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ABSTRACT

BACKGROUND AND PURPOSE: MR imaging is a useful and widely used evaluation for chordomas. Prior studies have classified chordomas into cell-dense type and matrix-rich type according to the ultrastructural features. However, the relationship between the MR imaging signal intensity and ultrastructural classification is unknown. We hypothesized that MR imaging signal intensity may predict both tumor ultrastructural classification and prognosis.

MATERIALS AND METHODS: Seventy-nine patients with skull base chordomas who underwent 95 operations were included in this retrospective single-center series. Preoperative tumor-to-pons MR imaging signal intensity ratios were calculated and designated as ratio on T1 FLAIR sequence (R_{T1}), ratio on T2 sequence (R_{T2}), and ratio on enhanced T1 FLAIR sequence (R_{EN}), respectively. We assessed the relationships among signal intensity ratios, ultrastructural classification, and survival.

RESULTS: Compared with the matrix-rich type group, the cell-dense type chordomas showed lower R_{T2} (cell-dense type: 1.90 \pm 0.38; matrix-rich type: 2.61 \pm 0.60 P < .001). The model of predicting cell-dense type based on R_{T2} had an area under the curve of 0.83 (95% CI, 0.75–0.92). In patients without radiation therapy, both progression-free survival (P = .003) and overall survival (P = .002) were longer in the matrix-rich type group than in the cell-dense type group. R_{EN} was a risk factor for progression-free survival (hazard ratio = 10.24; 95% CI, 1.73–60.79); R_{T2} was a protective factor for overall survival (hazard ratio = 0.33; 95% CI, 0.12–0.87); and R_{EN} was a risk factor for overall survival (hazard ratio = 4.76; 95% CI, 1.51–15.01).

CONCLUSIONS: The difference in MR imaging signal intensity in chordomas can be explained by electron microscopic features. Both signal intensity ratios and electron microscopic features may be prognostic factors.

ABBREVIATIONS: CDT = cell-dense type; MRT = matrix-rich type; OS = overall survival; PFS = progression-free survival; R_{TI} , R_{T2} , and R_{EN} = ratios of tumor-to-pons signal intensity in the TI FLAIR sequence, T2 sequence, and enhanced TI FLAIR sequence, respectively.

hordoma is a rare malignant tumor with an overall incidence of 0.08 per 100,000 in the United States.¹ These tumors occur mainly in the sacrum, followed by the skull base and spine.² Skull base chordomas account for 1% of all brain tumors and 4% of all primary bone tumors.³ Chordomas generally grow slowly; however, recurrence is common because en bloc resection is difficult in the

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skull base.³⁻⁵ The recurrence intervals are distinctively variable among patients due to biologic variability among and within patients. This difference also impacts the treatment plan; for example, some patients should receive radiation therapy as early as possible, and others can wait.⁶ Previous studies by our group and others⁵⁻¹² have demonstrated several risk factors related to long-term survival, such as the expression level of Ki-67, *Platelet-derived growth factor receptor* β (*PDGFR-* β) and *SNF5*, resection extent, age, postsurgery radiation therapy, and histopathologic findings. On the basis of our previous study on the electron microscopic ultra-structural characteristics of chordomas, we have classified chordomas into 2 different groups: cell-dense type (CDT) and matrix-rich

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Table 1: Key points for the differential diagnosis of cell-dense and matrix-rich types according to the ultrastructures of chordomas^a

	Cell-Dense Type	Matrix-Rich Type
Tumor cell content in a low-power field (×3000)	Densely arranged	Seldom or none
Extracellular matrix content	Relatively sparse	Abundant
Complex of rough endoplasmic reticulum and the mitochondria	Rich	Relatively seldom
Nuclei and chromatin	Abnormal nuclei with condensed, deeply staining heterochromatin	Regularly shaped nuclei, mainly with loosely arranged euchromatin
Cell junctions	Desmosomes are obvious	Few desmosomes
Collagen fibrils	Type III collagen fibrils can be found	Seldom
Exocytosis	Easily found	Seldom

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type (MRT).¹³ We also demonstrated the association between clinical outcomes and electron microscopic features of tumor specimens.¹³ We have shown that CDT is predictive of poorer outcomes, higher risk of recurrence, and shorter survival, though these findings were derived from a small sample size (n = 27).

MR imaging allows a differential diagnosis between chordomas and other tumors using signal intensity. 14,15 Recent studies have suggested that MR imaging signal intensity ratios on different sequences (ratio of tumor-to-pons signal intensity on T1 FLAIR [R_{T1}]; ratio of tumor-to-pons signal intensity on T2 [R_{T2}]; ratio of tumor-to-pons signal intensity on enhanced T1 FLAIR [R_{EN}]) are useful in predicting outcomes. Tian et al 16 revealed that higher $R_{\rm T2}$ predicted diminished tumor progression and higher $R_{\rm EN}$ predicted more rapid tumor progression; this correlation between $R_{\rm EN}$ and recurrence or progression was also shown by Lin et al 17 However, neither of these studies showed evidence for the possible explanation of signal intensity difference.

We hypothesized that ultrastructural morphologic characters could explain the mechanism of different signal intensities. Hence, this study aimed to explore the relationships among MR imaging signal intensity ratio, the electronic microscopic feature, and survival, which may provide meaningful preoperative prognostic information and help apply better individual-based treatment strategies.

MATERIALS AND METHODS

Patients

Data from 127 patients who underwent an operation by the same surgical team between July 2012 and July 2016 were reviewed retrospectively from the chordoma data base of the neuroendoscopic ward in Beijing Tiantan Hospital. The present series were classified into CDT and MRT according to the electronic features (Table 1).¹³ Overall, in contrast to the features in the MRT tumors, tumor density was high in CDT tumors; thus, the extracellular matrix was sparse. 13 Only the patients with both electron microscopy images and preoperative MR imaging available were included for further analysis; thus, 79 patients were enrolled in the present study. Among these patients, 27 cases were previously reported.¹³ The relevant clinical information was collected from the inpatient digital archives, including sex, age, tumor status (primary or recurrent tumor), presurgery radiation therapy, tumor blood supply (rich or not rich, with a rich blood supply being defined by having a tumorresection surface that bled easily and was difficult to suction cleanly, while a blood supply was that was not rich was defined by having a tumor resection surface that bled less easily and was easy to suction

cleanly), resection rate (≥90% or <90%), and histopathology (Ki-67 level, conventional chordoma, or chondroid chordoma). As a retrospective observational study of de-identified data without any additional therapy or monitoring procedures, this study did not need approval from the Ethics Committee of Beijing Tiantan Hospital.

MR Imaging Scans and Image Evaluation

The patients underwent preoperative MR imaging with a 3T MR imaging scanner (Siemens, Erlangen, Germany). Axial unenhanced T1 FLAIR and T2-weighted images were obtained with the following parameters, respectively: TR/TE range = 1850–2500/9.4–19.8 ms and 4500–6000/84–97 ms. Postcontrast scanning was performed 2 minutes after gadopentetate dimeglumine injection, with a concentration of 0.2 mL/kg (0.5 mol/L). The TR/TE ranges for the enhanced T1 FLAIR sequence were 1850–2500/9.4–19.8 ms. The section thickness was 3.0–50.5 mm.

The images were analyzed on our PACS. Operator-defined ROIs were placed in the tumor and the pons. The mean signal intensity of the tumor was normalized by dividing by the mean signal intensity of the corresponding pons on axial T1 FLAIR (R_{T1}), T2 (R_{T2}), and enhanced T1 FLAIR (R_{EN}), respectively.

Follow-Up

The follow-up period was calculated from the date of the first operation in Beijing Tiantan Hospital to the last follow-up in March 2019 or the date of death, whichever occurred first. The median follow-up time in this series was 52 months (range, 1–118 months), and 7 patients were lost to follow-up. Progression-free survival (PFS) was calculated from the date of the operation to the date of tumor recurrence or progression of residual tumor, which was confirmed by the imaging findings; and overall survival (OS) was defined as the time from diagnosis to disease-specific death or to the last follow-up time. Patients were censored at the last follow-up date if no death occurred or on the date of death if death was not disease-specific. ^{5,12}

Statistical Analysis

Statistical analyses were performed with SPSS software, Version 24 (IBM). T tests or Mann-Whitney U tests were used to analyze the continuous variables between groups. A χ^2 test was used in analyzing the distributions of categoric variables between groups. To evaluate the relationship between signal intensity ratio and electron microscopic classification, we included the signal intensity ratios (R_{T1} , R_{T2} , R_{EN}) in univariate analyses separately; multivariate analysis was performed using logistic regression. The area under the receiver operating characteristic curve was calculated to estimate the accuracy.

Table 2: Demographics of the 79 patients undergoing chordoma surgery in Beijing Tiantan Hospital between July 2012 and July 2016

	Total	MRT	CDT	Р
No. of cases	79	39	40	
Tumor status				
Primary	37	24	13	.01
Recurrent	42	15	27	
Sex				
Male	43	21	22	.918
Female	36	18	18	
Age (mean) (yr)				
Primary	47.8 ± 17.4	42.3 ± 18.2	58.2 ± 9.97	.006
Recurrent	47.1 ± 15.3	46.5 ± 16.5	47.6 ± 14.9	.828
Total	47.5 ± 16.2	43.9 ± 17.5	51.0 ± 14.3	.050
Histopathology				
Conventional	71	35	36	1.000
Chondroid	8	4	4	
Blood supply ^a				
Rich	21	7	14	.073
Not rich	55	31	24	
Resection grade				
≥90%	43	23	20	.423
<90%	36	16	20	
Survival status				
Alive	42	27	15	.018
Died	26	8	18	
NA ^b	11	4	7	
Presurgery radiation therapy				
Yes	17	5	12	.063
No	62	34	28	
Postsurgery radiation therapy				
Yes	30	14	16	.667
No	43	21	22	
NA ^c	6	4	2	
PFS (mo) ^d	40.3 ± 5.4	52.2 ± 5.7	18.7 ± 6.9	.003
OS (mo)	100.5 ± 7.1	113.9 ± 6.3	88.9 ± 9.3	.018

Note:—NA indicates not applicable.

OS and PFS were estimated using the Kaplan-Meier method, and survival curves for the 2 electron microscopic classifications were compared using the log-rank test. Statistically significant variables in the univariate analyses were included in the multivariate Cox proportional hazards regression model, using forward stepwise variable selection, to find significant predictors of recurrence and survival. A P value < .05 was considered statistically significant.

RESULTS

Clinical Characteristics and Electron Microscopic Classifications

Seventy-nine cases fit the inclusion criteria and were enrolled in the study (Table 2), which included 37 cases with newly diagnosed primary tumors and 42 cases with recurrent tumors or progression of residual tumors after prior operations. CDT chordomas were more common in recurrent tumors than in primary tumors (P=.01). There were 43 men and 36 women, and the mean age was 47.5 \pm 16.2 years. Seventy-one cases were confirmed as conventional chordomas, and 8 were chondroid chordomas. Among them, 47 patients

received adjuvant radiation therapy, while the other 32 patients received no radiation therapy. Because no medical treatment was approved for chordoma, no patients received any drug treatment.

According to the electron microscopic classification, 13 39 cases were classified as MRT and 40 cases were classified as CDT in the first operation. The MRT tumor cells were loosely arranged, the extracellular matrix was abundant, and the tumor cells showed bubble-like vacuoles. CDT tumors displayed relatively tightly arranged tumor cells, a less abundant matrix, and vacuoles not as rich as those in MRT cells (Fig 1). In the patients with primary tumors, patients with MRT tumors were younger than patients with CDT tumors (42.3 \pm 18.2 years versus 58.2 ± 9.97 years, P = .006).

Sixty-five patients underwent 1 operation during follow-up, and 14 patients underwent repeat operations due to tumor recurrence or progression of residual tumor. No patients underwent a repeat operation because of an incomplete initial resection. Among those 14 patients, 12 patients underwent 2 operations and 2 patients underwent 3 operations. Therefore, a total of 79 patients with 95 operations were included in the following analysis (Table 3). The median PFS for the patients who underwent a repeat operation was 10.0 months (range, 2–42

months). Among the patients who underwent 2 operations, 6 patients were diagnosed with CDT tumors in both operations, 3 patients were diagnosed with MRT tumors in both operations, 2 patients were diagnosed with a MRT tumor in the first operation and a CDT tumor in the second operation, and 1 patient was diagnosed with a CDT tumor in the first operation and an MRT tumor in the second operation. Among the 2 patients who had 3 operations, one was diagnosed each time with CDT tumors, while the other was diagnosed twice with CDT tumors and once with an MRT tumor.

Relationship between Signal Intensity Ratio and Electron Microscopic Classifications

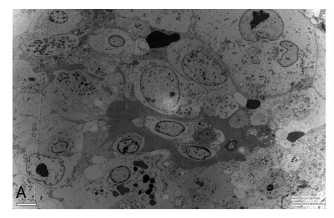
Presurgery MR images of 95 operations were pooled and analyzed. The CDT chordomas showed lower signal intensity than MRT chordomas on T2-weighted images (Table 4 and Fig 2). The R_{T1} values of CDT chordomas were higher than those of MRT chordomas. The R_{EN} value was also higher in patients with CDT than in those with MRT. When only 37 primary tumors were considered, the difference was still significant (Table 4).

^a Operative notes on 3 cases showed no specified blood supply.

^b Including both patients lost to follow-up and patients who did not die of chordomas.

^c Including the patients whose radiation therapy information was not available.

^d Patients without radiation therapy (n = 32).



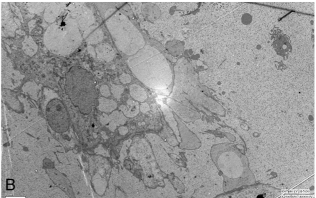


FIG 1. The electron microscopic features of the 2 classifications for skull base chordomas in low-power fields (\times 2500). *A*, In a cell-dense type chordoma, the physaliphorous cells are relatively tightly arranged and there is little extracellular matrix present. The vacuoles are not as rich as those observed in matrix-rich type tumor cells. The nuclei vary in size and condensed nuclei are found. *B*, In a matrix-rich type chordoma, the physaliphorous cells are sparse, and abundant extracellular matrix contents exist. The vacuoles are rich and apparent. The nuclei are slightly atypical.

Table 3: Details for the electron microscopic classification

Surgery No./Electron Microscopic Classification	Tumor Status ^a	No. of Cases
1 Operation $(n = 65)$	0.10.10.0	
CDT	Primary	6
	Recurrent	23
MRT	Primary	22
	Recurrent	14
2 Operations ($n = 12$)		
Both CDTs	Primary	3
	Recurrent	3
Both MRTs	Primary	3
	Recurrent	0
First MRT, second CDT	Primary	1
	Recurrent	1
First CDT, second MRT	Primary	1
	Recurrent	0
3 Operations $(n=2)$		
3 CDTs	Primary	1
	Recurrent	0
2 CDTs, 1 MRT ^b	Primary	0
	Recurrent	1

^a The tumor status when the first operation was performed in our hospital.

Table 4: Comparison of signal intensity ratios between CDT and MRT

	Electron Microscopic Classification		
Signal Intensity Ratio	CDT	MRT	P
Entire cohort ($n = 95$)			
R _{T2}	1.90 ± 0.38	2.61 ± 0.60	<.001
R _{T1}	0.63 ± 0.14	0.56 ± 0.13	.016
R _{EN}	1.19 ± 0.42	0.99 ± 0.41	.022
Primary tumors $(n = 37)$			
R _{T2}	1.99 ± 0.39	2.69 ± 0.53	<.001
R _{T1}	0.67 ± 0.18	0.55 ± 0.13	.019
R _{EN}	1.17 ± 0.42	0.89 ± 0.32	.024
Recurrent tumors ($n = 58$)			
R_{T2}	1.87 ± 0.38	2.50 ± 0.67	.001
R _{T1}	0.62 ± 0.13	0.58 ± 0.13	.307
R _{EN}	1.19 ± 0.43	1.11 ± 0.49	.526

However, when only recurrent tumors were analyzed, both R_{T1} and R_{EN} were no longer statistically different, while R_{T2} was still significantly different (Table 4).

Logistic regression analysis was then conducted to predict electron microscopic classification. A univariate analysis indicated that recurrent tumor ($P\!=\!.004$) and decreased R_{T2} ($P\!<\!.001$) and increased R_{T1} and R_{EN} ($P\!=\!.019$ and 0.027, respectively) were significant variables for predicting CDT, while presurgery radiation therapy and histopathology subtype were not significant variables ($P\!>\!.05$). From the multivariate analysis, R_{T2} was the only significant independent predictor of CDTs ($P\!<\!.001$). The area under the receiver operating characteristic curve was 0.83 (95% CI, 0.75–0.92) with only R_{T2} in the model.

Relationship between Electron Microscopic Classifications and PFS and OS

At follow-up, 26 patients (32.9%) had died of disease progression. Concerning the 32 patients without radiation therapy, 9 patients had died, including 2 patients in the MRT group and 7 patients in the CDT group. Among the 95 operations, 59 tumors (62.1%) had recurred.

When all 95 operations were analyzed together, the PFS was significantly longer in the MRT group than in the CDT group, with median PFS values of 36.0 months (interquartile range, 14–73 months) and 10 months (interquartile range, 6–42 months), respectively (P=.003). When the tumors without radiation therapy were considered, the PFS was 56 months (interquartile range, 34–undefined months) in the MRT group and 10 months (interquartile range, 5–24 months) in the CDT group (P=.003) (Fig 3A).

The OS was significantly longer in the MRT group than in the CDT group, with a median OS of 132.0 months (interquartile range, 82–undefined months) and 96.0 months (interquartile range, 48–114 months), respectively (n = 79; P = .018). When just the tumors without radiation therapy (n = 32) were considered, the MRT group showed a

^b The second operation was diagnosed as MRT.

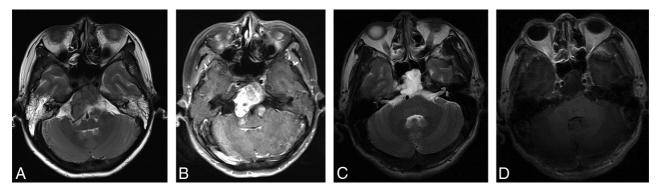


FIG 2. Representative MR images of the 2 classified chordomas. *A* and *B*, The cell-dense type chordomas. *A*, T2-weighted MR imaging shows a low signal intensity, recurrent chordoma in the clival region. *B*, Contrast-enhanced MR imaging shows the lesion with marked enhancement. *C* and *D*, Matrix-rich type chordomas with high signal intensity on T2-weighted images and low signal intensity on contrast-enhanced MR imaging, respectively.

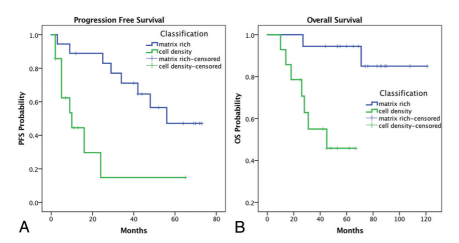


FIG 3. Kaplan-Meier analysis illustrating PFS and OS of 32 patients with chordomas without adjuvant radiation therapy. A, The patients with matrix-rich type chordomas have a significantly longer PFS than the patients with cell-dense type chordomas (P = .003). B, The OS of patients with matrix-rich type chordomas is significantly longer than that of those with cell-dense type chordomas (P = .002).

significantly longer OS than the CDT group, with 2 deaths in the MRT group compared with 7 deaths in the CDT group (P = .002) (Fig 3B).

Relationship between MR Imaging Signal Intensity Ratio and PFS and OS

To test for possible risk factors for recurrence, sex, tumor blood supply, and resection rate were included in the univariate analyses among patients not receiving radiation therapy ($n\!=\!32$). Blood supply and resection rate were associated with recurrence ($P\!=\!.015$ and .019, respectively). When these 2 parameters were combined with MR imaging signal intensity ratios and age in the multivariate analysis, R_{EN} and blood supply were significant risk factors for PFS (R_{EN} : hazard ratio = 10.24; 95% CI, 1.73–60.79; $P\!=\!.010$; blood supply: hazard ratio = 0.87; 95% CI, 1.15–13.01; $P\!=\!.029$).

Univariate Cox regression analyses indicated that postsurgery radiation therapy and resection rate (P = .023 and .036, respectively) were potentially significant prognostic factors

for OS in all 79 cases. In the multivariable analysis, R_{T2} and postsurgery radiation therapy were significant protective factors for OS (hazard ratio = 0.33; 95% CI, 0.12–0.87; P=.026; and hazard ratio = 0.12; 95% CI, 0.03–0.41; P=.001, respectively). R_{EN} was associated with significantly lower OS (hazard ratio = 0.76; 95% CI, 1.51–15.01; P=.008).

Relationship between MR Imaging Signal Intensity Ratio and Expression of Ki-67

The Ki-67 levels were available in 53 samples, which were collected from pathologic reports. The median Ki-67 level was 2% (range, 0%–20%) in 23 MRT chordomas, and the median Ki-67 level was 5% (range, 1%–20%) in 30 CDT chordomas. The expression of Ki-

67 was significantly higher in the CDT group than in the MRT group (P = .033).

DISCUSSION

Skull base chordoma can be an aggressive malignancy with poor prognosis. We previously demonstrated that electron microscopic features of chordomas were variable and could be classified into 2 morphologic types, CDT and MRT. In the present study, we show that electron microscopic classification is a valid predictor of survival. Patients with CDT had a significantly shorter PFS and OS than those with MRT, even after adjusting for the adjuvant radiation therapy. We further demonstrated that the different electron microscopic features of chordomas exhibited different MR imaging signal intensity ratios, especially $R_{\rm T2}$, and CDT chordomas had lower $R_{\rm T2}$ than MRT chordomas. In a recent radiomics study of the differentiation among primary chordoma, giant-cell tumor, and metastatic tumor of the sacrum, the radiomics model constructed on the basis of T2-weighted and contrast-enhanced T1-weighted MR imaging provided good predictive values. Another radiomics

study also found that T2-weighted and contrast-enhanced T1-weighted MR imaging provided more useful information in differentiating chordoma from chondrosarcoma, compared with T1-weighted MR imaging. An additional study reported that both T2-weighted and contrast-enhanced T1-weighted MR imaging are helpful in differentiating recurrent tumors from postoperative changes. In the present study, we additionally found that lower R_{T2} was a relatively accurate indicator for CDT (area under the receiver operating characteristic curve = 0.83; 95% CI, 0.75–0.92), which is applicable in both primary chordomas and recurrent chordomas.

In CDT chordomas, the tumor cells are rich and contact tightly with each other focally, and the extracellular matrix contents are relatively thin. Conversely, in MRT chordomas, the tumor cells are sparsely arranged and the extracellular matrix contents are abundant.¹³ These morphologic characteristics were also confirmed by hematoxylin-eosin staining under low-power microscopy.²⁰ There were numerous vacuoles of various sizes in the cytoplasm of MRT chordomas, while the vacuoles were absent or less obvious in CDT chordomas. Furthermore, the karyoplasm ratio increased occasionally in CDT tumor cells, 13 which led to reduced fluid content in tumor cells. Both high fluid content in the physaliphorous cells and extracellular matrix likely contribute to the high signal intensity in T2-weighted MR imaging in MRT chordomas. 3,14,15 Similar findings have been reported in liposarcoma and breast tumors, in which histopathologic components of loose myxoid stroma generate high T2 signal.^{21,22} To our knowledge, this is the first study focusing on the MR imaging signal intensity and electron microscopic features. The results partially-but-directly explain the pathologic basis of the different MR imaging signal intensities among chordomas.

Previous studies 15,23 demonstrated that poorly differentiated chordomas exhibited no extracellular myxoid stroma or very minimal matrix in pathologic slides and low signal intensity in T2-weighted MR imaging, and this subset of patients has decreased survival compared with those with conventional or chondroid chordomas. Although there were no poorly differentiated chordomas in our series, CDT chordomas do have a sparser extracellular matrix and higher tumor cell density, characteristics that make it reasonable to suggest that our results are comparable with those of these studies 15,23 and that R_{T2} is also qualified in poorly differentiated chordomas.

In the patients who underwent 1 operation by our team, we found a higher ratio of CDT in recurrent tumors (27/42, 64.3%) than in primary tumors (13/37, 35.1%) (Table 2). This finding indicates that recurrent chordomas are more prone to be CDT tumors than primary chordomas. In the patients who underwent >1 resection by our team, some tumors recurred as a different electron microscopic type from their previous surgical result. These findings may indicate that the tumor cells were heterogeneously arranged,²⁴ and the tiny amount of tissue in the electron microscopic sample may not exactly represent the whole tumor. With MR imaging analysis, the sampling error is greatly reduced because ROIs are selected by drawing the largest possible elliptic area within the tumor but avoiding areas of hemorrhage, necrosis, cyst, or calcification. 15,17,25 This feature may also indicate the emergence of new malignant cell populations within the recurrent tumor when the MRT tumors recurred as CDT tumors in some cases. Additionally,

we found that most of the tumors that underwent multiple resections eventually evolved into CDTs (10/14) (Table 3). A previous study reported that 2 primary sacral chordomas with high signal intensity in T2-weighted images recurred with poorly differentiated chordomas with intermediate signal intensity on T2-weighted images. ¹⁴ This occurrence suggests that decreasing T2 signal intensity may represent increasing malignancy. Recently, we confirmed that Ki-67 was associated with poor prognosis. ⁷ In the present study, we found higher Ki-67 levels in CDT tumors than in MRT tumors. The finding suggests that there is a more aggressive proliferation ability in CDT chordomas. ¹³ Furthermore, we found that CDT tumors were more likely to recur. ¹³ Taken together, we believe that the extent of malignancy increases in most chordomas when they recur, and this progression can be observed by both electron microscopic diagnosis and MR signal intensity.

MR imaging is important not only for diagnosing and differentiating disease but also in predicting prognosis. ^{3,16,24} Among the 32 patients who did not undergo adjuvant radiation therapy in the present series, R_{EN} and tumor blood supply were demonstrated as independent factors for PFS. Tian et al¹⁶ and Lin et al¹⁷ have also had similar results, which suggested that R_{EN} was a risk factor for PFS. Tian et al¹⁶ also found that tumors with higher R_{EN} had more tumor blood supply. A recent radiomic analysis of 148 skull base chordomas showed that both radiomic signature (which was based on preoperative axial T1 FLAIR, T2-weighted, and enhanced T1 FLAIR) and blood supply were independent markers of tumor progression. ²⁵ These 4 independent studies provide convincing evidence to conclude that the high R_{EN} and rich blood supply of chordomas can be used as prognostic indicators for rapid recurrence or progression.

We also demonstrated that R_{T2} and postsurgery radiation therapy are protective factors for survival and that R_{EN} is a risk factor for OS. Radiation therapy has been recommended as a standard procedure in chordoma treatment,² and many studies also showed that adjuvant radiation therapy improves outcomes.^{4,5,26} To our knowledge, the correlation between signal intensity ratio and OS in chordomas has not been studied previously, and the precise mechanism of our findings needs further research. However, considering that 79 cases are a relatively large sample size for the low incidence and prevalence of chordoma and the statistical significance of our findings, our results are meaningful. Therefore, MR imaging signal intensity ratios are useful clinical indicators for risk stratification of patients with chordomas or treatment-plan ranking.

Limitations

The main limitation of this study is that the time interval between the operation and adjuvant radiation therapy was not available in the present data base, and radiation therapy most likely positively influences recurrence intervals. Furthermore, radiation therapy has been shown to impact the MR imaging enhancement parameters. Therefore, we were unable to control for the effect of radiation therapy in the progression-free survival models. Another limitation is that this is a retrospective study for which some clinical information was missing; for example, the tumor blood supply of 3 tumors was not recorded in the archives. A study showed that lower ADC values may predict tumor

progression in postsurgery chordomas²⁷ and may also predict tumor-control probability in skull base chordomas treated with carbon ion therapy.²⁸ This probability suggests that the relationship between ADC values and electronic microscopic characteristics will also be worth investigating. However, diffusion-weighted MR imaging was previously not routinely performed in our hospital; therefore, ADC values were also not available in the present data base. These deficiencies will be remedied in our future study.¹⁵

CONCLUSIONS

MR imaging signal intensity ratio could distinguish CDT from MRT preoperatively, particularly $R_{\rm T2}$ values. The electron microscopic features may be the basis of the MR imaging signal intensity difference. Patients with CDT chordomas have significantly shorter PFS and OS than those with MRT chordomas. A higher $R_{\rm T2}$ correlates with an increased likelihood of MRT chordoma and longer survival, while $R_{\rm EN}$ is an independent adverse prognostic factor for both PFS and OS.

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