A numerical study on the effect of conductivity change in cell kill distribution in irreversible electroporation

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(Received 30 January 2020; accepted 4 May 2020)

Abstract
Introduction: irreversible electroporation (IRE) is a tissue ablation technique and physical process used to kill the undesirable cells. In the IRE process by mathematical modelling we can calculate the cell kill probability and distribution inside the tissue. The purpose of the study is to determine the influence of electric conductivity change in the IRE process into the cell kill probability and distribution.

Methods: cell death probability and electric conductivity were calculated with COMSOL Multiphysics software package. 8 pulses with a frequency of 1 Hz, pulse width of 100 µs and electric field intensity from 1000 to 3000 V/Cm with steps of 500 V/Cm used as electric pulses.

Results: significantly, the electrical conductivity of tissue will increase during the time of pulse delivery. According to our results, electrical conductivity increased with an electric field intensity of pulses. By considering the effect of conductivity change on cell kill probability, the cell kill probability and distribution will change.

Conclusion: we believe that considering the impact of electric conductivity change on the cell kill probability will improve the accuracy of treatment outcome in the clinic for treatment with IRE.

Key words: electroporation; Finite Element Analysis; electric conductivity; Peleg-Fermi; pulsed electric field.

Introduction

During electroporation, cells are exposed to the sufficiently high voltage electric field pulses that increase the transmembrane potential (TMP) of the cell membrane above a threshold [1-3]. The result of this increase in TMP is pore formation in the cell membrane. If the cell membrane recovers and pores were closed, reversible electroporation (RE) occurred. In the RE the cells remain viable. Reversible electroporation is used to transfer of macromolecules into the cells [4]. The combination of RE and chemotherapeutic agents has been used for cancer treatment which named electrochemotherapy (ECT) [5].

But if the cell membrane cannot recover and permanent pores were formed, irreversible electroporation (IRE) has occurred and the consequent is cell dies [6-9]. Recently, IRE with minimally invasive ablation and without joule heating effects is used for the treatment of tumors [10,11]. The IRE is used as a unique mode of cell death that does not rely on thermal damage from joule heating [8,12-14].

Successful IRE therapy depends on treatment planning parameters such as electric field intensity, frequency of the pulses, number of pulses and electrical tissue parameters, etc. Inducing maximum damage to the tumor and minimum damage to the surrounding healthy tissue is one of the most important parameters. This parameter is accomplished by numerical models. Several numerical models describing cell kill probability as a function of the electric field intensity [15,16]. These models are usually based on experimental data. The Peleg-Fermi model as a mathematical model is the most usable model for expression of cell death during IRE [17-19]. The Peleg-Fermi model in comparison with other models seems the most adequate because it includes dependency of pulse number and electric field intensity. For this reason, in this study Peleg-Fermi model was used.

The goal of IRE is inducing the cell death of undesirable cells. There are some factors that affect cell kill probability. One of the most important factors is electric field magnitude and distribution inside the tissue [20]. The electric field distribution and magnitude depends on the electric pulse and tissue properties and electrodes configuration [21]. Electrical conductivity is one of the tissue properties which can affect the electric field distribution and magnitude. Different studies have shown an increase in tissue conductivity in IRE [22-24]. The goal of this study was to describe the effects of electrical conductivity change during IRE on the cell kill distribution.
Material and methods

Finite element and parameter model

To calculate a finite element model, COMSOL Multiphysics 5.3 finite element analysis software was used. A cube geometry with dimensions 32*32*17 mm was modeled as liver tissue. A pair of stainless-steel needle electrodes with 0.43 mm diameter was used for simulation. The distance between the two electrodes is 8.66 mm. Figure 1 represented the electrodes and tissue geometries that were used in this study. In this study, a triangular meshing model which contained 56194 mesh nodes were used. The geometry was created in symmetric model, in order to reduce the simulation time (Figure 1).

8 pulses with a frequency of 1 Hz, pulse width of 100 µs and electric field intensity from 1000 to 3000 V/Cm with steps of 500 V/Cm used as electric pulses. The electrical and thermal properties of the liver and electrodes are listed in Table 1 [25,26].

Calculating method

The electric field and electric potential distribution inside the tissue were obtained by Laplace’s equation:

\[ \nabla \cdot (\sigma \nabla \phi) = 0 \quad \text{Eq. 1} \]

Where \( \sigma \) and \( \phi \) are tissue conductivity and electrical potential, respectively. Heat transfer in the tissue was estimated using Penne’s Bioheat equation:

\[ \nabla \cdot (k \nabla T) + \sigma |\nabla \phi|^2 + q''' - W_b \rho c_b T = \rho c \frac{\partial T}{\partial t} \quad \text{Eq. 2} \]

where \( T \) is temperature, \( \phi \) is electrical potential, \( q \) is the heat produced by metabolism, \( W_b \rho c_b T \) is the heat produced by perfusion, \( \rho \) is density, and \( c_b \) is specific heat capacity of the tissue. The conductivity changes in the IRE inside the tissue was calculated as [25,26]:

\[ \sigma = \sigma_0 \cdot (1 + \text{flc2hs}(E - E_{\text{delta}}, E_{\text{range}}) + \alpha \cdot (T - T_o)) \quad \text{Eq. 3} \]

where \( \sigma_0 \) is tissue initial conductivity, \( E \) is the electric field in the deserted point, \( E_{\text{delta}} \) is threshold electric field in electroporation, \( E_{\text{range}} \) is electric field range, \( \alpha \) is temperature coefficient, \( T \) is the temperature of the tissue, and \( T_o \) is the initial temperature of the tissue. Additionally, flc2hs is a smoothed Heaviside function in the COMSOL that changes from zero to one when \( E - E_{\text{delta}} = 0 \) over the range \( E_{\text{range}} \) to ensure the convergence of the numerical solution [25]. The parameters used in Equation 3, are listed in Table 2.

The electrical boundary condition at the active row of electrodes was set to be \( \phi = V(t) \). Where \( V(t) \) were pulses with time-varying voltage. Another row of electrodes was set at \( \phi = 0 \). The remaining boundaries were considered as electrical insulation.

The mathematical model used to calculate the cell kill due to IRE. The Peleg-Fermi model is given by:

\[ S(E, N) = \frac{N}{N_0} = \frac{1}{1 + e^{-k_1 N}} \quad \text{Eq. 4} \]

where \( S \) is the ratio of surviving cells, \( N \) is the number of surviving cells after IRE treatment, \( N_0 \) is the number of cells prior to IRE treatment, \( E \) is an electric field, \( E_c(N) \) is the critical electric field in which 50% of the cells are killed, and \( A(N) \) is a function of the pulse number. This model accounts for dependency on the number of pulses and electric field during IRE. In equation \( E_c(N) \) and \( A(N) \) is given by:

\[ E_c(N) = E_0 e^{-k_1 N} \quad \text{Eq. 5} \]
\[ A(N) = A_0 e^{k_2 N} \quad \text{Eq. 6} \]

where \( E_0 = 399,600 \text{ V/m}, \ A_0 = 144,100 \text{ V/m}, \ k_1 = 0.03 \) and \( k_2 = 0.06 \) [17].
These equations resulted from fit the experimental data to Equation 4 which performed by Globerg et al. [17]. Additionally, because we wanted to calculate the percentage cell kill, we converted the cell viability into a percentage cell kill given by:
\[ P_{\text{kill}} = 100(1 - S) \]
Eq. 7

Results
The results of this work are divided into two parts: conductivity change during IRE and percentage cell kill during IRE treatment.

Conductivity change during IRE
In this work, the conductivity changes were calculated in two points, tip and middle of electrodes. The conductivity changes of 8 pulses with 2500 V/cm electric field intensity during pulse transmission time are presented in Figure 2. When the intensity of the electric field was higher than the critical value, an increase in conductivity has occurred. As we can see in Figure 2, tissue conductivity in the transmission time of pulses was increased. This increase in the tip of the electrode was higher than in between electrodes. The conductivity changes at the time of last pulses (8th) with different electric field intensity are listed in Table 3.

Percentage cell kill during IRE treatment
To investigate the impact of conductivity change during IRE on the cell kill distribution two groups of simulations were done. One group of simulations is done with constant conductivity and other groups are done with variable conductivity which is given by Equation 3.

I have presented the cell kill probability histogram with different pulses in Figure 3. The cell kill probability histogram represents the frequency of occurrence of cell kill probability which lies within a volume fraction of simulated tissue. The cell kill probability histogram for 3000 V/cm pulses with variable and constant tissue electrical conductivity during IRE are shown in Figure 4.

We can use two-dimensional and three-dimensional images for a better showing of the percentage cell kill distribution inside the tissue. Figures 5 and 6 display the percentage cell kill distribution due to IRE around the electrode for different pulse electric field intensity. The percentage cell kill distribution in Figures 5 and 6 are calculated with variable and constant electrical conductivity during the IRE respectively. In order to visualize the percentage of cell kill due to IRE in 3D, the volume of the liver which has a cell kill probability of more than 80% are presented in Figures 7 and 8 for different pulses. Simulation results in Figures 7 and 8 are calculated with variable and constant electrical conductivity respectively.

![Figure 2](image-url) Conductivity changes used 8 pulses with the frequency of 1 Hz, pulse width of 100 μs, and the intensity of the electric fields of 2500 V/cm in the tip and between of electrode

![Figure 3](image-url) The cell kill probability histogram at last pulse (8th) for 1000-1500-2000-2500-3000 V/cm electric field intensity of pulses with constant tissue electrical conductivity during IRE

![Figure 4](image-url) The cell kill probability histogram at last pulse (8th) for 1000-1500-2000-2500-3000 V/cm electric field intensity of pulses with variable electrical conductivity during IRE

![Figure 5](image-url) Percentage cell kill distribution due to IRE around the electrode for different pulse electric field intensity.

![Figure 6](image-url) Percentage cell kill distribution in Figures 5 and 6 are calculated with variable and constant electrical conductivity during the IRE respectively.

![Figure 7](image-url) Volume of the liver which has a cell kill probability of more than 80%.

![Figure 8](image-url) Volume of the liver which has a cell kill probability of more than 80%.

Table 3. Conductivity change in the tip and between the electrodes at the time of the last pulse with 8 pulses with the frequency of 1 Hz, pulse width of 100 μs, and electric field intensity of 1000-3000 V/cm

| Electric field intensity (V/cm) | Tip (S/m) | Between (S/m) |
|-------------------------------|----------|---------------|
| 1000                          | 0.136    | 0.134         |
| 1500                          | 0.139    | 0.134         |
| 2000                          | 0.143    | 0.134         |
| 2500                          | 0.149    | 0.134         |
| 3000                          | 0.157    | 0.134         |
Figure 5. The percentage cell kill distribution due to IRE around the electrode in the plane perpendicular to the tip of electrodes at last pulse (8th) with variable electric conductivity for pulses with a) 1000 V/cm b) 1500 V/cm c) 2000 V/cm d) 2500 V/cm e) 3000 V/cm electric field intensity.

Figure 6. The percentage cell kill distribution due to IRE around the electrode in the plane perpendicular to the tip of electrodes at last pulse (8th) with constant electric conductivity for pulses with a) 1000 V/cm b) 1500 V/cm c) 2000 V/cm d) 2500 V/cm e) 3000 V/cm electric field intensity.
Figure 7. Volume of the liver which has a cell kill probability of more than 80% at last pulse (8th) with variable electric conductivity for pulses with a) 1000 V/cm b) 1500 V/cm c) 2000 V/cm d) 2500 V/cm e) 3000 V/cm electric field intensity.

Figure 8. Volume of the liver which has a cell kill probability of more than 80% at last pulse (8th) with constant electric conductivity for pulses with a) 1000 V/cm b) 1500 V/cm c) 2000 V/cm d) 2500 V/cm e) 3000 V/cm electric field intensity.
In order to better visualize the volume fraction of tissue killed by IRE, Figure 9 was calculated. Figure 9 displays the computed volume fraction of tissue which has cell kill probability of more than 80% as a function of the electric field intensity of electric pulses. Figure 9 provides the calculated volumes of tissue killed with constant and variable tissue electrical conductivity during the simulation.

Discussion

IRE is a new and invasive ablation technique with nonthermal effects. In order to visualize the cell kill probability distribution and maximizing damage to the tumor and minimizing damage to the surrounding healthy tissue, numerical models have been developed [27-30].

According to the Peleg-Fermi equation (Equation 4) some factors affecting the cell kill distribution such as electric field intensity, number of pulses, etc. Tissue electrical conductivity is one of the factors which can affect the electric field intensity inside the tissue and therefore can change cell kill probability. In this study, initially, the tissue electric conductivity change was calculated during the IRE for different pulses (Figure 2, Table 3). The remarkable result to emerge from the data is that the tissue electrical conductivity increased with an electric field intensity of pulses. This is in good agreement with [31,32]. The observed increase in tissue electrical conductivity could be attributed to being a result of pore creation in the cell membrane during pulse delivery. Which pore creation increased with an electric field intensity of pulses (Table 3). As expected, simulation results demonstrate that the change of tissue electrical conductivity because of greater electric field intensity in the tip of the electrode is more than in point of between electrodes. This confirms our previous findings [31,33].

As stated in the introduction, the main goal of this work was to describe the effects of different electric field intensity and tissue electrical conductivity change during IRE on the cell kill probability distribution. Figures 5 and 6 are showing the cell kill probability distribution for different pulses. It is important to note that the cell kill probability increased with the electric field intensity of pulses. Importantly, cell kill probability is higher in the vicinity of electrodes because of greater electric field intensity in this region. This finding highlights the usefulness of greater electric filed intensity of pulses for the treatment of big tumors. A reasonable explanation for this phenomenon is that, with an increase of electric field intensity of the pulse, the volume of tissue which has cell kill probability >80% become bigger, and high electric field intensity pulses can use for bigger tumor. These findings correlate fairly well with [17].

Histogram (Figures 3 and 4) analysis showed that, with an increase of electric field intensity of pulses, the volume fraction of tissue with low cell kill probability was reduced and volume fraction of tissue with high cell kill probability was increased. These results offer powerful evidence for tumor treatment with an adequate electric field intensity of pulses. The experiment in this work is in line with previous results [34,35]. Figure 4 is representing the cell kill probability histogram for 3000 V/cm pulses with constant and variable tissue electrical conductivity. According to the results, it is important to note that with variable tissue electrical conductivity during IRE, the volume fraction of tissue with low cell kill probability was reduced and the volume of tissue with high cell kill probability was increased in constant electric field intensity of pulse. These results match well with Figure 4. The possible reasons for this result are, during the transmission time of pulse, the pore was created in the cell membrane. The result of pore creation is an increase in the electric conductivity of the medium. The consequence of tissue electrical conductivity raise is an increase in electric current and electric field intensity in the far region of the electrode. According to the Peleg-Fermi equation (Equation 4) the cell kill probability is related to the electric field intensity. So because of an increase in electric field intensity in the far region of the electrode, the cell kill probability was increased. And change the cell kill probability distribution inside the tissue (Figures 5-8).

Conclusion

One of the goals of treatment planning in IRE is identification the treated volume of target tissue before the IRE process being started. The key to successful treatment is ensuring that the tumor treated with sufficient cell kill probability during the treatment. This is done by numerical models such as finite element methods. This paper has evaluated the importance of electric conductivity change and electric field intensity of pulse on cell kill probability. The results of this paper have demonstrated that, by considering the impact of electric conductivity change on cell kill probability, the cell kill
probability was changed. So, in order to accurate prediction of treatment in treatment planning, it has been suggested to considering the role of electric conductivity change on cell kill probability. The results have obtained satisfactory demonstrating that for treatment of the big tumor, we can use pulses with high electric field intensity. However, using the pulses with high electric field intensity increase the temperature of tissue [36] and must be considered. This is important for future research to survey the impact of electrical conductivity change on tissue temperature raise during IRE.

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