Altered corticospinal microstructure and motor cortex excitability in gliomas: an advanced tractography and transcranial magnetic stimulation study

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OBJECTIVE This prospective case-control study was conducted to examine whether spherical deconvolution (SD) can unveil microstructural abnormalities in the corticospinal tract (CST) caused by IDH-mutant gliomas. To determine the significance of abnormal microstructure, the authors investigated the correlation between diffusion parameters and neurophysiological data collected with navigated transcranial magnetic stimulation (nTMS).

METHODS Twenty participants (10 patients and 10 healthy controls) were recruited. Diffusion-weighted images were acquired on a 3-T MRI scanner using a cardiac-gated single-shot spin echo-planar imaging multiband sequence (TE 80 msec, TR 4000 msec) along 90 diffusion directions with a b-value of 2500 sec/mm² (FOV 256 × 256 mm). Diffusion tensor imaging tractography and SD tractography were performed with deterministic tracking. The anterior portion of the ipsilateral superior peduncle and the precentral gyrus were used as regions of interest to delineate the CST. Diffusion indices were extracted and analyzed for significant differences between hemispheres in patients and between patient and control groups. A navigated brain stimulation system was used to deliver TMS pulses at hotspots at which motor evoked potentials (MEPs) for the abductor pollicis brevis, first digital interosseous, and abductor digiti minimi muscles are best elicited in patients and healthy controls. Functional measurements such as resting motor threshold (rMT), amplitude of MEPs, and latency of MEPs were noted. Significant differences between hemispheres in patients and between patients and controls were statistically analyzed. The Spearman rank correlation was used to investigate correlations between diffusion indices and functional measurements.

RESULTS The hindrance modulated orientational anisotropy (HMOA), measured with SD tractography, is lower in the hemisphere ipsilateral to glioma (p = 0.028). The rMT in the hemisphere ipsilateral to a glioma is significantly greater than that in the contralateral hemisphere (p = 0.038). All measurements contralateral to the glioma, except for the mean amplitude of MEPs (p = 0.001), are similar to those of healthy controls. Mean diffusivity and axial diffusivity from SD tractography are positively correlated with rMT in the hemisphere ipsilateral to glioma (p = 0.02 and 0.006, respectively). The interhemispheric difference in HMOA and rMT is correlated in glioma patients (p = 0.007).

CONCLUSIONS SD tractography can demonstrate microstructural abnormality within the CST of patients with IDH1-mutant gliomas that correlates to the functional abnormality measured with nTMS.

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KEYWORDS low-grade gliomas; pathological excitability; tractography; spherical deconvolution; resting motor threshold; transcranial magnetic stimulation; corticospinal tract; oncology
can impact the extent of resection, occurrence of new motor deficits, and long-term recovery after surgery.\textsuperscript{5,7,8}

The CST can be radiologically assessed using diffusion MRI tractography. In tractography, information about the directional variance (anisotropy) of water diffusion is used to estimate mathematical reconstructions (trajectories) of white matter tracts. Diffusion imaging parameters include measures of the degree of anisotropy (fractional anisotropy [FA]), average displacement during diffusion (mean diffusivity [MD]), and average displacement in the radial and axial axes during diffusion (axial diffusivity [AD] and radial diffusivity [RD]). Traditional tractography is, however, limited by crossing fibers and the partial volume effects.\textsuperscript{9}

Traditional tractography parameters have yielded conflicting results when used to assess white matter connections in glioma patients. It is unclear, for example, whether these parameters represent axonal integrity,\textsuperscript{10} and there is inconsistent evidence about their variation between high-grade and low-grade gliomas.\textsuperscript{11,12} Finally, there is no consensus about the strength of their correlation to clinical motor function measures.\textsuperscript{13,14}

Advanced tractography methods such as spherical deconvolution (SD) estimate multiple fiber orientations per voxel. SD uses a continuous fiber orientation distribution function (fODF) to describe the amplitude (density) and orientation of multiple white matter fibers in each voxel. This function is derived from diffusion MRI data deconvolved with a model of white matter fiber (fiber response function). One of the widely used deconvolution methods is the damped Richardson-Lucy (dRL) SD.\textsuperscript{15} This method accurately resolves crossing fibers even in noisy data and preserves the angular resolution of main fiber orientations even in the presence of partial volume contamination.\textsuperscript{15,16} Moreover, advanced tractography parameters, such as measures of the hindrance modulated orientational anisotropy (HMOA), are tract specific and can be used as a proxy for fiber density and individual fiber anisotropy.\textsuperscript{16,17}

Navigated transcranial magnetic stimulation (nTMS) has been introduced in the last decade to noninvasively map the primary motor cortex for presurgical planning in patients with brain tumors.\textsuperscript{5,18–20} By stimulating the primary motor cortex, motor evoked potentials (MEPs) are elicited in peripheral muscles, and their latency and amplitude can be estimated. nTMS is also useful in defining the resting motor threshold (rMT) of the primary motor cortex. Differences in the rMT between the pathological and healthy hemisphere have been used to assess the presence of a “pathological excitability” of the motor cortex (defined as an rMT difference of at least 10% between the healthy and pathological brain hemisphere). Patients with a pathological excitability are at higher risk of developing motor deficits after surgery.\textsuperscript{5}

It is unclear whether any correlation exists between changes observed in the CST with tractography indices (including “traditional” and advanced parameters such as HMOA) and the presence of pathological excitability or altered MEP values. This is relevant, as at present we do not know whether changes seen with tractography in the CST have a neurophysiological correlate. In the current study, we assessed the microstructural quantification of the CST in healthy volunteers and in glioma patients using traditional diffusion tensor imaging (DTI) tractography and relative metrics (FA, MD, AD, RD) compared to SD tractography and tract-specific metrics (HMOA). We also correlated changes in diffusion measures with neurophysiology findings obtained with nTMS (MEP and rMT values and the presence or absence of pathological excitability of the motor cortex).

**Methods**

**Patient and Healthy Control Sample**

Twenty participants were recruited into this single-center prospective cohort study. Ten patients (4 females, 6 males; mean age 41 years, range 28–57 years) had a histologically confirmed diagnosis of infiltrative diffuse glioma involving or near the central lobule (precentral and postcentral gyri). Patients with previous surgery, multiple brain lesions, a WHO performance status of 2 or more, a histological diagnosis other than glioma, or an age below 18 or above 60 were not included. Ten healthy controls (4 females, 6 males; mean age 31 years, range 24–42 years) were closely matched to the patients with regard to sex and age. Healthy controls were screened for neurological and psychiatric history before inclusion. All patients and healthy controls were right-handed. Recruitment and data acquisition were approved by the South Yorkshire Research Ethics Committee, and written informed consent was obtained from all participants.

**MRI Study Acquisition**

A 3-T MR750 MRI scanner (General Electric) was used to acquire imaging data. Diffusion data were acquired using a cardiac-gated single-shot spin echo echoplanar imaging multiband sequence and 32-channel head coil (Nova Medical) with the following parameters: 75 axial slices, TE 80 msec, TR 5 R-R interval (approximately 4000 msec), acquisition matrix 128 × 128, and FOV 256 × 256 mm\textsuperscript{2}. Data were collected along 90 diffusion directions with a b-value of 2500 sec/mm\textsuperscript{2}. A total of 18 b = 0 diffusion weighting volumes were collected, 12 with anterior-to-posterior (AP) phase orientation and 6 with reversed posterior-to-anterior (PA) phase polarity for susceptibility correction. A multiband acceleration factor of 3 and an in-plane acceleration factor ARC (autocalibrating reconstruction for Cartesian imaging) of 2 were used. Diffusion data were de-noised\textsuperscript{21} and corrected for Gibbs ringing,\textsuperscript{22} motion,\textsuperscript{23} eddy currents,\textsuperscript{24} and susceptibility distortion.\textsuperscript{25} Outlier slices were then detected and replaced.\textsuperscript{24} Finally, intravolume slice motion was corrected for.\textsuperscript{25} Data from one patient were incomplete and therefore excluded from analysis.

**Tractography Reconstruction**

Neuroimaging analysis was completed by NatBrainLab members (A.S.M. and A.B.) under the supervision of the senior author (F.D.). StarTrack software was used for traditional single tensor tractography (DTI) and advanced SD tractography (www.natbrainlab.com). Whole-brain DTI tractography used deterministic tracking (Euler streamline propagation with a step size of 1 mm). Included
trajectory streamlines were no shorter than 20 mm and no
longer than 30 mm. Tracking was terminated if there was
development any greater than 25° or FA below 0.15. The SD
modeling was performed using the dRL algorithm (fiber
response parameter ALFA = 1.5, 300 iterations, regular-
ization η = 0.0015, ν = 16). Deterministic tracking was
performed with an angle threshold of 35°, an absolute
fODF amplitude threshold of 0.002, and a step size of 1
mm. Seeding was set to 1 seed per fiber orientation.

The CST was dissected using regions of interest (ROIs)
previously described: anterior portion of the ipsilateral
superior peduncle, precentral gyrus, and CST area in
the brainstem.26,27 Freehand ROIs were outlined on 2D
FA maps on TrackVis to select fibers from specific ana-
tomical regions through which the CST passes. Exclusion
ROIs were created based on prior anatomical knowledge
to eliminate artifactual streamlines. Trajectories were
checked by the senior author (F.D.).

Measures of microstructure were averaged across all
voxels through which the CST passed. For all CST tra-
jectories (both DTI and SD), mean FA, MD, AD, and RD
were extracted. In addition, for trajectories reconstructed
by SD, the HMOA (absolute amplitude of each lobe in the
fODF) was extracted. The maximum HMOA is averaged
across all voxels through which the CST passes and used
as a proxy for fiber density and FA in SD trajectories.

Transcranial Magnetic Stimulation

The eXimia navigated brain stimulation system (Nex-
stim) was used for preoperative motor mapping. A fig-
ure-8 coil was used to deliver a TMS pulse. The electric
field was estimated based on a dynamic spherical model
adjusted in real time. The lowest stimulation intensity
(percentage of maximum stimulator intensity) capable of
eliciting an MEP in 50% of trials for the hand area in both
brain hemispheres was used to calculate the rMT. MEPs
were recorded from the abductor pollicis brevis, first digi-
tal interosseous, and abductor digiti minimi muscles. The
mean amplitude and mean latency of MEPs of the tested
muscles were measured. TMS mapping (not blinded) was
performed by two operators (F.V. and H.H.). A pathologi-
cal excitability was considered to be present when the in-
temperisperic rMT ratio (irMTTr) was larger than 10% (irMTTr < 0.90 or > 1.10).22

Tumor Resection

Tumor removal was performed with the aim of maximal
resection according to anatomical and functional bound-
daries. Neuronavigation ( StealthStation S7, Medtronic) was
employed in all cases. The compound 5-aminolevulinic
acid (5-ALA) was used as an adjunct when enhancement
was present on preoperative MRI. Intraoperative motor
mapping with a constant current stimulator and a monop-
olar probe (ISIS Xpress system, Inomed GmbH) was per-
duced using train of 5 stimulations with a 0.5-msec pulse
width and interstimuli interval of 4.0 msec. Cortical map-
ing was performed with positive pulse form, and the sub-
cortical mapping was performed with positive pulse form.
Motor responses were recorded from the face, upper limb/
hand, and/or lower limb/foot, according to tumor location
and extent of motor cortex exposure. The monitoring of
the integrity of the CST was performed with continuous
MEPs elicited from a 4-contact subdural strip electrode
placed over the precentral gyrus (best contact of the 4).
Placement of the strip was guided by the cortical mapping
or by the nTMS preoperative mapping when the primary
motor cortex was not exposed. The extent of resection
was calculated as follows, based on the postoperative MRI
study obtained within 48 hours from surgery: gross total
(no residual tumor visible on postoperative MRI), subtotal
(residual volume 1–10 ml), and partial (residual volume
more than 10 ml).23 The residual volume was calculated
based on the residual hyperintensity visible on post opera-
tive T2/FLAIR sequences.

Statistical Analysis

SPSS software (version 25, IBM Corp.) was used for
statistical analysis. Prior to hypothesis testing, the distri-
bution of diffusion indices and rMTs were assessed with the
Shapiro-Wilk test. For all statistical tests, a significance
threshold of 0.05 was used. We tested the hypothesis that
microstructure and TMS indices are significantly differ-
ent between cerebral hemispheres ipsilateral and contra-
lateral to a glioma. Relative differences of indices from
trajectories ipsilateral and contralateral to the tumor were
calculated. The Wilcoxon signed-rank test was used to
determine if the difference is significant. This hypothesis
assumes that measurements contralateral to the glioma are
within the healthy range. The assumption was tested using the
Kolmogorov-Smirnov test. It was also used to test for asymmetry between the right and left hemispheres in the
healthy data set. This test was chosen because it does not
assume that the data are homoscedastic.

The hypothesis that functional (nTMS) and anatomical
(tractography) measurements correlate was tested us-
ing the Spearman rank order correlation, which was used
because it does not assume that the data are normally dis-
tributed.

Results

Patient Results

Histological analysis for the 10 patients was consistent
with WHO grade II (1 case) and III (9 cases) gliomas; all
10 cases were IDH1 mutant. There were 5 astrocytomas
and 5 oligodendrogliomas (Ip/19q codeletion confirmed).
Seventy percent of the patients had a glioma within 1 mm
of the CST as assessed on preoperative tractography. All
patients had a Medical Research Council (MRC) grade
5/5 preoperatively. Thirty percent of the patients had
gross-total resection, 50% had subtotal resection, and 20%
had partial resection. One patient suffered a postoperative
motor deficit (MRC grade 4/5 in the right upper limb) that
did not improve by the 6-month follow-up. This was pos-
sibly the result of a postoperative brain abscess that had
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rMT on TMS generally corresponding to a higher rMT intraoperatively), although the relationship was not significant. Mean subcortical stimulation was 5.4 mA. Fibers of the CST were identified in 7 patients using subcortical stimulation, which guided resection.

Tractography Results

Qualitative analysis of CST trajectories from DTI and SD revealed that in 60% of the cases, SD tractography reconstructs a thicker trajectory through the glioma/edematous regions. In all of the reconstructions, SD tractography reconstructed trajectories that more closely resembled the anatomical fanning shape of the CST at its cranial end (Fig. 1).

Diffusion indices ipsilateral and contralateral to a glioma are illustrated in Figs. 2 and 3. The differences in all traditional tractography measures (FA, MD, AD, and RD) between the hemisphere with a glioma and the contralateral hemisphere are not statistically significant using both SD and DTI (p = 0.759, 0.759, 0.400, 0.988, not applicable, respectively) and SD tractography (p = 0.988, 0.759, 0.400, 0.759, 0.759) in healthy controls.

Transcranial Magnetic Stimulation Results

In healthy controls, the median rMT in the left hemisphere (35%, IQR ± 10%) is statistically similar to that in the right hemisphere (35%, IQR ± 9%; p = 0.988).

In patients, the median rMT in the hemisphere contralateral to glioma was 40% (IQR ± 15%) of the maximum output stimulation. In brain hemispheres with glioma, the median rMT was 46% (IQR ± 17%). The rMT in the hemisphere ipsilateral to glioma was significantly greater than that in the contralateral hemisphere (median relative difference −21.1%, IQR ± 14.9%; p = 0.038). The rMT contralateral to a glioma was not statistically different from the rMT in healthy subjects (p = 0.073).

Nine of the 10 patients demonstrated a pathological excitability of the motor cortex, with the relative difference in rMT ranging from 12% to 136% between the affected and unaffected hemispheres. Only 2 healthy controls had a relative difference in the rMT larger than 10% between the two hemispheres, but this difference appeared to be modest (11.5% and 12%) and was not statistically significant (p = 0.988).

The median mean amplitude in healthy brains was 212 mV (IQR ± 115.65 mV). There was no right to left asymmetry in healthy brains (p = 0.699). In the patient group, the median mean amplitude ipsilateral to glioma was 76 mV (± 88 mV) and contralateral to glioma was 123 mV (± 46 mV). There was no significant difference in the mean amplitude of MEPs between brain hemispheres contralateral and ipsilateral to a glioma (p = 0.297).

The mean amplitude of MEPs contralateral to a glioma was statistically smaller than that in healthy controls (p = 0.001). Similarly, the mean amplitude of MEPs ipsilateral to a glioma was statistically smaller than those in healthy brains (p = 0.002).

The median mean latency of MEPs in a healthy brain was 23 msec (IQR ± 1 msec) on the left and 23 msec (± 1.5 msec) on the right. There was no asymmetry between left and right hemispheres in healthy brains (p = 0.336). The median mean latency of MEPs ipsilateral to a glioma was 24 msec (± 47 msec) and contralateral to a glioma was
24 msec (± 5 msec). There was no significant difference in the mean latency of MEPs between brain hemispheres ipsilateral and contralateral to a glioma (p = 0.518). The mean latencies contralateral and ipsilateral to a glioma were not statistically different from those of healthy brains (p = 0.928).

Correlation Between TMS and Tractography

The higher the rMT ipsilateral to glioma, the lower the HMOA ipsilateral to glioma. The correlation between CST microstructure, as defined by the relative difference between HMOA contralateral and ipsilateral to the glioma, and motor excitability, as defined by the relative difference between rMT contralateral and ipsilateral to the glioma, was also analyzed. The relative difference in HMOA is significantly related to the relative difference in the rMT for the motor cortex contralateral and ipsilateral to glioma. In other words, the degree of microstructure changes seen in the CST (with HMOA) correlates with the degree of functional changes (assessed with rMT; p = 0.007, r = −0.871; Fig. 4).

In brain hemispheres affected by glioma, MD and AD of the CST positively correlated with rMT when SD tractography was used (r = −0.518, p = 0.020 and r = 0.824, p = 0.006, respectively; Figs. 5 and 6). MD of the CST was also negatively correlated with latency in healthy brains (r = −0.518, p = 0.028).

Discussion

It is well known that gliomas can cause changes in the surrounding white matter through a combination of infiltration, edema, and mass effect. The interaction between gliomas and white matter tracts is of clinical relevance, as it can contribute to the development of neurological deficits. From a neurosurgical perspective, it is mandatory to spare functional white matter tracts, which often constitute the functional boundary of resection identified with intraoperative mapping. There is extensive literature to suggest that gliomas can change the diffusion profile of white matter tracts, as assessed on preoperative tractography. Several studies have reported that white matter microstructure can be affected by tumors. A recent meta-analysis on the relation between diffusion metrics and tumor grade in gliomas has suggested the significant effects of tumor grade on average MD and FA. These changes were more consistently observed in high-grade gliomas, supporting the concept that high-grade tumors are more destructive and infiltrative than low-grade tumors. However, the authors warned against a large heterogeneity in the existing...
literature, especially with regard to tractography methods, tract analysis, and patient populations.

In the present study, we focused on the changes observed in diffusion indices of the CST in a homogeneous population of IDH-mutant grade II and III gliomas in close relationship with the central lobule and CST. Diffusion parameters were assessed with both traditional (FA, MD, AD, and RD) and advanced (HMOA) indices, and results from the glioma population were compared to those of a healthy population. Our results illustrate that traditional diffusion indices, but not HMOA, are within the healthy range in brain hemispheres ipsilateral to a glioma (i.e., not significantly different compared to healthy subjects or to hemispheres contralateral to a glioma). Traditional diffusion indices (FA, MD, AD, and RD) are based on average voxel characteristics and are therefore susceptible to partial volume effects and crossing fibers. This is particularly relevant for the CST, where there is a significant amount of fiber crossing from callosal and superior longitudinal fasciculus fibers. Additionally, tractography results represent a mathematical reconstruction of the underlying white matter trajectories. Thus, tractography may show partial reconstructions in the presence of a strong partial volume effect due to edema or tumor infiltration. Nevertheless, for this study, we acquired state-of-the-art diffusion imaging data and applied advanced preprocessing and tractography methods. In particular, the use of high b-values and SD tractography helped to suppress these signal contaminations, resulting in reliable tractography reconstructions consistent across all subjects.

On the other hand, advanced tractography indices, such as HMOA, can be considered true tract-specific indices and can identify characteristics not illustrated by other diffusion indices. In addition, changes in tract-specific characteristics due to factors such as axonal loss, especially in regions of crossing fibers, are better characterized by HMOA. This may be the reason why only a significant decrease in HMOA is seen in CST trajectories ipsilateral to a glioma. A decrease in HMOA can therefore represent a microstructural change in the CST due to axonal loss. Although more commonly reported in glioblastomas, changes in traditional diffusion indices have also been found in grade II and III gliomas. However, the IDH status of gliomas was not reported in these papers; therefore, the patient populations are difficult to compare. It is well recognized that IDH-mutant and IDH-wildtype gliomas differ significantly in terms of biological behavior, aggressiveness, and prognosis. In our homogeneous series of IDH-mutant, predominantly grade III gliomas, traditional indices did not show any significant alteration: this may reflect the relatively more indolent behavior of these tumors, requiring advanced (and more sensitive) dif-

<table>
<thead>
<tr>
<th>Index</th>
<th>Median Value in Healthy Controls</th>
<th>Median Value Contralateral to Tumor</th>
<th>p Value (healthy controls vs CST contralateral to glioma)</th>
<th>Median Value Ipsilateral to Tumor</th>
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<td>FA</td>
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<tr>
<td>MD</td>
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<tr>
<td>AD</td>
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<td>0.413</td>
<td>0.915615</td>
</tr>
<tr>
<td>RD</td>
<td>0.3991535</td>
<td>0.404025</td>
<td>0.453</td>
<td>0.412706</td>
</tr>
<tr>
<td>HMOA</td>
<td>0.0243325</td>
<td>0.0247182</td>
<td>0.724</td>
<td>0.0231091</td>
</tr>
</tbody>
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FIG. 4. Correlation between abnormal excitability and abnormal microstructure in glioma patients.
fusion parameters to capture microstructural changes at the level of the CST. A larger sample with a more balanced group of grade II and grade III gliomas is needed to address the effect of increased grading on the observed results. It is possible that the observed changes on a microstructural level may be attributable to increased mitotic activity in grade III gliomas.

To further investigate the meaning of the microstructural changes observed in the CST, we attempted to establish whether there is a neurophysiological correlate between changes in the HMOA and measures of cortical excitability of the motor system. To this extent, we used preoperative nTMS as a noninvasive way to assess the motor system in patients and healthy subjects. The advantage of this approach is that, unlike other modalities of noninvasively mapping motor function (such as MEG or fMRI), nTMS has better spatial resolution and allows one to record MEPs for offline analysis. In addition, unlike direct cortical stimulation, nTMS can explore both hemispheres, permitting interhemispheric comparison in patients (to compare the motor excitability of the pathological versus nonpathological hemisphere) and allowing data to be obtained from healthy volunteers.

Different authors have attempted to establish a relationship between TMS and tractography-derived variables by studying different pathologies and age groups. Hübers et al. failed to demonstrate a relation between rMT and FA in healthy subjects in a voxel-wise analysis. Jang and colleagues established a correlation between rMT and fiber number in the hemispheres of patients with motor deficits after ischemic vascular injury but failed to show a correlation with the FA values of the CST. Papadelis et al. studied the evolution of rMT and DTI-derived metrics in children with hemiplegic cerebral palsy and found that the normal correlation of AD, MD, and rMT with age was disrupted in these patients. In the particular case of
gliomas, Sollmann et al. reported an inverse correlation between rMT and the distance between tumor and CST in patients with permanent or temporary motor deficits after surgery. However, a clear correlation between nTMS, tractography, and gliomas is missing in the literature.

In our series, the rMT of brain hemispheres with glioma was significantly higher than the rMT of hemispheres contralateral to glioma and that of healthy subjects. In addition, 9 of 10 patients demonstrated a pathological MT (defined as a difference of more than 10% between the rMT of the two hemispheres), whereas an rMT difference larger than 10% was found in only 2 of 10 healthy subjects and with very small and insignificant relative differences. (This latter finding is likely attributable to a margin of error intrinsic to nTMS.) The rMT changes observed in the brain hemisphere with glioma confirm the interaction between glial tumors and motor function, suggesting the presence of a pathological excitability of the motor system in patients.

When SD tractography was used, a significant relationship was found between diffusion indices and rMT. A significant correlation was found between an increase in MD and AD and an increase in the rMT in the affected brain hemisphere. In addition, a relation between changes in the CST microstructure and the presence of a pathological motor excitability was found. The higher the rMT ipsilateral to a glioma, the lower the HMOA ipsilateral to a glioma. These results point toward a direct relationship between structural changes in the CST and pathological excitability of the motor system. Different mechanisms can be hypothesized to explain this finding: a loss of axons in the CST, direct tumor invasion, or distortion and compression on the CST due to mass effect. However, in our series, the presence of peritumoral edema was minimal and diffusion weighting was high, making the latter hypothesis less likely.

It is important to note that all patients were neurologically intact prior to surgery, showing no motor deficits. A correlation between CST changes and pathological rMT may therefore suggest that preclinical changes affecting the motor system can be captured by combining SD tractography and TMS. Risk-stratification models combining imaging and TMS data have been designed with the aim to identify patients at high risk for developing new motor deficits following surgery. Our results support the incorporation of tract-specific indices such as HMOA and interhemispheric rMT difference in such risk-stratification models. Information about CST integrity could also be applied when planning the resection of gliomas close to the CST, helping the surgeon in deciding when to stop the resection at the subcortical level in order to prevent damage to the CST itself. Further research is needed to address these aspects.

Conclusions

Our results demonstrate that SD tractography reveals the presence of microstructural abnormality within the CST of patients with IDH1-mutant gliomas close to the motor cortex and CST. The microstructural CST changes, captured by the HMOA index, correlate with a pathological excitability of the motor cortex in neurologically intact patients. The combined use of SD tractography and TMS can provide valuable information on the anatomical-functional preoperative assessment of the motor system.

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