

Comparing Glioblastoma Surgery Decisions Between Teams Using Brain Maps of Tumor Locations, Biopsies, and Resections

Domenique M.J. Müller, MD¹; Pierre A.J.T. Robe, MD, PhD²; Roelant S. Eijgelaar, Msc³; Marnix G. Witte, PhD³; Martin Visser, Msc²; Jan C. de Munck, PhD²; Marieke L.D. Broekman, MD, PhD, LLM²; Tatjana Seute, MD, PhD²; Jeroen Hendrikse, MD, PhD²; David P. Noske, MD, PhD¹; William P. Vandertop, MD, PhD¹; Frederik Barkhof, MD, PhD^{2,4}; Mathilde C.M. Kouwenhoven, MD, PhD¹; Emmanuel Mandonnet, MD, PhD⁵; Mitchel S. Berger, MD⁶; and Philip C. De Witt Hamer, MD, PhD¹

PURPOSE The aim of glioblastoma surgery is to maximize the extent of resection while preserving functional integrity, which depends on the location within the brain. A standard to compare these decisions is lacking. We present a volumetric voxel-wise method for direct comparison between two multidisciplinary teams of glioblastoma surgery decisions throughout the brain.

METHODS Adults undergoing first-time glioblastoma surgery from 2012 to 2013 performed by two neuro-oncologic teams were included. Patients had had a diagnostic biopsy or resection. Preoperative tumors and postoperative residues were segmented on magnetic resonance imaging in three dimensions and registered to standard brain space. Voxel-wise probability maps of tumor location, biopsy, and resection were constructed for each team to compare patient referral bias, indication variation, and treatment variation. To evaluate the quality of care, subgroups of differentially resected brain regions were analyzed for survival and functional outcome.

RESULTS One team included 101 patients, and the other included 174; 91 tumors were biopsied, and 181 were resected. Patient characteristics were largely comparable between teams. Distributions of tumor locations were dissimilar, suggesting referral bias. Distributions of biopsies were similar, suggesting absence of indication variation. Differentially resected regions were identified in the anterior limb of the right internal capsule and the right caudate nucleus, indicating treatment variation. Patients with (n = 12) and without (n = 6) surgical removal in these regions had similar overall survival and similar permanent neurologic deficits.

CONCLUSION Probability maps of tumor location, biopsy, and resection provide additional information that can inform surgical decision making across multidisciplinary teams for patients with glioblastoma.

Clin Cancer Inform. © 2019 by American Society of Clinical Oncology

Creative Commons Attribution Non-Commercial No Derivatives 4.0 License 

INTRODUCTION

Standard treatment of glioblastoma consists of surgery and radiotherapy with concomitant and adjuvant chemotherapy. Nevertheless, the prognosis remains dismal, with a 1-year survival rate of 39%.¹ Initial neuro-oncologic treatment decisions include whether to undergo resective surgery or diagnostic biopsy or refrain from surgery, if the patient is unfit or unmotivated or the tumor is considered nonresectable.^{2,3} More extensive resections are associated with longer survival in glioblastoma, often recognized as a causal relation.⁴⁻⁷ The aim of neurosurgery is therefore to maximize tumor removal while preserving functional integrity. The surgical dilemma is that patient outcome depends on the decision of where to stop tumor removal. If stopped too early, more residual disease remains, with the risk of earlier tumor recurrence and shorter survival. If stopped too late, more tumor-infiltrated functional brain is removed, leading to

brain function deficits and reducing the patient's quality of life.⁸⁻¹¹

Neurosurgical decision making depends on several factors before surgery: the condition, age, and comorbidity of the patient; neurologic signs and symptoms; brain location of the tumor and derived resectability; expectations of the neurosurgeon regarding benefit and risk for the patient; and motivation of the patient after discussion of these expectations.^{5,12-15} Intraoperatively, neurosurgeons use various techniques to distinguish tumoral tissue from functional brain tissue, such as image guidance with neuro-navigation, magnetic resonance imaging (MRI), ultrasound or fluorescence,¹⁶ and intraoperative stimulation mapping.¹⁷ These factors and techniques add up to practice variation in neuro-oncologic decision making, which was captured by population-based cohort studies evaluating patterns of care in patients with glioblastoma.¹⁸⁻²² Biopsy percentages in

ASSOCIATED CONTENT

Data Supplement

Author affiliations and support information (if applicable) appear at the end of this article.

Accepted on October 29, 2018 and published at ascopubs.org/journal/cci on January 23, 2019; DOI <https://doi.org/10.1200/CCI.18.00089>

these reports range widely: 5%,²² 12%,²⁰ 24%,¹⁸ 34%,¹⁹ and 44%.²¹ The quality of glioblastoma resection is usually reported as the percentage of patients with a gross total resection, which also ranges considerably: 28%,²¹ 32%,¹⁸ 46%,²⁰ and 47%.²² Quality registries are being developed to evaluate the quality of neuro-oncologic care^{23,24}; however, no standards exist for the evaluation of neuro-oncologic and neurosurgical decision making among multidisciplinary teams. For comparison of teams, bias resulting from heterogeneity of tumor locations within the brain should be addressed; we proposed using a voxel-wise approach in nonenhancing glioma.²⁵

Voxel-wise methodologies were introduced for low-grade gliomas to evaluate surgical decision making with volumetric analysis of preoperative tumor and residual disease throughout the brain.^{26,27} The results of voxel-wise analyses can be plotted in standard brain space, which is called a probability map. Surgical decision making in non-enhancing gliomas has been compared among neurosurgical teams using resection probability maps (RPMs).²⁵

Ultimately, the best decision for a patient with glioblastoma regarding biopsy or resection, and in case of resection, the extent of tumor removal, should be based on a collaborative body of objective data rather than on the intuition of one or a few surgeons. A step in that direction is to compare and discuss the patterns of current care. In this study, we address the proof of concept for the use of probability maps to assess and compare the patterns of (intra)operative decision making between multidisciplinary teams in patients with glioblastoma.

METHODS

Patient Inclusion and Data Collection

Consecutive patients older than 17 years with a histopathologic diagnosis of supratentorial glioblastoma and first-time brain surgery in 2012 or 2013 were included from two brain tumor treatment centers: the Vrije Universiteit Medical Center in Amsterdam and the University Medical Center in Utrecht, the Netherlands. Patients underwent either resection with glioblastoma removal or biopsy for diagnosis. We defined tumor removal for histopathologic diagnosis only by an open or stereotactic approach as biopsy and more contrast-enhanced tumor removal than necessary for a histopathologic diagnosis by craniotomy as resection. The histopathologic diagnosis was based on WHO 2007 criteria. Written informed consent was obtained from all patients.

The two multidisciplinary neuro-oncologic teams had a similar approach to decision making in tumor board meetings on surgical indications for new patients by consensus of neurosurgeons based on patient and tumor characteristics. Intraoperative decision making was performed by neurosurgical teams under the supervision of a neuro-oncologic surgeon from a pool of two or three per center, respectively. Image guidance with neuronavigation

CONTEXT SUMMARY

Key objective:

To develop a method to compare patterns of surgical decisions without bias from tumor location in patients with glioblastoma across multidisciplinary teams.

Knowledge generated:

Probability maps of tumor location, biopsy and resection based on MRI before and after surgery enable comparison of patient selection and surgical decisions across teams. We demonstrate referral bias of patients with tumors in the left hemisphere and differential tumor removal in the right caudate nucleus between two teams.

Relevance:

Probability maps provide additional information that can inform patient counseling and surgical decision-making on biopsy or resection indications and on optimization of stimulation mapping. Our findings support the inclusion of standardized imaging in neuro-oncological registries. Probability maps may enhance knowledge transfer between neuro-oncological teams and serve as educational instrument.

was routinely applied. Functional MRI, DTI tractography, and intraoperative stimulation mapping for language and motor functions were used by indication. The Utrecht team had started using fluorescence-guided microscopy in 2013; the Amsterdam team had not.

Baseline characteristics of these patients were extracted from digital department databases, as well as from pre- and postoperative MRI. MRI acquisition protocols were comparable between the two teams (Data Supplement). A complication was defined as a new intervention or intensive care admission. A functional decline was defined as a drop in Karnofsky performance score of > 20 between admission and 6 weeks after surgery.

Brain Map Construction

The construction of brain maps is described in detail in the Data Supplement. Briefly, the enhancing portions of tumors were segmented on pre- and postoperative T1-weighted gadolinium-enhanced images. For patients who underwent biopsy, postoperative MRI was usually omitted, and we therefore considered the residual tumor to be identical to the preoperative tumor. Next, tumor volumes were nonlinearly registered to the Montreal Neurological Institute (MNI) 152 1-mm brain template to compare tumor locations between patients and teams.

Three types of probability maps were constructed by voxel-wise aggregations of the registered tumor volumes, each

reflecting clinical decision making. First, a tumor probability map (TPM) representing tumor burden before treatment was constructed by summation of preoperative tumors divided by the total number of patients treated by a team. A TPM difference between two teams would indicate referral or recruitment bias given a tumor location. Second, a biopsy probability map (BPM) representing the likelihood of a biopsy procedure over resection was constructed for each team by dividing the number of biopsies per location by the number of

resections. A BPM difference between two teams would indicate treatment variation in indications for biopsy given a tumor location. This can be considered a measure of multidisciplinary neuro-oncologic team decision making. Third, an RPM representing the likelihood of residual disease after surgery was constructed. A difference between RPMs of patients undergoing resection would indicate treatment variation in tumor removal given an indication for resection. This can be considered a measure of neurosurgical team decision making during surgery. In addition, a difference between RPMs based on all patients in a cohort, including those undergoing biopsy or resection, would then indicate treatment variation in tumor removal for all patients with newly diagnosed glioblastoma. This can be considered an overall measure of multidisciplinary neuro-oncologic decision making after first-time surgery for newly diagnosed glioblastoma.

TABLE 1. Baseline Patient and Treatment Characteristics and Outcomes

Characteristic	No. (%)		P
	Amsterdam Team (n = 101)	Utrecht Team (n = 174)	
Median age, years	63.2	66.1	.025
Female sex	37 (37)	68 (38)	.798
Karnofsky performance score			.160
100	9 (9)	5 (3)	
90	22 (22)	54 (31)	
80	23 (23)	40 (23)	
70	28 (28)	34 (20)	
60	12 (12)	19 (11)	
50	7 (7)	12 (7)	
40	0	3 (2)	
30	0	1 (1)	
Year of surgery			.617
2012	49 (49)	77 (44)	
2013	52 (52)	97 (56)	
Tumor in left brain hemisphere	39 (39)	95 (55)	.012
Median preoperative tumor volume, mL	28.7	37.9	.082
MRI timing			
Days between preoperative MRI and surgery	-4	-1	.000
Days between postoperative MRI and surgery	1	3	.000
Surgical strategy			.001
Biopsy	21 (21)	70 (40)	
Resection	80 (79)	104 (60)	
Median residual tumor volume, mL	1.0	0.3	.438
No. of patients undergoing resection with residue < 3 mL	59 (74)	44 (42)	.283
Median extent of resection, %	96.3	98.6	.206
No. of patients undergoing resection with extent of resection > 95%	45 (56)	43 (41)	.406
Outcomes			
Median OS, months	12	9	.187
Complications	3 (3)	7 (4)	.750
Functional decline	5 (5)	20 (11)	.083

Abbreviations: MRI, magnetic resonance imaging; OS, overall survival.

Subgroup Analysis

As an example of quality evaluation from treatment variation, we determined the subgroup of patients involved in the largest cluster of differentially resected voxels. Therefore, we extracted clinical follow-up and radiologic data from medical records to explore differences in outcome potentially related to treatment variation.

RESULTS

Patient and Treatment Characteristics

A total of 279 patients were treated, 102 by the Amsterdam team and 177 by the Utrecht team. For the probability map analysis, 101 (99%) and 174 patients (98%) were available, respectively. In two patients, the preoperative MR scan was unavailable from the referring hospital, and in two patients, the glioblastoma was nonenhancing. Patients were mostly comparable between teams in baseline characteristics (Table 1).

An MR scan within 3 days after resection was obtained in 77 patients (100%) who underwent resection in Amsterdam and in 68 (65%) of 104 in Utrecht. To adhere to current imaging standards,²⁸ we based the resection probabilities for evaluation of neurosurgical decisions on the patients with postoperative scans within 3 days. The quality of neurosurgical team decisions in patients undergoing resection was not significantly different between teams in terms of residual volume, extent of resection, or outcome (Table 1).

Referral Bias

A significant difference was indicated in relative tumor incidence by the TPM comparison between both teams (Fig 1). In several regions in the left hemisphere, regions with differential tumor incidence were identified, such as the caudate nucleus, anterior limb of the internal capsule, and temporal stem. These observations seemed to be based on systematic differences, such as referral or recruitment bias.

Variation in Biopsy Indications

No differentially biopsied regions were identified by the BPM comparison between teams (Fig 2). The BPMs

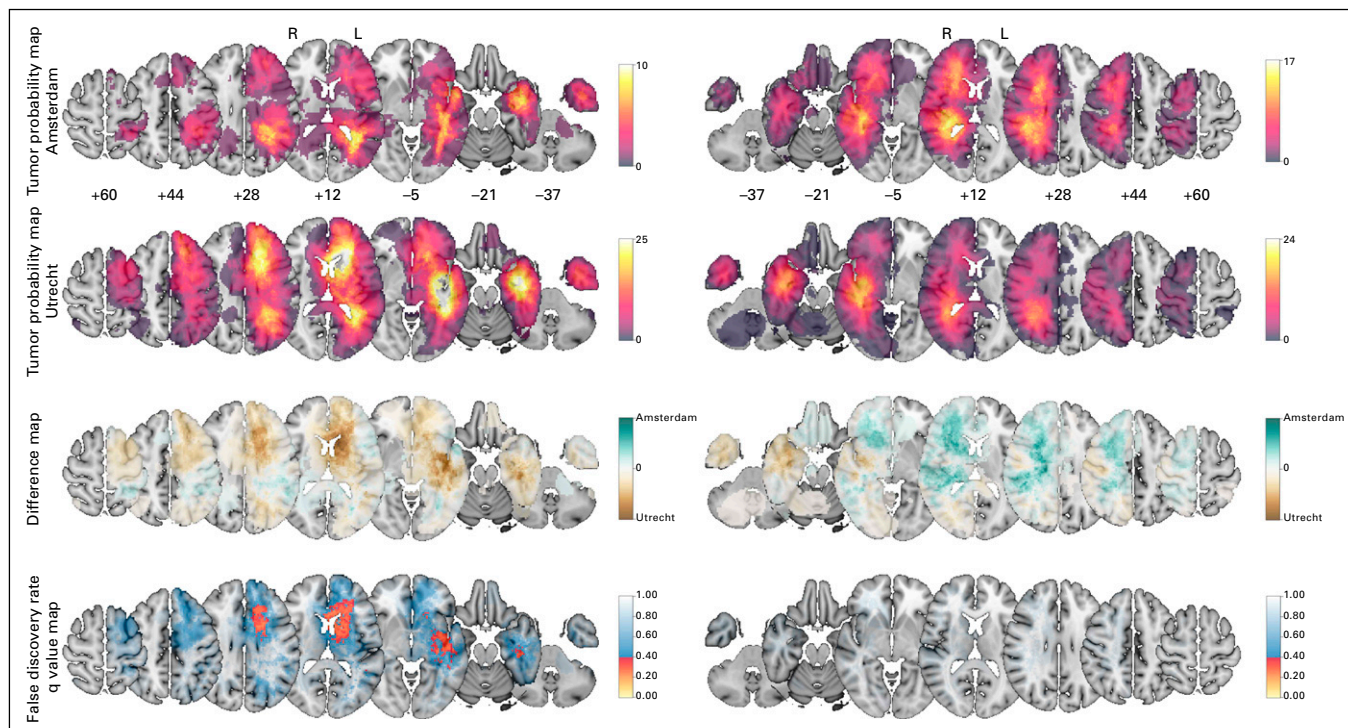


FIG 1. Tumor probability map (TPM) comparison of all 98 versus 174 patients per team to evaluate referral bias for the left (L; 38 v 95) and right (R) hemispheres (60 v 79). Plotted from top to bottom are the TPMs from Amsterdam, TPMs from Utrecht, difference maps, and false discovery rate q value maps. The color codes correspond with the adjacent legends. From left to right, the axial sections of the Montreal Neurological Institute (MNI) brain template are shown of the plotted z coordinates. The Data Supplement provides results over all axial sections, including the unadjusted P value map.

demonstrated that consensus existed regarding in which brain regions biopsy was preferred over resection, such as the thalamus, internal capsule, splenium of the corpus callosum, and periventricular region around the occipital horns, and conversely in which brain regions resection was preferred over biopsy, such as the anterior temporal lobe, right prefrontal cortex, and right inferior parietal lobe. The absence of differences in the BPMs suggests that brain location was not an important factor contributing to multidisciplinary decision making on performing a biopsy procedure. Nevertheless, the overall biopsy percentage differed, with 21 (21%) and 70 patients (40%) undergoing biopsy in Amsterdam and Utrecht, respectively ($P = .002$). Patients undergoing biopsy did not differ with regard to age (median, 63.1 and 65.9 years), condition (Karnofsky score < 70 in six [29%] and 16 [24%]), or tumor volume (median, 27 and 29 mL), respectively.

Variation in Tumor Removal

Tumor removal between teams was comparable in many brain regions. However, the Amsterdam team more often resected tumor in the right caudate nucleus and anterior limb of the right internal capsule than the Utrecht team (Fig 3). Common regions where tumor was generally removed included the temporal lobes, prefrontal cortices, and superior parietal lobules. Other tumor-infiltrated regions were commonly not removed, such as the anterior perforating

substance and corticospinal tract. The precision of these probabilities varied by location depending on the number of patients with information (Fig 1).

Subgroup Analysis of Patients With Different Tumor Removal

The brain region with the largest differential resection probability was the right caudate nucleus and surrounding white matter pathways, in which the Amsterdam team decided to remove the tumor, whereas the Utrecht team did not. This would probably have remained unnoticed without a probability map comparison. We identified 18 patients who underwent resection. No differences were noted in neurologic outcome, survival, postsurgical hydrocephalus, or ventricular dissemination of glioblastoma for patients with ($n = 12$) and without ($n = 6$) tumor removal (Data Supplement).

Variation in Oncologic Quality of Glioblastoma Surgery

The decisions for the whole cohort on whether to perform biopsy or resection and the observed extents of resection were similar between the two teams in both hemispheres (Fig 4). Therefore, overall decisions on biopsy and extent of resection do not depend on brain location.

DISCUSSION

The main finding of this study is that probability maps of tumor location, biopsy, and resection enable the evaluation

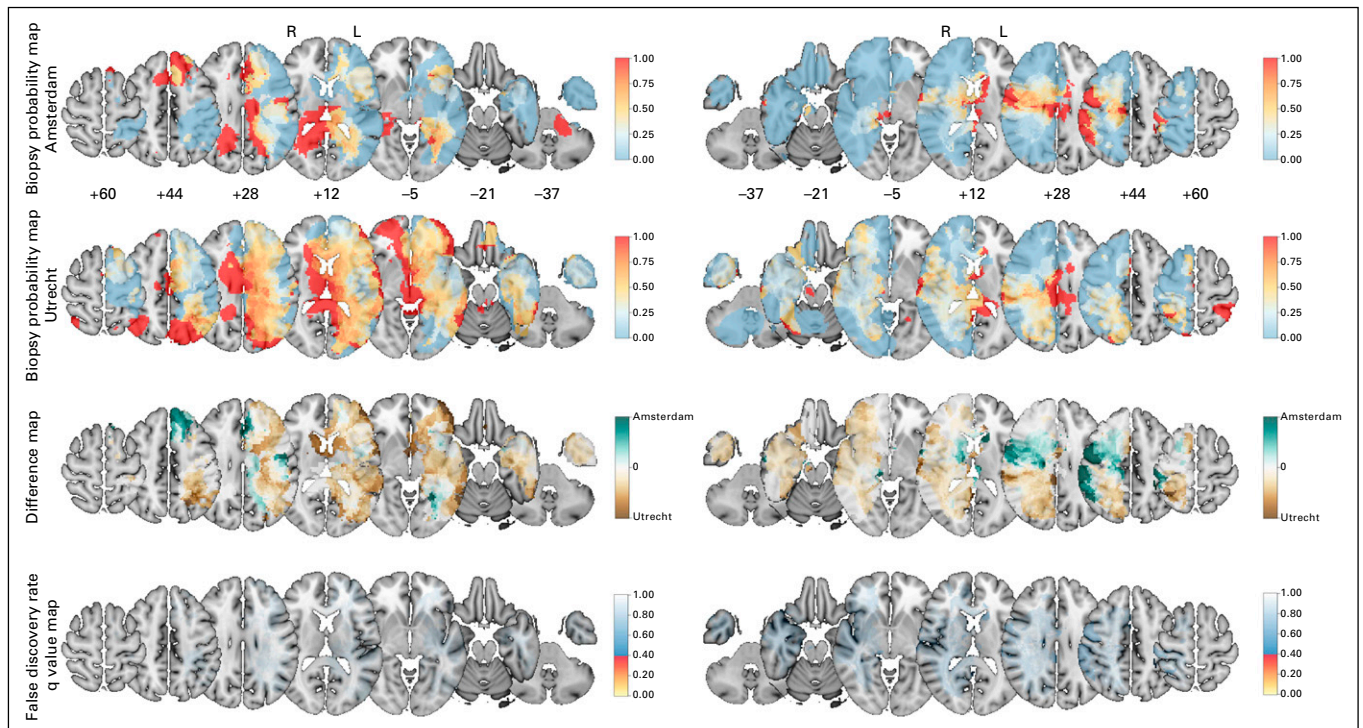


FIG 2. Biopsy probability map comparison of all 98 versus 174 patients per team to evaluate multidisciplinary decision making on biopsy indications for the left (L; 38 v 95) and right (R) hemispheres (60 v 79). A value of 0 represents a location where all patients have undergone resection, and a value of 1 represents a location where all patients have undergone biopsy. Note that the color code for biopsy probability in rows 1 and 2 is different from that in Figure 1. The color codes for difference maps in row 3, and false discovery rates in row 4 are identical in all figures. The Data Supplement provides the full results.

of surgical decision making among multidisciplinary teams in patients with glioblastoma.

Its potential use is illustrated by some examples. First, we identified a significant difference in tumor lateralization between two teams, and with TPM comparison, a more detailed specification of the brain regions involved could be made. This bias could be related to regional referral patterns of which we are as yet unaware. Because each center is considered to serve designated geographic regions, neurologists seldom cross-refer patients to other hospitals. Second, we identified a significant difference in biopsy percentage between two teams. This could not be attributed to specific brain locations according to the BPM analysis, nor to patient age, condition, or tumor volume. The difference in fluorescence guidance and application of stimulation mapping is not an explanation, because these techniques were used by the team performing more biopsies and fewer extensive resections. It seems other factors contributed to the differential overall biopsy preference, possibly related to expectations between the multidisciplinary teams regarding the benefits and risks of resection.

In addition, several of the observed differences in neurosurgical decision making were related to unresolved clinical dilemmas. Several plausible explanations were collected as

feedback from both neuro-oncologic teams for the difference in resection probability lateral of the frontal horn at the level of the caudate nucleus in the right hemisphere. As another example of the potential use of the probability map comparison, our findings highlight these dilemmas. First, opinions differed on the justification of surgical removal of the nondominant caudate nucleus. Arguments against removal included the functional anatomy of the caudate nucleus, which has been associated with behavior, sometimes resulting in severe cognitive impairment, such as abulia, as learned from observations in stroke²⁹⁻³² and other focal lesions.³³ In addition, the caudate nucleus is part of the negative motor network, connected to the supplementary motor area through the frontostriatal tract.³⁴ For supra-complete glioma resection, the caudate nucleus is considered a strict functional boundary by neurosurgeons who push the limits of resection.³⁵ In contrast, deficits were absent after caudate removal in low-grade glioma surgery.³⁶ In our data, no argument was raised against removal of the caudate to preserve the patient's functional integrity, although no systematic neuropsychological examinations were available for these patients. Second, the opinions also differed on justification of ventricular entry at the anterior horn of the lateral ventricle between the two teams. One team favored more extensive tumor removal, whereas the other team primarily intended to avoid adverse events resulting

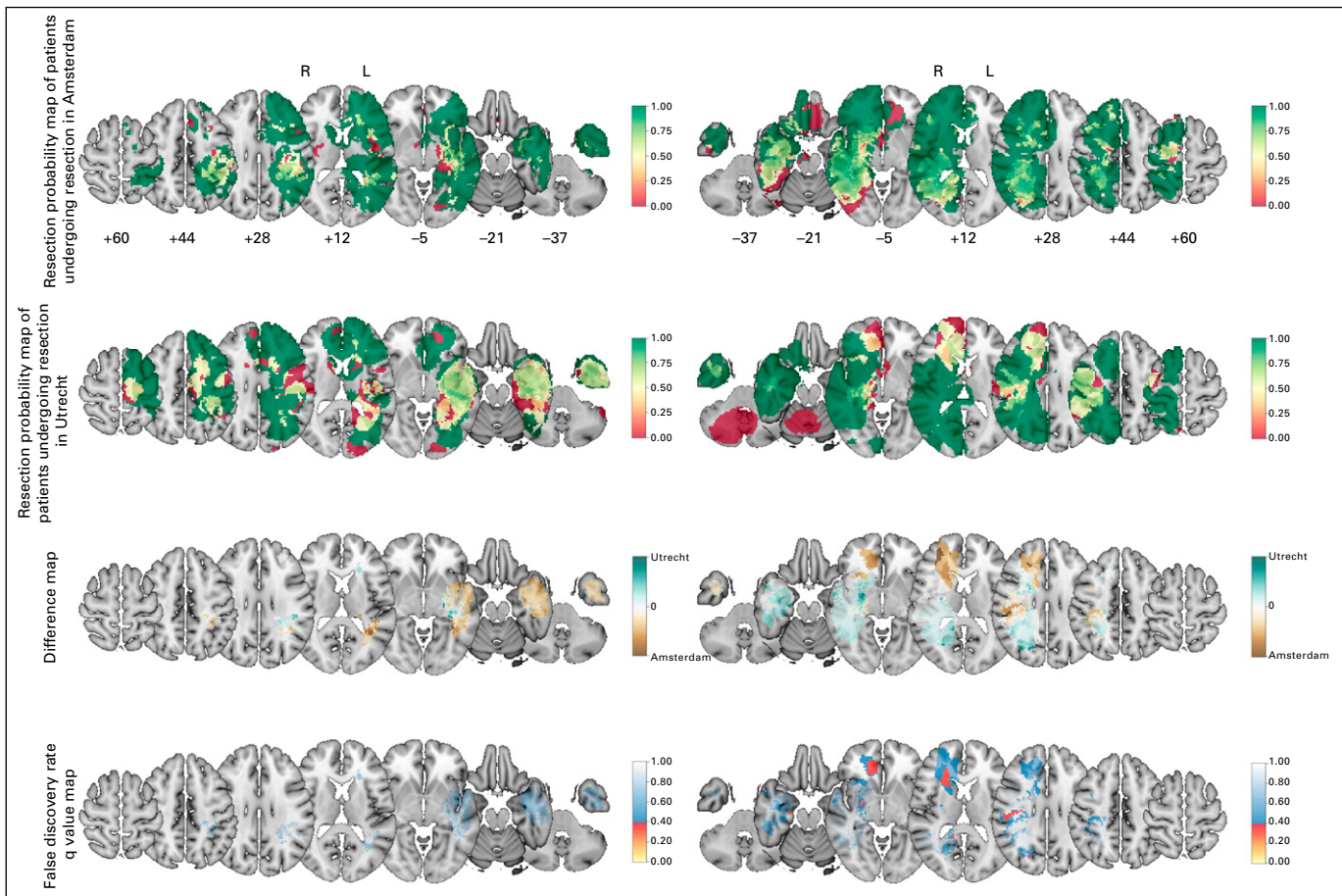


FIG 3. The resection probability map comparison of 77 versus 68 patients undergoing resection per team to evaluate neurosurgical decision making on residual disease for the left (L; 28 v 32) and right (R) hemispheres (49 v 36). Note that the color code for resection probability in rows 1 and 2 is different from those in Figures 1 and 2. The color codes for difference maps in row 3, and false discovery rates in row 4 are identical in all figures. The Data Supplement provides the full results.

from ventricular entry, such as hydrocephalus or tumor seeding. Hydrocephalus is an uncommon complication after glioma surgery, but it has been reported more frequently after ventricular entry.³⁷⁻⁴² Tumor seeding can be facilitated by wide ventricular opening. Although early publications did not find support for this,⁴³ later publications have indicated more frequent leptomeningeal glioblastoma dissemination after ventricular entry.^{38,44,45} In our data, tumor seeding only occurred in cases with opening of the ventricle, although this difference was not statistically significant with this limited number of patients. Altogether, our data do not support avoidance of ventricular entry to prevent these complications. Third, the opinions differed on the use of intraoperative stimulation mapping with cognitive tasks in the right frontal lobe. One team argued that cognitive impairment is an uncommon outcome and that cognitive mapping has not been validated and could impede tumor removal. The other team had started to develop a cognitive mapping protocol. More recently, awake mapping of nonlanguage functions has been proposed for low-grade glioma surgery, particularly in the right hemisphere.⁴⁶⁻⁵¹ On the basis of these

arguments and a small number of patients in our study, the dilemma over removal of a tumor-infiltrated caudate nucleus remains.

No standards exist for quality measurement of neuro-oncologic care. Often, for a patient cohort, a variety of outcome measures are reported. Examples of oncologic outcome measures include the percentage of patients undergoing biopsy, median extent of resection, percentage of patients undergoing gross total resection, median residual disease volume, time to progression, and overall survival time. Examples of functional outcome measures include the percentage of patients with neurologic deficits, average cognitive performance scores, and health-related quality-of-life indices. We postulate that in addition to these outcome measures, probability map comparison of tumors, biopsies, and resections aids in evaluating oncologic outcome without bias from brain location. Probability maps thus enable outcome comparison among hospitals treating patients with diverging complexity of tumor locations. Another advantage is that probability map information is

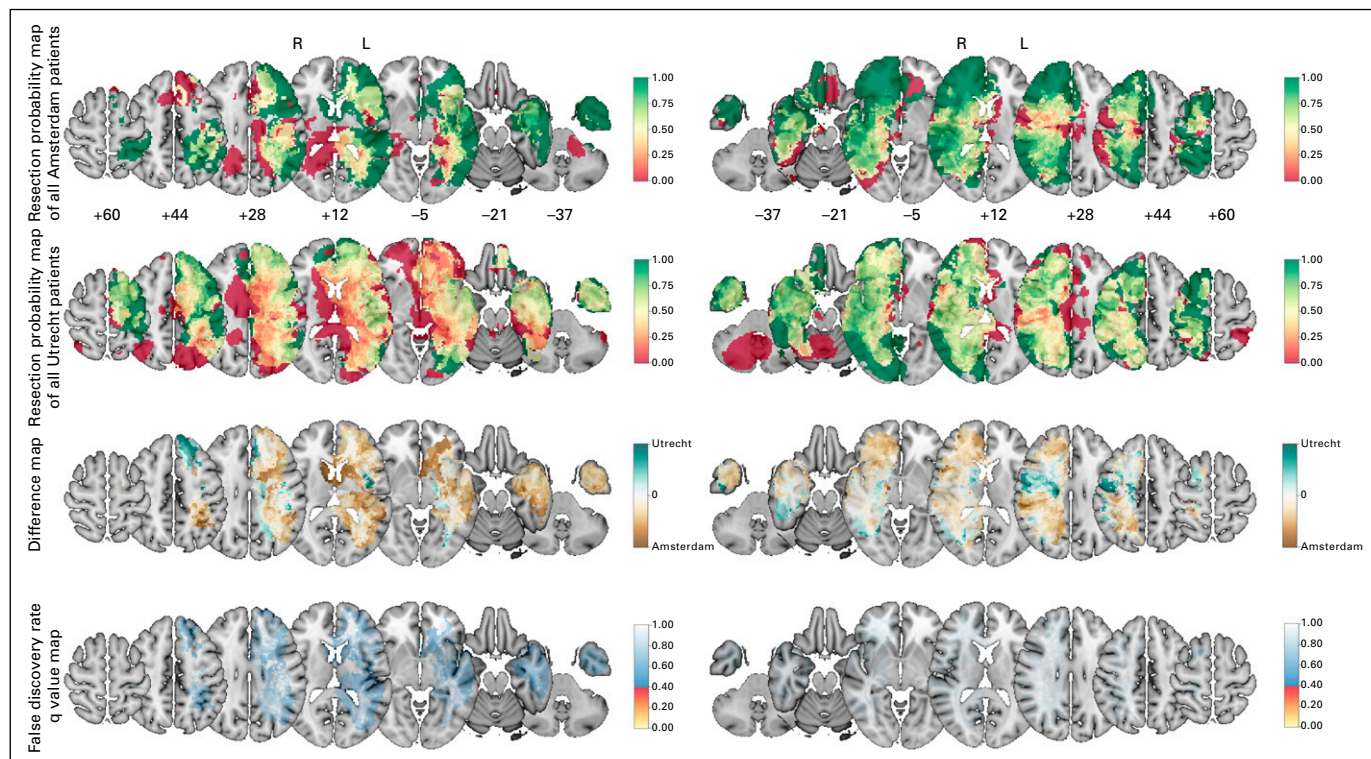


FIG 4. The resection probability map comparison of all 98 versus 174 patients per team to evaluate multidisciplinary decision making on tumor removal for the left (L; 38 v 95) and right (R) hemispheres (60 v 79). The layout and color codes are identical to those in Figure 3. The Data Supplement provides the full results.

available early in the course of disease, so quality assessment is not delayed until progression or survival. To optimize decision making and improve quality of care, a balance is required between oncologic and functional goals.⁵² That is, quality evaluation should not be narrowed to “less red is more quality” in BPMs and RPMs.

Several strengths of our study can be discerned. Volumetric analysis of residual disease was an important innovation compared with surgeon impression of resection completeness for an individual patient; now probability map comparison extends this by aggregating results for many patients per multidisciplinary team. New patterns of disease and treatment can be discovered, as demonstrated. By applying these methods at high resolution, we avoid assumptions on eloquence and interpretation of brain region classifications. Furthermore, an open-minded approach toward quality evaluation, learning, and improvement by changing current opinions was required from both teams. The well-documented patient cohorts were sizeable and seemed to represent clinical practice, which encourages extrapolation of our results.

Limitations to our findings include technical and inclusion issues. The technical observations on segmentation and registration are described in detail in the Data Supplement. Regarding inclusion, patients not undergoing surgery, and consequently without histopathologic diagnosis, were not included in our study. The impact of this potential selection bias is small, because we identified only three patients in

Amsterdam with a radiologically presumed glioblastoma without any surgery.

Our results have a number of practical implications. First, BPMs and RPMs can identify patients who will benefit most from intraoperative stimulation mapping and guide intraoperative stimulation mapping to the tumor regions with most function variability to potentially improve decision making. Instead of a static concept of functional and nonfunctional brain territories (ie, brain eloquence¹⁴), we now have a dynamic quantitative measurement per brain voxel of resectability that applies to a patient cohort treated by a team during a period of time. Second, neuro-oncologic registries have been initiated for the collection of patient data for quality evaluation.^{21,24,53} Our findings support the inclusion of standardized imaging as part of this collection. Third, our findings may serve patient counseling regarding decision making for biopsy or resection indications within multidisciplinary teams and for optimization of surgical strategies within neurosurgical teams. Fourth, probability maps may speed up knowledge transfer between neuro-oncologic teams. Fifth, the use of probability maps may enhance education and training of young neurologists, neurosurgeons, and radiologists by putting treatment decisions for individual patients into a broader perspective of aggregated patient cohorts and by providing intuitive and objective brain map visualization.

Future work could focus on larger patient populations to resolve the pictured clinical dilemmas and identify more patterns of treatment variation. The impact on patient outcome of identifying and resolving dilemmas resulting from treatment variation remains to be determined. Furthermore, RPMs could be used for individualized patient

care to predict resection results, contribute to image-guided surgical plans, and evaluate resection results.

In conclusion, probability maps of tumor location, biopsy, and resection provide additional information that can inform surgical decision making across multidisciplinary teams for patients with glioblastoma.

AFFILIATIONS

¹Vrije Universiteit Medical Center, Amsterdam, the Netherlands

²University Medical Center Utrecht, Utrecht, the Netherlands

³Netherlands Cancer Institute, Amsterdam, the Netherlands

⁴University College London, London, United Kingdom

⁵Hôpital Lariboisière, Paris, France

⁶University of California San Francisco, San Francisco, CA

CORRESPONDING AUTHOR

Philip C. De Witt Hamer, MD, VU University Medical Center, Department of Neurosurgery De Boelelaan 1117, 1081 HV Amsterdam, the Netherlands; e-mail: p.dewitthamer@vumc.nl.

SUPPORT

Supported in part by Project No. 10-10400-96-14003 of the Innovative Medical Devices Initiative, financed by the Netherlands Organisation for Scientific Research; by Research Grant No. VU2014-7113 from the Dutch Cancer Society; by the National Institute for Health Research of the University College London Hospitals Biomedical Research Centre; and by Brainlab, which provided segmentation software. This work was performed on the Dutch national e-infrastructure with the support of SURF Cooperative and the Translational Research IT project, an initiative from the Center for Translational Molecular Medicine.

AUTHOR CONTRIBUTIONS

Conception and design: Dominique M.J. Müller, Pierre A.J.T. Robe, Marnix G. Witte, Jan C. de Munck, Frederik Barkhof, Philip C. De Witt Hamer

Financial support: Jan C. de Munck, Frederik Barkhof, Philip C. De Witt Hamer

Administrative support: William P. Vandertop, Frederik Barkhof, Philip C. De Witt Hamer

Provision of study material or patients: Tatjana Seute, David P. Noske, William P. Vandertop, Mathilde C.M. Kouwenhoven, Philip C. De Witt Hamer

Collection and assembly of data: Dominique M.J. Müller, Pierre A.J.T. Robe, Marnix G. Witte, Marieke L.D. Broekman, Tatjana Seute, Frederik Barkhof, Emmanuel Mandonnet, Philip C. De Witt Hamer

Data analysis and interpretation: Dominique M.J. Müller, Pierre A.J.T. Robe, Roelant S. Eijgelaar, Marnix G. Witte, Martin Visser, Jan C. de Munck, Jeroen Hendrikse, David P. Noske, William P. Vandertop, Frederik Barkhof, Mathilde C.M. Kouwenhoven, Emmanuel Mandonnet, Mitchel S. Berger, Philip C. De Witt Hamer

Manuscript writing: All authors

Final approval of manuscript: All authors

Accountable for all aspects of work: All authors

AUTHORS' DISCLOSURES OF POTENTIAL CONFLICTS OF INTEREST

The following represents disclosure information provided by authors of this manuscript. All relationships are considered compensated.

Relationships are self-held unless noted. I = Immediate Family Member, Inst = My Institution. Relationships may not relate to the subject matter of this manuscript. For more information about ASCO's conflict of interest policy, please refer to www.asco.org/rwc or ascopubs.org/jco/site/ifc.

Pierre A.J.T. Robe

Honoraria: Medac

Marieke L.D. Broekman

Employment: Vertex (I)

Frederik Barkhof

Honoraria: Merck Serono, Biogen, Roche/Genentech

Consulting or Advisory Role: IXICO

Research Funding: Novartis (Inst), TEVA Pharmaceuticals Industries (Inst)

No other potential conflicts of interest were reported.

REFERENCES

- Ostrom QT, Gittleman H, Xu J, et al: CBTRUS statistical report: Primary brain and other central nervous system tumors diagnosed in the United States in 2009-2013. *Neuro Oncol* 18:v1-v75, 2016 (suppl 5)
- Young RM, Jamshidi A, Davis G, et al: Current trends in the surgical management and treatment of adult glioblastoma. *Ann Transl Med* 3:121, 2015
- Weller M, van den Bent M, Hopkins K, et al: EANO guideline for the diagnosis and treatment of anaplastic gliomas and glioblastoma. *Lancet Oncol* 15:e395-e403, 2014
- Chaichana KL, Jusue-Torres I, Navarro-Ramirez R, et al: Establishing percent resection and residual volume thresholds affecting survival and recurrence for patients with newly diagnosed intracranial glioblastoma. *Neuro-oncol* 16:113-122, 2014
- Marko NF, Weil RJ, Schroeder JL, et al: Extent of resection of glioblastoma revisited: Personalized survival modeling facilitates more accurate survival prediction and supports a maximum-safe-resection approach to surgery. *J Clin Oncol* 32:774-782, 2014
- Sanai N, Polley M-YY, McDermott MW, et al: An extent of resection threshold for newly diagnosed glioblastomas. *J Neurosurg* 115:3-8, 2011
- Brown TJ, Brennan MC, Li M, et al: Association of the extent of resection with survival in glioblastoma: A systematic review and meta-analysis. *JAMA Oncol* 2:1460-1469, 2016
- McGirt MJ, Mukherjee D, Chaichana KL, et al: Association of surgically acquired motor and language deficits on overall survival after resection of glioblastoma multiforme. *Neurosurgery* 65:463-469, discussion 469-470, 2009
- Gulati S, Jakola AS, Nerland US, et al: The risk of getting worse: Surgically acquired deficits, perioperative complications, and functional outcomes after primary resection of glioblastoma. *World Neurosurg* 76:572-579, 2011
- Rahman M, Abbatematteo J, De Leo EK, et al: The effects of new or worsened postoperative neurological deficits on survival of patients with glioblastoma. *J Neurosurg* 127:123-131, 2017

11. Jakola AS, Gulati S, Weber C, et al: Postoperative deterioration in health related quality of life as predictor for survival in patients with glioblastoma: A prospective study. *PLoS One* 6:e28592, 2011
12. Lamborn KR, Chang SM, Prados MD: Prognostic factors for survival of patients with glioblastoma: Recursive partitioning analysis. *Neuro Oncol* 6:227-235, 2004
13. Spena G, D'Agata F, Panciani PP, et al: Supratentorial gliomas in eloquent areas: Which parameters can predict functional outcome and extent of resection? *PLoS One* 8:e80916, 2013
14. Sawaya R, Hammoud M, Schoppa D, et al: Neurosurgical outcomes in a modern series of 400 craniotomies for treatment of parenchymal tumors. *Neurosurgery* 42:1044-1055, discussion 1055-1056, 1998
15. Sonabend AM, Zacharia BE, Cloney MB, et al: Defining glioblastoma resectability through the wisdom of the crowd: A proof-of-principle study. *Neurosurgery* 80:590-601, 2017
16. Barone DG, Lawrie TA, Hart MG: Image guided surgery for the resection of brain tumours. *Cochrane Database Syst Rev* 1:CD009685, 2014
17. De Witt Hamer PC, Robles SG, Zwinderman AH, et al: Impact of intraoperative stimulation brain mapping on glioma surgery outcome: A meta-analysis. *J Clin Oncol* 30:2559-2565, 2012
18. Gan HK, Rosenthal MA, Cher L, et al: Management of glioblastoma in Victoria, Australia (2006-2008). *J Clin Neurosci* 22:1462-1466, 2015
19. Graus F, Bruna J, Pardo J, et al: Patterns of care and outcome for patients with glioblastoma diagnosed during 2008-2010 in Spain. *Neuro Oncol* 15:797-805, 2013
20. Scoccianti S, Magrini SM, Ricardi U, et al: Patterns of care and survival in a retrospective analysis of 1059 patients with glioblastoma multiforme treated between 2002 and 2007: A multicenter study by the Central Nervous System Study Group of Airo (Italian Association of Radiation Oncology). *Neurosurgery* 67:446-458, 2010
21. Bauchet L, Mathieu-Daudé H, Fabbro-Peray P, et al: Oncological patterns of care and outcome for 952 patients with newly diagnosed glioblastoma in 2004. *Neuro Oncol* 12:725-735, 2010
22. Chang SM, Parney IF, Huang W, et al: Patterns of care for adults with newly diagnosed malignant glioma. *JAMA* 293:557-564, 2005
23. Sarnthein J, Stieglitz L, Clavien P-A, et al: A patient registry to improve patient safety: Recording general neurosurgery complications. *PLoS One* 11:e0163154, 2016
24. Hansen S, Nielsen J, Laursen RJ, et al: The Danish Neuro-Oncology Registry: Establishment, completeness and validity. *BMC Res Notes* 9:425, 2016
25. De Witt Hamer PC, Hendriks EJ, Mandonnet E, et al: Resection probability maps for quality assessment of glioma surgery without brain location bias. *PLoS One* 8:e73353, 2013
26. Ius T, Angelini E, Thiebaut de Schotten M, et al: Evidence for potentials and limitations of brain plasticity using an atlas of functional resectability of WHO grade II gliomas: Towards a "minimal common brain". *Neuroimage* 56:992-1000, 2011
27. Mandonnet E, Jbabdi S, Taillandier L, et al: Preoperative estimation of residual volume for WHO grade II glioma resected with intraoperative functional mapping. *Neuro-oncol* 9:63-69, 2007
28. Wen PY, Macdonald DR, Reardon DA, et al: Updated response assessment criteria for high-grade gliomas: Response assessment in neuro-oncology working group. *J Clin Oncol* 28:1963-1972, 2010
29. Caplan LR, Schmahmann JD, Kase CS, et al: Caudate infarcts. *Arch Neurol* 47:133-143, 1990
30. Kumral E, Eyyapan D, Balkir K: Acute caudate vascular lesions. *Stroke* 30:100-108, 1999
31. Grahn JA, Parkinson JA, Owen AM: The cognitive functions of the caudate nucleus. *Prog Neurobiol* 86:141-155, 2008
32. Pellizzaro Venti M, Paciaroni M, Caso V: Caudate infarcts and hemorrhages. *Front Neurol Neurosci* 30:137-140, 2012
33. Bhatia KP, Marsden CD: The behavioural and motor consequences of focal lesions of the basal ganglia in man. *Brain* 117:859-876, 1994
34. Kinoshita M, de Champfleury NM, Deverduin J, et al: Role of fronto-striatal tract and frontal aslant tract in movement and speech: An axonal mapping study. *Brain Struct Funct* 220:3399-3412, 2015
35. De Benedictis A, Sarubbo S, Duffau H: Subcortical surgical anatomy of the lateral frontal region: Human white matter dissection and correlations with functional insights provided by intraoperative direct brain stimulation: Laboratory investigation. *J Neurosurg* 117:1053-1069, 2012
36. Duffau H, Denvil D, Capelle L: Absence of movement disorders after surgical resection of glioma invading the right striatum. *J Neurosurg* 97:363-369, 2002
37. Marquardt G, Setzer M, Lang J, et al: Delayed hydrocephalus after resection of supratentorial malignant gliomas. *Acta Neurochir (Wien)* 144:227-231, discussion 231, 2002
38. Inamasu J, Nakamura Y, Saito R, et al: Postoperative communicating hydrocephalus in patients with supratentorial malignant glioma. *Clin Neurol Neurosurg* 106:9-15, 2003
39. Montano N, D'Alessandris QG, Bianchi F, et al: Communicating hydrocephalus following surgery and adjuvant radiochemotherapy for glioblastoma. *J Neurosurg* 115:1126-1130, 2011
40. Bulstrode HJCJ, Natalwala A, Grundy PL: Atypical presentation of delayed communicating hydrocephalus after supratentorial glioma resection with opening of the ventricles. *Br J Neurosurg* 26:222-226, 2012
41. Fischer CM, Neidert MC, Péus D, et al: Hydrocephalus after resection and adjuvant radiochemotherapy in patients with glioblastoma. *Clin Neurol Neurosurg* 120:27-31, 2014
42. John JK, Robin AM, Pabaney AH, et al: Complications of ventricular entry during craniotomy for brain tumor resection. *J Neurosurg* 127:426-432, 2016
43. Elliott JP, Keles GE, Waite M, et al: Ventricular entry during resection of malignant gliomas: Effect on intracranial cerebrospinal fluid tumor dissemination. *J Neurosurg* 80:834-839, 1994
44. Roelz R, Reinacher P, Jabbarli R, et al: Surgical ventricular entry is a key risk factor for leptomeningeal metastasis of high grade gliomas. *Sci Rep* 5:17758, 2015
45. Chan DTM, Hsieh SYP, Kam MKM, et al: Pattern of recurrence and factors associated with cerebrospinal fluid dissemination of glioblastoma in Chinese patients. *Surg Neurol Int* 7:92, 2016
46. Duffau H: Awake surgery for nonlanguage mapping. *Neurosurgery* 66:523-528, discussion 528-529, 2010
47. Wager M, Du Boisgueheneuc F, Pluchon C, et al: Intraoperative monitoring of an aspect of executive functions: Administration of the Stroop test in 9 adult patients during awake surgery for resection of frontal glioma. *Neurosurgery* 72:ons169-ons180; discussion ons180-ons181, 2013
48. Hulou MM, Cote DJ, Olubiye OI, et al: Awake right hemisphere brain surgery. *J Clin Neurosci* 22:1921-1927, 2015
49. Kinoshita M, Nakajima R, Shinohara H, et al: Chronic spatial working memory deficit associated with the superior longitudinal fasciculus: A study using voxel-based lesion-symptom mapping and intraoperative direct stimulation in right prefrontal glioma surgery. *J Neurosurg* 125:1024-1032, 2016
50. Herbet G, Moritz-Gasser S, Duffau H: Direct evidence for the contributive role of the right inferior fronto-occipital fasciculus in non-verbal semantic cognition. *Brain Struct Funct* 222:1597-1610, 2017

51. Vilasboas T, Herbet G, Duffau H: Challenging the myth of right nondominant hemisphere: Lessons from corticosubcortical stimulation mapping in awake surgery and surgical implications. *World Neurosurg* 103:449-456, 2017
52. Duffau H, Mandonnet E: The “onco-functional balance” in surgery for diffuse low-grade glioma: Integrating the extent of resection with quality of life. *Acta Neurochir (Wien)* 155:951-957, 2013
53. Dasenbrock HH, Yan SC, Smith TR, et al: Readmission after craniotomy for tumor: A national surgical quality improvement program analysis. *Neurosurgery* 80:551-562, 2017

