The Impact of MRI Features and Observer Confidence on the Treatment Decision-Making for Patients with Untreated Glioma

Paulina Due-Tønnessen¹,²*, Marco C. Pinho³, Kyrre E. Emblem⁴, John K. Hald¹, Masafumi Kanoto⁵, Andreas Abildgaard⁶, Donatas Sederevicius⁴, Inge R. Groote⁴, Otto Rapalino⁶ & Atle Bjørnerud⁴,⁷

In a blind, dual-center, multi-observer setting, we here identify the pre-treatment radiologic features by Magnetic Resonance Imaging (MRI) associated with subsequent treatment options in patients with glioma. Study included 220 previously untreated adult patients from two institutions (94 + 126 patients) with a histopathologically confirmed diagnosis of glioma after surgery. Using a blind, cross-institutional and randomized setup, four expert neuroradiologists recorded radiologic features, suggested glioma grade and corresponding confidence. The radiologic features were scored using the Visually AcceSAble Rembrandt Images (VASARI) standard. Results were retrospectively compared to patient treatment outcomes. Our findings show that patients receiving a biopsy or a subtotal resection were more likely to have a tumor with pathological MRI-signal (by T2-weighted Fluid-Attenuated Inversion Recovery) crossing the midline (Hazard Ratio; HR = 1.30 [1.21–1.87], P < 0.001), and those receiving a biopsy sampling more often had multifocal lesions (HR = 1.30 [1.16–1.64], P < 0.001). For low-grade gliomas (N = 50), low observer confidence in the radiographic readings was associated with less chance of a total resection (P = 0.002) and correlated with the use of a more comprehensive adjuvant treatment protocol (Spearman = 0.48, P < 0.001). This study may serve as a guide to the treating physician by identifying the key radiologic determinants most likely to influence the treatment decision-making process.

Gliomas remain the most common primary brain tumors in adults¹,² and are classified according to histopathological features using the World Health Organization (WHO) grading system³. While contrast-enhancement Magnetic Resonance Imaging (MRI) is a supplement for glioma characterization and monitoring⁴,⁵, surgery as the first-line defense makes MRI unlikely to replace histopathology as the benchmark for classification of tumor type and grade. Instead, MRI may better capture the heterogeneity and structural complexity of the disease, and presents an attractive source of complementary information for treatment decisions⁶.

However, which radiologic features are systematically associated with the subsequent patient treatment plan, and moreover, how observer confidence influences the decision-making process are underreported. Performing a study to address these issues is not straightforward because radiologic features of glioma grades, unlike histopathology, do not follow the same level of quantification, but are arguably subjective. Moreover, functional information from diffusion MRI and dynamic susceptibility contrast (DSC)-MRI, provide data that may not fit histopathologic classification nor traditional MRI evaluation. While perfusion MRI may influence a hypothetical management plan⁷, to our knowledge, there is a paucity of literature addressing the extent to which such functional MRI techniques affect treatment decisions in a real clinical setting⁸–¹⁰.

¹Department of Radiology, Division of Radiology and Nuclear Medicine, Oslo University Hospital, Oslo, Norway. ²Faculty of Medicine, University of Oslo, Oslo, Norway. ³Department of Radiology, University of Texas Southwestern Medical Center, Dallas, TX, USA. ⁴Department of Diagnostic Physics, Division of Radiology and Nuclear Medicine, Oslo University Hospital, Oslo, Norway. ⁵Department of Diagnostic Radiology, Faculty of Medicine, Yamagata University, Yamagata, Japan. ⁶Department of Radiology and Athinoula A. Martinos Center for Biomedical Imaging, Massachusetts General Hospital and Harvard Medical School, Boston, MA, USA. ⁷Department of Physics, Faculty of Mathematics and Natural Sciences, University of Oslo, Oslo, Norway. *email: pdue@ous-hf.no
To this end, using a large database of clinical and imaging data in a blind, cross-institutional, multi-observer setting, the purpose of our study was to perform a retrospective and unbiased evaluation on the impact of the radiologic assessment and observer confidence of MRI-based preoperative glioma characterization on subsequent treatment decisions.

**Materials and Methods**

**Ethical approval and informed consent.** Ethical approvals were obtained from institution A (Oslo University Hospital, Oslo, Norway) by the Norwegian Regional Committees for Medical and Health Research Ethics (reference number 2013/81) and for institution B (Massachusetts General Hospital, Boston, MA, USA) by the Partners Human Research Committee (PHRC) Institutional Review Board (2012P000303), respectively. According to the respective national laws, written informed consent was required and obtained for all subjects (patients) from institution A, whereas written informed consent was waived by the Institutional Review Board committee of institution B due to the retrospective nature of the study. All procedures performed in studies involving human participants were in accordance with the ethical standards of the institutional and/or national research committee and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards.

**Patients and treatment options.** We retrospectively included 220 previously untreated adult patients from two institutions on different continents. Of these, 94 patients came from institution A (49 males, median age 49 years, range 23–79 years), and 126 patients came from institution B (69 males, median age 56 years, range 21–87 years). Institution A is a national referral site for brain tumor patients and accounts for almost 50% of all patients in the country. All patients were referred to a diagnostic, contrast-enhanced MR exam between 2003 and 2012, before surgery and subsequent histopathological diagnosis of a WHO grade II–IV glioma (Fig. 1). Patient demographics, histopathological assessments, Karnofsky Performance Status (KPS) at the time of MRI are summarized in Table 1. Per study protocol, treatment options were reviewed up until the end of August 2013 using patient medical records and hospital registry systems. The treatment data included (Table 2); steroid use at the time of MRI, type of surgery (biopsy, subtotal resection <90% or gross total resection >90%), type of adjuvant therapy (fractionized radiotherapy, chemotherapy, combined chemoradiation and/or anti-angiogenic therapy with bevacizumab)\(^1\), and the number of reoperations (no/one/multiple) within the study period. Post-surgery treatment was decided using all information available to the treating physicians at the time, including MRIs, clinical information and histopathological diagnosis.

**Histopathologic analyses.** Tumor tissue from needle biopsies or surgical resections were routinely formalin-fixed and paraffin-embedded before diagnostic review by an experienced neuropathologist. Because of the retrospective nature of our study and the closure date of August 2013, all tumors were classified according to the WHO 2007 criteria (glioma grades II–IV). Only previously untreated patients with a newly diagnosed glioma by histopathology were included in the study.

![Figure 1. Flow chart of study inclusion criteria. Flow chart showing study inclusion and exclusion criteria. From May 2003 to July 2012, a total of 243 adult patients met the inclusion criteria from both institutions combined. Twenty-three patients were excluded because of undisclosed previous surgery or treatments before the MRI exam (upon re-exam), missing or inconclusive surgery and/or histopathology after the MRI exams, as well as corrupted MRI data. The final study sample for analysis was 220 patients.](https://www.nature.com/scientificreports/)

---

**Table 1: Patient demographics, histopathological assessments, Karnofsky Performance Status (KPS) at the time of MRI**

| Institution | Eligible patients | Excluded patients | Patients included in study |
|-------------|-------------------|-------------------|---------------------------|
| A           | 109               | 10                | 99                        |
| B           | 134               | 2                 | 132                       |

**Table 2: Treatment data**

| | Biopsy | Subtotal Resection | Gross Total Resection | Steroid Use | Adjuvant Therapy | Reoperations (No/O/Many) |
|---|---|---|---|---|---|---|
| 1 | Yes | No | No | Yes | Yes | No |
| 2 | No | Yes | Yes | No | No | Yes |

---

---

**Figure 1.** Flow chart showing study inclusion and exclusion criteria. From May 2003 to July 2012, a total of 243 adult patients met the inclusion criteria from both institutions combined. Twenty-three patients were excluded because of undisclosed previous surgery or treatments before the MRI exam (upon re-exam), missing or inconclusive surgery and/or histopathology after the MRI exams, as well as corrupted MRI data. The final study sample for analysis was 220 patients.
MR imaging. Imaging was performed with 1.5 Siemens scanners at institution A, and 3 Tesla Siemens and 1.5 Tesla GE (General Electric) scanners at institution B, with 8-, 12-, or 32-channels (Siemens) and 8-, or 16-channels (GE) head coils (Table 1). An important aspect of study protocol was the use of actual clinical data without any enforced cross-institutional standardization of imaging protocols.

Table 1. Patient demographics and DSC imaging parameters. Note. WHO = World Health Organization, KPS = Karnofsky performance status, GE = gradient echo, SE = spin echo, EPI = echo planar imaging, TR = repetition time, TE = echo time, FA = flip angle, Gd-DTPA = gadopentetate-dimeglumine, DA = Diffuse Astrocytoma, OA = Oligoastrocytoma, OD = Oligodendroglioma, EP = Ependymoma, CG = Chordoid glioma, AA = Anaplastic Astrocytoma, AOA = Anaplastic Oligoastrocytoma, AOD = Anaplastic Oligodendroglioma, GBM = Glioblastoma Multiforme.

Table 2. Histopathologic diagnoses and treatments used in the study. Note. WHO = World Health Organization, B = Biopsy, <90% = subtotal resection, >90% = gross total resection. C = Chemotherapy, R = Radiotherapy, CR = Chemoradiation, DA = Diffuse Astrocytoma, OA = Oligoastrocytoma, OD = Oligodendroglioma, EP = Ependymoma, CG = Chordoid glioma, AA = Anaplastic Astrocytoma, AOA = Anaplastic Oligoastrocytoma, AOD = Anaplastic Oligodendroglioma, GBM = Glioblastoma Multiforme. †incomplete data, *some patients receiving CR also received concomitant anti-angiogenic therapy.
For institution A, a standard imaging protocol included a 0.45 × 0.45 × 5 mm³ voxel size two-dimensional axial 19-slice fast spin-echo T₂-weighted (repetition time TR = 4000 ms, echo time TE = 104 ms), coronal 25-slice T₂-weighted fluid-attenuated inversion recovery (FLAIR) (TR = 9050 ms, TE = 114 ms; inversion time TI = 1500 ms) and 0.45 × 0.45 × 5 mm³ voxel size axial/coronal/sagittal 19-slice pre/post-contrast spin-echo T₁-weighted (TR = 500 ms, TE = 7.7 ms) images. Diffusion MRI was available in 83 of 94 subjects using axial 19-slice single-shot, spin-echo echo-planar imaging with TR = 3300 ms, TE = 95 ms, 1.8 × 1.8 × 5 mm³ voxel size, three averages, 1.5 mm inter-slice gap, 90° flip angle and b-values of 0, 500, 1000 s/mm² in three orthogonal directions. DSC-MRI was performed in all patients (sequence details shown in Table 1), including administration of a single dose of 0.2 mmol/kg gadobutrol (Gadovist®, Bayer Schering Pharma, Berlin, Germany) at a rate of 5 ml/sec followed by a 20 ml saline flush (BB Melsungen, Melsungen, Germany).

For institution B, the imaging protocol mirrored that of institution A, including concurrent diffusion MRI in 119 of 126 patients. All patients had DSC-MRI with either a single or a double dose of 0.1 mmol/kg gadopentate dimeglumine (Magnevist®, Bayer Schering Pharma, Berlin, Germany) at a rate of 5 ml/sec followed by 20 ml saline (sequence details shown in Table 1).

Image pre-processing. All image pre-processing was performed using nordicICE (NordicNeuroLab AS, Bergen, Norway) or the open source platform 3D Slicer. As previously described, a board-certified neuroradiologist at each institution identified tumor regions on conventional MRIs using manual outlining in nordicICE for institution A, and a semi-automatic approach in 3D Slicer for institution B. Using nordicICE, apparent diffusion coefficient (ADC) maps were estimated using Stejskal-Tanner diffusion approximation and DSC-MRI data were automatically processed to create relative cerebral blood volume (rCBV) maps using standard kinetic modelling and corrected for contrast agent extravasation. All rCBV maps were normalized to normal-appearing tissue and presented as semi-transparent color overlays on anatomical MRIs.

Observers. Two neuroradiologists with >10 and >15 years of clinical experience with brain MRI, were included from institution A. Two additional neuroradiologists were included to represent institution B, one with >10 years of clinical experience with brain MRI and working at institution B. The other observer representing institution B was a consultant neuroradiologist from a third institution with >5 years of clinical experience with brain MRI. The observers from institution A reviewed the MRI data of institution B, and vice versa.

First MRI reading. Using anatomical and diffusion MRI only, the observers recorded their scores from visual grading using the Visually AcceSAble Rembrandt Images (VASARI) standardized feature set through the joint National Institute of Health (NIH) and National Cancer Institute (NCI) initiated Annotation and Image Markup (AIM) scoring template for gliomas. The VASARI scoring system includes 19 semantic descriptors of imaging features of brain tumors (Supplementary Table 1). Tumor location was also assessed by inclusion (Yes/No) of the NIH and National Cancer Institute (NCI) initiated Annotation and Image Markup (AIM) scoring template for gliomas. The VASARI scoring system includes 19 semantic descriptors of imaging features of brain tumors (Supplementary Table 1). Tumor location was also assessed by inclusion (Yes/No) of

Statistical analyses. Any statistical analyses were performed using SPSS 22 (SPSS Inc., USA). A P-value of 0.05 was considered significant, and all tests with multiple comparisons were Bonferroni corrected (P = 0.05 number of tests).

Results

Institutional differences in patient data and treatment options. The ratio of low-grade to high-grade gliomas was significantly higher for institution A compared to institution B (49% versus 18%, P = 0.003). However, the distributions of WHO grades within the institution was similar (Table 1; P = 0.047;
Table 3. Resulting WHO grading and confidence scores. Note. †Data show average values (unitless), standard deviations and sample size. A = different from MRI #1 at the P < 0.001 level, B = different from MRI #2 at the P < 0.001 level, (>5–15 yrs) corresponds to years of clinical experience with brain MRI. WHO = World Health Organization, MRI #1 = first conventional MRI reading, MRI #2 = second conventional MRI reading, +CBV = with addition of CBV maps.

not passing Bonferroni). The average KPS was correspondingly higher at institution A (90.53% ± 11.20% versus 86.59% ± 9.48%, P = 0.005), and steroids during initial MRI were administered more frequent (53% versus 17%, P < 0.001). The number of gross-total resections in WHO grade II gliomas was lower for institution A compared to institution B (Table 2; 13% versus 58%, P = 0.018). Patients at institution A were also more likely to have repeated surgery (all patients: 26% versus 9%, P = 0.002, versus 38% versus 3% (P < 0.001), HR 1.37 [1.17–1.61]).

Observer outcome measures and intra-observer repeatability. An overview of the average WHO grade and corresponding confidence levels of all observers are shown in Table 3. At both institutions and for all observers, the proposed WHO grades did not change over the course of the study readings. In contrast, the confidence scores increased significantly for all observers with the addition of the CBV map (at the P < 0.001 level, all observers, Table 3). Moreover, for institution A, the ICCs when grading patients by WHO grades II-IV were 0.93 (P < 0.001, n = 91) and 0.88 (P < 0.001, n = 73) for the two observers, respectively. The ICCs of the corresponding confidence scores were lower at 0.7381 (P < 0.001, n = 91) and 0.58 (P < 0.001, n = 73), respectively. For institution B, the ICCs when grading patients by WHO grades II-IV were 0.69 (P < 0.001, n = 120) and 0.83 (P < 0.001, n = 124) for observer 1 and 2, respectively. Again, the ICCs of the corresponding confidence scores were lower at 0.35 (P < 0.001, n = 120) and 0.25 (P = 0.002, n = 124) for observer 1 and 2, respectively.

Institutional differences in MRI features. The average size of the tumors was smaller (37.63 ± 33.34 mL versus 68.42 ± 58.87 mL, P < 0.001), while the average peritumoral edema region was larger (53.43 ± 47.01 mL versus 34.38 ± 50.90 mL, P = 0.012) at institution A compared to institution B. Matched for WHO glioma grade, ependymal extension was the only imaging features from the first MRI reading separating patients of institution A from institution B (67% versus 17%, P < 0.001). For WHO grades II and IV only, the non-enhancing tumor margins were less well-defined at institution A compared to institution B (30% versus 73%, P < 0.003). There was no significant difference in the enhancement quality between 1.5 and 3 Tesla systems (Table 1) (P > 0.79; both observers), nor between a single - versus double-dose contrast agent administration (P > 0.79).

Associations between MRI features and subsequent neurosurgery. Table 4 highlights the pan-institutional imaging features associated with the choice of surgical procedure. In short, by VASARI, patients with a biopsy or a subtotal resection were more likely to have a tumor with pathologic FLAIR/T2 signal crossing the midline, and those receiving a biopsy sampling more often had multifocal or multicentric lesions. Patients with low-post-contrast enhancement quality (0–1) had longer time between the pre-surgical MRI and subsequent surgery (trimmed mean = 10.84 days versus 3.92 days, P < 0.01, Hazard Ratio [95% conf.int.]; HR = 1.21 [1.06–1.38]). For WHO grade IV glioblastomas only (N = 121), lack of FLAIR/T2 signal crossing the midline and no satellite enhancement foci were associated with a gross total resection (P < 0.001, HR = 1.37 [1.17–1.61]).

For Institution A, patients with a biopsy sampling or a subtotal resection more often had tumors with an infiltrative T1/FLAIR ratio (1.94 ± 0.71 and 1.91 ± 0.68 versus 1.41 ± 0.54, P = 0.008) and less pial invasion (16% and 41% versus 53%, P = 0.002) compared to those with a total resection. Findings of institution B mirrored the pan-institutional analyses (Table 4).

Finally, repeated surgery at both institutions was associated with a higher incidence of pial invasion (67% versus 38%, P = 0.001) and less restricted diffusion (1.52 ± 0.40 versus 1.70 ± 0.61, P = 0.002). For WHO grade IV glioblastomas, more peritumoral edema (74.94 ± 83.93 mL versus 44.71 ± 38.95 mL, P = 0.001) and lack of satellite enhancement foci (24% versus 43%, P = 0.001) were also associated with repeated surgery (N = 17 of 121).

Associations between MRI features and adjuvant therapy. Table 5 summarizes the associations relevant for both institutions. In short, patients receiving steroids at the time of MRI (N = 72) more frequently had tumors with poorly defined non-enhancing margins (65% versus 40%, P < 0.001), and pial invasion (34% versus 16%, P < 0.001). This association also included ependymal extension, but it did not pass Bonferroni correction.
Table 4. Associations between MRI features and subsequent neurosurgery (both institutions). Note. ***Significant features at the $P<0.001$ level (Bonferroni corrected). **Significant features at the $P<0.01$ level (Bonferroni corrected). #Observer scorings deemed indeterminate were excluded from analysis. Highest incidence/value highlighted in bold.

| MRI feature# | Biopsy | Subtotal resection | Gross total resection | Repeated surgery | Hazard ratio $[95\% \text{ conf.int}]$
|-------------|--------|-------------------|---------------------|-----------------|------------------|
|             | $\bar{x}$ ± st.dev | $\bar{x}$ ± st.dev | $\bar{x}$ ± st.dev | Yes | No |
| FLAIR/T2 signal cross midline | 45% (29/64) | 30% (30/99) | 6% (3/51) | — | — | 1.30 [1.21–1.87] *** |
| Enhancement quality | 1.96 ± 1.17 | 2.77 ± 0.90 | 1.97 ± 1.31 | — | — | 1.27 [1.11–1.56] *** |
| Multifocal lesions | 42% (27/64) | 27% (27/99) | 6% (3/51) | — | — | 1.30 [1.16–1.64] *** |
| Pial invasion | — | — | — | 43% (16/36) | 17% (31/187) | 1.30 [1.11–1.48] ** |
| Facilitated diffusion | — | — | — | 1.52 ± 0.40 | 1.70 ± 0.61 | 1.32 [1.09–1.37] ** |

Table 5. Associations between MRI features and adjuvant therapy (both institutions). Note. ***Significant features at the $P<0.001$ level (Bonferroni corrected). **Significant features at the $P<0.01$ level (Bonferroni corrected). #Observer scorings deemed indeterminate were excluded from analysis. Highest incidence/value highlighted in bold.

| MRI feature# | Steroids at MRI | Radiation | Chemo-radiation | Hazard ratio $[95\% \text{ conf.int}]$
|-------------|----------------|-----------|-----------------|------------------|
|             | $\bar{x}$ ± st.dev | $\bar{x}$ ± st.dev | $\bar{x}$ ± st.dev | Yes | No |
| Poor non-enhancing margins | 65% (46/70) | 40% (59/145) | — | — | 1.36 [1.19–1.66] *** |
| Pial invasion | 34% (24/70) | 16% (23/145) | — | — | 1.35 [1.18–1.71] *** |
| Cortical involvement | 74% (52/70) | 79% (115/145) | — | — | 1.29 [1.13–1.70] ** |
| Glioma grade (with DSC-MRI) | 3.96 ± 0.81 | 3.49 ± 0.74 | — | — | 1.29 [1.05–1.53] ** |
| Glioma grade (with DSC-MRI) | 3.64 ± 0.64 | 3.06 ± 0.84 | — | — | 1.38 [1.15–1.45] *** |
| Ependymal extension | 34% (47/141) | 48% (38/79) | — | — | 1.30 [1.12–1.51] *** |
| Involvement of basal ganglia | 24% (34/141) | 36% (29/79) | — | — | 1.26 [1.09–1.45] ** |

Cortical involvement was seen less frequent in patients receiving steroids (74% versus 79%, $P=0.002$). For patients receiving mono-radiotherapy ($N=24$), a lower WHO glioma grade when including DSC-MRI was the only associated variable ($2.96 ± 0.81 versus 3.49 ± 0.74, P=0.001$). For patients receiving chemotherapy only (Institution A; $N=8$), the T1/FLAIR ratio was more infiltrative ($2.06 ± 0.73 versus 1.52 ± 0.61, P=0.027$; Mann-Whitney) and the suggested grade with DSC-MRI was lower ($2.88 ± 0.64 versus 3.46 ± 0.76, P=0.014$; Mann-Whitney). Patients receiving combined chemo-radiation therapy ($N=141$) had a higher suggested WHO grade with DSC-MRI ($3.64 ± 0.64 versus 3.06 ± 0.84, P<0.001$) and lack of deep-brain involvement by less ependymal extension (34% versus 48%, $P=0.001$) and less involvement of the basal ganglia (24% versus 36%, $P=0.002$). For WHO grade IV glioblastomas, tumor with less frequent involvement of the basal ganglia (23% versus 54%, $P<0.001$) and more well-defined non-enhancing tumor margins (61% versus 35%, $P=0.002$) more often received combined chemo-radiation therapy.

Comparing patients on combined chemo-radiation with- or without additional anti-angiogenic therapy (institution B), patients receiving bevacizumab ($N=29$) were more likely to have a tumor in the deep-brain by more frequent involvement of the thalamus (28% versus 11%, $P=0.002$) and less frequent involvement of the temporal lobes (31% versus 56%, $P=0.002$).

Associations between observer confidence and treatment. Figure 2 shows representative MRI of two patients with low- and high observer confidence, respectively. Steroid use at the time of MRI ($N=72$) was associated with higher observer confidence when evaluating anatomical MRI only (2.79 ± 0.75 versus 2.43 ± 0.64, $P<0.001$). Interestingly, when adding DSC-MRI, the confidence level for the non-steroid group ($N=148$) increased significantly (from 2.43 ± 0.64 to 3.03 ± 0.80, paired t-test: $P<0.001$), and to a higher level than the steroid group (3.03 ± 0.80 versus 2.79 ± 0.75, $P<0.001$). Moreover, adding DSC-MRI reduced the number of VASARI features associated with low confidence (score 1–2) at the $P<0.001$ level (median 3 versus 0 features; Supplementary Table 1). For WHO glioma grades II and III ($N=99$), observer confidence was associated with the choice of surgical procedure. The lowest observer confidence was found in patients receiving a biopsy only (2.02 ± 0.44) compared to subtotal resection (2.25 ± 0.50) and gross total resection (2.40 ± 0.44, $P=0.002$).

Finally, for low-grade gliomas ($N=50$), the use of adjuvant therapy was inversely correlated with observer confidence (Spearman $=-0.48, P<0.001$). Patients not receiving any adjuvant therapy had the highest observer confidence (median 2.50, range 2.00–3.25, $N=22$), whereas patients on combined chemo-radiation with- or without anti-angiogenic therapy had the lowest observer confidence (median 2.00, range 1.75–3.5, $N=13$).
Discussion

In a blind and retrospective, dual-center, multi-observer study, we quantify pre-treatment radiologic features by MRI that are systematically associated with the outcome of subsequent treatment of adult patients with gliomas and therefore likely to play a key role in the clinical decision-making process. The choice of surgical intervention was associated with the complexity of tumor infiltration, and patients whose tumors had indiscernible contrast-enhancement patterns received a more conservative management by approximately 3 times longer time between the MRI exam and subsequent surgery. Tumors with a pathologic MRI-signal crossing the midline and/or multifocal disease were more likely to have sub-total resection or especially a biopsy. This finding was also associated with lower observer confidence. Tumor progression and the need for repeated surgery were associated with pial invasion and a more edematous signature of the peritumoral region19–21. Moreover, steroids were more often seen in invasive and infiltrative tumors with poorly defined non-enhancing margins. Still, use of steroids was also linked to higher observer confidence, where the tumors' overall appearance probably showed less ambiguous imaging features. For adjuvant therapy and corrected for grade, patients not receiving any additional therapy outside surgery had the highest observer confidence. Observer confidence also returned lower ICCs than those from glioma grading. Interestingly, this suggests that treating physicians are more likely to opt for additional adjuvant treatment options for tumors in which the imaging appearance is less typical, irrespective of histological grade. This fact is not entirely unexpected and likely reflects physicians understanding about the inherent limitations of the WHO grading system as the main or sole determinant of treatment decisions, a historical dogma which is being challenged and replaced by recent discoveries in glioma oncogenetics22,23. This finding

Figure 2. Example MRIs returning low- and high observer confidence. Illustration showing representative contrast-enhanced (CE) T1-weighted (T1w) and T2-weighted (T2w) MRIs and rCBV maps from DSC of two patients from institution A with WHO grade IV glioblastomas. (Left) MRIs of a 74-year old male returning low average observer confidence ('somewhat confident', 50–70% certainty), and typical appearance of a multifocal lesion with complex and diffuse contrast-agent enhancement patterns. (Right) MRIs of a 60-year old male returning high average observer confidence ('extremely confident', >90% certainty), and with typical appearance of a single lesion with well-defined contrast-enhancement patterns. Adding DSC-MRI reduced the number of VASARI features associated with low observer confidence at the P < 0.001 level. Both patients used steroids at the time of the MRI exam.
may also warrant the need for standardized disease registries in order to learn from the decisions made and the subsequent outcomes of previous decision-making processes. Patients receiving combined chemo-radiation had well-defined non-enhancing tumor margins and lack of deep-brain involvement. Because the choice of treatment is arguably linked to glioma location, these imaging features probably helped identify the target area for radiotherapy. Also, the lack of deep-brain involvement is consistent with the goal of minimizing radiation damage to basic functions of the brain. Instead, patients with tumors of the deep-brain where more likely to receive anti-angiogenic therapy.

Our study advocates the need for a high-quality, focused MRI protocol to complement the clinical and histopathologic data for pre-treatment assessment of patients with brain tumors. Novel imaging techniques are regularly introduced into oncologic research, with the ability to visualize new aspects of tumor pathophysiology, cellularity, metabolic profile and hemodynamic status. Glioma imaging protocols are therefore becoming increasingly comprehensive, time-consuming and costly, whilst quantification of any added impact on the decision-making process is still rarely performed. It can therefore be debated under what circumstances the diagnostic process is really improved in a cost-effective way by increasing the number of exams. In line with previous reports, adding DSC-MRI to a conventional imaging protocol improved the observers’ confidence of the glioma characterization in untreated (non-steroid) patients. Also, with DSC-MRI, observers suggesting a lower glioma grade was associated with adjuvant radiation- or chemo- monotherapy, whereas the suggestion of a higher glioma grade was associated with combined chemo-radiation. The reduced number of VASARI features associated with a low observer confidence could potentially be seen as a time-saving feature of DSC-MRI. While comparing subjective confidence scores across that observers with various levels of experience should be performed with care, our results indicate DSC-MRI may aid less experienced readers. For prospective studies, introducing machine learning alternatives may help confirm or identify other relevant imaging feature of the disease, and also reduce the inherent observer variations that follow complex diagnostic readings. By comparing the results of an artificial intelligence (AI) model to that of our current expert radiologic examination, we can reveal the added value of the AI model for assessment of disease. Finally, use of AI-based model interpretability may help generate more powerful radiomics signatures from the hidden layers of the neural network beyond the radiologist-labeled, classical VASARI features.

**Our study has some limitations.** Owing to inherent regional and national determinants, differences in patient demographics between the two institutions may have influenced our results. However, this difference is also welcomed in what makes a multi-center study stand out from a single-institution analogue, and introduce a compelling range in our findings beyond a certain demographic setting. Moreover, while taking measures to blind the observers, a study design of this nature will never truly mimic the dynamic and complex workup of oncologic practice. Undoubtedly, the treating physician will also include information from the histopathologic analyses in the treatment decision-making process. Therefore, our findings also include imaging analyses from patients of the same WHO type and grade. Also, owing to the retrospective nature of our study, the WHO grading system of 2007 was used in histopathological diagnosis, and neither we, nor the treating physicians at the time, had access to the molecular profiles of individual tumors. Both mutation status of isocitrate dehydrogenase (IDH) 1/2 and methylated O6-methylguanine DNA methyltransferase (MGMT) may influence the treatment decision process, and also potentially be determined by DSC-MRI. Furthermore, and unlike today, not all of the anatomical MRI data at the time used a 3D image readout. However, this should only to a limited extent affected the VASARI criteria as presented in our study.

To conclude, in a comprehensive study we identify the key radiographic determinants of glioma patients associated with the treatment decision-making process. The choice of surgical intervention was associated with the complexity of tumor infiltration and low observer confidence was associated with a more extensive adjuvant treatment protocol.

**Data availability**

The authors declare that all data supporting the findings of this study are available within the paper and its Supplementary Information. The datasets generated during and/or analyzed during the current study are available from the corresponding author on reasonable request.

**References**

1. Ostrom, Q. T. et al. CBTRUS Statistical Report: Primary Brain and Central Nervous System Tumors Diagnosed in the United States in 2008–2012. *Neuro Oncol* 17(Suppl 4), iv1–iv62, https://doi.org/10.1093/neuonc/nov189 (2015).
2. Crocetti, E. et al. Epidemiology of glial and non-glial brain tumours in Europe. *Eur. J. Cancer* 48, 1532–1542, https://doi. org/10.1016/j.ejca.2012.11.013 (2012).
3. Louis, D. N. et al. The 2016 World Health Organization Classification of Tumors of the Central Nervous System: a summary. *Acta Neuropathol* 131, 803–820, https://doi.org/10.1007/s00401-016-1545-z (2016).
4. Kalpathy-Cramer, J., Gerstner, E. R., Emblem, K. E., Andronesi, O. & Rosen, B. Advanced magnetic resonance imaging of the physical processes in human glioblastoma. *Cancer Res* 74, 4622–4637, https://doi.org/10.1158/0008-5472.CAN-14-0383 (2014).
5. Henson, J. W., Gaviani, P. & Gonzalez, R. G. MRI in treatment of adult gliomas. *Lancet Oncol* 6, 167–175 (2005).
6. Gutman, D. A. et al. MR imaging predictors of molecular profile and survival: multi-institutional study of the TCGA glioblastoma data set. *Radiology* 267, 560–569, radiol.13120118 [pii]; 10.1148/radiol.13120118 [doi] (2013).
7. Geer, C. P. et al. Does MR perfusion imaging impact management decisions for patients with brain tumors? A prospective study. *AJNR Am. J. Neuroradiol.* 33, 556–562, https://doi.org/10.3174/ajnr.A2811 (2012).
8. Park, J. K. et al. Scale to predict survival after surgery for recurrent glioblastoma multiforme. *J. Clin. Oncol* 28, 3838–3843 (2010).
9. Hilario, A. et al. The added value of apparent diffusion coefficient to cerebral blood volume in the preoperative grading of diffuse gliomas. AJNR Am. J. Neuroradiol. 33, 701–707, https://doi.org/10.3174/ajnr.A2846 (2012).

10. Verburg, N. et al. Diagnostic Accuracy of Neuroimaging to Delineate Diffuse Gliomas within the Brain: A Meta-Analysis. AJNR Am. J. Neuroradiol. 38, 1884–1891, https://doi.org/10.3174/ajnr.A5368 (2017).

11. Stupp, R. et al. Radiotherapy plus concomitant and adjuvant temozolomide for glioblastoma. N. Engl. J Med 352, 987–996, 352/10/987 [pii];10.1056/NEJMoa043330 [doi] (2005).

12. Emblem, K. E. et al. Machine learning in preoperative glioma MRI: survival associations by perfusion-based support vector machine outperforms traditional MRI. J Magen Reson. Imaging 40, 47–54, https://doi.org/10.1002/jmri.24390 (2014).

13. Emblem, K. E. et al. A generic support vector machine model for preoperative glioma survival associations. Radiology 275, 228–234, https://doi.org/10.1148/radiol.141140770 (2015).

14. Emblem, K. E. et al. Glioma grading by using histogram analysis of blood volume heterogeneity from MR-derived cerebral blood volume maps. Radiology 247, 808–817 (2008).

15. Fedorov, A. et al. 3D Slicer as an image computing platform for the Quantitative Imaging Network. Magn Reson. Imaging 30, 1323–1341, S0735-755X(12)00181-6 [pii];10.1016/j.mri.2012.05.001 [doi] (2012).

16. Helseth, R., Kaur, G. & Guilfoyle, D. N. Principles and limitations of NMR diffusion measurements. J. Med. Phys. 32, 34–42, https://doi.org/10.4103/0971-6203.31146 (2007).

17. Bjornerud, A., Sorensen, A. G., Mouridsen, K. & Emblem, K. E. T(1)- and T(2)(*)-dominant extravasation correction in DSC-MRI: Part I-theoretical considerations and implications for assessment of tumor hemodynamic properties. J. Cereb. Blood Flow Metab (2011).

18. Emblem, K. E. et al. Automatic glioma characterization from dynamic susceptibility contrast imaging: brain tumor segmentation using knowledge-based fuzzy clustering. J. Magen Reson. Imaging 30, 1–10 (2009).

19. Helseth, R. et al. Overall survival, prognostic factors, and repeated surgery in a consecutive series of 516 patients with glioblastoma multiforme. Acta Neuro. Scand 122, 159–167 (2010).

20. Gerstner, E. R. et al. Infiltrative patterns of glioblastoma spread detected via diffusion MRI after treatment with cediranib. Neuro Oncol 12, 466–472, doi:10.1093/neuonc/noq051 [doi] (2010).

21. McDonald, C. R. et al. Restriction spectrum imaging predicts response to bevacizumab in patients with high-grade glioma. Neuro Oncol 18, 1579–1590, https://doi.org/10.1093/neuonc/nox063 (2016).

22. Weller, M. et al. Personalized care in neuro-oncology coming of age: why we need MGMT and 1p/19q testing for malignant glioma patients in clinical practice. Neuro Oncol 14(Suppl 4), iv100–108, https://doi.org/10.1093/neuonc/nox206 (2012).

23. Weller, M. et al. Molecular neuro-oncology in clinical practice: a new horizon. Lancet Oncol 14, e370–379, https://doi.org/10.1016/s1470-2045(13)70168-2 (2013).

24. Liu, X. et al. Pathological factors contributing to crossed cerebellar diaschisis in cerebral gliomas: a study combining perfusion, diffusion, and structural MR imaging. Neuroradiology 60, 643–650, https://doi.org/10.1007/s00234-018-1015-3 (2018).

25. Aronen, H. J. et al. Cerebral blood volume maps of gliomas: comparison with tumor grade and histologic findings. Radiology 191, 41–51 (1994).

26. Hempel, J. M. et al. In Vivo Molecular Profiling of Human Glioma: Cross-Sectional Observational Study Using Dynamic Susceptibility Contrast Magnetic Resonance Perfusion Imaging. Clin Neuroradiol, https://doi.org/10.1007/s00062-018-0676-2 (2018).

27. Gazelle, G. S., McMahon, P. M., Siebert, U. & Reinfeld, M. T. Cost-effectiveness analysis in the assessment of diagnostic imaging technologies. Radiology 235, 361–370, https://doi.org/10.1148/radiol.2352040330 (2005).

28. Juan-Albarracin, J. et al. Glioblastoma: Vascular Habitats Detected at Preoperative Dynamic Susceptibility-Weighted Contrast-enhanced Perfusion MR Imaging Predict Survival. Radiology, 170845, https://doi.org/10.1148/radiol.2017170845 (2018).

29. Montavon, G., Samek, W. & Müller, K.-R. Methods for interpreting and understanding deep neural networks. Digital Signal Processing 73, 1–15, https://doi.org/10.1016/j.dsp.2017.10.011 (2018).

30. Anzalone, N. et al. Brain Gliomas: Multicenter Standardized Assessment of Dynamic Contrast-enhanced and Dynamic Susceptibility Contrast MR Images. Radiology 287, 933–943, https://doi.org/10.1148/radiol.2017170362 (2018).

Acknowledgements
We thank the patients and their relatives for involvement in the study. This work received funding from the Research Council of Norway grant 191088/F20 (AB) and FRIPRO grant 261984 (KEE), the SouthEastern Norway Regional Health Authority grants 2017073 (KEE) and 2013069 (KEE), and the European Research Council (ERC) under the European Union’s Horizon 2020 Programme (ERC Grant Agreement No. 758657-ImPRESS) (KEE).

Author contributions
P.D.-T., M.C.P., K.E.E., O.R. and A.B. conceived the project, P.D.-T. M.C.P., K.E.E., O.R. and A.B. performed the data analysis; A.A., I.R.G. and O.R. provided expertise on the neuroscience aspects of the experiments and manuscript text; P.D.-T. M.C.P., K.E.E. and O.R. provided MRI images and patients characteristics; O.R. and A.B. supervised the project and provided guidance on experimental design, data interpretation and writing of the manuscript; all authors contributed to discussions on crucial aspects of the project and reviewed the manuscript drafts; all authors read and approved the final version of the manuscript submitted for review.

Competing interests
Oso University Hospital, Oslo, Norway, has received payment from NordicNeuroLab AS, Bergen, Norway for patents and/or intellectual property rights of two authors (KEE) and (AB). One author has stock or stock options in NordicNeuroLab AS, Bergen, Norway (AB). The other authors have no disclosures to report.

Additional information
Supplementary information is available for this paper at https://doi.org/10.1038/s41598-019-56333-x.

Correspondence and requests for materials should be addressed to P.D.-T.

Reprints and permissions information is available at www.nature.com/reprints.

Publisher’s note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.
