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Cellular level studies and coil system design for transcranial magnetic stimulation

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Cellular level studies and coil system design for transcranial magnetic stimulation

by

Yiwen Meng

A thesis submitted to the graduate faculty
in partial fulfillment of the requirements for the degree of

MASTER OF SCIENCE

Major: Electrical Engineering

Program of Study Committee:
David.C.Jiles, Co-Major Professor
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Iowa State University
Ames, Iowa
2015

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DEDICATION

I would like to dedicate this thesis to my mother Li and my father Yulin for their unconditional and tremendous love and support for me to overcome any difficulty I met.
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ABSTRACT

Transcranial Magnetic Stimulation is a novel non-invasive neuromodulation technique to treat human brain disorders such as depression, Parkinson’s disease and PTSD. It uses pulsed currents in the coils to generate time varying magnetic field which induce eddy currents in the conductive tissues of the human brain. Recently, there have been many research publications in the field of TMS, specifically on coil designs, clinical trials and some in-vivo animal studies.

Even though FDA has approved TMS technique to treat depression, the basic mechanism or how the neural tissue reacts to TMS is still not well understood. Therefore, conducting in-vitro study on TMS will enable researchers to understand how TMS has influence on neural cells and neural tissue growth rate, morphology, axon length and other factors. In this work, I have conducted experiments on effect of TMS on N27 dopaminergic neural cells, an immortal cell line of rat, to investigate the effect on cell’s growth rate. Results will enable neuroscientists to understand the mechanism of TMS on neural cells.

As a part of TMS project, I have also worked on the development of a TMS helmet design. Due to the limitation of patient’s head size and rapid decay rate of magnetic field away from coil surface, designing an efficient and compact coil system is needed to treat deep brain regions. We have developed a variable coil system with combination of fixed single coil on top and variable Halo coil to realize deep brain stimulation with automatic control system and graphic user interface (GUI). In the meantime, I also conducted thermal and mechanical analysis of new coil configuration to investigate heating effect and electromagnetic force on the whole coil system. This system can be used by researchers or
clinicians with relative ease, maintaining the accuracy of coil position relative to the patients head.
CHAPTER 1. GENERAL INTRODUCTION

Transcranial Magnetic Stimulation (TMS)

Transcranial Magnetic Stimulation (TMS) is a non-invasive neuromodulation technique which has potential to treat various neurological disorders such as major depressive disorder, Parkinson’s disease, Post-traumatic stress disorder (PTSD) and migraine non-invasively and safely. It uses short pulses of time varying magnetic field to induce an electric field in the conductive tissues of the brain, hence, modulating the synaptic transmission of neurons. This neuromodulation technique can be used to excite or inhibit the firing rate of neurons by influencing the ion activities inside and outside of neuron’s plasma membrane [1-5].

Fig.1 The illustration of TMS treatment on human brain (from Laboratory for cognition and neural stimulation in school of medicine at University of Pennsylvania)

Fig.1 illustrates the basic mechanism of TMS and how it affects brain behavior of human. TMS uses different types of coils such as single coil, double coil, Halo coil and Helmholtz coil to generate different types of magnetic field in the human brain. In clinical trials, physicians put
certain type of coil according to the specific region of human brain like hippocampus and motor cortex to treat specific brain disorders [5].

**Research Motivation**

Since the US Food and Drug Administration (FDA) approved TMS as a treatment for depression in 2008, there has been an increasing research interest on TMS. As shown in Fig.2, the major fields are computer modeling and coil design, in-vitro and in-vivo studies and clinical trials [6-7]. More importantly, understanding the mechanism of TMS on brain or how TMS affects individual neurons or neural tissues would bring a big breakthrough to the current theory.

![Fig. 2 Illustration of research areas on TMS](image)

Therefore, doing in-vitro study on TMS is a good approach to investigate effect of TMS on several parameters like growth rate, soma size of individual neurons or neural networks. These kinds of work would be addressed into the study on growth rate, morphology and protein analysis of neurons. The motor symptoms of Parkinson's disease result from the death of dopamine generating cells in the midbrain region, substantial nigra.1RB3AN27 cell line is the immortalized dopamine neural cells from rat brain. This immortalized cell line has been carefully
characterized in studies of dopamine biosynthesis, neurotoxicity and used as a dopaminergic neuron model for in vitro and in vivo studies. Therefore, there have been numerous efforts to understand the basic mechanism of the degenerative process of dopaminergic neurons and to realize neural genesis in substantia nigra to cure Parkinson’s disease [8].

Besides, another aspect of research on TMS is designing efficient, compact coils or a coil system to treat different regions of the human brain especially the deep-lying regions. Thus, overcoming the fast decay rate of the magnetic field to induce enough electromagnetic fields in deep brain regions is one big challenge right now. Meanwhile, building a compact or even an automatic TMS coil system will make it easier for doctors to conduct TMS treatment in clinical trials.

**Thesis Organization**

Chapter 2 and Chapter 3 describe two parts of my work on TMS, focusing on in-vitro and coil design of TMS, respectively. Chapter 2 mainly shows the effect on different orientation of magnetic field on the proliferation rate of dopaminergic neurons with three different cell counting methods. It also gives a literature review of the beneficial effect of static and time varying electromagnetic field with different frequency ranges. Chapter 3 mainly describes work on computer modeling of thermal and mechanical analysis of variable TMS coil system and an illustration of graphic user interface (GUI) of the coil system.

Chapter 4 summarizes all of work in this thesis and some recommendations for future work on TMS like morphology study and protein analysis of neurons to understand the mechanism of TMS. References can be found at the end of each chapter. Appendix A lists all of my journal and conference publications during my master’s degree. Appendix B shows the results of computer modeling of electromagnetic field for the variable TMS coil system.
References


CHAPTER 2. DIFFERENTIAL EFFET OF MAGNETIC FIELD ORIENTATION ON THE PROLIFERATION RATE OF DOPARMINERGIC NEURONS DURING TRANSCRANIAL MAGNETIC STIMULATION

Modified from a paper submitted to *Neuroscience of Elsevier*

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Abstract

Transcranial magnetic stimulation (TMS) has been used to investigate possible treatments for a variety of neurological disorders. But the effect that magnetic fields have on neurons has not been well documented in the literature. Using a monophasic stimulator, we investigated the effect of different orientation of magnetic field generated by TMS coils on the proliferation rate of N27 neuronal cells cultured in flasks and multi-well plates. Exposing horizontally adherent N27 cells to a magnetic field pointing upward through the neuronal proliferation layer increased the proliferation of cells compared with the control group. On the other hand, proliferation rate decreased in cells exposed to a magnetic field pointing downward through the neuronal growth layer compared with the control group. The results were consistent across different methods of measuring proliferation and cell counting procedures. We confirmed results obtained from the Trypan-blue and automatic cell counting methods with those from the CyQuant and MTS cell viability assays. Our findings could have important implications for the preclinical development of TMS treatments of neurological disorders and represents a new method to control the proliferation rate of neuronal cells.

Key words: TMS; dopaminergic neurons; proliferation rate; orientation of magnetic field
Introduction

Transcranial Magnetic Stimulation is a non-invasive neuromodulation technique that uses time varying short pulses of magnetic field to induce an electric field in the conductive tissues of the brain thus, modulating the synaptic transmission of neurons. This neuromodulation technique can be used to excite or inhibit the firing rate of neurons which can then be used for treatment of various neurological disorders such as major depressive disorder, Parkinson's disease, Post-traumatic stress disorder and migraine [1-5]. Since the US Food and Drug Administration (FDA) approved TMS as a treatment for depression in 2008, there has been less focus on \textit{in vitro} and animal studies in the literature compared to \textit{in vivo} studies in humans [6-8]. The effects of TMS on individual neurons need to be thoroughly understood to fully utilize TMS as a neuromodulation tool for treating neurological disorders especially those originating from subcortical regions of the brain.

Few articles have reported the effect of time-varying magnetic fields, similar to those generated by TMS, on the proliferation rates of neurons. Bonmassar et al. designed micro TMS coils and showed that the direction of magnetic field affects the firing frequency of neurons, but the authors did not report the effect of magnetic field on the proliferation rate [9]. Meanwhile, some articles have reported the effect of static magnetic field on cell’s proliferation rate. Authors have used static magnetic fields from 1 to 10 tesla and did not find any significant effect on cell proliferation or on genetic toxicity, regardless of the length of treatment. However, there was a small effect on intracellular Ca\textsuperscript{2+} ion control [10]. Some articles have reported beneficial effects of DC electric field (EF) on neural proliferation and differentiation. The EF gradient affects morphology and phenotype of adult neural stem/progenitor cells (NPCs), which shows the potential of utilizing EF to control migration,
differentiation and alignment of stem cells transplanted to treat nervous system disorders [11]. Extremely low-frequency electromagnetic fields (ELF-EMFs) have been used therapeutically to drive cardiac-specific differentiation in adult human cardiac progenitor cells without any pharmacological or genetic manipulation of cells [12]. As far as we know, no one has published on the effect of TMS magnetic field on the proliferation rate of neurons or on the morphology of cells.

In this paper, we have presented the effect of magnetic field generated by TMS coils on the proliferation of N27 dopaminergic neurons. We have used different cell proliferation and cell counting procedures to confirm that directing a magnetic field downward or upward through the horizontal proliferation plane of adherent cell cultures decreased or increased cell proliferation rates, respectively. It is important to note that the direction of the induced electric current from the time varying TMS fields will be in clockwise or counterclockwise loops when the magnetic field is in up or down direction of the cell culture as shown in Fig. 4. This experimental set up is similar to the TMS treatment on human brain where the induced electric field from the TMS coils will be in clockwise or counterclockwise loops in the cortex.

**Experimental procedures**

*A. Magnetic Field Generated by TMS coils*
A Magstim Standard 70 mm double coil was used for treating N27 neurons. Magnetic field was measured on the surface of the coil using a gaussmeter and a Hall probe. The field was also calculated using finite element electromagnetic modeling software, SEMCAD X. The measured and calculated axial components of the magnetic field intensities are shown in Fig.3 and Fig.4. Magnetic field is negative in the negative x-axis and positive in the positive x-axis which is shown Fig.4. It also shows magnetic field values at 5mm above the coil surface where dopaminergic neurons are placed during TMS treatment after considering the thickness of flask and thermal insulation layer. According to these figures, the peak value of measured magnetic field intensity at 5mm above the coil surface is 0.55 MA/m which is reduced by approximately 0.1MA/m. Fig. 5 shows the top view of distribution of magnetic field intensity generated by double coil. Fig.6 shows the different orientations of magnetic field generated by the coil and directions of current in each circle of the double coil. The red
arrows on the left indicate the directions of supplied current (5000 A) in left circle as counterclockwise and clockwise in the right circle. The cross symbols indicate the magnetic flux pointing into the plane and the dot symbols indicate the magnetic flux pointing out of the plane.

![Fig. 5 Distribution of magnetic field intensity of double coil (2D top view).](image)

![Fig. 6 Orientation of magnetic flux lines generated by double coil, cross representing upward and dot representing downward field. The red arrows show the direction of supplied current with a peak magnitude of 5000 A. The two blue polygons represent the two flasks.](image)

According to Maxwell’s equation \( \nabla \times \mathbf{E} = -\frac{\partial \mathbf{B}}{\partial t} \), time-varying magnetic field will generate an electric field which induces eddy currents in the conducting neurons. The supplied current is a pulse wave which has a frequency of 2.5 kHz and magnitude of 5000 A, so its period is 0.4 ms. The stimulator sends only one pulse with a current amplitude of 5000 A.
A in clockwise and counterclockwise directions in left and right circles of the coil respectively as shown in Fig. 6. Thus, the supplied current in each circular coil will generate a time varying magnetic field changing from 0 to its peak value, during its first half period, which results in the corresponding induced eddy current in both areas shown in Fig. 7 and Fig. 8. According to Lenz’s law the induced current in the left circular coil was counterclockwise and it was clockwise in the right circular coil. Similarly, the value of the supplied current in both coils would change from its peak of 5000 A to 0 during the second half period. Thus, the induced eddy currents on the left and right flasks were clockwise and counterclockwise, respectively. Therefore, the difference between the two flasks was the sequence of the direction of the eddy currents.

B. Cell Culture

Immortalized rat mesencephalic 1RB3AN27 cells (N27) were grown in RPMI-1640 medium supplemented with 10% fetal bovine serum, 1% L-glutamine, 50 units penicillin and 50 µg/ml streptomycin and maintained at 37°C with a humidified atmosphere containing 5% CO₂, as described previously [13,14]. On Day 0, an equal number of N27 cells were seeded into each T-75 flask or 96-well plate. Groups were distinguished by culture time with two control and two TMS groups per time point and four replicate samples (flask or plate) per group. Control 1 was always kept in the incubator and was named Incubator in Table I. Control 2 was kept in the biosafety cabinet during the TMS treatment and was named Environmental in Table I. Table I shows culture time points and counting time points for the different sample locations and magnetic field orientations (“Field up” and “Field down”), which were used in a Trypan blue cytotoxicity assay. Table II shows cell culture samples with their culture time points as well as counting time points used in a CyQuant cell proliferation assay. Table III shows culture samples with their culture and counting time
points for an MTS cell viability assay and cell counting method. We performed a cell count for each sample of cells 24 hours after its TMS treatment to ensure that the cells had enough time to show any effects of TMS on their proliferation.

C. TMS experiment on dopaminergic neurons

We used a monophasic stimulator to treat N27 cell cultures. A set of 6 pulses with 4 seconds waiting time in between them was formed as one train and a waiting time of 10 seconds between each train was introduced, so the pulse repetition rate (TMS treatment frequency) we used is 0.25 Hz. This is a low frequency compared to usual clinical protocol frequency however, in order to obtain 100% power in the coil and avoid rapid heating up of the coil we have used this low frequency. It is not possible to operate at higher frequencies at full power with the existing set-up. A total of 60 trains with 360 pulses were delivered per 30-minute TMS treatment. An air-cooled double coil was used which has opposite current directions in each coil, generating magnetic fields on top side of each coil with opposite directions. Using air-cooled coils allowed us to induce magnetic fields without raising the temperature of the T-75 flasks placed on them. All TMS treatments on N27 cells were performed in a sterile biosafety cabinet (Fig. 9). The flask set above the left coil was designated “Field up” and the flask set above the right coil was designated “Field down”, corresponding to the orientation of the magnetic fields.
Fig. 9 Arrangement of the 30-minute TMS treatment delivered to two T-75 flasks. The directions of the two oppositely oriented magnetic fields were labeled on the double coil. Field orientation was upward on the left coil and downward on the right coil, as shown in the inset figure. We used two clamps to fix the coil in the cell culture cabinet and a layer of bubble wrap separated both flasks from the coils to maintain a thermal barrier.

D. Cytotoxicity assay

Cytotoxic cell death was measured as per Life Technologies’ Trypan blue exclusion cell counting method [15]. Briefly, after treatment cells were harvested with trypsin-EDTA and resuspended in 1X PBS, we then took 10 µl of cell suspension from one sample and added with 10 µl of 0.4% trypan blue solution (Life Technologies). Then, we put 10 µl of the mixture into the cell counting slide and place the slide into the automatic cell counter to count the concentration of cells in each sample. Finally, we extrapolated the total number of cells in each sample by multiplying its volume and concentration [15]. By using this method, we counted the number of cells in each of the four replicate samples according to the counting time points shown in Table. I. We studied the effect of TMS on 2 different initial cell densities, 1 million cells/flask and 0.5 million cells/flask (n=4).
E. CyQuant cell proliferation assay

We used Life technologies’ CyQuant cell viability assay to confirm our results from the Trypan blue cell counting procedures. On day 0, we seeded the N27 cells in 24-well plates (n=3) with the four rows per plate. In each plate, there were 4 rows and three columns. Row 1 to row 4 have different seeding densities; 20k, 50k, 80k, to 100k, respectively. Each row had three replicated samples in three columns to account for standard deviation. We had three groups with different culture times: Day 0, Day 1 and Day 2 (Table II). Briefly, after 24
hours post-treatment, we read the fluorescence with excitation maximum at 485 nm and the emission maximum at 530 nm using a Synergy 2 plate reader (BioTek) [16]. We pooled the groups designated as Incubator and Environmental in Table I, because the difference between them was insignificant.

F. MTS cell viability assay cell counting method

Cell viability was measured using Promega’s MTS assay to confirm the results from Trypan blue and CyQuant cell proliferation assays. Briefly, on day 0, we seeded the wells of 96-well plates with 15k for one row and half with 20k for another row of N27 cells in 200 µL of proliferation medium per well. Each row had 6 duplicated samples (n=6). The design of the experiment was according to Table III., TMS treatment was performed with “Field up” and “Field down” on 2 different well plates. After 24 hours post-treatment, 20µl MTS reagent (CellTiter 96® Aqueous One Solution Reagent) was added to each well and incubated at 37°C in a CO2 incubator for 90 min and absorbance was read at 490 nm and 670 nm in a Spectramax plate reader (Molecular Devices). We subtracted the baseline via Abs490-Abs670 prior to data analysis [17].

Table 3. Design of the TMS experiment with MTS cell viability assay cell counting method for cell densities of 15k and 20k per well.

<table>
<thead>
<tr>
<th>Culture Time Point</th>
<th>Counting Time Point</th>
<th>Sample Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Day 0</td>
<td>Day 1</td>
<td>Control Environmental Field up Field down</td>
</tr>
</tbody>
</table>

G. Statistical significance analysis

Statistical significance analysis was performed using Originlab 9.0 software (OriginLab Corporation, Northampton, MA, USA). Raw data analysis were analyzed using a
two unpaired t-test. Statistically significant differences are indicated by asterisks as follows: *p<0.05, **p<0.01 and ***p<0.001.

Results

After TMS treatment of N27 cells, we counted the number of viable cells using the Trypan blue method for initial seeding densities of 1 million (Fig. 10) and 0.5 million (Fig. 11) cells per flask. The culture time and counting time points are indicated in Table I. The results showed that the proliferation rate increased after TMS stimulation with the magnetic field oriented upward through the horizontal plane of adherent cells, compared to incubator and environmental samples. The proliferation rate decreased when the field was oriented downward through the horizontal growth plane compared to incubator and environmental samples. Also, environmental samples exhibited slower proliferation compared to the incubator condition. For the lower seeding density (Fig. 11), the difference of cell counting for each group became larger over time. The difference peaked on Day 3 when the number of cells in the “Field up” group was 23.57 ± 3.21% (mean ± STD, ***p<0.001) higher than that in the Environmental group, while in the Field down group, it was 11.45 ± 1.99% (**p<0.001) lower than in the Environmental group. Therefore, the total difference in cell’s proliferation rate attributable to TMS field direction was +35.02%.
To investigate the effect of different culture times on cell proliferation, we conducted another experiment expanding culture time from 2 days to 2.5 days. The seeding density was 0.5 million/flask and the new culture time points were Day 0.5, Day 1, Day 1.5, Day 2 and Day 2.5. Cells were counted 24 h after each treatment, so the corresponding counting time points were Day 1.5, Day 2, Day 2.5, Day 3 and Day 3.5 respectively. The effect of TMS and its direction on the proliferation of cells over time (Fig. 12) was similar to the previous results for this seeding density (Fig. 11). On Day 3.5, the number of cells in the “Field up” group was 13.53 ± 1.36% (**p<0.001) higher than in the Environmental group, whereas the number of cells in the “Field down” group was 12.61 ± 1.76% (**p<0.001) lower than in the Environmental group. Therefore, the total difference in cell’s proliferation rate attributable to TMS field direction was +26.14 %.
We used the CyQuant cell viability assay to confirm the results obtained with the Trypan blue cell counting method. This time we eliminated group 1 (Incubator) and we set four seeding densities. The design of this experiment was based on Table II. The effect of TMS field direction on cell proliferation obtained via the CyQuant method (Fig 13-16) was similar to the effect measured using the Trypan blue cell counting method. However, Fig.8, which had the seeding density of 100k per well did not follow the trend similar to other seeding densities i.e. the difference in the proliferation rate was not pronounced. It may be due to the fact that a large number of cells grew in the limited space so the cells might have attained 100% confluency earlier than Day 3.
Fig. 13 Cell proliferation after TMS treatment of plates with an initial seeding density of 20k/well. We used the CyQuant cell viability assay to count cells. On day 3, the number of cells in the “Field up” group was 22.06 ± 4.14% (mean ± STD as a percentage of the initial seeding density, *p<0.05) higher than in the Environmental group, whereas cell numbers in the “Field down” group were 28.77± 1.00% (**p<0.01) lower than in the Environmental group.

Fig. 14 Cell proliferation after TMS treatment of plates with an initial seeding density of 50k/well. We used the CyQuant cell viability assay to count cells. On day 3, the number of cells in the “Field up” group was 15.49 ± 7.26% (mean ± STD as a percentage of the initial seeding density, *p<0.05) higher than in the Environmental group, whereas cell numbers in the “Field down” group were 9.94 ± 2.47% (**p<0.05) lower than in the Environmental group.

Fig. 15 Cell proliferation after TMS treatment of plates with an initial seeding density of 80k/well. We used the CyQuant cell viability assay to count cells. On day 3, the number of cells in the “Field up” group was 15.57 ± 5.17% (mean ± STD as a percentage of the initial seeding density, *p<0.05) higher than in the Environmental group, whereas cell numbers in the “Field down” group were 11.62 ± 1.55% (**p<0.01) lower than in the Environmental group.

Fig. 16 Cell proliferation after TMS treatment of plates with an initial seeding density of 100k/well. We used the CyQuant cell viability assay to count cells. On day 3, the number of cells in the “Field up” group was 14.69 ± 5.74% (mean ± STD as a percentage of the initial seeding density, p>0.05) higher than in the Environmental group, whereas cell numbers in the “Field down” group were the same (0 ± 4.50%, p>0.05) as those in the Environmental group.
A third cell counting method, the MTS cell viability assay, was performed to confirm the results obtained with the Trypan blue and CyQuant cell counting methods. With an initial seeding of 15k (Fig. 17), the number of cells in the “Field up” group was 19.88 ± 4.56% (**p<0.001) higher than in the Environmental control group. Meanwhile, the number of cells in the “Field down” group was 8.88 ± 1.39% (**p<0.01) lower than in the Environmental group. Next, using an initial seeding of 20k (Fig. 18), the number of cells in the “Field up” group was 19.60 ± 4.57% (**p<0.01) higher than in the Environmental group, while the number of cells in the “Field down” group was 8.16 ± 0.09% (**p<0.01) lower than in the Environmental group. Therefore, the total difference in cell’s proliferation rate attributable to TMS field direction was +27.76%.

![Fig. 17 Cell proliferation after TMS treatment of plates with an initial seeding density of 15k/well. We used the MTS cell viability assay to count cells, reported here as a percentage of control group1 (Incubator).](image1)

![Fig. 18 Cell proliferation after TMS treatment of plates with an initial seeding density of 20k/well. We used the MTS cell viability assay to count cells, reported here as a percentage of control group1 (Incubator).](image2)

**Discussion**

We investigated the effect of magnetic field orientation on the proliferation rate of N27 dopaminergic neuronal cells using three different cell counting methods to cross-validate
the results. The MTS assay showed the highest difference in cell proliferation rate. It was also easy to replicate this counting procedure three times to obtain standard deviation. In the Trypan blue cell counting method, we used flasks to culture neuronal cells, which required more area to incubate replicate samples. Cell counting using the Trypan blue method was more time consuming because cell counting was performed one flask at a time, unlike the MTS method where cell counting was performed in groups. There were three replicate samples for each group (n=3) for Trypan blue method and for MTS and CyQuant cell viability assay, n=6, which is adequate to show statistical significance. In the CyQuant cell viability cell counting method, it was easy to replicate samples and we were also able to count cells of all groups at once, but the differences among groups were slightly smaller than those from the MTS cell counting method. Thus, the MTS cell viability assay cell counting method is recommended for investigating the proliferation rate of N27 dopaminergic neuronal cells under TMS treatment.

According to the design of all these experiments, each group of N27 dopaminergic neuronal cells received a 30 minutes TMS treatment each day. After experimenting with a one-hour treatment, we found that increasing the treatment time did not make much difference on cell proliferation rate. We used 0.25 Hz as the actual frequency because the minimum discharging and recharging time for the capacitor is 4 seconds when we set intensity of the monophasic stimulator at 100%. This time can be reduced by setting a lower intensity. However, we have used 100% intensity in order to have significant effect on the growth/proliferation rate. The temperature on the coil surfaces was measured by a thermal sensor which showed the temperature of the coil during stimulation. A temperature of 21.7 ± 0.1 °C was maintained in the flask and throughout the stimulation period. There was no
obvious vibration of coils during the stimulation discerned by visually since the coil was fixed by two stages. Therefore, the difference in neural proliferation rate was due to the different orientation of magnetic field generated by double coil. Since during TMS the corresponding electric field generated by time varying magnetic field can affect neurons firing rate [9], different orientation of magnetic field generated clockwise and counterclockwise electric fields and induced current in the brain. The difference in the sequence of clockwise and counterclockwise induced eddy current in the neurons is the reason for the different proliferation of neurons. Since, the interaction between magnetic field and neurons is not well established, further investigation of changes in neuron responses due to application of time varying magnetic fields such as TMS are warranted.

We plan to use different types of neuronal cells in future experiments to assess whether our results were cell-specific. We will also employ advanced imaging techniques to investigate any morphological changes in cells and cell components due to the effect of magnetic field orientation and stimulus parameters. Many factors potentially impact the proliferation of neuronal cells, such as BDNF, GDNF and NGF [18] so we will investigate the effects of TMS fields on these growth proteins.

**Conclusions**

The effect of magnetic field direction generated by TMS coils on the proliferation of N27 dopaminergic neuronal cells was investigated. Orienting the magnetic field upward through the horizontal plane of adherent cells increased their proliferation rate while orienting the magnetic field downward through the cell growth plane decreased their proliferation rate. The results obtained by the Trypan blue method of cell counting was verified by the CyQuant and MTS cell viability assay methods and all the results are
statistically significant. The changes in cell proliferation rate due to magnetic field direction is an important step forward in understanding the effect of magnetic fields on neuronal cell biology. Our findings could have important implications for the preclinical development of TMS treatments of neurological disorders and represents a new method to control the proliferation rate of neuronal cells.

References


CHAPTER 3. THERMAL AND MECHANICAL ANALYSIS OF VARIABLE TMS COIL SYSTEM

Modified from a paper published in *Journal of Applied Physics*, vol. 117, no. 17, p. 17B305, 2015

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**Abstract**

Transcranial Magnetic Stimulation (TMS) has the potential to treat various neurological disorders non-invasively and safely. The “Halo coil” configuration can stimulate deeper regions of the brain with lower surface to deep-brain field ratio compared to other coil configurations. The existing “Halo coil” configuration is fixed and is limited in varying the site of stimulation in the brain. We have developed a new system based on the current “Halo coil” design along with a graphical user interface (GUI) system that enables the larger coil to rotate along the transverse plane. The new system can also enable vertical movement of larger coil. Thus, this adjustable “Halo coil” configuration can stimulate different regions of the brain by adjusting the position and orientation of the larger coil on the head. We have calculated magnetic and electric fields inside an MRI-derived heterogeneous head model for various positions and orientations of the coil. We have also investigated the mechanical and thermal stability of the adjustable “Halo coil” configuration for various positions and orientations of the coil to ensure safe operation of the system.

**Introduction**

Transcranial magnetic stimulation (TMS) is a painless and non-invasive neuromodulation technique based on the principles of magnetic induction [1-2]. TMS has been used to study brain function and is being investigated as a possible treatment for
numerous brain disorders [3]. The technique already shows good efficacy for the treatment of major depressive disorder [4]. We have previously reported a “Halo coil” configuration which can stimulate deeper regions of the human brain, but the configuration was fixed so that only a single site in the brain has a lower surface to deep-brain field ratio compared to other coil configurations [5]. Now, we have built a variable “Halo coil” configuration with a circular coil fixed on top of the head and with vertical and rotational movement of the larger coil to selectively stimulate different regions of the brain. During the stimulation, we used two stimulators to send AC current signals to two coils. One stimulator sends an AC current with a frequency of 2.5 kHz and an amplitude of 2500 A to the circular coil. The other stimulator sends AC current with a frequency of 2.5 kHz and an amplitude of 5000 A to the larger coil.

We have also conducted thermal and mechanical analysis of the system to ensure its feasibility and stability. A GUI system has been built that accurately controls the movement and rotation of the larger coil using an Arduino microcontroller.

**Magnetic (Lorentz) Force Response**

COMSOL Multiphysics (Los Angeles, CA, USA) was used for magnetic force analysis. A 5000 A DC current was assigned in both coils to evaluate the maximum forces induced on the variable “Halo coil” system. Any forces experienced by the coils will be transferred to the insulation and thus the yield strength of insulation should be higher than the Lorentz forces exerted by the magnetic fields generated by the coils. The yield strength of copper is 70 MPa [6] and the ultimate tensile strength of the insulation, Nylon is 125 MPa [7]. The Lorentz force density in the coil can be calculated by equation (4), where $J [A/m]$ is the current density and $B [T]$ is the magnetic flux density. In this study, we used 3D models where $J [A/m^3]$ is the current density and $i = x, y, z$. 
The calculated Lorentz force density $f$ [N/m$^3$] is shown in Fig. 3 for two extreme conditions for our system. The larger coil is rotated $+30^\circ$ and the distance between the centers of two coils is 5 cm and 15 cm. Fig. 19 (a) is the top view, (b) is the side view for a distance of 5 cm between the coils. In Fig. 19 (c) and (d), the distance between the centers of two coils is 15 cm, (c) is the top view, (d) is the side view.

Fig. 19 The result of force analysis between the two coils with larger coil rotated at $+30^\circ$. In (a) and (b), the distance between the centers of two coils is 5 cm, while (a) is the top view, (b) is the side view. In (c) and (d), the distance between the centers of two coils is 15 cm, while (c) is the top view, (d) is the side view.

Fig. 20 The result of thermal analysis of two coils with larger coil rotated at $+30^\circ$. In (a) and (b), the distance between the centers of two coils is 5 cm, while (a) is the top view, (b) is the side view. In (c) and (d), the distance between the centers of two coils is 15 cm, while (b) is the top view, (d) is the side view.

The arrows in the picture show the direction of Lorentz forces and their lengths indicate the magnitude of that force density. In the side view, the majority of Lorentz force density was parallel to the direction of vertical movement of the large coil. The maximum
Lorentz force was 335.01 MN/m³. The equivalent stress on the larger coil was 3.35 MPa. The coils are made of copper with a yield strength of 70 MPa and covered with Nylon, which has a yield/ultimate tensile strength of 125 MPa. Thus, the stress due to the Lorentz forces in the larger coil was significantly smaller than the yield strength of copper and Nylon. The system is therefore be mechanically stable and can withstand the expected Lorentz forces. However, the cyclic loading conditions which can occur in repetitive TMS have not been analyzed.

**Thermal Analysis**

The temperature in the coils was another important factor in the system because of the high amplitude of the current induced in the coil. This can generate a large amount of heat in the coil due to Joule’s law \((Q = I^2 \cdot R \cdot t)\). The limit of surface temperature for electrical medical equipment has been specified by General Standard IEC 60601-2-37, which is 50°C in air and 43°C at the surface of the body [8]. Thus, the modeling of heat was focused on the duration of the stimulation when either of the coils reached 50°C. The incompressible Navier-Stokes heat equation from the COMSOL Heat Transfer module was used to model the thermal changes in the coil system under TMS therapy conditions, as shown equation (5),

\[
\rho C_p \frac{\partial T}{\partial t} + C_p \mathbf{u} \cdot \nabla T = \nabla \cdot (k \nabla T) + Q \tag{5}
\]

where \(\rho\) is the fluid density, \(C_p\) is the fluid heat capacity, \(T\) is the temperature, \(\mathbf{u}\) is the velocity field of the fluid, \(k\) is the thermal diffusivity of the material, and \(Q\) is external source heating [9]. According to the modeling results shown in Fig. 20, after stimulation for 231 seconds, the small circular coil which is placed on the top on patient’s head, reached 50.04 °C. Additionally, the vertical position and the rotational movement of the larger coil did not
have a significant effect on the heat generated in the two coils as all positions and orientations demonstrated similar results of approximately 50 °C in the smaller coil after 231 seconds of stimulation.

**GUI system**

A graphical user interface (GUI) was developed in Java to control the movement and rotation of the larger coil with a computer via an Arduino microcontroller as shown in Fig. 21. The left portion of the interface is the control panel which has two buttons to control the vertical movement of large coil by a linear actuator. The range of vertical movement is -5 cm to +5 cm compared to its origin with a step size of 1 cm. It also has two buttons to control the rotation by a servo motor. The range of rotation is -30° to +30° compared to its origin with a step size of 5°. The right portion of the interface shows the modeling results of electric and magnetic field for the selected position of the large coil. These images will show the distribution of magnetic and electric field which will indicate the site of stimulation with a field larger than the threshold or peak field for the selected position of the large coil.

**Conclusion**

TMS is a novel non-invasive and safe treatment for various neurological disorders. In our present work, we have designed and developed a variable “Halo coil” system that can
achieve deep brain stimulation at specific treatment areas with the vertical and rotational movement of the larger coil in the “Halo coil” system. We have also developed a GUI system to control the movement precisely via a computer. The modeling results of magnetic and electric field confirm that our design can stimulate different parts of human brain. The modeling result of Lorentz forces show that the magnetic forces in coils do not exceed the yield strengths of the coil material and casing in the system. The modeling results of Joule heating showed that the treatment time of 231 seconds will heat the coil to a temperature of 50.04 °C. Thus for longer treatment times, an active cooling system using external air or water circulation should be considered.

References


CHAPTER 4. GENERAL CONCLUSIONS

General Discussion

The study in Chapter 2 has investigated the effect of different orientation of magnetic field for TMS on the proliferation rate of N27 dopamine cells. Basically, the proliferation rate of N27 cells would increase after being treated with the magnetic field orienting upward through the horizontal plane of adherent cells while the rate would decrease after being treated with the magnetic field orienting downward through the cell growth plane. It also compared the advantages of three different cell counting methods which could give a suggestion to related studies in future. The result of this study would bring more attention on the in-vitro and in-vivo study on TMS to understand its mechanism or its effect on neural cells or tissues.

The result in Chapter 3 has demonstrated the stability and feasibility of the Variable TMS coil system. It shows that the maximum electromagnetic (Lorentz) force was much smaller than the yield strength of the coil and its cover materials. Meanwhile, it points out the maximum time for treatment using this system is 231 seconds or 3 minutes and 51 seconds because of one of coils in the system would reach 50 °C at that time, which is the highest temperature for electrical medical equipment according to General Standard IEC 60601-2-37. Moreover, the vertical and rotational movement of the Halo coil does not affect both the mechanical and thermal properties of the whole system, which shows its stability.

The results in chapter 2 and chapter 3 are addressed two major research areas in-vitro study and computer modeling and coil design, respectively. The results could help us to investigate more of the basic mechanism of TMS and bring more efficient and compact tools in clinical trials. Therefore, there are many promising research opportunities in TMS in the
three major areas shown in Fig. 2, which would eventually bring TMS as a major tool to treat different kinds of human brain diseases.

**Recommendation for Future Research**

Although the results in chapter 2 have shown the effect of magnetic field on proliferation rate of neurons, the basic mechanism of how TMS influences the growth of neurons is still not understood. In the meantime, there are many factors that could affect cell’s growth such as temperature, electromagnetic force, gravity and certain proteins like BDNF, GDNF and NGF [1]. Since we have monitored and demonstrated that temperature and electromagnetic force did not contribute much to the results in the experiment. Future work can be done on analysis those proteins like BDNF, GDNF and NGF. Besides, some articles have shown effect of magnetic field on the soma size and axon length of neural cells and using magnetic field to direct axon growth [2-3]. Thus, morphology studies on neurons by TMS treatment would be another approach to investigate its mechanism. Moreover, magnetic nanoparticles have shown possibilities for cell recognition, isolation, purification and enhancing the effect of magnetic field [3-4], so using magnetic nanoparticles in the experiment could show more significant results.

The heat issue of TMS coil is one main challenge for long time repetitive TMS (rTMS) treatment. In our variable TMS coil system, the maximum treating time is less than 4 minutes, which is quite small compared to treating time of regular repetitive TMS treatment. Although this issue can be resolved by adding water-cooling or air-cooling to the coil [5], the cooling component highly increases the weight of the coil which makes it difficult to build enough support structure for vertical and rotational movement of the coil. In our current design, we used polymer and 3D printing technology to make the support structure. However,
it cannot support the heavy cooling component of the coil, so future work can be done on
design a more firm support structure to hold the air-cooling coil for long time TMS treatment
with the variable TMS coil system.

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APPENDIX A. MODELING OF ELECTRIC AND MAGNETIC FIELD FOR VARIABLE TMS COIL SYSTEM

We have used SEMCAD X (SPEAG, Swiss) finite element software to calculate the electric and magnetic fields generated by the fixed circular coil positioned at the vertex of the head with different orientations of the larger coil. An AC current with a frequency of 2.5 kHz and an amplitude of 2500 A was applied to the circular coil which is comparable to the pulse signal generated by a biphasic commercial TMS stimulator with 50% power intensity, and a current signal of the same frequency but with an amplitude of 5000 A was applied to the large coil [1]. We have used an anatomically realistic human head model with different electrical properties assigned to each tissue of the brain [2]. These parameters are shown in Table I [3].

<table>
<thead>
<tr>
<th>Tissue</th>
<th>εr (Relative Permittivity)</th>
<th>σ (Electric Conductivity) [S/m]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brain (grey matter)</td>
<td>7.81 × 10^4</td>
<td>1.04 × 10^-1</td>
</tr>
<tr>
<td>Brain (white matter)</td>
<td>3.43 × 10^4</td>
<td>6.45 × 10^-2</td>
</tr>
<tr>
<td>Cerebellum</td>
<td>7.84 × 10^4</td>
<td>1.24 × 10^-1</td>
</tr>
<tr>
<td>Cerebrospinal fluid</td>
<td>1.09 × 10^2</td>
<td>2.00</td>
</tr>
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<td>1.14 × 10^3</td>
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</tr>
<tr>
<td>Skull</td>
<td>1.44 × 10^3</td>
<td>2.03 × 10^4</td>
</tr>
</tbody>
</table>

The calculation of magnetic field was based on the Biot-Savart law as shown in equation (1) [4].

\[
A_\theta(r) = \frac{\mu_0}{4\pi} \int \frac{J_\phi(r')}{|r-r'|} dr' \quad (1)
\]

The vector potential \( A \) is decoupled from the electric field \( E \) which is calculated using equation (2) where \( \nabla \cdot E_s = 0 \) (solenoidal) and \( \nabla \times E_i = 0 \) (irrotational).

\[
E = -j\omega A + \nabla \varphi = E_s + E_i \quad (2)
\]

The magneto-quasi-static calculation is described by equation (3).

\[
\nabla \cdot \sigma \nabla \varphi = j\omega \nabla \cdot (\sigma A_\theta) \quad (3)
\]
Fig. 22 (a), (b), (c) and (d) show the difference in electric and magnetic fields generated in the head for different vertical positions of the large coil. When comparing Fig. 22(b) and Fig. 22(d), it shows that the electric field in Fig. 22(b) is higher than that in Fig. 22(d).

Fig. 22 (a) and (b) show the magnetic field (a and c) and electric field (b and d) generated in the anatomically realistic human head model for different vertical positions of the large coil. In figures a and b, the distance between two coils is 5 cm. In figures c and d, the distance between two coils is 15 cm.

Fig. 23 Magnetic field (a and c) and electric field (b and d) generated in the anatomically realistic human head model for different rotational angles of the large coil. In figures a and b, the coil is rotated +30 degrees. In figures c and d, the coil is rotated -30 degrees.

However, the electric field in lower part of head model is higher in Fig. 22(b) than in Fig. 22(d), which is enhanced by the position of the large coil. These modeling results show the evidence that the larger coil enhances the electric and magnetic fields at the deeper regions of the brain by reducing the decay of field generated by the smaller circular coil which is fixed on the top of head. Thus, different positions of larger coil enables stimulation
of different deeper regions of the human brain and helps clinicians to vary the site of stimulation according to the disorder that is being treated.

Fig. 23 shows the induced electric field in the anatomical heterogeneous head model with the rotational movement of the larger coil. According to Fig. 23 (b) and (d), the position of the peak value of the electric field was different according to different positions of the larger coil and the peak value of electric field was approximately 250 V/m which is larger than the threshold electric field of 150 V/m reported by March et al. [5] and 120 V/m by Rosanova et al. [6]. Therefore, rotation of the larger coil also reduces the decay of electric and magnetic field generated by the small circular coil similar to vertical movement.

References


APPENDIX B. LIST OF PUBLICATIONS AND REPRINTS OF PUBLICATIONS DERIVED FROM THE WORK PERFORMED FOR THIS THESIS

Archived Journal Publications:


Peer Reviewed Conference Publications:


Deep brain transcranial magnetic stimulation using variable “Halo coil” system

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Department of Electrical and Computer Engineering, Iowa State University, Ames, Iowa 50011, USA

(Submitted 4 March 2015; published online 4 March 2015)

Transcranial Magnetic Stimulation has the potential to treat various neurological disorders non-invasively and safely. The “Halo coil” configuration can stimulate deeper regions of the brain with lower surface to deep-brain field ratio compared to other coil configurations. The existing “Halo coil” configuration is fixed and is limited in varying the site of stimulation in the brain. We have developed a new system based on the current “Halo coil” design along with a graphical user interface system that enables the larger coil to rotate along the transverse plane. The new system can also enable vertical movement of larger coil. Thus, this adjustable “Halo coil” configuration can stimulate different regions of the brain by adjusting the position and orientation of the larger coil on the head. We have conducted magnetic and electric fields inside a MRI-derived heterogeneous head model for various positions and orientations of the coil. We have also investigated the mechanical and thermal stability of the adjustable “Halo coil” configuration for various positions and orientations of the coil to ensure safe operation of the system. © 2015 AIP Publishing LLC.

[http://dx.doi.org/10.1063/1.4913937

INTRODUCTION

Transcranial magnetic stimulation (TMS) is a painless and non-invasive neuromodulation technique based on the principles of magnetic induction.1,2 TMS has been used to study brain function and is being investigated as a possible treatment for numerous brain disorders.3 The technique already shows good efficacy for the treatment of major depressive disorder.4 We have previously reported a “Halo coil” configuration which can stimulate deeper regions of the human brain, but the configuration was fixed so that only a single site in the brain has a lower surface to deep-brain field ratio compared to other coil configurations.5 Now, we have built a variable “Halo coil” configuration with a circular coil fixed on top of the head and with vertical and rotational movement of the larger coil to selectively stimulate different regions of the brain. During the stimulation, we used two stimulators to send AC current signals to two coils. One stimulator sends an AC current with a frequency of 2.5 kHz and an amplitude of 2500 A to the circular coil. The other stimulator sends an AC current with a frequency of 2.5 kHz and an amplitude of 5000 A to the larger coil.5 We have used an anatomically realistic human head model with different electrical properties assigned to each tissue of the brain.6 These parameters are shown in Table I.7

The calculation of magnetic field was based on the Biot-Savart law as shown in the following equation:8

\[ A_0(r) = \frac{\mu_0}{4\pi} \int_{\Omega} \frac{J_0(r')}{|r - r'|} d\Omega . \] (1)

The vector potential A is decoupled from the electric field E which is calculated using Eq. (2) where \( \nabla \cdot E_s = 0 \) (solenoidal) and \( \nabla \times E_s = 0 \) (irrotational).

\[ E = -j\omega A + \nabla \varphi = E_s + E_i. \] (2)

The magneto-quasi-static calculation is described by the following equation:

\[ \nabla \cdot \sigma \nabla \varphi = j\omega \nabla \cdot (\sigma A_0). \] (3)

TABLE I. Values of dielectric properties at 2.5 kHz.

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</table>

a)Author to whom correspondence should be addressed. Electronic mail: hadimani@iastate.edu.
Figs. 1(a)–1(d) show the difference in electric and magnetic fields generated in the head for different vertical positions of the large coil. When comparing Figs. 1(b) and 1(d), it shows that the electric field in Fig. 1(b) is higher than that in Fig. 1(d). However, the electric field in lower part of head model is higher in Fig. 1(b) than in Fig. 1(d), which is enhanced by the position of the large coil. These modeling results show the evidence that the larger coil enhances the electric and magnetic fields at the deeper regions of the brain by reducing the decay of field generated by the smaller circular coil which is fixed on the top of head. Thus, different positions of larger coil enable stimulation of different deeper regions of the human brain and helps clinicians to vary the site of stimulation according to the disorder that is being treated.

Fig. 2 shows the induced electric field in the anatomical heterogeneous head model with the rotational movement of the larger coil. According to Figs. 2(b) and 2(d), the position of the peak value of the electric field was different according to different positions of the larger coil and the peak value of electric field was approximately 250 V/m which is larger than the threshold electric field of 150 V/m, reported by March et al. and Rosanova et al. Therefore, rotation of the larger coil also reduces the decay of electric and magnetic field generated by the small circular coil similar to vertical movement.

**MAGNETIC (LORENTZ) FORCE RESPONSE**

COMSOL Multiphysics (Los Angeles, CA, USA) was used for magnetic force analysis. A 5000 A DC current was assigned in both coils to evaluate the maximum forces induced on the variable “Halo coil” system. Any forces experienced by the coils will be transferred to the insulation and thus the yield strength of insulation should be higher than the Lorentz forces exerted by the magnetic fields generated by the coils. The yield strength of copper is 70 MPa (Ref. 11) and the ultimate tensile strength of the insulation, Nylon is 125 MPa. The Lorentz force density in the coil can be calculated by Eq. (4), where \( J \) (A/m) is the current density and \( B \) (T) is the magnetic flux density. In this study, we used 3D models where \( J \) (A/m\(^3\)) is the current density and \( i = x, y, z \)

\[
f_i = J \times B.
\] (4)

The calculated Lorentz force density \( f \) (N/m\(^3\)) is shown in Fig. 3 for two extreme conditions for our system. The larger coil is rotated \(+30^\circ\) and the distance between the centers of two coils is 5 cm and 15 cm. Fig. 3(a) is the top view and Fig. 3(b) is the side view for a distance of 5 cm between the coils. In Figs. 3(c) and 3(d), the distance between the centers of two coils is 15 cm, i.e., Fig. 3(c) is the top view and Fig. 3(d) is the side view.

The arrows in the picture show the direction of Lorentz forces and their lengths indicate the magnitude of that force density. In the side view, the majority of Lorentz force density was parallel to the direction of vertical movement of the large coil. The maximum Lorentz force was 335.01 MN/m\(^3\). The equivalent stress on the larger coil was 3.35 MPa. The coils are made of copper with a yield strength of 70 MPa and covered with Nylon, which has a yield/ultimate tensile strength of 125 MPa. Thus, the stress due to the Lorentz forces in the larger coil was significantly smaller than the yield strength of copper and Nylon. The system is therefore be mechanically stable and can withstand the expected Lorentz forces. However, the cyclic loading conditions, which can occur in repetitive TMS, have not been analyzed.
THERMAL ANALYSIS

The temperature in the coils was another important factor in the system because of the high amplitude of the current induced in the coil. This can generate a large amount of heat in the coil due to Joule’s law ($Q = I^2 R t$). The limit of surface temperature for electrical medical equipment has been specified by General Standard IEC 60601-2-37, which is 50°C in air and 43°C at the surface of the body. Thus, the modeling of heat was focused on the duration of the stimulation when either of the coils reached 50°C. The incompressible Navier-Stokes heat equation from the COMSOL Heat Transfer module was used to model the thermal changes in the coil system under TMS therapy conditions, as shown in the following equation:

$$\rho C_p \frac{\partial T}{\partial t} + \rho C_p u \cdot \nabla T = \nabla \cdot (k \nabla T) + Q, \quad (5)$$

where $\rho$ is the fluid density, $C_p$ is the fluid heat capacity, $T$ is the temperature, $u$ is the velocity field of the fluid, $k$ is the thermal diffusivity of the material, and $Q$ is external source heating. According to the modeling results shown in Fig. 4, after stimulation for 231 s, the small circular coil, which is placed on the top on patient’s head, reached 50.04°C. Additionally, the vertical position and the rotational movement of the larger coil did not have a significant effect on the heat generated in the two coils as all positions and orientations demonstrated similar results of approximately 50°C in the smaller coil after 231 s of stimulation.

GUI SYSTEM

A GUI was developed in Java to control the movement and rotation of the larger coil with a computer via an Arduino microcontroller, as shown in Fig. 5. The left portion of the interface is the control panel, which has two buttons to control the vertical movement of large coil by a linear actuator. The range of vertical movement is $-5$ cm to $+5$ cm compared to its origin with a step size of 1 cm. It also has two buttons to control the rotation by a servo motor. The range of rotation is $-30^\circ$ to $+30^\circ$ compared to its origin with a step size of 5°. The right portion of the interface shows the modeling results of electric and magnetic field for the selected position of the large coil. These images will show the distribution of magnetic and electric field, which will indicate the site of stimulation with a field larger than the threshold or peak field for the selected position of the large coil.
CONCLUSION

TMS is a novel non-invasive and safe treatment for various neurological disorders. In our present work, we have designed and developed a variable “Halo coil” system that can achieve deep brain stimulation at specific treatment areas with the vertical and rotational movement of the larger coil in the “Halo coil” system. We have also developed a GUI system to control the movement precisely via a computer. The modeling results of magnetic and electric field confirm that our design can stimulate different parts of human brain. The modeling result of Lorentz forces show that the magnetic forces in coils do not exceed the yield strengths of the coil material and casing in the system. The modeling results of Joule heating showed that the treatment time of 231 s will heat the coil to a temperature of 50.04 °C. Thus, for longer treatment times, an active cooling system using external air or water circulation should be considered.

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Differential Effect of Magnetic Field Orientation on the Proliferation Rate of Dopaminergic Neurons during Transcranial Magnetic Stimulation

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Abstract--Transcranial magnetic stimulation (TMS) has been used to investigate possible treatments for a variety of neurological disorders. But the effect that magnetic fields have on neurons has not been well documented in the literature. Using a monophasic stimulator, we investigated the effect of different orientation of magnetic field generated by TMS coils on the proliferation rate of N27 neuronal cells cultured in flasks and multi-well plates. Exposing horizontally adherent N27 cells to a magnetic field pointing upward through the neuronal proliferation layer increased the proliferation of cells compared with the control group. On the other hand, proliferation rate decreased in cells exposed to a magnetic field pointing downward through the neuronal growth layer compared with the control group. The results were consistent across different methods of measuring proliferation and cell counting procedures. We confirmed results obtained from the Trypan-blue and automatic cell counting methods with those from the CyQuant and MTS cell viability assays. Our findings could have important implications for the preclinical development of TMS treatments of neurological disorders and represents a new method to control the proliferation rate of neuronal cells.

Key words: TMS; dopaminergic neurons; proliferation rate; orientation of magnetic field

INTRODUCTION

Transcranial Magnetic Stimulation is a non-invasive neuromodulation technique that uses time varying short pulses of magnetic field to induce an electric field in the conductive tissues of the brain thus, modulating the synaptic transmission of neurons. This neuromodulation technique can be used to excite or inhibit the firing rate of neurons which can then be used for treatment of various neurological disorders such as major depressive disorder, Parkinson's disease, Post-traumatic stress disorder and migraine (Barker
et al., 1985; George et al., 2010; Vonloh et al., 2013; Rosenberg, 2002; Dodick et al., 2010). Since the US Food and Drug Administration (FDA) approved TMS as a treatment for depression in 2008, there has been less focus on in vitro and animal studies in the literature compared to in vivo studies in humans (Meng et al., 2015; Crowther et al., 2013; Wassermann and Zimmermann, 2012; March et al., 2014). The effects of TMS on individual neurons need to be thoroughly understood to fully utilize TMS as a neuromodulation tool for treating neurological disorders especially those originating from subcortical regions of the brain.

Few articles have reported the effect of time-varying magnetic fields, similar to those generated by TMS, on the proliferation rates of neurons. (Bonmassar et al., 2009) designed micro TMS coils and showed that the direction of magnetic field affects the firing frequency of neurons, but the authors did not report the effect of magnetic field on the proliferation rate. Meanwhile, some articles have reported the effect of static magnetic field on cell’s proliferation rate (Miyakoshi, 2009). Authors have used static magnetic fields from 1 to 10 tesla and did not find any significant effect on cell proliferation or on genetic toxicity, regardless of the length of treatment. However, there was a small effect on intracellular Ca^{2+} ion control. Some articles have reported beneficial effects of DC electric field (EF) on neural proliferation and differentiation. The EF gradient affects morphology and phenotype of adult neural stem/progenitor cells (NPCs), which shows the potential of utilizing EF to control migration, differentiation and alignment of stem cells transplanted to treat nervous system disorders (Ariza et al., 2010). Extremely low-frequency electromagnetic fields (ELF-EMFs) have been used therapeutically to drive cardiac-specific differentiation in adult human cardiac progenitor cells without any pharmacological or genetic manipulation of cells (Gaetani et al., 2009). As far as we know, no one has published on the effect of TMS magnetic field on the proliferation rate of neurons or on the morphology of cells.

In this paper, we have presented the effect of magnetic field generated by TMS coils on the proliferation of N27 dopaminergic neurons. We have used different cell proliferation and cell counting procedures to confirm that directing a magnetic field downward or upward through the horizontal proliferation plane of adherent cell cultures decreased or increased cell proliferation rates, respectively. It is
important to note that the direction of the induced electric current from the time varying TMS fields will be in clockwise or counterclockwise loops when the magnetic field is in up or down direction of the cell culture as shown in Fig. 4. This experimental set up is similar to the TMS treatment on human brain where the induced electric field from the TMS coils will be in clockwise or counterclockwise loops in the cortex.

EXPERIMENTAL PROCEDURES

A. Magnetic Field Generated by TMS coils

A Magstim Standard 70 mm double coil was used for treating N27 neurons. Magnetic field was measured on the surface of the coil using a gaussmeter and a hall probe. The field was also calculated using finite element electromagnetic modeling software, SEMCAD X. The measured and calculated axial components of the magnetic field intensities are shown in Fig.1 and Fig.2. Magnetic field is negative in the negative x-axis and positive in the positive x-axis which is shown Fig. 2. It also shows magnetic field values at 5mm above the coil surface where dopaminergic neurons are placed during TMS treatment after considering the thickness of flask and thermal insulation layer. According to these figures, the peak value of measured magnetic field intensity at 5mm above the coil surface is 0.55 MA/m which is reduced by approximately 0.1MA/m. Fig. 3 shows the top view of distribution of magnetic field intensity generated by double coil. Fig.4 shows the different orientations of magnetic field generated by the coil and directions of
current in each circle of the double coil. The red arrows on the left indicate the directions of supplied current (5000 A) in left circle as counterclockwise and clockwise in the right circle. The cross symbols indicate the magnetic flux pointing into the plane and the dot symbols indicate the magnetic flux pointing out of the plane.

Fig. 3 Distribution of magnetic field intensity of double coil (2D top view).

According to Maxwell’s equation, time-varying magnetic field will generate an electric field which induces eddy currents in the conducting neurons. The supplied current is a pulse wave which has a frequency of 2.5 kHz and magnitude of 5000 A, so its period is 0.4 ms. The stimulator sends only one pulse with a current amplitude of 5000 A in clockwise and counterclockwise directions in left and right circles of the coil respectively as shown in Fig. 4. Thus, the supplied current in each circular coil will generate a time varying magnetic field change from 0 to its peak value, during its first half period, which results in the corresponding induced eddy current in both areas shown in Fig. 5 and Fig. 6. According to
Lenz’s law the induced current in the left circular coil was counterclockwise and it was clockwise in the right circular coil. Similarly, the value of the supplied current in both coils would change from its peak of 5000 A to 0 during the second half period. Thus, the induced eddy current on the left and right flasks was clockwise and counterclockwise, respectively. Therefore, the difference between the two flasks was the sequence of the direction of the eddy currents.

B. Cell Culture

Immortalized rat mesencephalic 1RB3AN27 cells (N27) were grown in RPMI-1640 medium supplemented with 10% fetal bovine serum, 1% L-glutamine, 50 units penicillin and 50 µg/ml streptomycin and maintained at 37°C with a humidified atmosphere containing 5% CO₂, as described previously (Anantharam et al., 2002; Prasad et al., 1998). On Day 0, an equal number of N27 cells were seeded into each T-75 flask or 96-well plate. Groups were distinguished by culture time with two control and two TMS groups per time point and four replicate samples (flask or plate) per group. Control 1 was always kept in the incubator and was named Incubator in Table I. Control 2 was kept in the biosafety cabinet during the TMS treatment and was named Environmental in Table I. Table I shows culture time points and counting time points for the different sample locations and magnetic field orientations (“Field up” and “Field down”), which were used in a Trypan blue cytotoxicity assay. Table II shows cell culture samples with their culture time points as well as counting time points used in a CyQuant cell proliferation assay. Table III shows culture samples with their culture and counting time points for an MTS cell viability assay and cell counting method. We performed a cell count for each sample of cells 24 hours after its TMS treatment to ensure that the cells had enough time to show any effects of TMS on their proliferation.

C. TMS experiment on dopaminergic neurons

We used a monophasic stimulator to treat N27 cell cultures. A set of 6 pulses with 4 seconds waiting time in between them was formed as one train and a waiting time of 10 seconds between each train was introduced, so the pulse repetition rate (TMS treatment frequency) we used is 0.25 Hz. This is a low frequency compared to usual clinical protocol frequency however, in order to obtain 100% power in the
coil and avoid rapid heating up of the coil we have used this low frequency. It is not possible to operate at
higher frequencies at full power with the existing set-up. A total of 60 trains with 360 pulses were
delivered per 30-minute TMS treatment. An air-cooled double coil was used which has opposite current
directions in each coil, generating magnetic fields on top side of each coil with opposite directions. Using
air-cooled coils allowed us to induce magnetic fields without raising the temperature of the T-75 flasks
placed on them. All TMS treatments on N27 cells were performed in a sterile biosafety cabinet (Fig. 7).
The flask set above the left coil was designated “Field up” and the flask set above the right coil was
designated “Field down”, corresponding to the orientation of the magnetic fields.

![Image of TMS treatment setup](image)

Fig. 7 Arrangement of the 30-minute TMS treatment delivered to two T-75 flasks. The directions of the two
 oppositely oriented magnetic fields were labeled on the double coil. Field orientation was upward on the left coil and
downward on the right coil, as shown in the inset figure. We used two clamps to fix the coil in the cell culture
cabinet and a layer of bubble wrap separated both flasks from the coils to maintain a thermal barrier.

D. Cytotoxicity assay

Cytotoxic cell death was measured as per Life Technologies’ Trypan blue exclusion cell counting
method (Xu et al.,2008). Briefly, after treatment cells were harvested with trypsin-EDTA and
resuspended in 1X PBS, we then took 10 µl of cell suspension from one sample and added with 10 µl of
0.4% trypan blue solution (Life Technologies) and triturated this mixture. Then, we put 10 µl of the
mixture into the cell counting slide and place the slide into the automatic cell counter to count the
concentration of cells in each sample. Finally, we extrapolated the total number of cells in each sample
by multiplying its volume and concentration (Xu et al.,2008). By using this method, we counted the
number of cells in each of the four replicate samples according to the counting time points shown in Table I. We studied the effect of TMS on 2 different initial cell densities, 1 million cells/flask and 0.5 million cells/flask (n=4).

Table I. Design of the TMS experiment with Trypan blue cell counting method for cell densities of 1 and 0.5 million cells/flask.

<table>
<thead>
<tr>
<th>Culture Time Point</th>
<th>Counting Time Point</th>
<th>Sample Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Day 0</td>
<td>Day 1</td>
<td>Incubator Environmental Field up Field down</td>
</tr>
<tr>
<td>Day 0.5</td>
<td>Day 1.5</td>
<td>Incubator Environmental Field up Field down</td>
</tr>
<tr>
<td>Day 1</td>
<td>Day 2</td>
<td>Incubator Environmental Field up Field down</td>
</tr>
<tr>
<td>Day 1.5</td>
<td>Day 2.5</td>
<td>Incubator Environmental Field up Field down</td>
</tr>
<tr>
<td>Day 2</td>
<td>Day 3</td>
<td>Incubator Environmental Field up Field down</td>
</tr>
</tbody>
</table>

Table II. Design of the TMS experiment with CyQuant cell viability assay cell counting method for cell densities of 100k, 80k, 50k and 20k per well.

<table>
<thead>
<tr>
<th>Culture Time Point</th>
<th>Counting Time Point</th>
<th>Sample Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Day 0</td>
<td>Day 1</td>
<td>Environmental Field up Field down</td>
</tr>
<tr>
<td>Day 1</td>
<td>Day 2</td>
<td>Environmental Field up Field down</td>
</tr>
<tr>
<td>Day 2</td>
<td>Day 3</td>
<td>Environmental Field up Field down</td>
</tr>
</tbody>
</table>

E. CyQuant cell proliferation assay

We used Life technologies’ CyQuant cell viability assay to confirm our results from the Trypan blue cell counting procedures. On day 0, we seeded the N27 cells in 24-well plates (n=3) with the four rows per plate. In each plate, there were 4 rows and three columns. Row 1 to row 4 have different seeding densities; 20k, 50k, 80k, to 100k, respectively. Each row had three replicated samples in three columns to account for standard deviation. We had three groups with different culture times: Day 0, Day 1 and Day 2 (Table II). Briefly, after 24 hours post-treatment, we read the fluorescence with excitation maximum at 485 nm and the emission maximum at 530 nm using a Synergy 2 plate reader (BioTek) (Jones et al., 2001). We pooled the groups designated as Incubator and Environmental in Table I, because the difference between them was insignificant.

F. MTS cell viability assay cell counting method

Cell viability was measured using Promega’s MTS assay to confirm the results from Trypan blue and CyQuant cell proliferation assays. Briefly, on day 0, we seeded the wells of 96-well plates with 15k for
one row and half with 20k for another row of N27 cells in 200 µL of proliferation medium per well. Each row had 6 duplicated samples (n=6). The design of the experiment was according to Table III., TMS treatment was performed with “Field up” and “Field down” on 2 different well plates. After 24 hours post-treatment, 20µl MTS reagent (CellTiter 96® Aqueous One Solution Reagent) was added to each well and incubated at 37°C in a CO2 incubator for 90 min and absorbance was read at 490 nm and 670 nm in a Spectramax plate reader (Molecular Devices). We subtracted the baseline via Abs$^\text{490}$-Abs$^\text{670}$ prior to data analysis (Mahon et al., 2000).

Table III. Design of the TMS experiment with MTS cell viability assay cell counting method for cell densities of 15k and 20k per well.

<table>
<thead>
<tr>
<th>Culture Time Point</th>
<th>Counting Time Point</th>
<th>Sample Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Day 0</td>
<td>Day 1</td>
<td>Control</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Environmental</td>
</tr>
<tr>
<td>Field up</td>
<td></td>
<td>Field down</td>
</tr>
</tbody>
</table>

G. Statistical significance analysis

Statistical significance analysis was performed using Originlab 9.0 software (OriginLab Corporation, Northampton, MA, USA). Raw data analysis were analyzed using a two unpaired t-test. Statistically significant differences are indicated by asterisks as follows: *p<0.05, **p<0.01 and *p<0.001.

RESULTS

After TMS treatment of N27 cells, we counted the number of viable cells using the Trypan blue method for initial seeding densities of 1 million (Fig. 8) and 0.5 million (Fig. 9) cells per flask. The culture time and counting time points are indicated in Table I. The result showed that the proliferation rate increased after TMS stimulation with the magnetic field oriented upward through the horizontal plane of adherent cells, compared to incubator and environmental samples. The proliferation rate decreased when the field was oriented downward through the horizontal growth plane compared to incubator and environmental samples. Also, environmental samples exhibited slower proliferation compared to the incubator condition. For the lower seeding density (Fig.9), the difference of cell counting for each group became larger over time. The difference peaked on Day 3 when the number of cells in the “Field up” group was 23.57 ± 3.21%
(mean ± STD, ***p<0.001) higher than that in the Environmental group, while in the Field down group, it was $11.45 \pm 1.99\%$ (**p<0.001) lower than in the Environmental group. Therefore, the total difference in cell’s proliferation rate attributable to TMS field direction was $+35.02\%$.

To investigate the effect of different culture times on cell proliferation, we conducted another experiment expanding culture time up from 2 days to 2.5 days. The seeding density was 0.5 million/flask and the new culture time points were Day 0.5, Day 1, Day 1.5, Day 2 and Day 2.5. Cells were counted 24 h after each treatment, so the corresponding counting time points were Day 1.5, Day 2, Day 2.5, Day 3 and Day 3.5 respectively. The effect of TMS and its direction on the proliferation of cells over time (Fig. 10) was similar to the previous results for this seeding density (Fig. 9). On Day 3.5, the number of cells in the “Field up” group was $13.53 \pm 1.36\%$ (**p<0.001) higher than in the Environmental group, whereas the number of cells in the “Field down” group was $12.61 \pm 1.76\%$ (**p<0.001) lower than in the Environmental group. Therefore, the total difference in cell’s proliferation rate attributable to TMS field direction was $+26.14\%$. 

![Fig. 8 Cell densities in the TMS experiment, derived using the Trypan blue cell counting method with an initial seeding density of 1 million cells/flask. Counting time is indicated in Table. I.](image1)

![Fig. 9 Cell densities in the TMS experiment, derived using the Trypan blue cell counting method with initial seeding density of 0.5 million cells/flask. Counting time is indicated in Table. I.](image2)
We used the CyQuant cell viability assay to confirm the results obtained with the Trypan blue cell counting method. This time we eliminated group 1 (Incubator) and we set four seeding densities. The design of this experiment was based on Table II. The effect of TMS field direction on cell proliferation obtained via the CyQuant method (Fig 11-14) was similar to the effect measured using the Trypan blue cell counting method. However, Fig.8, which had the seeding density of 100k per well did not follow the trend similar to other seeding densities i.e. the difference in the proliferation rate was not pronounced. It may be due to the fact that a large number of cells grew in the limited space so the cells might have attained 100% confluency earlier than Day 3.
Fig. 11 Cell proliferation after TMS treatment of plates with an initial seeding density of 20k/well. We used the CyQuant cell viability assay to count cells. On day 3, the number of cells in the “Field up” group was 22.06 ± 4.14% (mean ± STD as a percentage of the initial seeding density; *p<0.05) higher than in the Environmental group, whereas cell numbers in the “Field down” group were 28.77± 1.00% (**p<0.01) lower than in the Environmental group.

Fig. 12 Cell proliferation after TMS treatment of plates with an initial seeding density of 50k/well. We used the CyQuant cell viability assay to count cells. On day 3, the number of cells in the “Field up” group was 15.49 ± 7.26% (mean ± STD as a percentage of the initial seeding density; *p<0.05) higher than in the Environmental group, whereas cell numbers in the “Field down” group were 9.94 ± 2.47% (*p<0.05) lower than in the Environmental group.

Fig. 13 Cell proliferation after TMS treatment of plates with an initial seeding density of 80k/well. We used the CyQuant cell viability assay to count cells. On day 3, the number of cells in the “Field up” group was 15.57 ± 5.17% (mean ± STD as a percentage of the initial seeding density; *p<0.05) higher than in the Environmental group, whereas cell numbers in the “Field down” group were 11.62 ± 1.55% (*p<0.05) lower than in the Environmental group.

Fig. 14 Cell proliferation after TMS treatment of plates with an initial seeding density of 100k/well. We used the CyQuant cell viability assay to count cells. On day 3, the number of cells in the “Field up” group was 14.69 ± 5.74% (mean ± STD as a percentage of the initial seeding density, p>0.05) higher than in the Environmental group, whereas cell numbers in the “Field down” group were the same (0 ± 4.50%, p>0.05) as those in the Environmental group.
A third cell counting method, the MTS cell viability assay, was performed to confirm the results obtained with the Trypan blue and CyQuant cell counting methods. With an initial seeding of 15k (Fig. 15), the number of cells in the “Field up” group was 19.88 ± 4.56% (***p<0.001) higher than in the Environmental control group. Meanwhile, the number of cells in the “Field down” group was 8.88 ± 1.39% (**p<0.01) lower than in the Environmental group. Next, using an initial seeding of 20k (Fig. 16), the number of cells in the “Field up” group was 19.60 ± 4.57% (**p<0.01) higher than in the Environmental group, while the number of cells in the “Field down” group was 8.16 ± 0.09% (**p<0.01) lower than in the Environmental group. Therefore, the total difference in cell’s proliferation rate attributable to TMS field direction was +27.76 %.

**DISCUSSION**

We investigated the effect of magnetic field orientation on the proliferation rate of N27 dopaminergic neuronal cells using three different cell counting methods to cross-validate the results. The MTS assay showed the highest difference in cell proliferation rate. It was also easy to replicate this counting procedure three times to obtain standard deviation. In the Trypan blue cell counting method, we used flasks to culture neuronal cells, which required more area to incubate replicate samples. Cell counting using the Trypan blue method was more time consuming because cell counting was performed one flask at a time, unlike the
MTS method where cell counting was performed in groups. There were three replicate samples for each group (n=3) for Trypan blue method and for MTS and CyQuant cell viability assay, n=6, which is adequate to show statistical significance. In the CyQuant cell viability cell counting method, it was easy to replicate samples and we were also able to count cells of all groups at once, but the differences among groups were slightly smaller than those from the MTS cell counting method. Thus, the MTS cell viability assay cell counting method is recommended for investigating the proliferation rate of N27 dopaminergic neuronal cells under TMS treatment.

According to the design of all these experiments, each group of N27 dopaminergic neuronal cells received a 30 minutes TMS treatment each day. After experimenting with a one-hour treatment, we found that increasing the treatment time did not make much difference on cell proliferation rate. We used 0.25 Hz as the actual frequency because the minimum discharging and recharging time for the capacitor is 4 seconds when we set intensity of the monophasic stimulator at 100%. This time can be reduced by setting a lower intensity. However, we have used 100% intensity in order to have significant effect on the growth/proliferation rate. The temperature on the coil surfaces was measured by a thermal sensor which showed the temperature of the coil during stimulation. A temperature of 21.7 ± 0.1 °C was maintained in the flask and throughout the stimulation period. There was no obvious vibration of coils during the stimulation discerned by visually since the coil was fixed by two stages. Therefore, the difference in neural proliferation rate was due to the different orientation of magnetic field generated by double coil. Since during TMS was the corresponding electric field generated by time varying magnetic field can affect neurons firing rate (Bonmassar et al., 2012), different orientation of magnetic field will generate clockwise and counterclockwise electric fields and induced current in the brain. The difference in the sequence of clockwise and counterclockwise induced eddy current in the neurons is the reason for the different proliferation of neurons. Since, the interaction between magnetic field and neurons is not well established, further investigation of changes in neuron responses due to application of time varying magnetic fields such as TMS are warranted.
We plan to use different types of neuronal cells in future experiments to assess whether our results were cell-specific. We will also employ advanced imaging techniques to investigate any morphological changes in cells and cell components due to the effect of magnetic field orientation and stimulus parameters. Many factors potentially impact the proliferation of neuronal cells, such as BDNF, GDNF and NGF (Allen et al., 2013) so we will investigate the effects of TMS fields on these growth proteins.

CONCLUSIONS

The effect of magnetic field direction generated by TMS coils on the proliferation of N27 dopaminergic neuronal cells was investigated. Orienting the magnetic field upward through the horizontal plane of adherent cells increased their proliferation rate while orienting the magnetic field downward through the cell growth plane decreased their proliferation rate. The results obtained by the Trypan blue method of cell counting was verified by the CyQuant and MTS cell viability assay methods and all the results are statistically significant. The changes in cell proliferation rate due to magnetic field direction is an important step forward in understanding the effect of magnetic fields on neuronal cell biology. Our findings could have important implications for the preclinical development of TMS treatments of neurological disorders and represents a new method to control the proliferation rate of neuronal cells.

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REFERENCE


