TITLE

Magnetic Resonance Imaging Features of Gemistocytic Astrocytoma

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IRB Statement

Ethics approval was obtained from both institutions (Royal Melbourne Hospital, Human Research Ethics Committee; Austin Hospital, Office for Research).

Short Running Title

MRI Features of Gemistocytic Astrocytoma **Ianusc** \geq Auth

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- 7
- 8 ABSTRACT
- 9 Introduction

10 Gemistocytic astrocytoma is the second most common subtype of World Health Organization

- 11 grade 2 astrocytoma, but has a worse prognosis than other grade 2 lesions. We aim to
- 12 describe the MR imaging features of histopathologically proven gemistocytic tumours.
- 13
- 14 Methods

Ethics approval was obtained from both institutions. Patient consent was not required for this
retrospective study. We reviewed MR imaging findings of 16 consecutive cases of

17 histopathologically proven gemistocytic astrocytoma and anaplastic astrocytoma with

- 18 gemistocytic features.
- 19

20 **Results**

Average patient age was 48 years, with a 3:1 male to female ratio. Based on our series, the typical appearance of a gemistocytic astrocytoma is a large, heterogeneous mass most commonly supratentorial and lobar. Regions of cyst formation, partial signal suppression on FLAIR images and contrast enhancement are all common features. Additionally, contrary to previous literature that describes gemistocytic astrocytoma as a purely supratentorial lesion, we present two cases of gemistocytic astrocytoma involving the brainstem.

27

28 Conclusions

29 The possibility of gemistocytic astrocytoma should be considered in patients presenting with

- 30 large heterogeneous tumours that have regions of cyst formation, partial FLAIR suppression
- and contrast enhancement. This may be especially useful in reconciling a lesion with high-

- 32 grade MR imaging features with low grade histopathology. An infratentorial location does
- 33 not preclude the diagnosis of gemistocytic astrocytoma.
- 34

35 **KEYWORDS**

Astrocytoma; Brain Neoplasms; Gemistocytic Astrocytoma; Magnetic Resonance Imaging;
Radiology

38 INTRODUCTION

Although gemistocytic astrocytomas are the second most common histological subtype of
diffuse astrocytoma behind fibrillary astrocytoma, they are relatively uncommon, accounting
for 8 to 29% of all such tumours.¹⁻⁶ They are histologically characterized by a significant
population of gemistocytic astrocytes on a background of fibrillary astrocytes. Initially
thought to be reactive cells, it has since been proven that the gemistocytes are neoplastic,
sharing a p53 mutation with other neoplastic astrocytes.^{7, 8}

- The current World Health Organization (WHO) classification for gemistocytic 45 astrocytoma is grade 2. However, a 1991 study by Krouwer et al. showed that prognosis for 46 astrocytoma with a gemistocytic component was the same irrespective of a background of 47 fibrillary (WHO grade 2) or anaplastic (WHO grade 3) cells, and that as little as 5% 48 gemistocyte tumour cells is sufficient to negatively impact prognosis.⁹ It has thus been 49 proposed that all gemistocytic astrocytoma be clinically managed as grade 3 anaplastic 50 astrocytoma, and potentially even reclassified.⁹ Given the well-known limitations of sampling 51 error in the histological assessment of astrocytomas, being able to prospectively suggest the 52 diagnosis based on imaging could alter management.¹⁰ 53
- While there have been several studies analysing histopathological features of these tumours, there is a paucity of literature on the radiological appearance.¹¹ What literature is available is limited to single case reports and one clinico-pathologically focused study that briefly describes only some imaging findings.^{4, 12, 13} These publications, as well as anecdotal experience, suggest that gemistocytic astrocytomas have significantly different imaging features to the more common diffuse fibrillary astrocytomas.
- We aim to describe the MR imaging features of histopathologically provengemistocytic astrocytomas, and identify common or specific findings.
- 62
- 63 METHOD

64 **Patient Population**

Ethics approval was obtained from both institutions. Patient consent was not required for this 65 retrospective study. Patients were identified by a search of histopathology reports at two 66 tertiary hospitals for the keywords "gemistocyte" and "gemistocytic". The search 67 encompassed from 2007 (the introduction of the PACS at these institutions) to 2014. All 68 grade 2 or grade 3 astrocytomas with a diagnosis of gemistocytic astrocytoma were included. 69 Astrocytomas with a reported component of gemistocytic astrocytes were also considered for 70 review, regardless of whether they were classified as gemistocytic astrocytomas in the final 71 diagnosis. 27 cases were identified for initial analysis. Patient demographic information was 72 73 also collected.

74

75 Pathology Review

Cases were reviewed independently by two neuropathologists with 32 and 12 years 76 experience, who were aware of the purpose of the study. Tumour grade using WHO criteria 77 78 was also assessed. The pathology specimens were assessed to confirm the presence of 79 gemistocytes and to quantify the percentage of gemistocytic, fibrillary and protoplasmic 80 astrocytes. Disagreement was resolved by consensus. It was determined that any tumour with <5% gemistocytes could not be considered a gemistocytic astrocytoma, and would be 81 82 excluded. No cases were excluded for this reason. From the initial 27 cases, 16 were found to consist entirely of gemistocytes and fibrillary astrocytes. Of the remaining 11 cases, 10 were 83 84 determined to be mixed gemistocytic/protoplasmic astrocytoma, and one was felt, in retrospect, to represent a WHO grade 4 tumour. As both these entities are known to have 85 86 heterogeneous imaging appearances, these patients were excluded.

87

88 Imaging Sequences

Seven patients were scanned on 1.5T Signa Horizon Lx (General Electric, Milwaukee, WI). 89 90 Six patients were scanned on a 1.5T Magnetom Avanto (Siemens Medical Solutions, Erlangen, Germany). Two patients were scanned on a 1.5T Signa Horizon Echospeed Plus 91 (General Electric, Milwaukee, WI). One patient was scanned on a 3T TIM TRIO (Siemens 92 Medical Solutions, Erlangen, Germany). Exact protocols varied by machine, and evolved 93 94 over time. Two patients had only post-contrast T1 weighted images (T1WI) volumetric studies available, one patient did not have any post contrast sequences and one patient had 95 dual-echo imaging rather than T2 weighted images (T2WI) and fluid attenuation inversion 96 recovery (FLAIR). The remaining 15 patients all had T2WI (repetition time (TR)/echo time 97 (TE) = 3360-6400 / 93.62-163), FLAIR (TR/TE = 8000-9002 / 87 - 137.50) and pre- and 98

- post-contrast T1WI (TR/TE = 440-640 / 7.38-20). T2* was available for 15 patients and
 diffusion weighted imaging (DWI) was available for 13 patients.
- 101

102 Imaging Review

103 All cases were reviewed independently by two neuroradiologists with 21 and 7 years

- 104 experience. The reviewers were aware of the purpose of the study but blinded to
- 105 histopathological details and patient identity. Disagreement was resolved by consensus. The
- 106 cases were assessed for tumour size (axial T2/FLAIR signal abnormality), location, midline
- and cortex involvement, signal characteristics, FLAIR suppression, contrast enhancement,
- 108 apparent diffusion coefficient (ADC) of enhancing and non-enhancing regions, and
- 109 susceptibility.

FLAIR suppression was defined as signal reduction compared to the T2 images, but less than cerebrospinal fluid (CSF). Tumour regions that suppress to the same degree as CSF were described as cystic regions. Midline involvement was determined if any part of the tumour extended into the corpus callosum, anterior or posterior commissures, or brainstem on FLAIR or T1 post-contrast images. Susceptibility was defined as loss of signal on T2* images.

116

117 **RESULTS**

118 **Patient Population**

The mean age of our patients was 50 years (48 years for grade 2, 51 years for grade 3), with
an age range of 29 to 70 years (37 to 55 years for grade 2, 29 to 70 years for grade 3). Grade
2 tumours had a male predominance, with 6/8 (75%) cases, while in grade 3 tumours, only
3/8 (38%) cases were in male patients.

123

124 Pathology Review

Half (8/16) of the tumours were grade 2, compatible with a classical diagnosis of

- 126 gemistocytic astrocytoma. The remaining 8 tumours were grade 3 astrocytomas, classically
- 127 defined as anaplastic astrocytoma with gemistocytic features. The average percentage of
- gemistocytes present was 67% (range 20% to 90%) for grade 2 tumours, and 62% (range
- 129 10% to 88%) for grade 3 tumours. As mixed tumours (those with protoplasmic components)
- 130 were excluded, all included tumours were on a background of fibrillary astrocytes.
- 131

132 Imaging Review

The imaging characteristics are presented in Table 1. Grade 2 tumour size ranged from 32 to 133 73 mm (average of axial diameter), with a similar size range for grade 3 tumours. 6/8 (75%) 134 of grade 2 tumours were supratentorial and lobar, with 2/8 (25%) involving the brainstem, 135 one as the sole site of disease and the other multifocal with supratentorial lesions. All grade 3 136 tumours were supratentorial, and 7/8 (88%) were lobar with a single grade 3 tumour arising 137 in the thalamus. Both brainstem lesions were diagnosed on biopsy only and histopathological 138 review assigned both as grade 2. Grade 2 tumours showed midline involvement in 5/8 (63%) 139 cases, and 6/8 (75%) showed cortical involvement. For grade 3 tumours, midline involvement 140 141 was present in 3/8 (38%) cases, and cortex was involved in 7/8 (88%) cases.

All lesions were T1 hypointense. Most (14 patients) had T2WI and FLAIR imaging available. These all showed high T2 signal, the majority of which (6/7 grade 2; 4/7 grade 3) was heterogeneous. In 3/7 (43%) grade 2 tumours and 2/7 (29%) grade 3 tumours, welldefined cystic regions were demonstrated. This usually constituted only a small percentage of the whole tumour (5-25%), although one grade 2 case was predominantly cystic (>75%). 6/7 (86%) grade 2 tumours and 3/7 (43%) grade 3 tumours showed partial FLAIR suppression of ill-defined markedly T2 hyperintense regions.

Enhancement was present in 75% (6/8) grade 2 cases and 57% (4/7) grade 3 cases.
The most common enhancement pattern was predominantly peripheral and nodular. Mild or
avid enhancement was seen in equal numbers.

Diffusion restriction was infrequently seen, present in only 2/8 (25%) grade 2 cases,
and 2/7 (29%) grade 3 cases. ADC was lower in enhancing, solid regions of tumour
compared to non-enhancing areas. No susceptibility was seen on T2* images to indicate
presence of blood or calcification in any lesion.

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157 DISCUSSION

We present the imaging characteristics of 16 cases of gemistocytic tumours: 8 grade 2 gemistocytic astrocytomas, and 8 anaplastic astrocytomas with gemistocytic features. This retrospective case series is, to the best of our knowledge, the largest and first dedicated attempt to document the MR imaging features of this tumour. While imaging features have been touched upon in other papers, the literature is limited to either case reports or clinicopathological papers where only some imaging features are discussed.¹²⁻¹⁴

164

165 Background

The literature reports that gemistocytic astrocytoma accounts for between 8% and 166 29% of WHO grade 2 astrocytomas.¹⁻⁶ The variability in reported incidence likely results 167 from absence of clear diagnostic criteria. The percentage of gemistocytes required to give the 168 diagnosis has not been clearly defined. Historically, an arbitrary limit of at least 20% 169 gemistocytes has been used to define this lesion, with some studies further subcategorizing 170 tumours with more than 60% gemistocytes as, somewhat non-intuitively, "pure gemistocytic 171 astrocytomas".⁹ It is not known whether the percentage of gemistocytes is prognostically 172 important, with two studies showing no difference in progression-free survival or overall 173 survival between tumours with greater or less than 20% gemistocytes.^{8, 13} On the other hand, 174 a study by Watanabe et al has shown more rapid progression in tumours with as little as 5% 175 gemistocytes.¹⁵ 176

Gemistocytic astrocytomas are currently considered a WHO grade 2 lesion. The 177 presence of anaplasia precludes the classical diagnosis, with tumours demonstrating anaplasia 178 being defined as anaplastic astrocytoma regardless of the percentage of gemistocytes.¹¹ 179 Despite this WHO classification, gemistocytic astrocytomas have a worse prognosis than 180 other low-grade gliomas, with several studies showing higher recurrence and progression 181 rates^{2, 5} and worse overall survival.^{1, 4, 6} Conversely, grade 2 gemistocytic astrocytomas and 182 grade 3 anaplastic astrocytoma with gemistocytic features have been shown to have similar 183 survival.9 184

Given the questions surrounding impact on survival in the different classifications of 185 gemistocytic tumours, we chose to examine the imaging features of both grades of tumour. 186 Our results show no difference in MR characteristics between grade 2 and grade 3 lesions, 187 which may be seen as further evidence of blurring of the distinction between grade 2 188 gemistocytic astrocytomas and anaplastic astrocytoma with gemistocytic features.⁹ 189 Regardless, the poorer prognosis of gemistocytic astrocytoma relative to other grade 2 lesions 190 highlights the value of prospective diagnosis on imaging for treatment planning, especially 191 given the well-known limitations of sampling error and inter-observer variability in 192 histological assessment of gliomas.¹⁰ 193

194

195 Patient Population

Patients diagnosed with gemistocytic astrocytomas have been reported to be older
than those with other diffuse astrocytomas, with an average age at diagnosis in the 40s,
approximately 10 years older than those with other diffuse low-grade gliomas.^{1,4} The
demographics of patients in our study conforms to previous literature, with an average age of

48 years. While some authors have postulated that this older age may contribute to the worse
prognosis in gemistocytic astrocytomas, as age is a strong independent poor prognostic factor
in astrocytoma,⁴ other studies show that the prognosis remains worse than for fibrillary

203 tumours even when stratified for age. 1

Gemistocytic astrocytomas also have a higher male predominance than other diffuse astrocytomas, with a 3:1 male to female ratio, compared with 1.5:1 for fibrillary astrocytoma.^{4,9} This is confirmed with our gemistocytic astrocytomas, though we showed a female predominance in anaplastic astrocytoma with gemistocytic features, possibly due to small sample size.

209

210 Imaging Features

Gemistocytic astrocytoma has historically been described as exclusively 211 supratentorial and most commonly found in the frontal lobes.⁹ This lobar distribution follows 212 many supratentorial neoplasms, and likely reflects nothing more than the larger size of the 213 frontal lobe compared to other lobes. We also describe two cases where gemistocytic 214 astrocytoma involved the brainstem (Figures 1 and 2). In one case, this was the sole location 215 of disease, and in the second there was also multifocal frontal lobe disease. Both cases were 216 217 grade 2 lesions (classic gemistocytic astrocytoma). Search of the literature reveals no previous cases of brainstem gemistocytic astrocytoma. 218

WHO grade 2 diffuse fibrillary astrocytomas are described in the literature as relatively homogeneous T2 hyperintense lesions involving both grey and white matter.¹⁶⁻¹⁸ In contrast, all but one of our gemistocytic astrocytomas showed signal heterogeneity on T2 weighted imaging.

While enhancement is not a feature of grade 2 fibrillary astrocytomas, it has been 223 noted in some grade 2 protoplasmic astrocytomas and grade 2 oligoastrocytomas.¹⁶⁻¹⁹ When 224 present in these lesions, however, it is usually only mild.^{19, 20} Our finding of frequent (75%) 225 enhancement (Figure 3), which is as often avid as it is mild, could result in gemistocytic 226 astrocytoma being mistaken for a higher grade lesion on imaging, and if not recognized, 227 potentially lead to the belief that lower histological grade was due to under-sampling.¹⁰ As 228 such, knowledge that the presence of gemistocytes is frequently associated with significant 229 enhancement will help in reconciliation of imaging and histological findings. 230 In contrast, grade 4 glioblastomas frequently have complete peripheral nodular 231 contrast enhancement, surrounding central areas of necrosis.^{21, 22} Although contrast 232

enhancement was common in gemistocytic astrocytomas, it is often mild, and when morepronounced did not have a continuous peripheral ring.

- Our finding of 43% of cases showing well-defined cystic regions (Figure 4) supports the study of Yang et al. (2003), which demonstrated regions of cystic change in 44% of their 25 selected cases. This is a feature more commonly associated with higher grade tumours,²²
- but we show it to be a common finding in gemistocytic astrocytoma.
- Partial FLAIR suppression of ill-defined markedly T2 hyperintense regions (Figures 4
 and 5) absent in fibrillary astrocytomas, has been described in protoplasmic astrocytomas and
 dysembryoplastic neuroepithelial tumours (DNETs).^{19, 23} It is not a described feature in
 glioblastomas, even when the tumour has a significant non-enhancing component.²⁴
- Histopathologically, gemistocytes are described as cells with "voluminous,
 homogeneous, slightly eosinophilic cytoplasm with few branching processes and an eccentric
 nucleus"²⁵ (see Figure 6 for an example). Although this cellular feature may in part account
 for FLAIR suppression, this imaging feature is encountered in other lesions without
 gemistocytes, so it is likely the other factors are contributing, likely within the extracellular
 milieu. Future studies aimed at determining a histopathological-radiological correlation for
 FLAIR suppression would be instructional.
- Diffusion restriction was seen in less than half of cases. This features has previously been ascribed to increased cellularity in gliomas²², and we noted lower ADC values in enhancing regions of tumour compared to non-enhancing regions, suggesting the enhancing regions have higher cellularity.
- Despite this study being the largest series of gemistocytic tumours reported in the radiology literature, a limitation is the small number of cases. Our finding of no difference on MR imaging between grade 2 and 3 lesions, for example, may merely represent small sample size. A minority of our cases were diagnosed on biopsy rather than resection, and the possibility of non-representative biopsies must also be acknowledged. This is particularly important with regard to the two brainstem gemistocytic astrocytomas we report, both diagnosed on biopsy alone.
- 261

262 **Conclusions**

Gemistocytic astrocytoma is a tumour with a worse prognosis than other WHO grade 264 2 astrocytomas. Prospective diagnosis may affect management, and help reconcile differences 265 between histopathological grade and radiological appearance. Based on our series of both 266 classical and grade 3 gemistocytic lesions, the possibility of gemistocytic tumour should be

- 267 considered in patients presenting with large, heterogeneous lesions that have regions of cyst
- 268 formation, partial FLAIR suppression and contrast enhancement.
- 269

270 COMPETING INTERESTS

- 271 None to disclose.
- 272
- 273 FUNDING INFORMATION

- 274 None to disclose.
- 275 **REFERENCES**

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Fig. 1 – 72-year-old-female with WHO grade 2 gemistocytic astrocytoma. T2 (a), FLAIR (b) and T1WI post-contrast axial (c) and coronal (d) images, showing an enhancing mass in the midbrain and pons. Note central FLAIR suppression. This patient also had lesions in both frontal lobes (not shown).

Fig. 2 – 38-year-old female with WHO grade 2 gemistocytic astrocytoma. T2 (a), FLAIR (b) and T1WI post-contrast axial (c) and coronal (d) images, showing a small, enhancing lesion in the brainstem (medulla oblongata, arrow).

Fig. 3 – T1-weighted images post contrast, indicating different patterns of enhancement (arrows). (a) Peripheral nodular enhancement in a 37-year-old male with WHO grade 2 gemistocytic astrocytoma. (b) Heterogeneous region of enhancement in a 29-year-old male with WHO grade 3 anaplastic astrocytoma with gemistocytic features.

Fig. 4 – 37-year-old male with WHO grade 2 gemistocytic astrocytoma. T2WI (a) and FLAIR (b) images, showing regions of cyst formation (arrow) and ill-defined partial FLAIR suppression (asterisk).

Fig. 5 – 29-year-old male with WHO grade 3 anaplastic astrocytoma with gemistocytic features. T2WI (a) and FLAIR (b) images, showing regions of cyst formation (arrow) and ill-defined partial FLAIR suppression (asterisk).

Fig. 6 – 32-year-old female with WHO grade 3 anaplastic astrocytoma with gemistocytic features. 400 x magnification hematoxylin and eosin (H&E) stain showing the typical appearance of neoplastic gemistocytes, with large volume pink-staining cytoplasm and eccentric nuclei (arrows).

Feature	Grade 2 Gemistocytic astrocytoma (n=8)	Grade 3 Anaplastic astrocytoma with gemistocytic features (n=8)
Location	Frontal (5/8), temporal (1/8),	Frontal (3/8), frontotemporal

Table 1: Imaging features by tumour WHO grade.

	brainstem (1/8), multifocal*	(3/8), parietal (1/8),
	(1/8)	thalamus (1/8)
Cortex	Involved (6/8), uninvolved	Involved (7/8), uninvolved
	(2/8)	(1/8)
Midline	Involved (5/8), uninvolved	Involved (3/8), uninvolved
	(3/8)	(5/8)
Size - mean diameter (range)	53 mm (32 to 73 mm)	56 mm (36 to 70 mm)
T1 signal	hypointense (8/8)	hypointense (7/7)
T2 signal	hyperintense (7/7)	hyperintense (7/7)
T2 characteristics	heterogeneous (6/7),	heterogeneous (4/7),
	homogeneous (1/7)	homogeneous (3/7)
Cystic regions (percentage	None (4/7), 5-25% (2/7),	None (5/7), 5-25% (2/7)
of total tumour)	>75% (1/7)	
Ill-defined FLAIR	Yes (6/7), No (1/7)	Yes (3/7), No (4/7)
suppression		
Enhancement	None (2/8), Mild (3/8),	None (3/7), Mild (2/7),
	Significant (3/8)	Significant (2/7)
Pattern of enhancement	Peripheral nodular (4/6),	Heterogeneous region (2/4),
	Solid region (1/6), Wispy	Peripheral Nodular (1/4),
	(1/6)	Wispy (1/4)
Diffusion restriction	No (4/6), Yes (2/6)	No (5/7), Yes (2/7)
ADC of non-enhancing	1590 (1535 to 1663)	1107 (554 to 1434)
region – mean (range)		
ADC of enhancing region –	1099 (900 to 1310)	983 (850 to 1115)
mean (range)		· · · /

Note. *multifocal case involved both frontal lobe and brainstem; ADC are expressed as 10⁻⁶ mm²/s







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