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Analysis of physical characteristics of Tumor Treating Fields for human glioblastoma

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Abstract

Tumor Treating Fields (TTFields) therapy is an approved treatment that has known clinical efficacy against recurrent and newly diagnosed glioblastoma. However, the distribution of the electric fields and the corresponding pattern of energy deposition in the brain are poorly understood. To evaluate the physical parameters that may influence TTFields, postacquisition MP-RAGE, T1 and T2 MRI sequences from a responder with a right parietal glioblastoma were anatomically segmented and then solved using finite-element method to determine the distribution of the electric fields and rate of energy deposition at the gross tumor volume (GTV) and other intracranial structures. Electric field–volume histograms (EVH) and specific absorption rate–volume histograms (SARVH) were constructed to numerically evaluate the relative and/or absolute magnitude volumetric differences between models. The electric field parameters \( E_{\text{AUC}} \), \( V_{\text{E150}} \), \( E_{300\%} \), \( E_{500\%} \), and \( E_{1000\%} \) as well as the SAR parameters \( \text{SAR}_{\text{AUC}} \), \( V_{\text{SAR-150}} \), \( \text{SAR}_{300\%} \), \( \text{SAR}_{500\%} \), and \( \text{SAR}_{1000\%} \) facilitated comparisons between models derived from various conditions. Specifically, TTFields at the GTV were influenced by the dielectric characteristics of the adjacent tissues as well as the GTV itself, particularly the presence or absence of a necrotic core. The thickness of the cerebrospinal fluid on the convexity of the brain and the geometry of the tumor were also relevant factors. Finally, the position of the arrays also influenced the electric field distribution and rate of energy deposition in the GTV. Using EVH and SARVH, a personalized approach for TTFields treatment can be developed when various patient-related and tumor-related factors are incorporated into the planning procedure.

Keywords
Conductivity, disposition, glioblastoma, malignant gliomas, multiphysics, tumor geometry, Tumor Treating Fields

Introduction

Tumor treating fields (TTFields) are alternating electric fields, tuned to a frequency between 100 and 300 kHz, that have antimitotic properties against rapidly dividing cancer cells. When properly applied, these fields disrupt macromolecular protein structures thought to possess large dipole moments, such as Tubulin and Septin, which are critical for proper cytokinesis [1–3]. In past randomized clinical trials, TTFields was shown to have comparable efficacy against recurrent glioblastoma when compared to chemotherapies and, when added to maintenance temozolomide, it had a superior benefit in newly diagnosed glioblastoma patients when compared to maintenance temozolomide alone [4, 5]. Although the clinical efficacy against glioblastoma is apparent, the intracranial distribution of the electric fields and to what extent the dose of these fields retard tumor growth remain largely unknown.

TTFields are applied to the shaved scalp via two pairs of orthogonally positioned transducer arrays. Clinical placement of the arrays is determined by the proprietary NovoTAL™ software that generates an array layout diagram [6]. Each array has nine ceramic disks acting like disk sources for the electric fields. However, unlike high-energy ionizing radiation that can penetrate intracranial structure in a straight beam path, the intensity and directionality of electric fields are heavily influenced by the local dielectric properties of various structures in the head, particularly the electric conductivity and the relative permittivity of brain tissue [7]. As the electric conductivity and relative permittivity of tissues vary, the absorption...
and attenuation of TTFields will change and thus the distribution of these fields will be distorted as they permeate throughout the brain. In addition, there is currently no standardized electric conductivity and relative permittivity values for glioblastoma. As each individual tumor will vary in size, geometry, location within the brain, cellular composition, and presence or absence of necrosis, all of which may influence the electric field and energy absorption distribution within the glioblastoma. Therefore, to properly evaluate how tissue dielectric properties and physical characteristics of the various tissues influence the applied TTFields in the glioblastoma patient, a computer simulated model, currently best solved by the finite-element method, is needed. This model takes into account the normal brain structures, the gross tumor volume (GTV), the presence or absence of a necrotic core within the tumor, and local tissue electric conductivity and relative permittivity values. Here, we performed detailed finite-element modeling of a patient who responded to TTFields treatment. We found that there is heterogeneity in the electric field intensity and the rate of energy absorbed at the GTV depending on the placement of the transducer arrays, the presence or absence of a necrotic core within the glioblastoma, the thickness of the cerebrospinal fluid on the convexity of the brain and tumor geometry. The intensity of the electric fields and the specific absorption rate (SAR) were also represented graphically by the finite-element mesh. The introduction or elimination of certain intracranial structures, and the expansion or contraction of cerebrospinal fluid volume, only isotropic electric conductivity values were used in order to simplify the modeling process. Data postprocessing of the solved models was performed using Microsoft Excel, including generation of volume histograms EVH and SARVH. Area under the curve was computed using Simpson’s Rule integration in MATLAB 2016 for electric fields (E_AUC) and SAR (SAR_AUC). The EVH was used in the comparison of electric field strength between different models and was referenced to (1) the percentage volume of a particular structure receiving at least 150 V/m (V_E150), (2) the magnitude of electric field strength encompassing 95% of a particular structure’s volume (E_95%), (3) the magnitude of electric field strength encompassing 50% of a particular structure’s volume (E_50%), and (4) the magnitude of electric field strength encompassing 20% of a particular structure’s volume (E_20%). Similarly, the comparison of the rate of energy absorbed in different SARVH models was referenced to (1) the percentage volume of a particular structure receiving at least 7.5 W/kg (V_SAR5), (2) the magnitude of SAR encompassing 95% of a particular structure’s volume (SAR_95%), (3) the magnitude of SAR encompassing 50% of a particular structure’s volume (SAR_50%), and (4) the magnitude of SAR encompassing 20% of a particular structure’s volume (SAR_20%).

Materials and Methods

Critical neuroanatomical structures such as the scalp, skull, dura, cerebrospinal fluid, white matter, gray matter, brainstem, cerebellum, orbits, and bilateral ventricles were segmented based on the postacquisition MP-RAGE image dataset from a responder with a glioblastoma in the right parietal brain. The segmentation of these masks was performed using ScanIP (Simpleware LTD., UK) where grayscale thresholding methods were applied to initially segmented tissues, followed by manual correction on these masks. Additionally, T1 and T2 MRI sequences were also imported into ScanIP, and then coregistered with the segmented masks for delineation of the GTV and necrotic core; the remaining unsegmented tissue was labeled as unspecified tissue and given material properties that of muscle. Structures such as orbits, brainstem, and cerebellum were segmented and their associated physical properties were included mainly for spatial reference in the model. The GTV and necrotic core were both manually segmented by the treating physician based on the visible enhancement shown on the coregistered postgadolinium T1- and T2-weighted image datasets. Transducer arrays were manually placed on the surface of the scalp in the model, approximating as closely as possible to the standard display as shown in the United States Food and Drug Administration’s communication on the NovoTTF-100A system [8]. Upon completion of segmentation, a three-dimensional finite-element mesh was generated within ScanIP. This mesh was then imported into COMSOL Multiphysics (COMSOL, Burlington, MA), where material properties, boundary conditions, and appropriate physics parameters were assigned and applied.

For the purpose of demonstrating the sensitivity of varying physical properties, such as electric conductivity and relative permittivity, tumor geometry, array disposition, the introduction or elimination of certain intracranial structures, and the expansion or contraction of cerebrospinal fluid volume, only isotropic electric conductivity values were used in order to simplify the modeling process. Data postprocessing of the solved models was performed using Microsoft Excel, including generation of volume histograms EVH and SARVH. Area under the curve was computed using Simpson’s Rule integration in MATLAB 2016 for electric fields (E_AUC) and SAR (SAR_AUC). The EVH was used in the comparison of electric field strength between different models and was referenced to (1) the percentage volume of a particular structure receiving at least 150 V/m (V_E150), (2) the magnitude of electric field strength encompassing 95% of a particular structure’s volume (E_95%), (3) the magnitude of electric field strength encompassing 50% of a particular structure’s volume (E_50%), and (4) the magnitude of electric field strength encompassing 20% of a particular structure’s volume (E_20%).

Results

Electric field distribution of a responder treated with TTFields

A retrospective analysis of one glioblastoma patient, who responded to TTFields treatment, was performed by modeling the intracranial electric field distribution using finite-element analysis. The responder had a glioblastoma in the right parietal lobe extending toward the bilateral ventricles with roughly 1.5 cm between the GTV and the lateral border of the right lateral ventricle. As expected,
the highest electric field intensity was seen within the sulci on the surface of the brain and it is associated with a high-to-low gradient from the surface to the deeper regions (Fig. 1). In particular, the lowest intensity was seen in the inferior portion of the frontal (Fig. 1C and I) and the temporal (Figs. 1B and H) lobes. Furthermore, the body of the corpus callosum had high electric field intensity, particularly in the regions between (Fig. 1D) and above the lateral ventricles (Fig. 1E and F). This is likely due to the relatively higher electric conductivity of the cerebrospinal fluid located next to either side of the lesser conductive white matter, which probably created a higher capacitive reactance similar to a capacitor, helped to retain a higher electric field intensity within the white matter. Lastly, the medial portion of GTV in the right parietal lobe also possessed a higher electric field intensity. This could be a result of a relatively higher electric conductivity of the necrotic core within the GTV on one side and the cerebrospinal fluid within the right lateral ventricle on the other side, both of which also likely contributed to a higher reactive capacitance within the right parietal tissue.

To investigate the strength of electric field and the rate of energy deposited into the GTV and various intracranial structures, EVH and SARVH were generated for the comparison between models that use the primary position for transducer array placement as outlined in Figure 3E and incorporate the isotropic electric conductivity and relative permittivity values listed in Table 1 [9, 10]. As expected, the highest E_{AUC} was found at the scalp and skull, whereas the lowest was located at the orbits, bilateral ventricles, and brainstem (Fig. 2A and C). In the GTV, 95% of the volume had an electric field intensity of >50 V/m, whereas 50% had >80 V/m and 20% has >150 V/m (Fig. 2A). Because at least 20% of the GTV volume had a minimum of 150 V/m of electric field

Figure 1. Distribution of TTFields within a patient brain. The electric field intensity is higher in the supratentorial than the infratentorial brain. Within the supratentorial brain, various parts of the sulci, the body of the corpus callosum, and the medial surface of the GTV appear to have the highest electric field intensity. Axial images from inferior to superior slices are shown in A, D and G. Coronal images from anterior to posterior slices are shown in B, E and H. Sagittal images from right to left slices are shown in C, F and I. TTFields, Tumor Treating Fields; GTV gross tumor volume.
coverage (Table 2), the $V_{E150}$ volume and $E_{50\%}$ magnitude were used as means of comparing different modeling outcomes in subsequent analyses. Similarly, the highest SAR $AUC$ was found at the skull, GTV, and the layer of cerebrospinal fluid between cortex and dura, whereas the lowest were located in the orbits, cerebellum, and brainstem (Fig. 2B and D). Because at least 15% of the GTV volume had a SAR of at least 7.5 W/kg (Table 2), the $V_{SAR7.5}$ volume and SAR $50\%$ magnitude were used as means of comparing different modeling outcomes in subsequent analyses.

### Disposition analysis of transducer array placement

As the placement of transducer arrays can be shifted during each array exchange on treatment, we investigated whether or not the disposition of the arrays can alter electric field coverage and rate of energy deposition in the GTV. The effects of shifting array positions at different locations on the scalp were modeled according to an aggregate 2-cm deviation from the primary position; each of the nine disks in each array were manually drawn to scale and placed in full contact with the surface of the scalp in the configuration of the primary position as shown in Figure 3E. The lateral arrays were then shifted in clockwise or counterclockwise configurations, whereas the anterior–posterior arrays were shifted in the forward or backward positions, resulting in eight additional configurations (Fig. 3, except for Fig. 3E [primary position]). The posterior array in all cases was not moved inferiorly due to the fact that the patient’s image dataset was truncated at the occiput, and thus had insufficient occipital anatomy to shift the posterior array inferiorly. Still, there was high variance in electric field coverage of the GTV between 100 and 150 V/m, ranging from only 20% volume having 100 V/m with clockwise rotation of the lateral arrays and no displacement of the anterior–posterior arrays (Fig. 4A, red curve and Table 3) to >40% volume having at least 150 V/m when the lateral arrays were rotated in a counterclockwise fashion and the anterior array was moved backward (Fig. 4A, green curve). However, the variability in SAR, as represented by the magnitude of SAR $50\%$, was low and it ranged between 3 and 6 W/kg (Fig. 4B). For the necrotic core of the tumor, there was also a similar but smaller variance in the electric field coverage (Fig. 4C) and its corresponding SAR magnitude (Fig. 4D).

### Sensitivity analysis of the conductivity of the GTV and the necrotic core

The strength of the electric field penetrating the GTV most likely depends on its dielectric properties, specifically the electrical conductivity and relative permittivity. Our previous modeling found that the electric field intensity at the GTV is more sensitive to changes in its electric conductivity than its relative permittivity characteristics [7]. However, glioblastoma frequently has a necrotic component that contains liquefied cellular products or exudates from adjacent highly permeable vasculature. The fluid component of this necrotic core can potentially influence the electric field strength within the GTV. To investigate the relationship between the field coverage at both GTV and necrotic core, the electric conductivity of the GTV was altered from 100 to 0.001 S/m, whereas the electric conductivity of the necrotic core was kept constant. The mean electric field strength within the necrotic core rose 600% from 5 to

---

Table 1. Physical parameters required as inputs for computer modeling.

<table>
<thead>
<tr>
<th>Tissue structure</th>
<th>Volume (cc)</th>
<th>Electric conductivity $\sigma$ (S/m)</th>
<th>Relative permittivity $\varepsilon_r$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gross tumor volume (GTV)</td>
<td>5.813874</td>
<td>2.50E-01</td>
<td>1.00E+04</td>
</tr>
<tr>
<td>Necrotic core</td>
<td>2.421458</td>
<td>1.00E+02</td>
<td>1.00E+00</td>
</tr>
<tr>
<td>Scalp</td>
<td>524.5453</td>
<td>1.05E-03</td>
<td>1.10E+03</td>
</tr>
<tr>
<td>Skull</td>
<td>463.5451</td>
<td>2.11E-02</td>
<td>2.04E+02</td>
</tr>
<tr>
<td>Dura</td>
<td>216.8171</td>
<td>5.02E-01</td>
<td>2.90E+02</td>
</tr>
<tr>
<td>Cerebrospinal fluid</td>
<td>238.8805</td>
<td>2.00E+00</td>
<td>1.09E+02</td>
</tr>
<tr>
<td>White matter</td>
<td>593.1396</td>
<td>8.68E-02</td>
<td>1.29E+03</td>
</tr>
<tr>
<td>Gray matter</td>
<td>261.5665</td>
<td>1.41E-01</td>
<td>2.01E+03</td>
</tr>
<tr>
<td>Bilateral ventricle</td>
<td>51.38429</td>
<td>2.00E+00</td>
<td>1.09E+02</td>
</tr>
<tr>
<td>Brainstem</td>
<td>28.7721</td>
<td>1.61E-01</td>
<td>2.30E+03</td>
</tr>
<tr>
<td>Orbits</td>
<td>12.89734</td>
<td>1.50E+00</td>
<td>9.66E+01</td>
</tr>
<tr>
<td>Cerebellum</td>
<td>44.55224</td>
<td>1.61E-01</td>
<td>2.30E+03</td>
</tr>
<tr>
<td>Unspecified tissue/muscle</td>
<td>133.3064</td>
<td>3.84E-01</td>
<td>6.38E+03</td>
</tr>
<tr>
<td>Electrodes</td>
<td>N/A</td>
<td>1.00E-05</td>
<td>1.10E+04</td>
</tr>
<tr>
<td>Titanium wires</td>
<td>N/A</td>
<td>1.28E+06</td>
<td>5.00E+01</td>
</tr>
</tbody>
</table>

The volume, electric conductivity and relative permittivity values for GTV, necrotic core, scalp, skull, dura, cerebrospinal fluid, white matter, gray matter, bilateral ventricles, brainstem, orbits, cerebellum, unspecified tissue/muscle, electrodes, and titanium wires that were used in the analysis.
30 V/m when the electric conductivity of the GTV decreased from 100 to 1 S/m, but further increase was markedly attenuated when the electric conductivity was lowered from 1 to 0.001 S/m (Fig. 5A). Similarly, the mean SAR increased from 0 to 1.8 W/kg when the electric conductivity of the GTV decreased from 100 to 1 S/m, but further increase was negligible when the electric conductivity was lowered from 1 to 0.001 S/m (Fig. 5C). In contrast, when the electric conductivity of the necrotic core was varied from 100 to 1 S/m, there was negligible change in the mean electric field strength in the GTV; however, there was up to a 10% increase in mean electric field strength of the GTV when the electric conductivity of the necrotic core was lowered from 1 to 0.001 S/m (Fig. 5B). But the change in mean SAR of the GTV was insignificant when the electric conductivity of the necrotic core was varied from 100 to 0.001 S/m (Fig. 5D).

To further investigate the electric field coverage of GTV as influenced by the conductivity of the necrotic core, both EVH and SARVH were constructed as the GTV was

**Figure 2.** Volume histograms EVH and SARVH. The EVH (A), SARVH (B), electric field map (C), and SAR map (D) were generated using the transducer array placement as outlined in Figure 3E. The highest $E_{\text{max}}$ was found at the scalp and skull, whereas the lowest was detected at the orbits, bilateral ventricles, and brainstem. The highest $SAR_{\text{max}}$ was found at the skull, GTV, and the layer of cerebrospinal fluid between cortex and dura, whereas the lowest was found in the orbits, cerebellum, and the orbits. EVH, electric field–volume histogram; SARVH, specific absorption rate–volume histogram; SAR, specific absorption rate.
modeled with or without the necrotic core. When highly conductive fluid in the necrotic core was replaced with electric conductivity and relative permittivity of white matter, the $E_{\text{AUC}}$ of the GTV shifted from 20% to 30% (Figs 6A and B) and the corresponding $V_{\text{SAR7.5}}$ increased progressively from +0.5 mm to −1.0 mm on the convexities of the brain. Specifically, $E_{\text{AUC}}$ increased from 94.0 V/m at +0.5 mm to 115.1 V/m at −0.5 mm, and to 122.1 V/m at −1.0 mm (Fig. 7C). Similarly, the rate of energy deposited in the GTV increased progressively, as represented by $SAR_{\text{AUC}}$, $V_{\text{SAR7.5}}$, and $SAR_{20\%}$ when the cerebrospinal fluid space was narrowed from +0.5 to −1.0 mm on the convexities of the brain. In particular, $SAR_{\text{AUC}}$ increased from 5.1 W/kg at +0.5 mm, to 6.8 W/kg at −0.5 mm, and to 7.4 W/kg at −1.0 mm (Fig. 7C). Therefore, increased cerebrospinal fluid space at the convexity shunts electric field and energy away from the brain, whereas decreased cerebrospinal fluid space allows a higher intensity of electric field and SAR to penetrate the brain.

The influence of cerebrospinal fluid on GTV

As cerebrospinal fluid is a highly conductive medium compared with ordinary brain tissue and it is in close proximity to the GTV, especially in our patient, we sought to investigate whether the cerebrospinal fluid along the surface of the brain and inside the ventricles could influence the electric field and SAR coverage of the GTV. First, 0.5 mm of cerebrospinal fluid was added on the surface or 0.5 mm to 1.0 mm of cerebrospinal fluid was subtracted from the surface of the original brain, corresponding to a respective volume of 275 cc, 196 cc and 179 cc compared to the initial volume of 240 cc. Indeed, $E_{\text{AUC}}$, $V_{\text{E150}}$ and $E_{\text{95\%}}$ of the GTV all increased progressively when the cerebrospinal fluid space was narrowed progressively from +0.5 mm to −1.0 mm on the surface of the original brain, or 0.5 mm to 1.0 mm of cerebrospinal fluid was subtracted from the surface of the original brain, corresponding to a respective volume of 275 cc, 196 cc and 179 cc compared to the initial volume of 240 cc. Indeed, $E_{\text{AUC}}$, $V_{\text{E150}}$ and $E_{\text{95\%}}$ of the GTV all increased progressively when the cerebrospinal fluid space was narrowed progressively from +0.5 mm to −1.0 mm on the convexities of the brain. Specifically, $E_{\text{AUC}}$ increased from 94.0 V/m at +0.5 mm to 115.1 V/m at −0.5 mm, and to 122.1 V/m at −1.0 mm (Fig. 7C). Similarly, the rate of energy deposited in the GTV increased progressively, as represented by $SAR_{\text{AUC}}$, $V_{\text{SAR7.5}}$, and $SAR_{20\%}$ when the cerebrospinal fluid space was narrowed from +0.5 to −1.0 mm on the convexities of the brain. In particular, $SAR_{\text{AUC}}$ increased from 5.1 W/kg at +0.5 mm, to 6.8 W/kg at −0.5 mm, and to 7.4 W/kg at −1.0 mm (Fig. 7C). Therefore, increased cerebrospinal fluid space at the convexity shunts electric field and energy away from the brain, whereas decreased cerebrospinal fluid space allows a higher intensity of electric field and SAR to penetrate the brain.

The influence of the material properties of tumor, necrotic core, and bilateral ventricles on EVH and SARVH

By replacing the material properties of the segmented volume representing the original tumor (GTV + necrotic core) with that of white matter, the electric field strength in terms of $E_{\text{AUC}}$, $V_{\text{E150}}$, $E_{\text{95\%}}$, $E_{\text{50\%}}$, and $E_{\text{20\%}}$ were all remarkably higher compared with the model containing the original tumor (Fig. 7C); proportionally the SAR was also increased dramatically. The $E_{\text{AUC}}$ for the original tumor was 104 V/m in comparison to 153.9 V/m after the change, which is an increase of about 48%. The $SAR_{\text{AUC}}$ also increased by 120%, from 5.9 to 13.0 W/kg. Similarly, but not as dramatically, replacing the material properties of the segmented necrotic core with that of white matter increased the electric field strength and SAR within the GTV. Specifically, the $E_{\text{AUC}}$ of the replaced vestibulocerebral area.

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**Table 2.** The values for electric field and rate of energy deposition parameters at the GTV and other structures in the head as shown in EVH and SARVH from Figure 2.

<table>
<thead>
<tr>
<th>Tissue structure</th>
<th>$E_{\text{AUC}}$</th>
<th>$V_{\text{E150}}$ (%)</th>
<th>$E_{\text{95%}}$</th>
<th>$E_{\text{50%}}$</th>
<th>$E_{\text{20%}}$</th>
<th>$SAR_{\text{AUC}}$</th>
<th>$V_{\text{SAR7.5}}$ (%)</th>
<th>$SAR_{95%}$</th>
<th>$SAR_{50%}$</th>
<th>$SAR_{20%}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bilateral ventricle</td>
<td>14.7</td>
<td>0.0</td>
<td>5.7</td>
<td>9.8</td>
<td>25.2</td>
<td>5.0</td>
<td>5.1</td>
<td>2.6</td>
<td>2.6</td>
<td>5.3</td>
</tr>
<tr>
<td>Brainstem</td>
<td>35.7</td>
<td>0.0</td>
<td>23.6</td>
<td>36.3</td>
<td>44.0</td>
<td>1.2</td>
<td>0.0</td>
<td>1.0</td>
<td>2.1</td>
<td>2.6</td>
</tr>
<tr>
<td>Cerebellum</td>
<td>50.4</td>
<td>0.1</td>
<td>32.0</td>
<td>49.0</td>
<td>62.6</td>
<td>1.0</td>
<td>0.0</td>
<td>0.8</td>
<td>2.5</td>
<td>2.5</td>
</tr>
<tr>
<td>Cerebrospinal fluid</td>
<td>67.3</td>
<td>0.1</td>
<td>21.0</td>
<td>53.5</td>
<td>90.2</td>
<td>1.9</td>
<td>0.0</td>
<td>58.6</td>
<td>2.2</td>
<td>9.2</td>
</tr>
<tr>
<td>Dura</td>
<td>108.3</td>
<td>0.1</td>
<td>25.4</td>
<td>86.2</td>
<td>169.8</td>
<td>8.2</td>
<td>32.3</td>
<td>4.9</td>
<td>17.1</td>
<td>20.3</td>
</tr>
<tr>
<td>Gray matter</td>
<td>108.1</td>
<td>0.0</td>
<td>37.7</td>
<td>87.2</td>
<td>164.7</td>
<td>2.5</td>
<td>5.6</td>
<td>0.6</td>
<td>2.6</td>
<td>4.3</td>
</tr>
<tr>
<td>Gross tumor volume (GTV)</td>
<td>104.0</td>
<td>0.1</td>
<td>27.5</td>
<td>89.7</td>
<td>153.5</td>
<td>5.9</td>
<td>13.9</td>
<td>2.1</td>
<td>4.1</td>
<td>6.2</td>
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<tr>
<td>Necrotic core</td>
<td>54.1</td>
<td>0.0</td>
<td>38.8</td>
<td>53.8</td>
<td>65.2</td>
<td>3.5</td>
<td>5.6</td>
<td>2.3</td>
<td>4.1</td>
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<td>Orbits</td>
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<td>6.2</td>
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<td>0.0</td>
<td>0.5</td>
<td>0.8</td>
<td>0.9</td>
</tr>
<tr>
<td>Scalp</td>
<td>596.0</td>
<td>0.0</td>
<td>1.0</td>
<td>1.0</td>
<td>1000</td>
<td>8.7</td>
<td>21.0</td>
<td>0.1</td>
<td>0.9</td>
<td>8.3</td>
</tr>
<tr>
<td>Skull</td>
<td>537.8</td>
<td>0.0</td>
<td>24.6</td>
<td>437.2</td>
<td>1000</td>
<td>31.8</td>
<td>38.1</td>
<td>0.6</td>
<td>2.5</td>
<td>42.8</td>
</tr>
<tr>
<td>White matter</td>
<td>126.9</td>
<td>0.0</td>
<td>48.7</td>
<td>137.0</td>
<td>169.8</td>
<td>2.2</td>
<td>2.4</td>
<td>0.9</td>
<td>2.7</td>
<td>4.3</td>
</tr>
</tbody>
</table>

GTV, gross tumor volume; EVH, electric field–volume histogram; SARVH, specific absorption rate–volume histogram; $E_{\text{AUC}}$, electric field area under the curve; $V_{\text{E150}}$, volume covered with electric field intensity of 150 volts per meter; $E_{\text{95\%}}$, the electric field intensity encompassing 95% of volume; $E_{\text{50\%}}$, the electric field intensity encompassing 50% of volume; $E_{\text{20\%}}$, the electric field intensity encompassing 20% of volume; $SAR_{\text{AUC}}$, SAR area under the curve; $V_{\text{SAR7.5}}$, volume covered with specific absorption rate of 7.5 watts per kilogram; $SAR_{95\%}$, the magnitude of specific absorption rate encompassing 95% of volume; $SAR_{50\%}$, the magnitude of specific absorption rate encompassing 50% of volume; $SAR_{20\%}$, the magnitude of specific absorption rate encompassing 20% of volume.
necrotic core and the original tumor was 111.8 and 104.0 V/m, respectively. In addition, the SAR_AUC of the replaced necrotic core and the original tumor was 6.1 and 5.9 W/kg, respectively. Interestingly, when the material properties of the bilateral ventricle, which was considered a highly conductive medium, were replaced with the properties of gray matter, neither the electric field strength nor SAR changed significantly. This is possibly due to the bulk of the incident electric fields, which permeate first through cerebrospinal fluid at the convexity of the brain, has already deposited most of the energy at the convexity interface as the cerebrospinal fluid in this region is highly conductive compared with adjacent tissues. In contrast, the intensity of the source fields was not easily altered in regions deep within the center of the brain where the bilateral ventricles are located.

**The influence of tumor geometry on EVH and SAR/VH**

The influence of tumor geometry on the electric field and rate of energy deposition was also investigated. The initial GTV was kept intact as segmented in the patient’s model, and was used to compare with GTVs of other standard geometric solids, while keeping the centroid of each GTV at the same location. All other parameters such as array position and electric conductivity of tissue remained the same (Fig. 3E and Table 1). Standard geometric solids, including cube, cylinder, sphere, icosahedron, and cone, were used to represent the shape of the tumor for studying changes in electric field distribution and energy deposition in the GTV (Fig. 8). To simplify the comparisons, the necrotic core properties across all models, including the original brain model, were set equal to the GTV; thus, the GTV studied in this section is essentially GTV and necrotic core combined into one entity. In addition, we were interested to determine how the electric fields are distributed depending on the orientation of the conical solid. Therefore, the conical solid was rotated about the geometric centroid of the original patient’s GTV, with the tip of the solid pointing in six different directions according to the patient’s anterior, posterior, left, right, superior, or inferior head positions.

Compared to the original tumor geometry, symmetric, less angulated geometries had a tendency to associate with a lower basal level of electric fields, as shown by lower values in $E_{AUC}$ and $E_{30\%}$, and diminished energy absorption, as shown by SAR_AUC and SAR_{20\%} (Fig. 9).

![Figure 3. Array displacement for disposition analysis.](image-url)
In particular, the cylindrical and spherical GTVs had lower $E_{\text{AUC}}$ and $E_{\text{20\%}}$ as well as $\text{SAR}_{\text{AUC}}$ and $\text{SAR}_{\text{20\%}}$ than that for the cube and icosahedron, and these changes are likely due to the fact that the latter ones had more angulated corners. For lesser symmetric geometric shapes, such as a cone, the orientation in three-dimensional space of the brain was important. The $E_{\text{AUC}}$ and $E_{\text{20\%}}$, as well as $\text{SAR}_{\text{AUC}}$ and $\text{SAR}_{\text{20\%}}$, had the lowest values when the cone was pointing superiorly but highest when pointing anteriorly (Figs. 8B and 9C). The highest values of $E_{\text{AUC}}$ 122.0 V/m and $E_{\text{20\%}}$ 163.9 V/m, as well as $\text{SAR}_{\text{AUC}}$ 6.5 W/kg and $\text{SAR}_{\text{20\%}}$ 7.8 W/kg, were found when the conical GTV was pointing anteriorly, with the flat surface facing orthogonal to the lateral ventricle.

**Discussion**

As it is being used to treat glioblastoma, TTFIELDS at 200 kHz permeate the patient brain according to the laws of physics, including Gauss’ law, Coulomb’s law, and the principles of continuity and capacitance. Our modeling of these fields using the finite-element method reveals that the intensity of TTFIELDS is highest at the sulci, genu of the corpus callosum, and at the medial surface of the

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1293
GTV facing the lateral ventricle. To better assess quantitatively the electric field distribution and rate of energy deposition within the brain, volume histograms were constructed for the electric fields and SAR. To our knowledge, this is the first comprehensive usage of both EVH and SARVH to quantify and compare, respectively, TTFields distribution and rate of energy deposition within the patient brain and tumor in various models. Both EVH and SARVH can also facilitate the quantitative comparison of intracranial TTFields between patients, using prespecified values at the GTV and other segmented structures, such as $E_{\text{AUC}}$, $V_{\text{E150}}$, $E_{95\%}$, $E_{50\%}$, and $E_{20\%}$ for TTFields distribution as well as $\text{SAR}_{\text{AUC}}$, $V_{\text{SAR7.5}}$, $\text{SAR}_{95\%}$, $\text{SAR}_{50\%}$, and $\text{SAR}_{20\%}$ for rate of energy deposition. $E_{\text{AUC}}$ represents the aggregate electric fields as represented by the area under the curve within any segmented structures and $V_{\text{E150}}$ is the volume of distribution for the electric field intensity at least 150 V/m on the histogram. Similarly, $\text{SAR}_{\text{AUC}}$ also represents the aggregate rate of energy deposited as represented by the area under the curve within any segmented structures and $V_{\text{SAR7.5}}$ is the volume of distribution for SAR at 7.5 W/kg on the histogram. In addition, $E_{95\%}$, $E_{50\%}$, and $E_{20\%}$ are data points on the EVH curve that represent the electric field intensity that covers 95%, 50%, and 20% of the GTV, whereas $\text{SAR}_{95\%}$, $\text{SAR}_{50\%}$, and $\text{SAR}_{20\%}$ are data points on the SARVH that represent the rate of energy deposited in 95%, 50%, and 20% of the GTV. Collectively, these parameters quantitatively represent the intrinsic characteristics of EVH and SARVH and therefore allow the comparison of TTFields among different patients.

Our TTFields modeling is based on MP-RAGE, T1, and T2 MRI sequences obtained from a patient with glioblastoma in the right parietal brain. Using patient-based MRI for finite-element modeling has several distinct advantages compared to modeling based on atlas-based human head and/or generalized tumor model [10, 11]. First, there are intrinsic factors that are unique to the individual human head that may influence the electric field and rate of energy deposition in the GTV. A prime example is the volume of cerebrospinal fluid within the subarachnoid space at the convexity. Because cerebrospinal fluid is a good electric conductor, a thicker layer tends to shunt TTFields away from the brain while a thinner layer allows more fields to penetrate the brain and eventually to the GTV. Indeed, as brain atrophy occurs in the general population spanning the entire age continuum in adulthood, there is a simultaneous increase in cerebrospinal fluid space from age 18 to 80 [12]. Furthermore, the rate of atrophy is quicker for men than woman, particularly in the left hemisphere, suggesting neuroanatomical differences between genders [12].
Glioblastoma also can influence the distribution of TTFields at the GTV [12]. The necrotic core is primarily consisted of a collection of thick fluid from broken-down cellular debris containing various proteins and metabolites [13, 14]. Because this fluid is ionic and highly conductive, it may direct electric fields toward the more cellular portion of the GTV. Indeed, our modeling has shown that the necrotic core can influence both the electric field strength and SAR within the adjacent GTV as the electric conductivity of the GTV varied from 100 to 1 S/m. Lastly, glioblastoma is an infiltrative tumor [15] and the accumulation of the tumor cells in the adjacent gyri and sulci may change the geometry of the tumor. Our geometric analysis has shown that electric field and SAR values grossly increase as the tumor geometry becomes more angulated, as seen in the cube and icosahedron models compared to spherical and cylindrical models. This is likely due to the fundamental basis of electric force and surface charge distribution, where charges bunched up at sharp corners experience a very high internal electric force exerted upon adjacent charges, thereby creating a higher electric field. Likewise, smoother surfaces enable electric charges to distribute more uniformly, generating a lower electric field. An extreme form of tumor geometric asymmetry is represented by a cone and modeling revealed that the orientation of this structure is important, resulting in highest electric field intensity and SAR in this particular patient model when the vertex is pointing anteriorly and the flat base is facing orthogonally to the lateral ventricle. Taken together, atlas-based modeling may not accurately

Figure 5. Sensitivity analysis of electric conductivity of GTV and necrotic core. When the electric conductivity of the necrotic core was held constant, the mean electric field strength within the necrotic core rose 600% from 5 to 30 V/m when the conductivity of the GTV decreased from 100 to 1 S/m, but further increase was markedly attenuated when the conductivity was lowered from 1 to 0.001 S/m (A). The mean SAR increased from 0 to 1.8 W/kg when the conductivity of the GTV was decreased from 100 to 1 S/m, but further increase was negligible when the conductivity was lowered from 1 to 0.001 S/m (C). When the electric conductivity of the GTV was held constant and the electric conductivity of the necrotic core was varied from 100 to 1 S/m, there was negligible change in the mean electric field strength in the GTV; but there was up to a 10% increase in mean electric field strength of the GTV when the electric conductivity of the necrotic core was lowered from 1 to 0.001 S/m (B). However, the change in mean SAR of the GTV was insignificant when the necrotic core conductivity was varied from 100 to 0.001 S/m (D). GTV, gross tumor volume; V/m, volt per meter; S/m, Siemens per meter; W/kg, watt per kilogram; SAR, specific absorption rate.
Figure 6. Sensitivity analysis for electric field strength and rate of energy deposition with or without the necrotic core. EVH (A) and SARVH (C) were modeled with the necrotic core, which consisted of highly conductive fluid. When the electric properties of the necrotic core were replaced with poorly conductive tissue, such as white matter, the electric field coverage and rate of energy deposition were increased as shown in the EVH (B) and SARVH (D), respectively. The electric field diagrams showed differences in the electric field coverage at the GTV with (E) and without (F) the necrotic core. EVH, electric field–volume histogram; SARVH, specific absorption rate–volume histogram.
incorporate patient-related differences in brain volume, CSF space and physical characteristics of the tumor, all of which influence TTFields distribution in the brains of glioblastoma patients.

Transducer array positioning also influences TTFields distribution within the brain. In accordance to the data present by Wenger et al. [6], we also noted changes in TTFields at the GTV depending on array positioning, with

Figure 7. Influence of cerebrospinal fluid on the electric field strength and rate of energy deposition at the GTV and necrotic core. The layer of cerebrospinal fluid was altered by −1.0 mm, −0.5 mm, and +0.5 mm at the convexity of the brain and the respective EVH (A) and SARVH (B) at the GTV were generated. The electric field parameters $E_{AUC}$, $V_{E150}$, and $E_{50\%}$, as well as parameters for the rate of energy deposition $SAR_{AUC}$, $V_{SAR7.5}$, and $SAR_{50\%}$, all increased progressively when the cerebrospinal fluid space was narrowed progressively from +0.5 mm to −0.5 mm, and then to −1.0 mm on the convexity of the brain. GTV, gross tumor volume; EVH, electric field–volume histogram; SARVH, specific absorption rate–volume histogram; $E_{AUC}$, electric field area under the curve; $V_{E150}$, volume covered with electric field intensity of 150 volts per meter; and $E_{50\%}$, the electric field intensity encompassing 50% of volume; SAR, specific absorption rate; $SAR_{AUC}$, SAR area under the curve; $V_{SAR7.5}$, volume covered with specific absorption rate of 7.5 watts per kilogram; $SAR_{50\%}$, the magnitude of specific absorption rate encompassing 50% of volume.
significant variability in the electric field strength of GTV between 100 and 150 V/m, ranging from only 20% volume having 100 V/m with clockwise rotation of the lateral arrays and no displacement of the anterior–posterior arrays to >40% volume having at least 150 V/m when the lateral arrays are rotated in a counterclockwise fashion and the anterior array is moved backward. SAR variability in the rate of energy deposition was less dramatic, as the magnitude of SAR$_{50\%}$ changes between 3 and 6 W/kg. These results suggest that array positioning is important to maximize TTFields distribution at the GTV and to determine if the maximal electric field distribution correlates with survival in the patient population treated with TTFields.

Our results have limitations. First, our modeling only used isotropic values but there are local fields generated by the electrical activities of neurons and the flow of cerebrospinal fluid that influence regional electric conductivity [16, 17]. Therefore, anisotropic mapping may delineate a more accurate distribution of TTFields. However, for comparing various structural components that may influence TTFields, isotropic modeling still allows a relatively accurate comparison among different dielectric-, geometric-, and tumor-related parameters. Second, the GTV is delineated by the leakage of gadolinium across permeable tumor vasculature. But this vascular permeability can be increased or decreased by radiation and drugs, as in, respectively,
pseudoprogression after concurrent radiation and temozolomide treatment for newly diagnosed glioblastoma or psuedoresponse after the administration of an antiangiogenic agent like bevacizumab [18]. Therefore, the GTV can be overestimated or underestimated depending on the biological response of the glioblastoma to treatment, and the resolution of this issue is limited by the currently available MRI technology. However, methionine positron emission tomography can help to confirm the presence of active tumor within the gadolinium-enhanced GTV, whereas diffusion-weighted MRI and multivoxel MR spectroscopy may detect nonenhancing tumor without hyperpermeable vasculature [19, 20]. Third, we did not account for infiltrative nonenhanced disease in the model because the purpose of this study is to address bulk changes in distribution of the TTFields and energy deposition by varying the physical aspects of the treatment delivery. Lastly, we only performed modeling on one patient. It will require comparison of multiple patients, in particular between responders and nonresponders, in order to determine the relevant parameters that influence TTFields in the glioblastoma population.

In conclusion, we constructed volume histograms EVH and SARVH to facilitate future comparisons between
glioblastoma patients undergoing treatment with TTFields. TTFields at the GTV are influenced by the dielectric characteristics of the adjacent tissues as well as the GTV itself, particularly the presence or absence of a conductive necrotic core. The amount of cerebrospinal fluid at the convexity of the brain and the geometry of the tumor are additional relevant factors. Finally, the position of the arrays also can influence the electric field distribution and rate of energy deposition within the GTV.

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Conflict of Interest

None declared.

References