>
</tr>
<tr>
<td>Temporal lobe</td>
<td>6 (16.2)</td>
</tr>
<tr>
<td>Occipital lobe</td>
<td>4 (10.8)</td>
</tr>
<tr>
<td>Other (including sellar lesion)</td>
<td>12 (32.5)</td>
</tr>
</tbody>
</table>

RIOH, radiation-induced organizing hematoma; SD, standard deviation; AVM, arteriovenous malformation; GKS, gamma knife surgery; RTx, radiation therapy.
showed significantly shorter latency in the GKS group than in the conventional RT group (5.85 ± 4.06 years vs. 11.15 ± 8.27 years, p = .046). In addition, the RIOH in the GKS group had significantly thicker tumor wall than that in the conventional RT group (693.7 ± 565.7 μm vs. 406.9 ± 519.7 μm, p = .049) (Fig. 2). Significant differences were not observed in age, tumor size of primary tumor, or RIOH. The primary tumor size did not differ between groups. However, the size of RIOH lesions tended to be larger in the GKS group than in the conventional RT group (p = .055). The multiplicity of RIOH was not significantly different between the groups. In terms of original pathology, the four groups based on preceding pathology were compared, and no significant differences were found in any of the clinicopathological parameters.

The results of the radiologic findings are summarized in Table 4. All patients with GKS-induced RIOH and 11 patients with RT-induced RIOH showed perilesional edema on T2-weighted images. Seven patients in the GKS-induced RIOH group showed subacute stage hemorrhage on T1-weighted images, and 21 showed hemosiderin rim deposit on T2-weighted images. Similarly, nine patients in the RT-induced RIOH group showed subacute stage hemorrhage, and 11 cases showed hemosiderin rim. Significant difference in radiologic image findings including perilesional edema (p = .223), subacute stage hemorrhage (p = .059) and hemosiderin deposit in MR image (p = .679), was not observed between the GKS and RTx groups.

No significant differences were found in other parameters. In the GKS group, the mean age at the time of RIOH diagnosis was 46.9 years; the mean sizes of primary tumor and RIOH were 3.35 cm and 2.01 cm, respectively; and size difference between primary tumor and RIOH was 1.34 cm. In the conventional RT group, the mean age was 45.8 years; the sizes of primary tumor and RIOH measured 3.04 cm and 1.51 cm on average, respectively; and the size difference between primary tumor and RIOH was 1.53 cm. However, the size of RIOH lesions tended to be larger in the GKS group than in the conventional RT group (p = .055).

In terms of original pathology, the four groups based on preceding pathology were compared. The mean age in the AVM, brain tumor, metastasis, and other tumors groups was 43.1 years, 43.5 years, 57.2 years, and 52.7 years, respectively. The mean sizes of the primary tumor and RIOH were 3.87 cm and 2.06 cm in the AVM group, 2.85 cm and 1.50 cm in the brain tumor group, 2.92 cm and 2.20 cm in the metastasis group, and 2.85 cm and 1.75 cm in the other tumors group, respectively. The size difference was 1.81 cm, 1.35 cm, 0.72 cm, and 1.10 cm in the AVM, brain tumor, metastasis, and other tumors groups, respectively. The mean tumor wall thickness was 714.2 μm in the AVM group, 560.0 μm in the brain tumor group, 362.5 μm in the metastasis group, and 538.5 μm in the other tumors group. The average latency in the AVM, brain tumor, metastasis, and other tumors groups was 46.9 years, 43.5 years, 57.2 years, and 52.7 years, respectively. The mean sizes of the primary tumor and RIOH were 3.35 cm and 2.01 cm, respectively; and size difference between primary tumor and RIOH was 1.34 cm. In the conventional RT group, the mean age was 45.8 years; the sizes of primary tumor and RIOH measured 3.04 cm and 1.51 cm on average, respectively; and the size difference between primary tumor and RIOH was 1.53 cm. However, the size of RIOH lesions tended to be larger in the GKS group than in the conventional RT group (p = .055).

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Three patients in the GKS group showed multiplicity, two in the conventional RT group, 1 in the AVM group, 2 in the brain tumor group, and two in the metastasis group; however, statistical difference was not found between the groups.

Table 2. Clinicopathological differences between 37 radiation-induced organizing hematomas

<table>
<thead>
<tr>
<th>Variable</th>
<th>Radiation treatment</th>
<th>Original pathology</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>GKS (n = 24)</td>
<td>RTx (n = 13)</td>
<td>AVM (n = 14)</td>
</tr>
<tr>
<td>Tumor size (cm)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Primary tumor</td>
<td>3.35 ± 1.22</td>
<td>3.04 ± 1.08</td>
<td>.387</td>
</tr>
<tr>
<td>RIOH</td>
<td>2.01 ± 0.80</td>
<td>1.51 ± 0.55</td>
<td>.055</td>
</tr>
<tr>
<td>(Size differences)</td>
<td>1.34 ± 1.33</td>
<td>1.53 ± 1.28</td>
<td>.695</td>
</tr>
<tr>
<td>Tumor wall thickness (μm)</td>
<td>693.7 ± 565.7</td>
<td>406.9 ± 519.7</td>
<td>.049</td>
</tr>
<tr>
<td>Latency (yr)</td>
<td>5.85 ± 4.06</td>
<td>11.15 ± 8.27</td>
<td>.046</td>
</tr>
</tbody>
</table>

Values are presented as mean±SD.
GKS, gamma knife surgery; RTx, radiation therapy; AVM, arteriovenous malformation; RIOH, radiation-induced organizing hematoma; SD, standard deviation.

Table 3. Correlation between multiplicity and treatment/original pathology

<table>
<thead>
<tr>
<th>Multiplicity</th>
<th>Total</th>
<th>No</th>
<th>Yes</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>GKS</td>
<td>24</td>
<td>21 (87.5)</td>
<td>3 (12.5)</td>
<td>.586</td>
</tr>
<tr>
<td>RTx</td>
<td>13</td>
<td>11 (84.6)</td>
<td>2 (15.4)</td>
<td></td>
</tr>
<tr>
<td>AVM</td>
<td>14</td>
<td>13 (92.8)</td>
<td>1 (7.2)</td>
<td>.155</td>
</tr>
<tr>
<td>Brain tumor</td>
<td>12</td>
<td>10 (83.3)</td>
<td>2 (16.7)</td>
<td></td>
</tr>
<tr>
<td>Metastasis</td>
<td>4</td>
<td>2 (50.0)</td>
<td>2 (50.0)</td>
<td></td>
</tr>
<tr>
<td>Others</td>
<td>7</td>
<td>7 (100)</td>
<td>0</td>
<td></td>
</tr>
</tbody>
</table>

GKS, gamma knife surgery; RTx, radiation therapy; AVM, arteriovenous malformation.
DISCUSSION

RIOH has been considered a sporadic form of de novo CH occurring as a late complication of cerebral radiation. RIOH occurs in children mainly after the use of cerebral radiation to treat medulloblastoma, glioma, or AVM or for prophylactic treatment of hematological malignancies such as acute lymphoblastic leukemia [4,5,14,15]. However, RIOH is rare in adults [14,16]. In a previous study with 84 cases of RIOH, the average age at diagnosis was 20.6 years, the median was 17 years, and the average latency to development of RIOH was 10.3 years, with a median of 8 years [11]. One rare case of RIOH has been diagnosed at 52 years after RT [17].

The pathophysiology of RIOH is not well-known; however, two hypotheses have been suggested: occult CHs that were previously present respond to radiation and become apparent; de

Table 4. Radiological differences between GKS-induced and conventional RTx-induced RIOH

<table>
<thead>
<tr>
<th></th>
<th>Perilesional edema</th>
<th>p-value</th>
<th>Subacute stage hemorrhage (T1 high)</th>
<th>p-value</th>
<th>Hemosiderin deposit (T2 dark rim)</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Total</td>
<td>Absent</td>
<td>Present</td>
<td>Total</td>
<td>Absent</td>
<td>Present</td>
</tr>
<tr>
<td>GKS</td>
<td>24</td>
<td>0</td>
<td>23 (95.8)a</td>
<td>9 (37.5)</td>
<td>7 (29.1)b</td>
<td>.223</td>
</tr>
<tr>
<td>RTx</td>
<td>13</td>
<td>2 (15.3)</td>
<td>11 (84.6)</td>
<td>4 (30.7)</td>
<td>9 (69.2)</td>
<td></td>
</tr>
</tbody>
</table>

GKS, gamma knife surgery; RTx, radiation therapy; RIOH, radiation-induced organizing hematoma; MRI, magnetic resonance imaging.

aOne case was undetectable on MRI; bEight cases were excluded that did not include the T1 series in preoperative MRI.

Fig. 1. The histology of radiation-induced organizing hematoma (RIOH) and de novo cavernous hemangioma (CH); Microscopically, RIOH shows a hematoma-like area composed of hyalinized vessels with fibrin and infiltrating foamy macrophages (A, B), and de novo CH consists of clusters of well-formed vascular lumens (C).

Fig. 2. Gamma knife surgery–induced radiation-induced cavernous hemangioma (RIOH) shows relatively thicker tumor walls (A) compared with conventional radiation therapy–induced RIOH (B) (Masson’s trichrome).
novel formation of the lesion in response to radiation [10]. The
devolvement mechanisms might include a series of processes, such as
vascular injury, proliferation of the vascular wall, necrosis, and
ischemia due to narrowing of the lumen. Previous studies showed
increased vascular endothelial growth factor after exposure to
radiation in rats, supporting this hypothesis [18].

Patients with de novo CH can be treated with antiepileptic
drugs and regular follow-up but are usually recommended to under-
go surgery if possible in case the symptoms worsen or the size
changes due to the risk of bleeding. In RIOH, which is regarded
dlephan form of de novo CM, surgical treatment is considered
the standard treatment option following the treatment algorithm
for de novo CM. However, if RIOH is considered a separate dis-
ease entity with different pathogenesis and clinical course than
de novo CM, conservative treatment modalities could be con-
sidered in addition to invasive surgical treatments, which could
cause neurological side effects. Therefore, several studies have
been conducted to compare the pathogenesis of RIOH and de
novo CH, and several notable results were reported. Compared
with de novo CH, RIOH was found to develop at a younger age,
symptoms at the time of diagnosis were milder, and tended to
be more multifocal. However, the prevalence of a hemorrhagic
event, the most fatal complication, was not significantly different
[11-13].

Considering the hematoma-like area and infiltration of foamy
macrophage, RIOH appears more likely to be an inactive hem-
angioma-like lesion, which might be closer to a recanalized cavi-
tary hemato ma induced by high-dose radiation than to vascular
malformation [6]. Therefore, we suggest the use of the term
RIOH rather than RICH.

In the present study, we hypothesized that RIOH would show clinical and histological differences depending on treat-
ment type or primary pathology. Based on the treatment type,
the GKS group showed shorter latency. In previous studies, sev-
eral factors affecting the duration of the latency period have
been reported, including radiation dose > 30 Gy and RT before
10 years of age [11,14,15,19], with increased risk of hemor-
rhagic event [9]. Hypothetically, the shorter latency observed in
the GKS group might be due to its requirement of a higher dose
in a smaller target area compared with conventional RT or fo-
cused damage that could accelerate tissue necrosis and tumori-
genesis.

The present study had several limitations. First, the small sam-
ple size might have led to skewed statistical results. Second, al-
though all available clinical, radiologic, and pathologic data were
collected, some parameters might be inconsistent in retrospec-
tive analysis. Third, the subjects were older in the present study.
As previously stated, RIOH occurs primarily in younger patients,
however, in the present study, the mean age of the subjects was
46.6 years, which is an older age group compared with other re-
ports (median 31.1 years [12]; mean age at the time of radiation
10.4 years with mean latency of 10.3 years [11]). Histopatho-
logical differences can exist in the RIOH of older subjects com-
pared with younger subjects and could affect the results. How-
ever, only 37 patients were included in this study, most of whom
were in their 30s or older, explaining why the results differ from
those of previous studies. Further research with a larger cohort
is needed to verify the results.

We hypothesized that the size of RIOH after conventional
RT would be larger than after GKS because conventional RT is
applied for larger lesions. However, the average size of the le-
sion tended to be larger in the GKS group, although the differ-
ce was not statistically significant. Furthermore, we hypothe-
sized that the size difference between primary tumor and RIOH
would be greater in the GKS group than in the RT group; how-
ever, significant difference was not observed. The repeated treat-
ment applied in the same lesion, the error in the measurement
of the radiological/pathological size, and the limitations of the
present study described above might have affected these results.

In summary, RIOH after GKS tended to occur earlier and
had thinner tumor wall than RIOH after conventional RT. The
original pathology of RIOH had no effect on the histologic and
clinical features of RIOH. These results suggest that the clinical
course of RIOH differs based on type of treatment. Understand-
ing the unique pathophysiology of RIOH, one of the most well-
known complications of cerebral radiation, has become more
important as the survival rate of brain tumor patients increases.
Therefore, further investigation in the form of prospective stud-
iess with larger cohorts is required to elucidate more detailed clini-
cal and histologic features of patients to provide appropriate med-
ical management and predict the clinical course more accurately.

Ethics Statement
This retrospective study was approved by the Institutional Review Board of
Severance Hospital (4-2020-0186), and patient informed consent was
waived. All procedures were performed in accordance with the 1964 De-
claration of Helsinki and its later amendments or comparable ethical stan-
dards.

Availability of Data and Material
All data generated or analyzed during the study are included in this pub-
ished article (and its supplementary information files).

Code Availability
Not applicable.
Author Contributions
Conceptualization: YJC, SHK. Data curation and interpretation: MSK, YJC, MP, SHK. Supervision: SHK, JHC, YJC. Writing—original draft: MSK, YJC. Writing—review & editing: YJC, JHC, SHK. Approval of final manuscript: all authors.

Conflicts of Interest
The authors declare that they have no potential conflicts of interest.

Funding Statement
No funding to declare.

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