Reconstruction of the Corticospinal Tract in Patients with Motor-Eloquent High-Grade Gliomas Using Multilevel Fiber Tractography Combined with Functional Motor Cortex Mapping


ABSTRACT

BACKGROUND AND PURPOSE: Tractography of the corticospinal tract is paramount to presurgical planning and guidance of intraoperative resection in patients with motor-eloquent gliomas. It is well-known that DTI-based tractography as the most frequently used technique has relevant shortcomings, particularly for resolving complex fiber architecture. The purpose of this study was to evaluate multilevel fiber tractography combined with functional motor cortex mapping in comparison with conventional deterministic tractography algorithms.

MATERIALS AND METHODS: Thirty-one patients (mean age, 61.5 [SD, 12.2] years) with motor-eloquent high-grade gliomas underwent MR imaging with DWI (TR/TE = 5000/78 ms, voxel size = 2 × 2 × 2 mm³, 1 volume at b = 0 s/mm², 32 volumes at b = 1000 s/mm²). DTI, constrained spherical deconvolution, and multilevel fiber tractography—based reconstruction of the corticospinal tract within the tumor-affected hemispheres were performed. The functional motor cortex was enclosed by navigated transcranial magnetic stimulation motor mapping before tumor resection and used for seeding. A range of angular deviation and fractional anisotropy thresholds (for DTI) was tested.

RESULTS: For all investigated thresholds, multilevel fiber tractography achieved the highest mean coverage of the motor maps (eg, angular threshold = 60°; multilevel/constrained spherical deconvolution/DTI, 25% anisotropy threshold = 71.8%, 22.6%, and 11.7%) and the most extensive corticospinal tract reconstructions (eg, angular threshold = 60°; multilevel/constrained spherical deconvolution/DTI, 25% anisotropy threshold = 26,485 mm², 6308 mm², and 4270 mm²).

CONCLUSIONS: Multilevel fiber tractography may improve the coverage of the motor cortex by corticospinal tract fibers compared with conventional deterministic algorithms. Thus, it could provide a more detailed and complete visualization of corticospinal tract architecture, particularly by visualizing fiber trajectories with acute angles that might be of high relevance in patients with gliomas and distorted anatomy.

ABBREVIATIONS: ADT = angular deviation threshold; CSD = constrained spherical deconvolution; CST = corticospinal tract; FAT = fractional anisotropy threshold; FOD = fiber orientation distribution; MLFT = multilevel fiber tractography; nTMS = navigated transcranial magnetic stimulation

Gliomas are the most prevalent malignant brain tumors in adults, and particularly anaplastic astrocytoma and glioblastoma as representatives of high-grade gliomas have poor prognoses. Contemporary treatment combines neurosurgical tumor resection with extended focal radiation therapy and adjuvant chemotherapy. Specifically, a maximum extent of resection correlates to

From the Department of Biomedical Engineering (A.Z., J.P.), Eindhoven University of Technology, Eindhoven, the Netherlands; Department of Diagnostic and Interventional Neuroradiology (N.S.), University Hospital ULM, Ulm, Germany; Helmholtz Zentrum München, Munich, Germany; Department of Diagnostic and Interventional Neuroradiology (N.S., F.K., B.W., C.Z., J.S.K.), Klinikum rechts der Isar, Image-Based Biomedical Modeling (F.K., B.M.), Department of Informatics, TranslATUM (F.K., B.W.), Central Institute for Translational Cancer Research, and Department of Neurosurgery (J.G., A.S., S.M.K.), School of Medicine, Klinikum rechts der Isar, TUM-Neuroraming Center (N.S., C.Z., J.S.K., S.M.K.), Central Institute for Translational Cancer Research, and Department of Neurosurgery (J.G., A.S., S.M.K.), School of Medicine, Klinikum rechts der Isar, Technical University of Munich, Munich, Germany; Department of Radiology and Biomedical Imaging (N.S.), University of California, San Francisco, San Francisco, California; Department of Imaging and Pathology (A.R., S.S.), Translational MR, and Department of Neurosciences (A.R., S.S.), Leuven Brain Institute, Katholieke Universiteit Leuven, Leuven, Belgium; Image Sciences Institute (A.D.L., A.L.) and Neurology Department (A.D.L.), University Medical Center Utrecht Brain Center, University Medical Center Utrecht, Utrecht, The Netherlands; and Department of Quantitative Biomedicine (B.M.), University of Zurich, Zurich, Switzerland.

Please address correspondence to Andrey Zhylka, MD, Eindhoven University of Technology, Biomedical Engineering, Rondom 70, Eindhoven, 5612AP, Netherlands; e-mail: a.zhylka@tue.nl; @AnjensonZ; @NSollmann

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prolonged survival and improved quality of life. However, a maximum extent of resection needs to be weighed against the risk of surgery-related functional decline such as persistent paresis or aphasia, which may arise from tumor resection in or near functionally eloquent structures such as the motor or language cortex. Additionally, subcortical WM pathways such as efferent fibers that interconnect certain brain areas or course down to the periphery may need to be respected to avoid lasting functional deficits.

Imaging and mapping of brain function are essential for a maximum safe resection combined with a high extent of resection and a low-risk profile for permanent functional deterioration. In the preoperative setup, DWI with tractography is used for delineation of WM architecture, which can then be visualized and respected during tumor resection. Particularly, DTI-based tractography is commonly applied to reconstruct certain fiber tracts, relating to its comparatively wide availability and low false-positive rate. However, despite its frequent application in the preoperative work-up of patients with glioma, the technique has relevant shortcomings that render the accuracy of the method questionable. Specifically, major issues relate to potential underrepresentation of fiber branching or crossing fibers, which are difficult to resolve by DTI-based tractography. Alternatives to DTI-based tractography exist, including methods such as constrained spherical deconvolution (CSD)—tractography, which has shown improved specificity compared with DTI-based tractography, given that CSD-based tractography is determined by higher angular resolution and the possibility of also disentangling more complex fiber configurations. In CSD, multiple fibers passing through a voxel with distinct orientations can be estimated, depending on the fiber orientation distribution (FOD).

Multilevel fiber tractography (MLFT) has been developed from CSD-based tractography and similarly propagates fiber pathways on the basis of FOD peaks, with the main advancement that MLFT proposes that FOD peaks do not solely reflect crossing fibers but may also reflect high angular deviation of fibers or their branching. A previous study has demonstrated that MLFT improved reconstruction of the corticospinal tract (CST) in patients with motor-eloquent high-grade gliomas by generating fiber bundles with higher radial extent (ie, delineation of CST fanning with a wider range) compared with DTI as well as CSD-based deterministic CST tractography, thus potentially showing a more complete picture of the actual CST architecture. Yet, without optimal seeding of the ROI for tractography, MLFT would be at considerable risk of reconstructing false-positive WM pathways, given that it can also include acute angles of fiber courses, which might be particularly frequently observed among patients with brain tumors due to the space-occupying effect that may lead to fiber diversion and compression.

The issue of optimal ROI seeding for CST reconstruction may be addressed by combining MLFT with preoperative functional mapping, such as navigated transcranial magnetic stimulation (nTMS). In essence, nTMS can target neurostimulation to the brain with subcentimeter precision and enables the spatial identification and demarcation of the motor cortex in relation to a lesion. Particularly, nTMS-based motor mapping has been frequently used in the preoperative setup and for guidance of intraoperative resection in patients with motor-eloquent brain tumors. The approach has been shown to result in cortical motor maps similar to those generated by intraoperative direct electrical stimulation as the reference standard for functional cortical mapping. More recently, combinations of tractography with nTMS have been used to establish function-based tractography of the CST, which is based on the definition of the nTMS-derived cortical motor map as an ROI.

Against this background, the purpose of the present study was to combine MLFT for reconstruction of the CST with nTMS for encasing the functional motor cortex in patients with motor-eloquent gliomas. We hypothesized that MLFT may show higher coverage of the nTMS motor map (ie, the highest percentage of fibers of the CST being connected to the motor cortex) compared with deterministic DTI-based and CSD-based tractography.

**MATERIALS AND METHODS**

**Study Design and Patient Cohort**

This study was approved by the local institutional review board (Ethikkommission Technische Universität München) and was conducted in accordance with the Declaration of Helsinki. The requirement for written informed consent for this study was waived due to the retrospective design.

Eligible patients were identified by chart review, covering the time interval from February 2019 to February 2020. Inclusion criteria were the following: 1) older than 18 years of age, 2) availability of preoperative 3T MR imaging, including diffusion-weighted sequences, 3) diagnosis of a high-grade glioma (based on imaging findings and later confirmation by histopathologic evaluation of biopsy probes or tumor tissue harvested during surgical resection), 4) suspected motor-eloquent tumor location according to preoperative MR imaging, and 5) availability of preoperative nTMS-based motor mapping of the tumor-affected hemisphere.

Overall, 31 patients (mean age, 61.5 [SD, 12.2] years; age range, 34.4–85.1 years; 12 women) matched the inclusion criteria and were considered for this study. Three patients were diagnosed with World Health Organization grade III gliomas, and 28 patients with World Health Organization grade IV gliomas, and the right hemisphere was affected by the tumor volume in 19 patients.

**Data Acquisition**

**MR Imaging.** Preoperative MR imaging was performed on two 3T scanners (Achieva dStream or Ingenia; Philips Healthcare) using a 32-channel head coil. The imaging protocol included a 3D FLAIR sequence (TR/TE = 4800/277 ms, 1-mm³ isotropic voxels covering the whole head), an axial T2-weighted sequence (TR/TE = 3396/87 ms, voxel size = 0.36 × 0.36 × 4 mm³), a diffusion-weighted sequence (TR/TE = 5000/78 ms, voxel size = 2 × 2 × 2 mm³, 1 volume at b=0 s/mm², 32 volumes at b=1000 s/mm²), and a 3D T1-weighted turbo field echo sequence (TR/TE = 9/4 ms, 1-mm³ isotropic voxels covering the whole head) without and with intravenous injection of a contrast agent using a dose of 0.2 mL per kg body weight of gadoteric acid (Dotagraf, 0.5 mmol/mL; Jenapharm).

**nTMS.** Preoperative motor mapping of the tumor-affected hemisphere was performed using nTMS (NBS system 4.3 or 5.0; Nexstim). For neuronavigation, the preoperatively acquired
contrast-enhanced 3D T1-weighted turbo field echo sequence was used. Motor mapping was performed according to a standardized protocol using a figure-of-eight stimulation coil and a biphasic pulse wave application. Both representations of upper and lower extremity muscles were mapped within the tumor-affected hemisphere using an intensity of 105%–110% of the individual resting motor threshold for the upper extremity and an intensity of at least 130% of the resting motor threshold for the lower extremity muscles. Motor-positive nTMS points were identified during post hoc analysis, which were required to show an amplitude of motor-evoked potentials of ≥50 μV, with motor-evoked potential onset latencies within the typical range.30,41

**Image Segmentation and Processing of DWI Data.** Before image segmentation, all MR imaging data were transferred to Montreal Neurological Institute space (with an isotropic voxel size of 1 mm³). Given that commonly used parcellation pipelines may not produce robust segmentations in the presence of brain tumors, lesion filling was performed using the Virtual Brain Grafting toolkit (https://github.com/KUL-Radneuron/KUL_VBG/).44,45 This approach replaces the tumor volume with synthetic healthy tissue.45 As a prerequisite for lesion filling, lesion segmentation was performed using the BraTS toolkit (https://github.com/neuronflow/BraTS-Toolkit).46,47 The BraTS toolkit was provided with T1-weighted images, both noncontrast and contrast-enhanced, as well as FLAIR and T2-weighted images, to perform the lesion segmentation.46,47 The segmentation differentiates between the tumor core (necrotic center and contrast-enhancing tumor parts) and FLAIR-hyperintense zones (edema/tumor infiltration).

All diffusion-weighted data sets were preprocessed by performing motion and eddy current corrections.48 To estimate FODs, we used recursive calibration of the response function.49 Given the number of acquired diffusion-weighted volumes, the order of spherical harmonics describing FODs was set to 6. All processing was performed in ExploreDTI (Version 4.8.6; http://www.exploredti.com/).

Because nTMS was performed as a separate acquisition, nTMS points had to be transferred to the space of the T1-weighted and diffusion imaging data sets, achieved by performing registration of the contrast-enhanced T1-weighted images to the masks containing all motor-positive nTMS points of the tumor-affected hemisphere. All nTMS points were enlarged using a hull of 2-mm radius by default to provide the final motor cortex seed mask.36,37 All image coregistrations and segmentations were visually inspected for quality and manually corrected, when necessary, by a trained neuroradiologist.

**Tractography Algorithms.** Three tractography algorithms were used for CST reconstruction: DTI-based tractography, CSD-based tractography, and MLFT. MLFT has been recently introduced and is a bundle-specific algorithm that reconstructs fiber bundles as multilevel structures.27,28 The algorithm requires seed and target regions (ie, the ipsilateral anterior pontine brainstem level and the nTMS-derived cortical motor map) and a maximum number of levels.27,28 At each iteration, streamline propagation is performed using a deterministic CSD-based algorithm. At the end of each iteration, FOD peaks of the points along pathways that did not reach the target region are used as initial directions at the following iteration. Thus, the potential of the peaks to represent branches is explored.

The angular deviation threshold (ADT) is aimed at maintaining smoothness and physical plausibility of the fiber pathways because it prevents propagation into directions highly deviating from the previous step direction. However, the threshold may impact the accuracy of the reconstruction, depending on the voxel and angular resolution of the data. Thus, each tractography algorithm was run with 3 different ADTs, namely 20°, 45°, and 60°. For DTI-based tractography, 3 levels of the fractional anisotropy threshold (FAT) were used by setting the individual fractional anisotropy to a maximum value that enables reconstructing a minimum fiber course (ie, 100% FAT). For CSD-based tractography as well as MLFT, the FOD peak threshold was set to 0.08.27,28 The tractography step size was set to half the voxel size, and the number of iterations for MLFT was set to 2.

**Statistical Analysis.** Statistical testing was performed using SPSS (Version 26.0; SPSS Statistics for Windows; IBM) and the SciPy library (Version 1.3.1; https://www.scipy.org). The threshold for statistical significance was set at α = 0.05.

To assess and compare reconstructions of the CST, we computed the ratio of the visited voxels of the nTMS mask to the reconstructed fibers for each parameter configuration (ADT = 20°, 45°, and 60°; FAT = 25%, 50%, and 75%) and algorithm (DTI-based tractography, CSD-based tractography, and MLFT). This computation provides an estimate of how much of the nTMS mask (ie, cortical motor-positive nTMS points) is covered and, consequently, how complete the reconstruction is according to the function-based reference (nTMS mask). Additionally, volumes of the reconstructed bundles were computed as the accumulated volume of all the voxels visited by at least 1 fiber pathway. Only the bundle part between the seed and target region was taken into account. To compare the nTMS coverage as well as the volumes of the reconstructions as obtained by the different algorithms depending on the FAT as well as ADT, we performed paired 1-sided Wilcoxon tests. Furthermore, the nTMS coverage achieved by the same algorithm at different ADT settings was compared using paired 2-sided Wilcoxon tests.

**RESULTS**

Tractography using MLFT was capable of reconstructing fiber bundles with higher volumes (on average 10,367, 19,567, and 26,485 mm³ for ADTs of 20°, 45°, and 60°) than what was achieved by the DTI- and CSD-based algorithms, true for all used ADTs and FATs (Fig 1 and Online Supplemental Data). DTI-based tractography reconstructed statistically significant smaller bundles than CSD-based tractography at 25% (P = .54, .99, and 1 for 20°, 45°, and 60° ADT), 50% (P = .97, 1, and 1 for 20°, 45°, and 60° ADT) and 75% FAT (P = 1 for all ADTs) based on 1-sided Wilcoxon tests.

**RESULTS**

Compared with the other approaches, MLFT reconstructions of the CST achieved the highest coverage of the nTMS motor map, which amounted to 38.7%, 60.8%, and 71.8%, on average, for ADTs of 20°, 45°, and 60°, respectively (Fig 2 and Online Supplemental Data). MLFT also achieved higher coverage in
case of tumor-induced bundle deformations (Fig 1). CSD-based tractography achieved higher coverage of the nTMS motor map than DTI-based tractography, regardless of the settings for ADT and FAT (Fig 2). This finding was also confirmed by comparing reconstructions obtained with the same ADTs with 1-sided Wilcoxon tests ($P .95$ for all comparisons, Online Supplemental Data).

The change of the ADT led to statistically significant changes in nTMS motor map coverage ($P .01$) based on 2-sided Wilcoxon tests, which can also be observed in an exemplary patient case (Fig 3). Only when using 75% FAT for DTI-based reconstruction was a statistically nonsignificant difference observed between reconstructions obtained with 20° and 60° ADT as well as 45° and 60° ADT.

**DISCUSSION**

We combined MLFT with motor mapping for CST reconstruction based on nTMS motor maps used for ROI placement in patients with motor-eloquent gliomas. Tractography results were compared against deterministic DTI-based and CSD-based tractography, given that these techniques are commonly used for clinical tractography in patients with brain tumors. The main findings are as follows: 1) MLFT enabled CST reconstruction with higher bundle volumes, and 2) MLFT yielded higher coverage of the nTMS motor map (ie, a higher percentage of the nTMS points reached by the reconstructed CST).

The MLFT algorithm we used was developed from CSD-based tractography, and it similarly reconstructs fiber pathways on the basis of FOD peaks. However, in contrast to CSD-based tractography, MLFT suggests that FOD peaks may also be reflective of high angular deviation of fibers or their branching and may not only stem from fiber crossings.27,28 Hence, reconstruction of the CST with MLFT may produce bundles with higher radial extent; thus, the delineation of CST fanning with a wider range by also reconstructing fiber courses with acute angles becomes possible.27,28 Zhylka et al27 have shown this feature in a previous study among patients with motor-eloquent high-grade gliomas, but the motor cortex mask was defined using a parcellation mask of the precentral, postcentral, and paracentral gyri combined with a cross-section of the brainstem at the pontine level.

Given that MLFT may naturally reconstruct bundles with higher fiber count due to inclusion of fibers that course with acute angles, the potential risk of increasing the false-positive rate (ie, proportion of fibers that are visualized but do not connect to the actual functional motor cortex) is present. Specifically, to avoid a high false-positive rate, MLFT would require well-defined ROI seeding, and if certain pathways do not connect to the ROI, the algorithm checks to see if any deviation at the previous reconstruction points would allow connection to the ROI, thus providing some control over specificity while potentially improving sensitivity.27,28 In this regard, motor maps from preoperative nTMS were used as the target ROIs, given that nTMS motor mapping has shown high agreement with intraoperative direct electrical stimulation as the reference standard for functional mapping in patients with brain tumor.33-35 Furthermore, nTMS has also been effectively combined with DTI-based tractography in the past, allowing DTI-based reconstruction of the CST using functional data for ROI seeding, which could improve tracking of fibers for preparation and guidance of tumor resection and stratification for perioperative functional deficits.36-39 In this context, compared with conventional seeding without functional data (eg, manual delineation of the brainstem for ROI generation), nTMS-based tractography of the CST has been shown to result in a lower number of aberrant tracts (ie, tracts not belonging to the CST), and it changed the surgical strategy in more than twice as many patients.38,50 Furthermore, detailed somatotopic CST reconstructions might become possible when using nTMS motor maps as ROIs, with greater spatial overlap between the motor cortex and the cortical end region of the CST compared with conventional anatomic seeding for tractography.40

Reconstructions of the CST using MLFT showed the highest bundle volume for the CST, combined with the highest coverage of the nTMS motor map compared with deterministic DTI- or
CSD-based tractography. This result may indicate that a more complete reconstruction of the CST can be achieved with MLFT, which is most likely due to reconstruction of fibers with higher radial extents. Glioma may cause considerable deviation of CST fibers, which may, in part, be lost to reconstruction when using DTI- or CSD-based approaches (Fig 1). The higher coverage of nTMS maps when applying MLFT could increase confidence because the fibers reconstructed from MLFT are actually...
representative of the motor system (Fig 2), given that higher nTMS coverage is suggestive of more fibers being connected to the actual functional motor cortex. Of note, preoperatively enclosing the functional motor cortex with nTMS allows definition of its individual extent and location, which can be drastically aberrant to the structural landmark anatomy due to shifts related to the space-occupying effects and, most notably, due to plastic reshaping of functional motor representations in response to the presence and growth of glioma.51-53

A more complete reconstruction of fibers belonging to the CST is of clinical merit when achieving an optimal onco-functional balance because their visualization could help avoid surgery-induced damage and, thus, occurrence of lasting perioperative paresis. Specifically, CST reconstruction with broader fanning by MLFT could also generate laterally coursing and marginal fibers, which could be at particular risk of damage when approaching a maximized extent of resection. The higher CST bundle volumes may likely also result in pathways with a smaller lesion-to-tract distance, having potential impact on resection planning. Previous work using nTMS motor mapping combined with DTI-based tractography of the CST has proposed the lesion-to-tract distance as a parameter for presurgical stratification of the risk of permanent motor function decline, with lower lesion-to-tract distances being associated with a higher risk for lasting deficits.36,37,39 Application of MLFT instead of DTI-based tracking might potentially allow refined results for lesion-to-tract distances, with a potential definition of more realistic lesion-to-tract distances that may facilitate improved surgical outcome in terms of the extent of resection and the patient’s functional status.

A major limitation of this study is that tractography results were not confirmed by intraoperative direct electrical stimulation as the reference standard for functional mapping. Due to the retrospective character of this study, this confirmation was not possible but may be achieved in future studies. Hence, potential overrepresentations of fibers when using the MLFT algorithm cannot be fully excluded. Another limitation of MLFT relates to FOD accuracy, given that a high number of diffusion directions and high b-values are not routinely acquired for clinically used diffusion-weighted MR imaging sequences. As a consequence, the accuracy of the fitted diffusion models could be restricted because FODs have to be represented by lower-order spherical harmonics. Furthermore, the FOD algorithm used does not estimate separate response functions for different tissues.49 However, using acquisitions with multiple diffusion-weights (eg, multi-shell imaging) over the sequence we used would facilitate applying FOD algorithms that can differentiate multiple tissues.54

**CONCLUSIONS**

Compared with routinely used deterministic DTI-based and CSD-based tractography of the CST, MLFT may enable CST reconstructions with a higher bundle volume paired with higher coverage of the functional motor cortex. Thus, MLFT could provide a more detailed visualization of CST architecture by also visualizing fiber courses with acute angles, which might be of particular relevance in patients with gliomas and distorted anatomy of the motor system. However, prospective confirmation of tractography results from MLFT by intraoperative direct electrical stimulation as the reference standard for functional mapping is required for validation purposes.

Disclosure forms provided by the authors are available with the full text and PDF of this article at www.ajnr.org.

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