Multi-Modal Glioblastoma Segmentation: Man versus Machine

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Abstract

Background and Purpose: Reproducible segmentation of brain tumors on magnetic resonance images is an important clinical need. This study was designed to evaluate the reliability of a novel fully automated segmentation tool for brain tumor image analysis in comparison to manually defined tumor segmentations.

Methods: We prospectively evaluated preoperative MR Images from 25 glioblastoma patients. Two independent expert raters performed manual segmentations. Automatic segmentations were performed using the Brain Tumor Image Analysis software (BraTumIA). In order to study the different tumor compartments, the complete tumor volume TV (enhancing part plus non-enhancing part plus necrotic core of the tumor), the TV+ (TV plus edema) and the contrast enhancing tumor volume CETV were identified. We quantified the overlap between manual and automated segmentation by calculation of diameter measurements as well as the Dice coefficients, the positive predictive values, sensitivity, relative volume error and absolute volume error.

Results: Comparison of automated versus manual extraction of 2-dimensional diameter measurements showed no significant difference (p = 0.29). Comparison of automated versus manual segmentation of volumetric segmentations showed significant differences for TV+ and TV (p<0.05) but no significant differences for CETV (p>0.05) with regard to the Dice overlap coefficients. Spearman’s rank correlation coefficients (ρ) of TV+, TV and CETV showed highly significant correlations between automatic and manual segmentations. Tumor localization did not influence the accuracy of segmentation.

Conclusions: In summary, we demonstrated that BraTumIA supports radiologists and clinicians by providing accurate measures of cross-sectional diameter-based tumor extensions. The automated volume measurements were comparable to manual tumor delineation for CETV tumor volumes, and outperformed inter-rater variability for overlap and sensitivity.


Introduction

Gliomas are the most frequent primary brain tumors in adults, accounting for 70% of adult primary cerebral malignancies. Glioblastoma (GBM), the most common malignant primary brain tumor in humans, exhibits very rapid infiltrative growth and a poor prognosis, with an average survival time after GBM diagnosis of one year [1]. In recent years, the overall survival of glioblastoma patients has increased due to more extensive treatment strategies such as concomitant radio- and chemotherapy [2] and advanced surgical techniques [3] like fluorescence-guided surgery with 5-aminolevulinic acid (5-Ala) or advanced mapping methods [4,5].

Prognostic biomarkers for improved survival include imaging of the extent of resection based on the amount of postoperatively enhancing tissue as determined by structural MRI. Depending on the study populations, gross resection of 78% to 98% of the enhancing tumor volume is associated with an improved survival in patients with previously untreated or recurrent GBM [6,7]. Preoperative imaging characteristics associated with better survival are: i) absence of necrosis, ii) amount of contrast enhancing tumor, and iii) extent of solid tumor exceeding the enhancement [8,9]. Whole tumor and sub-compartment segmentation is performed manually in most centers and is still considered the “gold standard” procedure. Several studies reported failures of individual expert rater segmentations because structural MRI may
obscure the precise visual delineation of glioma boundaries [10–12]. In addition, manual segmentation is time-consuming and is a rate-determining step for further treatment planning [13]. Fully automated user-independent segmentation tools are available for research purposes. However, up to now they were considered limited with regard to ease of use, accuracy and speed, and therefore not adequate for clinical applications [14].

There is a large body of literature available on automatic brain tumor segmentation [14], but only recently the focus is shifting towards segmentation of individual tumor sub-compartments from multi-modal images. Most of the relevant fully automatic brain tumor segmentation methods employ a supervised or unsupervised tissue classification, that assigns labels based on voxel-wise or regionally extracted features. Corso et al. [15] segmented tumor core and edema using Bayesian tissue classification fused with affinity assignment by weighted aggregation. Verma et al. [16] used a support vector machine classifier to segment necrotic, active and edema tumor tissues. Bauer et al. [17] followed a similar approach, but added regularization constraints to increase robustness. Menze et al. [18] combined an atlas of normal individuals with a latent tumor atlas to employ a generative model for segmenting tumor compartments in different modalities. Zikic et al. [19] used context-aware voxel features as input for a decision forest classifier to segment 3 different tumor compartments from multi-modal images. A comparison of the performance of many recent methods on a standardized research dataset can be found in [20].

Almost all evaluations so far focus on the assessment of overlap measures like Dice coefficient, which are well established for comparing segmentation algorithms. However, other clinically more relevant measures like the diameter-based RANO metrics [21], volume error or the tendency to over- or underestimate are hardly ever reported. Moreover, there are only very few methods, for which the source code is publicly available (e.g. [22,23]), and to the best of our knowledge there exists not a single publicly available fully automatic segmentation tool with a graphical user interface that can be used by non-specialists and that can be applied to clinical images directly.

This study was designed to compare a novel clinically-oriented fully automated segmentation tool for brain tumor image analysis with manually defined tumor segmentations by two independent raters. Specifically, we aimed to evaluate: i) whether the fully automated tool can reproduce current 2D diameter-based criteria for brain tumor assessment [21]; ii) whether volumetric criteria can be reliably estimated by fully automated segmentation, including susceptibility to failure within dedicated sub compartiments. Furthermore, we also report iii) whether tumor localization has an effect on the quality of automated segmentations; iv) how improved image acquisition protocols impact the segmentation result.

### Methods

#### Study population

Patients with newly diagnosed and histologically confirmed glioblastoma pre-operatively admitted to our institution between October 2012 and July 2013 were eligible for this prospective study. Exclusion criteria were: incomplete image acquisition, Karnofsky performance status <70%, abnormal hematologic, renal or hepatic function, and previous cranial neurosurgery. The study was approved by the Local Research Ethics Commission (Kantonale Ethikkommission Bern). All patients provided written informed consent.

MR Imaging Protocol

MR images were acquired on two different 1.5 TMR scanners (Siemens Avanto and Siemens Aera, Siemens, Erlangen/Germany). Every patient underwent a standardized MRI protocol including: i) 3D-T1w-MPR in sagittal acquisition, 1 mm isotropic resolution; ii) post-contrast 3D-T1w-MPR in sagittal acquisition, 1 mm isotropic resolution; iii) 3-D T2w (SPC) in sagittal acquisition, 1 mm isotropic resolution; iv) fluid-attenuated inversion recovery (FLAIR) (TIR 2D) in axial acquisition. The sequence parameters were as follows: for pre-contrast 3D-T1w-MPR sequences TE = 2.67 ms, TR = 1580 ms, FOV = 256 × 256 mm², FA = 8°, with an isotropic voxel resolution of 1 mm × 1 mm × 1 mm; for post-contrast T1w TE = 4.37 ms, TR = 2070 ms, FOV = 256 × 256 mm², FA = 15°, using isotropic 1 mm × 1 mm × 1 mm voxels; for 3D T2w (SPC) in sagittal acquisition TE = 300 ms, TR = 3000 ms, FOV = 256 × 256 mm², FA = 120°, using isotropic 1 mm × 1 mm × 1 mm voxels; for the 2D fluid-attenuated inversion recovery sequence TE = 80 ms, TR = 9000 ms, FOV = 256 × 256 mm², FA = 120°, using a non-isotropic voxel size 1 mm × 1 mm × 3 mm.

Re-evaluation of pooled datasets from the MICCAI BraTS challenge

In order to compare the impact of the modified imaging protocols on the segmentation quality, we additionally analyzed a dataset composed of MR images from the MICCAI 2012 Brain Tumor Segmentation (BraTS) challenge consisting of in-silico and research datasets of brain tumor patients [20]. This dataset is different from the dataset in our clinical study described above. The BraTS images included the same MRI sequences with lower resolutions and various parameter settings. Quality of the automated segmentation was tested compared to the manually generated ground truth using the online evaluation tool (http://www2.imm.dtu.dk/projects/BRATS2012).

Two-dimensional measurements

In order to compare the accuracy of automated versus manual segmentation, we computed the product of two maximum diameters for 2-dimensional measurements, as recommended for reporting of gross tumor volume by the WHO and the refined RANO guidelines [21,24]. The automated 2-dimensional measurements were extracted from the largest perpendicular diameters of the contrast-enhancing lesions of the automatically segmented tumor expansion. These were compared to the measurements of our two expert raters using the product of maximal cross-sectional enhancing diameters. In case of multiple lesions, the sum of products of diameters (SPD) of all lesions was calculated [25].

Manual volumetric segmentation

The manual segmentations were performed by two independent expert raters: an expert in brain tumor imaging (SB) and a neurosurgeon experienced in brain tumor analysis (NP). Both raters were supervised independently by two neuroradiologists with more than 10 years of experience in brain tumor imaging (RW and RKV). Manual segmentation was performed with the open source software 3D Slicer Version 4.2.2.3 (www.slicer.org) [26]. Every patient (n = 25) was segmented manually slice by slice. Segmentation was performed on T1w, T1wGd, T2w and FLAIR sequences according to the Vasari MR feature guide v.1.1 (https://wiki.nci.nih.gov/display/CIP/VASARI). Four different tumor compartments were classified: i) non-enhancing tumor, ii) enhancing tumor, iii) necrosis and iv) edema. Hence, non-enhancing tumor was defined on FLAIR, T2w and T1w.
Enhancing tumor parts were classified on post-contrast T1 weighted images compared to pre-contrast T1-weighted images excluding hemorrhage. Necrosis was defined as a region within the tumor that did not enhance and had a hyper-intense signal on T2w and FLAIR. Edema was classified on FLAIR, T1w, T2w and contrast-enhanced T1w [27]. The average time for manually segmenting all subcomponents of one study patient was approximately one hour.

Automated Segmentation

Automatic segmentations were performed using the Brain Tumor Image Analysis (BraTumIA) software (available under this link http://www.istb.unibe.ch/content/research/medical_image_analysis/software/index_eng.html). The software offers a completely integrated segmentation pipeline, where the user only has to load the original Dicom stacks of the four relevant MRI modalities (T1w, contrast-enhanced T1w, T2w, FLAIR). Then, the images are processed in a fully automatic way, including skull stripping [28] and subsequent rigid co-registration [29] to ensure voxel-to-voxel correspondence between the different MRI sequences. Based on the registered images, segmentation into unaffected tissue and tumor tissue, encompassing four different sub-compartments, is performed based on combined supervised classification and regularization. The algorithmic core of the segmentation evolved out of [17] and the basics of the current approach have been recently described in a conference paper [30]. Briefly, the segmentation problem is formulated as an energy minimization task in a conditional random-field context. From each voxel, a high-dimensional feature vector is extracted, consisting of multi-modal intensities, texture and gradient statistics from local image patches, multi-scale symmetry features across the mid-sagittal plane and location features. Based on this high-dimensional feature vector, each voxel is assigned a tissue label by a decision forest classifier [31]. Spatial constraints and prior knowledge are considered by a conditional random field regularization to increase robustness. More details are given in Data S1 and in [30].
Figure 3. Box and whisker plots (min-max) of TV+, TV and CETV between manual raters (inter-rater (IR)), automatic segmentation versus expert rater 1 (AE 1); and automatic segmentation versus expert rater 2 (AE 2) for A: Dice coefficients B: PPV = Positive predictive values C: Sensitivity for the sub-compartment segmentations.
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Figure 4. Scatter plot of the absolute volume measurement for TV+, TV and CETV (from left to right). Volumes measured automatically are shown on the x-axis, volumes measured manually by the two expert raters are shown on the y-axis.
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The segmentation method requires training on manually segmented images of brain tumor patients. Training was performed on 36 separate patients that were not part of the study. Computation time for the complete automatic processing was less than 5 minutes. The algorithmic core of the software was among the best performing methods at the MICCAI Brain Tumor Segmentation (BRATS) challenge (http://www2.imm.dtu.dk/projects/BRATS2012/). A screenshot of the user interface of the integrated software is shown in figure 1.

Statistical methods
We evaluated the results primarily using the Dice coefficient [32]. The Dice coefficient measures the overlap of two regions (i.e. region 1 given by a manual segmentation, region 2 given by the automatic segmentation). It can range from 0 to 1, with 0 indicating no overlap and 1 indicating complete overlap. Additionally, we also report the positive predictive value (PPV) [33], sensitivity [34] and volume error between manual and automatic segmentations. The Wilcoxon signed rank test [35] was used for the statistical analysis of the difference between manual and automated volumetric segmentation in terms of the metrics mentioned above. For inter-observer comparisons and automatic versus manual comparisons, we also used Spearman's rank correlation coefficients [36]. For the comparison of 2-dimensional measures and investigation of the impact of localization, non-parametric analysis of variance [37,38] was performed. Statistical analysis was done using Graphpad Prism version 5.

Results
Study population
The mean patient age at pre-operative MR imaging (+/− standard deviation (SD)) was 67.75 years (+/−6.191 (range 53–79 years), mean pre-operative Karnofsky performance status (+/−SD) was 84.38% (+/−6.292 (range 70–90%), and mean pre-operative NIHSS (+/−SD) was 0.25+/−1.12 (range 0–3). Of the 25 patients, 10 were female and 15 male. Five patients underwent stereotactic biopsy, 11 subtotal extirpations and nine complete resections of enhancing tumor (CRET). All diagnoses were confirmed by histopathology.

Comparison of automated versus manual segmentation
Two-Dimensional measurements. Comparison between automated and manual 2-dimensional measurements by two expert raters was done using the product of maximal cross-sectional enhancing diameters (SPD in mm²+/−SD). The mean absolute difference in SPD of automatic diameter versus expert rater 1 was 570 mm² (+/−569), (95% CI [335, 805]); the mean absolute difference in SPD of automatic diameter versus expert rater 2 was 495 mm² (+/−438), (95% CI [314, 676]). The mean absolute difference in SPD of expert rater 1 versus expert rater (+/−SD) was 355 mm² (+/−380), (95% CI [198, 511]). There was no significant difference between the three different measurements (Friedman test; p = 0.29). The results are depicted in figure 2.

Volumetric Segmentation. To evaluate the different tumor compartments, we separately investigated the complete tumor volume (TV) encompassing the enhancing part of the tumor, the non-enhancing part of the tumor plus the necrotic core, the TV (+/−SD) was 84.38% (+/−6.292 (range 70–90%), and mean pre-operative NIHSS (+/−SD) was 0.25+/−1.12 (range 0–3). Of the 25 patients, 10 were female and 15 male. Five patients underwent stereotactic biopsy, 11 subtotal extirpations and nine complete resections of enhancing tumor (CRET). All diagnoses were confirmed by histopathology.

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Table 1. Statistics of Dice coefficients from the MICCAI BraTS 2012 testing dataset compared to the Dice coefficients on our clinical study dataset (CS).

<table>
<thead>
<tr>
<th>Dice</th>
<th>TV+</th>
<th>TV</th>
<th>CETV</th>
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<tbody>
<tr>
<td>BraTS median</td>
<td>0.69</td>
<td>0.55</td>
<td>0.58</td>
</tr>
<tr>
<td>BraTS mean</td>
<td>0.71</td>
<td>0.55</td>
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<tr>
<td>BraTS SD</td>
<td>0.15</td>
<td>0.19</td>
<td>0.24</td>
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<tr>
<td>CS median</td>
<td>0.84</td>
<td>0.70</td>
<td>0.66</td>
</tr>
<tr>
<td>CS mean</td>
<td>0.80</td>
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<td>0.63</td>
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<tr>
<td>CS SD</td>
<td>0.12</td>
<td>0.14</td>
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<tr>
<td>p-value</td>
<td>0.06</td>
<td>0.06</td>
<td>0.01</td>
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Table 2. Spearman’s rank correlation coefficients (p) of TV+, TV and CETV.

<table>
<thead>
<tr>
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<th>AE1 versus AE2</th>
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<tr>
<td>TV+</td>
<td>0.85 0.00</td>
<td>0.74 0.00</td>
<td>0.79 0.00</td>
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<tr>
<td>TV</td>
<td>0.90 0.00</td>
<td>0.68 0.00</td>
<td>0.75 0.00</td>
</tr>
<tr>
<td>CETV</td>
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<td>0.84 0.00</td>
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doi:10.1371/journal.pone.0096873.t002
coefficients for the automatic segmentation were 0.8 for TV+, 0.66 for TV and 0.63 for CETV (compared to expert 1 who defined the ground truth, see table 1). The average absolute volume error of the automatic segmentation was 20.4 ml for TV+, 14.5 ml for TV and 7.2 ml for CETV (compared to expert 1 who defined the ground truth). The relationship between automatically and manually calculated volumes of individual tumor sub-compartments is shown with scatterplots in figure 4. Details can be found in Data S1.

Using the Wilcoxon signed ranks test, we observed significant differences for TV+ and TV (p < 0.05) but no significant differences for CETV (p > 0.05) with regard to the Dice coefficients. Spearman’s rank correlation coefficients (p) of the Dice coefficient for TV+, TV and CETV showed highly significant correlations (p-value) between automatic and manual segmentations (see Table 2).

Figure 5 illustrates results for a patient where manual and automatic segmentation match well, whereas figure 6 depicts a patient where only moderate agreement between manual and automatic segmentation could be achieved, mostly due to a mismatch of the CETV compartment.

**Tumor localization and impact on accuracy of segmentation.** Of the 25 analyzed data sets, 8 tumors were in the frontal lobes, 8 temporal, 8 parietal and one occipital (the latter was excluded from group analyses). In order to identify a potential impact of the tumor localization in different lobes of the brain on the accuracy of automated segmentation, we analyzed the influence of the tumor localization (frontal, temporal, parietal) on the Dice overlaps. A Kruskal-Wallis test revealed no differences related to the localization or the sub-compartments (see Table 3).

**Segmentation results on non-standardized imaging protocols.** Comparison of the Dice coefficients for automatic segmentation between the non-standardized MICCAI BraTS 2012 test data and the standardized high resolution datasets in our study patients revealed improved Dice coefficients for the standardized protocol in all sub-compartments (see Table 1). However the improvement was not statistically significant in all compartments (p = 0.06) due to the small number of patients in the BraTS dataset (11 patients).

**Discussion**

In this prospective study we validated an automatic multimodal segmentation software (BraTumIA) in clinical practice. We delineated different tumor compartments, i.e. TV, TV+ and CETV in de novo GBM. Our goal was to evaluate how BraTumIA performed in our clinical setting in comparison to manual ratings. We focused on two metrics for clinical evaluation: the SPD according to the WHO classification [24,39], and the volumes and overlaps of TV, TV+ and CETV. It was shown that automatic volumetry offers significant time gains compared to manual volumetry.

Two points stand out among the study results: i) computation of cross sectional SPD using automated 2-dimensional tumor extensions is comparable to manual tumor delineation; ii) estimation of TV, TV+ and CETV by the automatic method reaches a sensitivity, which is comparable to the inter-observer
variability between two experts performing manual segmentation. For CETV, BraTumIA had a better overlap with the ground-truth than Dice overlap between manual inter-rater segmentations.

Furthermore, we observed that classification of automated segmentation is insensitive to tumor localization and individual compositions of the sub-segments of GBM. Comparison of the quality of the automatic segmentations on our new prospective patient dataset of standardized images with the quality obtained on the non-standardized BraTS 2012 images, showed an average improvement of more than 10% (table 1). This indicates that a standardized high-resolution isotropic imaging protocol such as the one used in this study has the potential to optimize the quality of automatic segmentations.

A previous study demonstrated that an extent of resection (EOR) of the TV ≥ 78% impacts patient outcome, and that the positive relationship between EOR and patient outcome can be observed even at the highest levels of resection [8]. For the pre-operative baseline assessment, we observed good concordance of linear SPD assessment between manual raters and the automated approach, and a higher sensitivity for automated volumetry of the CETV compared to two manual raters (72% vs. 53%, figure 3). Although the linear SPD measures provided a good concordance between automatic and manual measurements, these measures are problematic in cases of non-solid GBMs [40]. GBMs usually appear with a complex morphology, thus necessitating volumetric analyses. However, diameter-based SPD measures only consider one tumor compartment (necrosis+enhancing tumor), usually in low resolution datasets.

Most segmentation techniques focus on reporting of Dice similarity coefficients of the gross tumor volume instead of individual tumor sub-compartments. A semi-automatic approach using Slicer recently reported an average Dice similarity coefficient of 0.80 [41]. Semi-automatic segmentations tend to produce better accuracies compared to fully automatic methods but previous studies suggest that semi-automatic methods are less objective [42]. While previous approaches aimed to assess only one tumor

Table 3. Regional Differences in Dice overlap according to localization.

<table>
<thead>
<tr>
<th>Dice</th>
<th>Frontal</th>
<th>Temporal</th>
<th>Parietal</th>
<th>p-Values</th>
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<tbody>
<tr>
<td>TV+ mean</td>
<td>0.84</td>
<td>0.81</td>
<td>0.85</td>
<td>0.62</td>
</tr>
<tr>
<td>TV mean</td>
<td>0.65</td>
<td>0.64</td>
<td>0.74</td>
<td>0.13</td>
</tr>
<tr>
<td>CETV mean</td>
<td>0.57</td>
<td>0.65</td>
<td>0.68</td>
<td>0.2</td>
</tr>
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</table>

doi:10.1371/journal.pone.0096873.t003
In summary, we demonstrated that BraTumIA supports radiologists and clinicians with accurate measures of cross sectional 2-dimensional tumor extensions that are equivalent to manual tumor delineation, and provides CETV tumor volumes that have lower variability than manual ratings.

Supporting Information

Data S1 This file includes Method S1 and Tables S1 to S6. Method S1: Details of the segmentation algorithm. Table S1: Automated versus manual segmentation of expert 1. Table S2: Automated versus manual segmentation of expert 2. Table S3: Inter-observer agreement of manual segmentations of expert 1 and expert 2. Table S4: Statistical analysis (Wilcoxon signed rank test) of agreement between automatic and manual segmentations. Table S5: Spearman rank correlation coefficients of automatic and manual segmentations. Table S6: Kappa coefficient of mutual agreement between different automatic and manual segmentations in all tumor sub-compartments. (PDF)

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Author Contributions

Conceived and designed the experiments: SB JS MR RW. Performed the experiments: NP SB RKV RW. Analyzed the data: NP SB. Contributed reagents/materials/analysis tools: SB AP PS JB. Wrote the paper: NP SB MR RW.

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