Electrosensitization Increases Antitumor Effectiveness of Nanosecond Pulsed Electric Fields In Vivo

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Abstract
Nanosecond pulsed electric fields are emerging as a new modality for tissue and tumor ablation. We previously reported that cells exposed to pulsed electric fields develop hypersensitivity to subsequent pulsed electric field applications. This phenomenon, named electrosensitization, is evoked by splitting the pulsed electric field treatment in fractions (split-dose treatments) and causes in vitro a 2- to 3-fold increase in cytotoxicity. The aim of this study was to show the benefit of split-dose treatments for in vivo tumor ablation by nanosecond pulsed electric field. KLN 205 squamous carcinoma cells were embedded in an agarose gel or grown subcutaneously as tumors in mice. Nanosecond pulsed electric field ablations were produced using a 2-needle probe with a 6.5-mm interelectrode distance. In agarose gel, splitting a pulsed electric field dose of 300, 300-ns pulses (20 Hz, 4.4-6.4 kV) in 2 equal fractions increased cell death up to 3-fold compared to single-train treatments. We then compared the antitumor effectiveness of these treatments in vivo. At 24 hours after treatment, sensitizing tumors by a split-dose pulsed electric field exposure (150 + 150, 300-ns pulses, 20 Hz, 6.4 kV) caused a 4- and 2-fold tumor volume reduction as compared to sham and single-train treatments, respectively. Tumor volume reduction that exceeds 75% was 43% for split-dose–treated animals compared to only 12% for single-dose treatments. The difference between the 2 experimental groups remained statistically significant for at least 1 week after the treatment. The results show that electrosensitization occurs in vivo and can be exploited to assist in vivo cancer ablation.

Keywords
nanosecond pulsed electric fields (nsPEF), irreversible electroporation, tumor ablation, nanoporation, electrosensitization

Abbreviations
DT, doubling time; EMEM, Eagle’s minimum essential medium; IRE, irreversible electroporation; nsPEF, nanosecond pulsed electric fields; PBS, phosphate-buffered saline; PEF, pulsed electric fields; PI, propidium iodide; Pr, propidium; PRR, pulse repetition rate; ROI, region of interest; ROS, reactive oxygen species

Received: February 15, 2017; Revised: April 24, 2017; Accepted: May 04, 2017.

Introduction
Several local ablative therapies have been explored as innovative treatments to fight cancer. The advantages of these procedures include increased safety, less scarring, fast recovery, and decreased length of hospital stay.¹ Irreversible electroporation (IRE) is a promising new minimally invasive technique for tumor ablation. Irreversible electroporation uses high-intensity pulsed electric fields (PEF) of 100 µs duration to cause irreparable cell damage and destroy tissues.² Compared to other conventional ablation methods, such as radiofrequency heating and cryoablation, IRE treatments preserve vital structures and major blood vessels within the ablated zone.³ Irreversible electroporation was found particularly efficient at treating tumors less than 3.0 cm.² However, larger lesions

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Electrosensitization can potentially enable the use of lower electric field amplitudes to achieve the same or better ablation effect than standard single-train protocols. Attaining the therapeutic effect at much lower exposure doses translates into a reduction in the thermal damage due to the Joule heating and could potentially minimize IRE known side effects such as pain, muscle contraction, and cardiac arrhythmias when treatments are done in the proximity of the heart.

Materials and Methods

Cell Line and Media

Experiments were performed using a mouse squamous cell carcinoma KLN 205 cell line (ATCC, Manassas, Virginia). Cells were cultured in Eagle’s minimum essential medium with l-glutamine (ATCC), supplemented with 10% (v/v) fetal bovine serum (Atlanta Biologicals, Norcross, Georgia), 100 U/mL penicillin and 0.1 mg/mL streptomycin (Mediatech Cellgro, Herdon, Virginia).

Pulsed Electric Field Exposure

The same PEF exposure system was used for both 3D cultures and tumors in mice. Trapezoidal pulses of 300 ns duration were produced by a custom pulse generation system with an output impedance of 100 Ω, adjustable pulse amplitude (up to 15 kV), duration (150-2000 ns), and frequency (1-100 Hz; Tangers Electronics, Norfolk, Virginia; Figure 1A). Pulsed electric field was delivered by a PEF-delivering 2-needle electrode probe made from 0.5 mm diameter hollow syringe needles. The edge-to-edge distance between the needles was 6.5 mm with a maximum penetration capacity in agarose culture or the mouse skin of 5 mm (Figure 1B). To produce pulse trains of predetermined duration at selected repetition rates, the generator was triggered externally from a model S8800 stimulator (Grass Instrument Co, Quincy, Massachusetts). Pulse amplitude was monitored in all experiments, using a 200 MHz, 1 GS/s DSO5202B digital oscilloscope (Antek, Qingdao City, China). The shape of the electric pulse at 6.4 kV is reported in Figure 1C.

Pulsed electric field delivery to cells embedded in agarose gel was accomplished by mounting the electrodes on a micro-manipulator to enable accurate and steady positioning of the needles within the gel with cells in a 35-mm Petri dish. For accurate comparison, different PEF treatments and sham exposure (no pulses delivered) were performed in the same cell sample for a maximum of 6 exposures per dish. We previously reported that electrosensitization develops only if the electric field is above the threshold for electroporation. In our experiments, each exposure was spaced 1 cm from the previous one to avoid “electroporation-preconditioning” of the area. In fact, for 300 ns pulses applied at 6.4 kV (which is the highest electric field used in the in vitro experiments shown in Figure 4), the electric field at 1 cm from the electrodes is 0.8 kV/cm. Based on the electroporation thresholds found, for...
200 and 600 ns pulses, which are 1.5 to 1.9 and 1 kV/cm, respectively.\textsuperscript{43,44} 0.8 kV/cm gives a good safety margin being about 2-fold below the expected threshold for 300 ns pulses.

For tumor PEF treatments, mice were anesthetized by inhalation of 3\% isoflurane in air (Patterson Veterinary, Devens, Massachusetts). The 2-needle electrode was inserted at the opposite margins of the tumor by lifting the overlying skin. Ultrasound conductive gel (Parker Laboratories, Fairfield, New Jersey) was used to ensure an efficient electrical continuity. During PEF treatments, the probe was held by hand and readjusted to the proper position between trains if needed. Animals in the sham control group underwent anesthesia and the probe insertion procedure but no PEF delivery. Tumors were subjected to a single-PEF treatment with 1 electrode insertion.

**Electric Field Simulation and Temperature Measurements**

To quantify the electric field distribution, we carried out a 3D numerical simulations using the commercial finite element method solver COMSOL Multiphysics, Release 5.0. Figure 2A shows the electric field at the cell location between the electrodes, 3.8 mm above the plane of the Petri dish. The dish was modeled as a 2-mm thick layer of polystyrene. To match the experimental conditions, the electrodes (modeled as stainless steel) were positioned perpendicular to the Petri dish, 1 mm above it. The electric currents interface was used to solve Maxwell equations under the assumption of steady-state conditions, for which:

\[
\nabla \cdot (-\sigma \nabla V) = 0
\]

This equation is solved for the voltage field, \(V\), which is used to compute the electric field, \(E = -\nabla V\), and the current, \(J = \sigma E\), where \(\sigma\) is the material conductivity.

Since, the calculations were based on an electrostatic model that disregards the dispersive properties of the medium: this was possible since the dielectric relaxation time for a buffer conductivity of 1.4 S/m would be on the order of few nanoseconds, much shorter than the nsPEF duration considered.\textsuperscript{45-47} The tetrahedral mesh chosen to discretize the domain of simulation (sphere of air with radius of 21.8 mm) resulted in a total of 668 169 elements, with a minimum size of 0.065 mm and a maximum size of 1.530 mm. Quadratic elements were used throughout the solution domain, giving \(0.9 \times 10^6\) degrees of freedom. Under these conditions, the mean electric field and its standard deviation (SD) in the region of interest (ROI), defined as a 1 mm \(\times\) 1 mm square between the electrodes, was 0.082 (0.001) kV/cm for 100 V of input. The coefficient of variation in percent, namely the ratio of...
the SD over the mean value of the electric field, was 1.2% showing the high homogeneity of the electric field in the ROI.

In the 3D cultures, the local heating was measured using a fiber optic ReFlex-4 thermometer (Nortech Fibronic, Quebec City, Canada). The fiber optic probe was inserted into the agarose culture in the center of the gap between the electrodes (Figure 2B). By the end of PEF exposure, the measured values for 300, 300-ns pulses at 4.4, 5.4, and 6.4 kV delivered at 20 Hz averaged 0.5, 0.8, and 1.1°C, respectively (Figure 2C). In practice, PEF trains that were used to study electrosensitization effects (up to 300 pulses) did not raise the temperature to potentially damaging levels.

**Cell Culture and PEF Cytotoxicity in 3D Cultures**

Three-dimensional cultures in agarose were described in detail previously. Briefly, the bottom of a 35-mm dish was coated with 3 mL of 2.5\% low-gelling-temperature agarose (Sigma-Aldrich, St Louis, Missouri) in the growth medium. Cells were resuspended at 5 × 10⁶ cell/mL in 1% agarose in the growth medium, and 1.5 mL of this suspension was pored over the presolidified 2.5% agarose base layer. The samples were incubated at 4°C for 5 minutes to speed up agarose jellification thus avoiding cell sedimentation, covered with 0.5 mL of media, and kept in the incubator for 30 minutes before PEF treatment.

The agarose cultures were analyzed 2 hours after exposure. Dead cells were stained using 4 mg/mL of propidium iodide (PI; Sigma-Aldrich) in phosphate-buffered saline (PBS). Thirty minutes before the analysis, the growth medium was replaced with 1 mL of PI solution. Images of the ablation zone were acquired using an Olympus SZX16 fluorescence stereomicroscope (Olympus America, Hamden, Connecticut) equipped with a Hamamatsu C9100 EM-CCD camera using a 0.9×, 0.44 NA objective. Images were analyzed with MetaMorph version 7.5.2 software (Molecular Devices, Foster City, California). Propidium (Pr) signal was quantified in the center of the gap between the electrodes within a region of 1 mm² with almost uniform electric field (Figure 2A). For each image, the fluorescence intensity of the exposed area was corrected for the background fluorescence.

**Murine Tumor Model**

Seven- to 8-week-old DBA/2 J female mice (Jackson Laboratory, Bar Harbor, Maine) were anesthetized by inhalation of 3% isoflurane and inoculated subcutaneously in the left dorsolateral
flank region with 1 × 10^6 KLN 205 cells in 50 μL of PBS. Mice were housed in individually ventilated cages in groups of 5 under pathogen-free conditions. Tumors were allowed to grow to a volume of approximately 35 to 45 mm^3 before PEF treatment. Tumor growth was measured at 24 hours posttreatment and twice weekly using a digital caliper, and volumes (v) were calculated using a standard formula \( v = \frac{ab^2\pi}{6} \), where \( a \) is the longest diameter, and \( b \) is the next longest diameter perpendicular to \( a \).

Mice were humanely euthanized when the tumor reached 800 mm^3. Tumors were considered eliminated when no recurrence was detected within 100 days after treatment. Tumor doubling time (DT) was calculated using the Schwartz formula.48

This experiment protocol was approved by the Old Dominion University Institutional Animal Care and Use Committee (permit number: 15-009).

**Statistical Analysis**

Data are presented as mean ± standard error for \( n \) independent experiments. Statistical analyses were performed using a 2-tailed \( t \) test where \( P < .05 \) was considered statistically significant. Statistical calculations, including data fits, and data plotting were accomplished using Grapher 11 (Golden Software, Golden, Colorado).

**Results**

**Quantification of the Electrosensitization Phenomenon In Vitro in 3D Agarose Cultures**

In our previous publications, the electrosensitization phenomenon has been investigated by using exposure system with 1 mm distance between the PEF-delivering electrodes (ie, 1-mm gap cuvettes or a 1-mm gap 2-needle probe).28,35,37 However, because our in vivo model envisages the treatment of tumors of about 5 mm in diameter, here, we studied the effect of dose fractionation on the size of ablation zone generated in a 3D agarose culture by applying a voltage across a 6.5-mm gap.

As a first step, we established the PEF conditions needed to kill in 2 hours 100% of cells between the PEF-delivering electrodes. For consistency with our previous research,37 we used pulses of 300 ns duration. The effect of increasing number of pulses 300 ns (6.4 V, 20 Hz) on KLN 205 cells embedded in 1% agarose was visualized by the Pr fluorescence in the lethally damaged cells. Figure 3 shows that significant cell killing started after 200 or more pulses. To quantify this effect, we measured the Pr fluorescence intensity in a region of 1 mm^2 in the center of the gap between the electrodes where cells experienced nearly uniform electric field (Figures 3A and 4). Propidium fluorescence intensity reached a plateau at 400 to 800 pulses indicating that at these doses, 100% of the cells in the quantified area were killed. Therefore, in the subsequent experiments, a parallel control exposure of 600 pulses was used as a reference point for Pr fluorescence that corresponds to 100% cell death.

To investigate the electrosensitization phenomenon, 300 pulses (300 ns, 20 Hz) were delivered either as a single train or as 2 trains with a 2-minute interval (also referred to as single- and split-dose treatments below), and the voltage applied was varied from 4.4 to 6.4 kV. Figure 4A shows for a representative experiment that fractionated treatments produced larger Pr-positive region. Maximum sensitization effect was seen at 4.4 kV (3.6 kV/cm in the center between the electrodes) where

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**Figure 3.** Analysis of the cytotoxic effect of 300 ns, 6.4 kV pulses in KLN 205 cells embedded in agarose. A, The ablation area between and around the 6.5-mm gap pulsed electric field (PEF)–delivering electrodes (arrows) was visualized by propidium (Pr) uptake by dead cells. Images were taken 2 hours after exposure to the indicated number of pulses. Scale bar: 1 mm. B, Quantification of the cytotoxic effect by the mean intensity of Pr fluorescence (as measured within the region of interest shown in (A)). Mean ± standard error (SE), \( n = 3 \). Note signal saturation at 400 to 800 pulses, which indicates killing of 100% of cells within the studied region.
the Pr signal increased 3 times and lethality increased from 12%-16% to 41%-46% (Figure 4B). At the next higher amplitude (5.4 kV), fractionated doses were significantly more efficient as well. However, at 6.4 kV where the single dose already killed 80% of the cells, approaching the fluorescence saturation point, splitting the PEF train in fractions had no additional effect.

Overall, in these in vitro experiments, we established the conditions necessary to achieve efficient sensitization effect using a large gap PEF-delivering probe.

Electrosensitization Assists Tumor Ablation by nsPEF In Vivo

The lethal electric field threshold for cells in medium may considerably differ from that of tumors in vivo.49,50 Hence, the PEF conditions to see electrosensitization in vivo needed to be tested in preliminary experiments. Our in vitro data showed that splitting in fractions a PEF dose that kills 20% to 50% of the cells, causes maximum electrosensitization effect. To identify the PEF dose that matches these conditions in vivo, we investigated the antitumor efficacy of 300 ns pulses over a range of electric field amplitudes. At 24-hour posttreatment, 300, 300-ns pulses (20 Hz) at 6.4 kV caused 50% tumor reduction, and therefore this PEF dose was selected to study the electrosensitization phenomenon in vivo (data not shown).

At the time of treatment, tumors were on average 40.2, 41.2, and 39.2 mm³ for sham exposure, single-dose, and split-dose groups, respectively (Table 1). Tumor size was assessed 24 hours after the treatment and then monitored twice weekly for 3 weeks. At 24 hours after the treatment, split-dose protocols (150 + 150 pulses) with intertrain interval of 2 minutes, caused a 4- and 2-fold tumor volume reduction as compared to sham and single-dose exposure controls, respectively (Figure 5A). In the split-dose group, 43% of the animals exceeded the 75% level of tumor volume reduction, whereas only 12% of single-dose treated animals achieved a similar ablation level (Figure 5B). The difference between single- and split-dose treatments remained statistically significant for at least 1 week after the treatment (Figure 5A). No differences in tumor DT were observed between the different experimental groups suggesting that the tested nsPEF did not trigger any tumor growth inhibition besides their lethal effect at the time of treatment (Figure 5C). Finally, none of the animals were cured with the single-dose protocol, whereas 1 split-dose treatment resulted in complete tumor regression with no recurrence (tumor-free animal for 100 days).

Discussion

In this study, we show, for the first time, that split-dose nsPEF treatments engaging electrosensitization facilitate tumor
destruction in vivo as well as in vitro. Despite delivering equivalent doses, splitting a PEF train in 2 fractions with a 2-minute interval caused a 2-fold tumor volume reduction compared to single-train treatments. All mice treated with the split-dose treatment experienced a significant reduction in tumor volume, and this difference remained statistically significant for a week. However, with the PEF dose used in the study (300, 300-ns pulses, 20 Hz at 6.4 kV), engaging electrosensitization did not cause persistent tumor regression. Maximum electrosensitization effect may develop within a certain range of pulse amplitudes, durations, frequencies, number of pulses, and trains. It remains to be fully established how various PEF parameters affect the development of electrosensitization. In this study, because the needles penetrated the body of the mouse, the range of PEF doses that could be tested was limited by the risk of harming internal organs. For instance, 6.4 kV was the highest PEF amplitude that was also well tolerated by the animals. We therefore believe that the results of this pilot study can be further improved by a careful design of the PEF-delivering electrodes thus allowing to optimize the PEF parameters in vivo.

We previously reported that the time interval between trains affects cell killing by electrosensitization. In KLN 205 embedded in 3D agarose, dose fractionation enhanced Pr uptake when the intertrain interval reached 100 seconds. However, the time course of electrosensitization in vitro and in vivo in tumors may differ. In this study, we used 2-minute intervals that might have not been optimal to cause maximum facilitation of cell killing by engaging electrosensitization. Future work will focus on the optimization of the intertrain interval to fully characterize the potential of electrosensitization for tumor ablation.

Other groups have introduced time intervals between trains in their PEF protocols but with the aim of either allow for heat dissipation or to recharge the pulse generator. We are aware of only 1 study that investigated how intertrain intervals affect IRE efficiency. Indeed, by splitting fifty-one 50-μs pulses in 3 trains of 17 pulses each with 30-second intervals, Jiang et al found increased tumor destruction. In reality, these authors exploit split-dose protocols without being aware of the electrosensitization phenomenon. Moreover, they did not use this term themselves.

A critical question that needs to be addressed is what physiological mechanisms are responsible for electrosensitization. One possible mechanism of electrosensitization may involve the influx of calcium (Ca^{2+}) through the nanopores. Ca^{2+} changes impact nearly every aspect of cellular life, and therefore its intracellular concentration and localization are highly regulated. Split-dose treatments prolong the time intervals when the internal Ca^{2+} is elevated, and this loss of homeostasis

Figure 5. In vivo split-dose treatments engaging electrosensitization increase the antitumor effectiveness of nsPEF. A, Tumor growth curves in sham exposure, single-dose 300 pulses (300 ns, 6.4 kV, 20 Hz), and split-dose 150 + 150 pulses with 2-minute interval experimental groups. For each animal, the data were normalized to the tumor volume measured immediately before treatment. B, The tumor reduction frequency histogram at 24 hours after the treatment. C, The tumor doubling time. Mean ± standard error (SE), n = 15, 17, and 14 for sham, split- and single-dose groups, respectively. *P < .05 for the difference between single- and split-dose groups.
may cause cell death through both necrosis and apoptosis. The prolonged PEF treatment may also aggravate the adenosine triphosphate (ATP) demand to sustain membrane repair and ion pumps activated to restore ion gradients. Therefore, one can speculate that a prolonged high demand for ATP combined with the ATP loss through pores could be a factor responsible for electrosensitization.

Future search for mechanisms responsible for electrosensitization should also focus on how PEF “prime” the cells making them hypersensitive to electroporation. The first train of pulses may increase the level of reactive oxygen species (ROS). A recent study reported that oxidative damage to the membrane increases its susceptibility to electroporation. Hence, the PEF-induced ROS production may increase the cell membrane sensitivity to subsequent treatments. Electrosensitization may be explained by the PEF-induced colloid osmotic cell swelling phenomenon. Permeabilization of cells leads to water uptake and cell swelling due to the so-called colloid osmotic mechanism. The increase of cell diameter translates into a higher PEF-induced transmembrane potential and therefore increases electroporation. Swelling takes tens of second, which may explain the time dynamic of the electrosensitization onset.

As of today, none of these mechanisms have been established as a cause of electrosensitization. We expect that a better understanding of the electrosensitization phenomenon will help to develop more effective PEF treatments.

The results of this in vivo study warrant further exploration of electrosensitization as a facilitating factor for tumor ablation by IRE. Contrary to other approaches used in combination with PEF, engaging electrosensitization does not require chemotherapeutic or cytotoxic agents, and it can be easily integrated in existing IRE protocols. Electrosensitization may allow to reduce the pulse amplitude or to enlarge the distance between the PEF-delivering electrodes without losing the ablation efficiency. More research is needed to establish the optimal PEF conditions as well as to reveal the mechanisms underlying electrosensitization.

Acknowledgements
The authors would like to thank Dr. C. Jiang, Dr. M.A. Malik, and Mr. E. Yang for manufacturing the two-needle probe and Dr. R. Heller, Ms. C. Edelblute, Dr. S. Guo for the valuable discussion and help provided during the in vivo study.

Declaration of Conflicting Interests
The author(s) declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

Funding
The author(s) received the following financial support for the research, authorship, and/or publication of this article: This work was supported by a grant from Pulse Biosciences, Inc. (to O.N.P.) and by a 2015 AFOSR MURI grant (to AGP) on Nanoelectropulse-Induced Electromechanical Signaling and Control of Biological Systems, administered through Old Dominion University.

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