Electrochemotherapy by pulsed electromagnetic field treatment (PEMF) in mouse melanoma B16F10 in vivo

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Introduction. Pulsed electromagnetic field (PEMF) induces pulsed electric field, which presumably increases membrane permeabilization of the exposed cells, similar to the conventional electroporation. Thus, contactless PEMF could represent a promising approach for drug delivery.

Materials and methods. Noninvasive electroporation was performed by magnetic field pulse generator connected to an applicator consisting of round coil. Subcutaneous mouse B16F10 melanoma tumors were treated with intravenously injection of cisplatin (CDDP) (4 mg/kg), PEMF (480 bipolar pulses, at frequency of 80 Hz, pulse duration of 340 µs) or with the combination of both therapies (electrochemotherapy – PEMF + CDDP). Antitumor effectiveness of treatments was evaluated by tumor growth delay assay. In addition, the platinum (Pt) uptake in tumors and serum, as well as Pt bound to the DNA in the cells and Pt in the extracellular fraction were measured by inductively coupled plasma mass spectrometry.

Results. The antitumor effectiveness of electrochemotherapy with CDDP mediated by PEMF was comparable to the conventional electrochemotherapy with CDDP, with the induction of 2.3 days and 3.0 days tumor growth delay, respectively. The exposure of tumors to PEMF only, had no effect on tumor growth, as well as the injection of CDDP only. The antitumor effect in combined treatment was related to increased drug uptake into the electroporated tumor cells, demonstrated by increased amount of Pt bound to the DNA. Approximately 2-fold increase in cellular uptake of Pt was measured.

Conclusions. The obtained results in mouse melanoma model in vivo demonstrate the possible use of PEMF induced electroporation for biomedical applications, such as electrochemotherapy. The main advantages of electroporation mediated by PEMF are contactless and painless application, as well as effective electroporation compared to conventional electroporation.

Key words: pulsed electromagnetic field; bipolar pulses; contactless electroporation; CDDP; electrochemotherapy; platinum determination; mouse melanoma

Introduction

Electroporation is a physical method enabling delivery of impermeable drugs, macromolecules, proteins and genetic material (plasmid DNA, siRNA, miRNA) into cells.1 Electroporation is related to the induced transmembrane voltage which if sufficiently high, enables the formation of tempo-
Electrochemotherapy mediated by pulsed electromagnetic field treatment


Electrochemotherapy is used in treatment of human cutaneous tumors of different histology, and has been translated also in treatment of deep seated tumors. In parallel, electrochemotherapy is being used for treatment of tumors in veterinary oncology. The main chemotherapeutics used in electrochemotherapy are nonpermeable bleomycin and poorly permeable cisplatin (CDDP), via systemic or intratumoral administration route. Electric pulses can be delivered to the tumors via systemic or intratumoral administration route. Electric pulses can be delivered to the tumors via noninvasive plate electrodes, which embrace the tissue, or invasive needle electrodes, which are inserted into the tumor.

In the past the effects of externally applied pulsed electromagnetic fields (PEMF) on the cells were studied extensively. It was demonstrated that externally applied PEMF can influence intracellular signal transduction, affect the cytoskeletal proteins involved in cell shape modification, induce changes in mitochondrial membrane potential, and besides that increase transmembrane molecular transport (electroporation). Since then, a few studies actually defined the PEMF parameters that enabled successful electroporation of cells; i.e. large number of 25 up to 800 the µs long magnetic field pulses applied at frequencies from 25 Hz up to 40 Hz and strength from 725 V/m up to 160 kV/m. Furthermore, its use as electroporation tool was shown in an approach for drug delivery in vivo induced by PEMF. If feasible and effective, electroporation induced by PEMF would have the advantage over “conventional” electroporation, since it is noninvasive, contactless and does not induce pain during electroporation. We assessed the feasibility and antitumor effectiveness of electroporation induced with PEMF as drug delivery system for CDDP to murine melanoma B16F10 subcutaneous tumors. To prove the underlying mechanism of electroporation we measured the platinum (Pt) bound to DNA in tumors. Electroporation induced by PEMF proved to facilitate drug uptake in tumors, such as CDDP, thus providing evidence of its feasibility and effectiveness.

Materials and methods

Drug

CDDP, a chemotherapeutic drug used in electrochemotherapy protocol in human and veterinary clinic, was chosen in the study to test the application of induced electroporation mediated with magnetic field. The stock solution of the chemotherapeutic drug used in the study, CDDP (5 mg/mL, Cysplatyl, Aventis Laboratory, Paris, France) was dissolved in aqua pro injection and frozen in aliquots of 1 mL. In order that each animal received a dose of 80 µg of CDDP, a fresh solution at appropriate concentration of CDDP (1 mg/mL) was prepared in 0.9% sodium chloride solution daily before each experiment.

Mouse tumor model

Female C57Bl/6 mice were purchased from Charles River Laboratories Italy s.r.l. (Calco, Italy) and were maintained in an adaptation period for 14 days.
They were kept at a constant room temperature with a 12 hours light cycle in a conventional animal facility. Eight-week old animals weighing 20–22 g were used in the experiments. Tumors in C57Bl/6 mice were implanted subcutaneously in the right flank of the mice by inoculation of suspension $1 \times 10^6$ B16F10 melanoma cells prepared in 100 µL of phosphate-buffered saline (PBS) for electrochemotherapy experiments. All animal experimental manipulations were conducted in accordance with the principles and procedures outlined with the guidelines for animal experiments of the EU directives and the permission from The administration of the Republic of Slovenia for food safety, veterinary and plant protection (permission No.: 34401–4/2012/2).

**In vivo electrochemotherapy protocol using noninvasive electroporation induced by PEMF or conventional electroporation**

Seven days after subcutaneously induction of B16F10 melanoma tumors (40 mm³) mice were randomly divided into the experimental groups as follows: intravenously injection of saline solution alone (Control) or combined with electroporation induced pulsed electromagnetic field (PEMF), intravenously injection of CDDP (CDDP) or combined with electroporation induced PEMF (PEMF + CDDP). Noninvasive electroporation was performed 3 minutes after intravenous injection of chemotherapeutic drugs by magnetic field pulse generator (TESLA Stym, Iskramedical, Slovenia) connected to an applicator consisted of round coil with 72 turns. The generator supplied the applicator with pulses of electric current that generated time-varying magnetic field around the coil, which in turn induced an electric field in the treated tissue (Figure 1).

In order to obtain precise application of electroporation mice were initially anesthetized with inhalation anaesthesia in the induction chamber with 2% (v/v) of isoflurane (Isoflurane; Piramal Healthcare UK Limited, London, UK) and afterwards the mouse muzzle was placed under inhalation tube to remain anesthetized during experiment. The applicator for electroporation was positioned over the tumor so that the tumor was in the middle of the applicator (Figure 1A).

In previous experiments, groups of positive controls, such as conventional electroporation (EP) and the combination of EP with CDDP (ECT), were obtained. The growth of untreated tumors (tumor doubling time (the time in which tumor reaches twice of the initial volume, DT) in control was $1.4 \pm 0.2$, $n = 6$) and of CDDP treated tumors alone ($2.3 \pm 0.1$, $n = 4$) in that independent experiment (data previously not published) were comparable.

Supplementary Figure 1), the most promising sequence of bipolar pulses (Supplementary Figure 1) was used in the combination with CDDP. Briefly, the sequence had 480 bipolar pulses with duration of $t_p = 340$ µs, with a peak of $I_p = 400$ A, repetition frequency ($f_p$) of 80 Hz and duration of each sequence ($t_s$) of 6 s (Figure 2). Electric pulses were measured using an oscilloscope (WavePro 7300A, LeCroy, Chestnut Ridge, NY) and current probe CWT Rogowski Current Transducer (Powertek, UK).

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to the experiment performed with PEMF treatment). In the conventional electrochemotherapy protocol three minutes after intravenously injection of CDDP eight square wave electric pulses at 1300 V/cm voltage to distance ratio, 100 µs long and 1 Hz (Cliniporator™, IGEA s.r.l., Carpi, Italy) were applied by plate electrodes (d = 8 mm) to the tumors. Electric pulses were delivered in perpendicular orientation (4 + 4) and good contact between the electrodes and tumor was assured using conductive gel.

**Determination of magnetic and electric field in the tumor**

Time-varying magnetic field and induced electric field of PEMF in the tumor was determined by means of numerical modelling. Numerical model of the applicator was modelled as multi-turn coil node which is a lumped model for tightly wound 72 wires separated by electrical insulator. Numerical model of the tumor was represented by an ellipsoid (Figure 1). Since volume of mice tumors varied from 30 to 40 mm³ an average volume, i.e. 35 mm³, was used in the numerical model of the tumor. Bipolar pulse (Figure 2) was used as electric current in the numerical model of the applicator. Calculations of time-varying magnetic field and induced electric field were performed using finite element method on a desktop PC (Windows 8.1, 3.50 GHz, 32 GB RAM) using commercial finite element software package COMSOL Multiphysics 5.1 (COMSOL AB, Stockholm, Sweden).

**Treatment evaluation**

The muscle contraction during PEMF treatment and conventional EP, tumor growth after therapy, skin area above the tumor and 2 cm in diameter around the tumor exposed to PEMF or conventional EP and the general well-being of animals (consumption of water and food, weight loss) were monitored during the experiment. Tumor growth was followed by measuring three mutually orthogonal tumor diameters (a, b, and c) with a Vernier caliper, every day. The tumor volumes were calculated by the formula:

\[ V = \frac{abc}{6} \]

The arithmetic mean of the tumor volumes and the standard error of the mean (SE) were calculated for each experimental group for each measurement day. The tumor growth delay was determined for each individual tumor by subtracting the average DT of the control group from the DT of each individual tumor.

**Platinum determination in the serum and tumors**

The measurements of platinum accumulation in the serum, tumors, platinum bound to the DNA in the cells and in extracellular fraction were performed by inductively coupled plasma mass spectrometry (ICP-MS, Agilent Technologies, model 7700x, Tokyo, Japan). 195Pt isotope was monitored. At optimized instrumental parameters, instrumental limit of detection (LOD) was 0.005 ng Pt/mL (3σ of the blanks). The linearity of the signal was confirmed from LOD to 10 µg Pt/mL. Repeatability of the measurements was better than 3%.

The platinum uptake in tumors and its total concentration in the serum were measured 1 hour after the treatment of mice with intravenously injection of CDDP, electroporation induced by PEMF or the combination of those therapies (PEMF + CDDP). The blood was collected with glass capillary from inorbital sinus (3–8 samples per group) and was coagulated at room temperature for two hours. Thereafter the blood was centrifuged at 3000 rpm for 10 minutes and serum was collected and stored at the temperature -20°C. On the day of measurements total amount of serum samples were digested in 1 mL of 1 : 1 mixture of 65% nitric acid (MERCK KgaA, Dermstadt, Germany) and 30% hydrogen peroxide (MERCK KgaA, Dermstadt, Germany) by incubation at 90°C for 48 hours. Obtained clear solutions were diluted with Milli-Q water before analysis.

For platinum determination in tumors, animals were sacrificed after the blood collection. The tumors (3–8 tumors per group) were excised and removed from the overlying skin. Each tumor was weighed, and placed into a 15 mL graduated polyethylene tube. For tumors digestion, the same procedure as for serum was applied, with the exception that 2 mL instead of 1 mL of 1 : 1 mixture of 65% nitric and 30% hydrogen peroxide was used. Before analysis samples were diluted with Milli-Q water.

**Determination of platinum bound to the DNA in the tumor cells and the extracellular fraction (fluid)**

The tumors were obtained as described in the chapter above, weighed and immediately mechani-
ically disintegrated. The sample was washed with 3 mL of freshly prepared PBS and filtered through the cell strainer with pore size of 40 µm (Corning Incorporated, Life Sciences, Durham, USA). The obtained cells in suspension were centrifuged at 1500 rpm for 10 minutes. Collected cells were used for the fast DNA isolation by salting-out protocol. Briefly, cells were lysed with lysis buffer (10 mM Tris-HCl, 1 mM ethylenediaminetetraacetic acid [EDTA], 1% sodium dodecyl sulfate [SDS]) with proteinase K (20 µg) for 30 minutes at 55°C by constant shaking. After the samples were cooled down proteins were precipitated by adding of 120 µL 4 M NaCl and shaken for 15 seconds. Precipitated proteins were centrifuged at 13000 rpm for 6 minutes. In addition, DNA was precipitated with 1 mL of ethanol (70%) for 2 min by gentle mixing of tube and centrifuged at 13000 rpm for 2 minutes. Precipitated DNA was washed with additional 1 mL of ethanol (70%) and centrifuged at 13000 rpm for 2 minutes. The pellet of DNA was dried out, resuspended in 100 µL of distilled water, digested under the same procedure as serum and the concentration determined in diluted samples by ICP-MS.

The rest of two fractions, supernatant and the interstitial fraction on the top of the cell strainer, named as extracellular fraction, were collected and stored at -20°C till the digestion with the mixture of nitric acid and hydrogen peroxide (see the section above).

### Statistical analysis

All data were tested for normal distribution with the Shapiro–Wilk test. A t-test and one-way analysis of variance followed by a Holm–Sidak test were used for evaluation of the differences between the experimental groups. A p value less than 0.05 was considered significant. SigmaPlot Software (Systat Software, Chicago, IL, USA) was used for statistical analysis and graphical representation.

### Results

#### Antitumor effectives of electrochemotherapy mediated by PEMF

Exposure of tumors to PEMF or conventional EP, performed 3 minutes after intravenous injection of CDDP, resulted in significant tumor growth delay, up to 3 days compared to untreated tumors, as well as compared to monotherapies. Nevertheless significantly higher antitumor effect was obtained after conventional electrochemotherapy compared to electrochemotherapy mediated by PEMF. Treatment of tumors with CDDP alone or exposure to PEMF or conventional EP had no significant effect on tumor growth (Table 1, Figure 3).

CDDP = intravenously injection of cisplatin (4 mg/kg); PEMF = pulsed electromagnetic field treatment; PEMF + CDDP = PEMF after intravenously injection of CDDP; EP = electric pulses treatment; ECT = electrochemotherapy, EP after intravenously injection of CDDP; DT = tumor doubling time; GD = tumor growth delay; p < 0.05 statistically significant difference; Data pooled from separate experiments after checking that DT in control and CDDP treatment alone were comparable.

<table>
<thead>
<tr>
<th>Group</th>
<th>n</th>
<th>DT (Mean ± SE)</th>
<th>GD</th>
<th>P (&lt;0.05)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control*</td>
<td>12</td>
<td>1.5 ± 0.1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>CDDP* 4 mg/kg</td>
<td>12</td>
<td>2.2 ± 0.2</td>
<td>0.7</td>
<td></td>
</tr>
<tr>
<td>PEMF</td>
<td>9</td>
<td>1.9 ± 0.1</td>
<td>0.4</td>
<td></td>
</tr>
<tr>
<td>PEMF + CDDP</td>
<td>10</td>
<td>3.8 ± 0.1</td>
<td>2.3</td>
<td>&lt;0.001 (to PEMF)</td>
</tr>
<tr>
<td>EP*</td>
<td>12</td>
<td>2.2 ± 0.3</td>
<td>0.7</td>
<td></td>
</tr>
<tr>
<td>ECT CDDP*</td>
<td>8</td>
<td>4.5 ± 0.2</td>
<td>3.0</td>
<td>&lt;0.009 (to PEMF + CDDP)</td>
</tr>
</tbody>
</table>

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**FIGURE 3.** Antitumor effectiveness of electrochemotherapy with CDDP mediated by PEMF in mouse melanoma B16F10. Data were collected from two individual experiments and each point on graph represents mean and standard error of the mean (AM ± SE). Each group consisted at least of 8 animals.

CDDP = intravenously injection of cisplatin (4 mg/kg); ECT = electrochemotherapy; EP = electric pulses treatment; PEMF = pulsed electromagnetic field treatment; PEMF + CDDP = PEMF after intravenously injection of CDDP; EP = electric pulses treatment; PEMF = pulsed electromagnetic field treatment; PEMF + CDDP = PEMF after intravenously injection of CDDP; DT = tumor doubling time; GD = tumor growth delay; p < 0.05 statistically significant difference; Data pooled from separate experiments after checking that DT in control and CDDP treatment alone were comparable.
Overall, the antitumor effect CDDP, i.e. the chemotherapeutic drug used, in the electrochemotherapy performed by PEMF was significantly increased in comparison to monotherapies, presumably due to facilitated transport of CDDP. However, electrochemotherapy after conventional EP was more effective than PEMF mediated electroporation.

**Determination of platinum in the serum and tumor after electroporation induced by PEMF**

In order to determine whether electroporation induced by PEMF facilitates drug delivery into the cells, as the underlying antitumor mechanism, Pt accumulation in the serum and tumors with plasma mass spectrometry was determined. First, Pt was measured as indicator of CDDP in whole tumors and plasma of the blood in mice. Intravenous CDDP injection demonstrated the drug accumulation in tumors, and electroporation induced by PEMF as successful method for increasing Pt accumulation in whole tumors, one hour after the drug administration (Figure 4). A statistically significant increase in platinum content in the tumors treated by electroporation induced by PEMF was observed, however this measurement does not indicate whether the electroporation induced by PEMF in fact facilitated drug delivery into the cells.

To prove that electroporation induced by PEMF facilitates transmembrane transport of CDDP, the extracellular and intracellular Pt amounts were measured. After mechanically disintegration of tumors, the suspension of tumor cells and extracellular fractions were obtained, and Pt was measured in both fractions. The concentration of the Pt detected in extracellular fraction statistically significantly decreased after electroporation induced by PEMF (Figure 5). Moreover, the Pt bound to DNA as indicator of the drug bound to the intracellular target was significantly increased after the electroporation induced by PEMF (Figure 5). Approximately 2 times higher values of Pt bound to DNA were obtained.

**Estimation of PEMF in the tumor**

Simulation results of PEMF in the tumor are presented in Figure 6. Electric field distribution in the tumor was linearly decreasing from the boundary of the tumor towards the center with a peak value of 8.6 V/m on the tumor boundary. Magnetic flux density had a peak value of 0.3 T and its distribu-
Discussion

This study demonstrated the use of contactless pulsed electromagnetic field (PEMF) treatment as an approach to achieve electroporation of melanoma tumor tissue, which increases drug uptake in vivo. We evaluated for the first time the antitumor effectiveness of electrochemotherapy obtained by PEMF after systemic injection of chemotherapeutic drug, CDDP, which is used in conventional electroporation protocol. Furthermore, we proved that the antitumor effect was related to increased drug uptake into the electroporated tumor cells, demonstrated by increased amount of Pt bound to the DNA. Thus PEMF treatment can be used (once optimized) for noninvasive drug delivery in vivo, which may be important for research where delicate tissues and organs needs to be avoided and for clinical applications, since it is noninvasive, contactless and painless compared to the classical electroporation using different electrodes.

Potential use of strong time-varying magnetic field which induced electric field to increase transmembrane molecular transport (i.e. electroporation), was already suggested before. In order to expand the use of contactless PEMF induced electroporation as drug delivery method the melanoma B16F10 tumor model in vivo was chosen, as it represents a great challenge in the treatment of human melanoma. We have shown that magnetic field generated by round coil which induced the 480 bipolar pulses, at frequency of 80 Hz, pulse duration of 340 µs, significantly improved the antitumor effectiveness of electrochemotherapy with CDDP. Our results are in accordance with in vitro study, where much stronger PEMF (6.1 T) was indicated as delivery method for therapeutic molecules in human pathogenic fungi. Namely, the synergistic effect of simultaneous treatment of 200 applied magnetic field pulses at frequency of 35 Hz and drug was observed. However, the application of stronger magnetic field (up to 16.4 T) did not result in better membrane permeabilization. The membrane permeabilization seems to be dependent more on the shape, number and frequency of generated pulses in addition to the amplitude of induced electric field, similar to the conventionally generated bipolar pulses. In addition, bipolar pulses were demonstrated two times more effective as monopolar and almost equally effective as conventional square wave pulses. In fact, for the almost equal membrane permeabilization larger number of short duration induced bipolar electric pulses at higher frequency has to be delivered. Similarly, in our study the obtained antitumor effectiveness of cisplatin by using short and larger number of induced electric pulses at higher frequency in comparison to conventional electric pulses was comparable. Furthermore, it is known that simple round coils induce less focused and lower peak electric field than figure-of-eight coils. It was also demonstrated in vitro that by using figure-of-eight coils the increased transmembrane molecular transport could be obtained by pulses of lower frequencies and larger number. However, we and others have shown that by increasing the number of pulses at the same frequency the effect of electroporation can be improved (supplementary data).

Presently, in conventional electrochemotherapy protocols mainly square wave or monotonically decreasing electric pulses are delivered through plate or needle electrodes to the cells or tissues. On the contrary, only a few studies were performed with bipolar electric pulses for the electroporation of cells in vitro and tissues in vivo. In general, the lower pulse amplitudes were needed for effective electroporation of cells in vitro with respect to unipolar pulses. Besides that, the pulse shape played important role in electroporation of cells as well. It was shown that electroporation, cell death and the uptake of Lucifer Yellow occurred by using the rectangular bipolar pulses at lowest, the sine bipolar pulses at medium and the triangular bipolar pulses at the highest pulses amplitudes. Bipolar pulses were already applied successfully in electrochemotherapy for human and veterinary clinic.

In fact, the combination of PEMF induced electroporation with CDDP had significant antitumor effectiveness, whereas the application of PEMF or the drug alone had none. The antitumor effectiveness of electrochemotherapy of applied PEMF (480 bipolar pulses, at frequency of 80 Hz, pulse duration of 340 µs) was presumably due to improved membrane permeabilization of cells in the tissue, since the monotherapies alone had very little but no significant effect on tumor growth in comparison to control. Similarly, preclinical and clinical studies performed in conventional scheme of electrochemotherapy with CDDP have shown great antitumor effectiveness of electrochemotherapy on different tumor types, mostly due to direct cytotoxic effect on tumor cells. It has been demonstrated...
that after conventional electroporation the cytotoxicity of CDDP could be improved by 70-times. Nevertheless, in our study sufficient antitumor effectiveness on melanoma B16F10 was obtained with electrochemotherapy after PEMF induced electroporation, despite the effect was significantly lower compared to that obtained after conventional electrochemotherapy with CDDP. However, calculations suggest that levels of electric field are 4 orders of magnitude lower than those associated to classical electroporation, but tend to be high enough to induce electroporation. On the other side, while using electric field bellow 0.09 V/cm at the position of tumor site, we suspect there was no possible occurrence of irreversible electroporation or thermal effect, as obtained at much higher electric field induced by PEMF (up to 40 V/cm). Even more, we speculate that PEMF could be improved by positioning of tumor towards the edge of the coil, where based on calculations the highest strength of magnetic field could be obtained, which consequently could induce higher electric field strength and thus, even more cells could be successfully electroporated at deeper parts of tumor tissue. Therefore, further studies to optimize the PEMF are warranted.

Even though it was previously known that the time varying magnetic field could induce electroporation, affect the cytoskeleton and intracellular signal transduction, the mechanisms of its action in the combination with CDDP have not been studied yet. Therefore, to clarify the antitumor effectiveness of electrochemotherapy the measurements of platinum amount after electroporation induced by PEMF were performed. Observed significant 1.6-fold increased platinum uptake into melanoma B16F10 tumors after electroporation induced by PEMF indirectly confirmed membrane permeabilization of the tumor cells and thus, its correlation with the antitumor effectiveness of electrochemotherapy. Our results are in accordance with results reported in other studies, where up to two times higher platinum uptake was obtained in sarcoma SA-1 and fibrosarcoma LPB tumors after conventional electrochemotherapy with CDDP. Even though lower increase of platinum amount in tumors was obtained after PEMF in comparison to conventional electrochemotherapy, the final amount of platinum in the tumors was comparable. The difference of platinum amount in tumors treated only with CDDP, might be tumor type dependent, since melanoma tumors are well vascularized and contained large spherical cells, with less surrounding extracellular matrix component in comparison to stiff SA-1 and LPB tumors with small spindle-shaped cells and high content of extracellular matrix component. On the other hand, the amount of platinum in the serum was significantly up to 6-times lower compared to tumor tissue. Therefore we could assume that excess of the drug which was not entrapped in the tumors after electroporation was washed out with similar kinetic as in nonelectroporated tumors. Moreover, our results indicated that CDDP in the cells reached its main intracellular target DNA, in fact significantly two times higher amount of Pt was bounded to the DNA in PEMF and CDDP treated tumors than in CDDP only treated tumors. At the same time, as expected the pool of Pt amount in the extracellular fraction of these tumors was lowered, up to 1.4-times. Thus, the increased Pt uptake in the cells and its binding to the DNA could indicate the main reason for antitumor effectiveness of electrochemotherapy mediated by PEMF.

Presently, it is not clear if the membrane permeabilization obtained after application of time varying magnetic field occurs only due to induced electric field as in conventional electroporation or due to direct effect of magnetic field with the plasma membrane and surrounding ions. Thus, the precise mechanism of cisplatin uptake after electroporation mediated by PEMF into the cells remains unclear. Obtained Pt amount in the tumor cells after treatment with CDDP only could be ascribed to passive diffusion and active transport mechanisms of cisplatin through the membrane, which are carrier-mediated through formed pores or via endocytosis. In addition, it has been demonstrated that exposure of cells to train of unipolar pulsed low electric fields at strength from 1.2 up to 20 V/cm can induce endo-endocytosis. Thus, we suspect that generated bipolar electric field of just below 0.09 V/cm by PEMF might also trigger endocytosis besides membrane permeabilization which enables the internalization of cisplatin in the cell and contributes partially to the increase of platinum amount.

In conclusion, our results show that PEMF at magnetic field below of 1 T was sufficient to achieve membrane permeabilization of tumor cells, thus, small molecules such as drug (CDDP) improved delivery and cellular uptake in solid tumors was enabled. Due to simple, contactless, painless, focused local application of PEMF, better field distribution irrespective of tissue type and thus, achieving electric field strength for membrane permeabilization can be established in deeper parts of tissue. However this approach has the limitation that the strength of the magnetic field decreases rapidly.
with distance from the coil which has to be taken into account by designing coils in order to achieve successful permeabilization at a greater tissue depth. PEMF might thus represent an alternative to conventional electroporation with electric fields in electrochemotherapy. However further studies are needed to improve the equipment, to optimize and establish precise protocols of drug application and PEMF parameters, as well as to reveal the effects of PEMF on variety of normal and tumor tissues.

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