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Safety parameter considerations of anodal transcranial Direct Current Stimulation in rats

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Abstract

A commonly referenced transcranial Direct Current Stimulation (tDCS) safety threshold derives from tDCS lesion studies in the rat and relies on electrode current density (and related electrode charge density) to support clinical guidelines. Concerns about the role of polarity (e.g. anodal tDCS), sub-lesion threshold injury (e.g. neuroinflammatory processes), and role of electrode montage across rodent and human studies support further investigation into animal models of tDCS safety. Thirty-two anesthetized rats received anodal tDCS between 0 and 5 mA for 60 min through one of three epicranial electrode montages. Tissue damage was evaluated using hemotoxylin and eosin (H&E) staining, Iba-1 immunohistochemistry, and computational brain current density modeling. Brain lesion occurred after anodal tDCS at and above 0.5 mA using a 25.0 mm² electrode (electrode current density: 20.0 A/m²). Lesion initially occurred using smaller 10.6 mm² or 5.3 mm² electrodes at 0.25 mA (23.5 A/m²) and 0.5 mA (94.2 A/m²), respectively. Histological damage was correlated with computational brain current density predictions. Changes in microglial phenotype occurred in higher stimulation groups. Lesions were observed at and above 0.5 mA using a 25.0 mm² electrode (electrode current density: 20.0 A/m²), which is below the previously reported safety threshold of 142.9 A/m² using cathodal tDCS. The lesion area is not simply predicted by electrode current density (and so not by charge density as duration was fixed); rather computational modeling suggests average brain current density as a better predictor for anodal tDCS. Nonetheless, under the assumption that rodent epicranial stimulation is a hypersensitive model, an electrode current density of 20.0 A/m² represents a conservative threshold for clinical tDCS, which typically uses an electrode current density of 2 A/m² when electrodes are placed on the skin (resulting in a lower brain current density).

1. Introduction

Transcranial direct current stimulation (tDCS) is a non-invasive method of brain stimulation used to modulate cortical excitability (Nitsche and Paulus, 2000). Conventional tDCS applies a small amount (1–2 mA) of direct current to the scalp using large (25–35 cm²) electrodes (Brunoni et al., 2012; Woods et al., 2016). Computational and animal models have shown that only a fraction of the applied current reaches the cortex, leading to neuronal polarization and excitability changes in the cortex (Datta et al., 2009a; Marquez-Ruiz et al., 2012; Rahman et al., 2013; Rohan et al., 2015) and hippocampus (Rohan et al., 2015; Kronberg et al., 2017). Given its ability to affect the function of cortical neurons, tDCS has been investigated for a variety of medical and augmentative applications, such as depression (Brunoni et al., 2011; Loo et al., 2012), motor rehabilitation (Edwards et al., 2009), speech rehabilitation (Baker et al., 2010; Fridriksson et al., 2011; Galletta et al., 2015), pain control (Fregni et al., 2006; Dasilva et al., 2012; Castillo-Saaedera et al., 2016), and working memory (Brunoni and Vanderhasselt, 2014). tDCS is considered a safe and well tolerated technique when proper protocols are followed (Bikson et al., 2009; Kasschau et al., 2015; Nitsche and Paulus, 2015; Gbadeyan et al., 2016; Palm et al., 2016; Woods et al., 2016). Nonetheless, as the application of tDCS becomes increasingly commonplace and indications for its use more widespread, additional work on tDCS safety is warranted for supporting basic dosing guidelines (Peterchev et al., 2012; Bikson et al., 2016; Jackson et al., 2016).

Initial safety limitations for tDCS were based upon literature from other electrical brain stimulation techniques. Nitsche et al. discussed safety of tDCS (Nitsche et al., 2003) by referencing safety standards in which pulsating current was applied directly to brain...
been reported at an anodal current intensity of 0.6 mA (47.8 A/m²) – corresponding in their electrode montage to 143 A/m² – along with electrode charge density (C/m²), which multiplies current density by time, as two generalized safety parameters for dosing guidelines (Bikson et al., 2009).

Building upon this framework for rodent safety studies, other tDCS paradigms were evaluated for lesion induction in the mouse (Rueger et al., 2012; Pikhovych et al., 2016a, b), where the lowest current intensity that produced detectable cortical damage was 0.5 mA (220 A/m² electrode current density) for both anodal and cathodal tDCS groups (Pikhovych et al., 2016b). However, damage was not consistently observed for 0.5 mA cathodal and 1.0 mA anodal stimulation groups (Pikhovych et al., 2016a, b), which indirectly suggests a role for polarity. More recently, lesions have been reported at an anodal current intensity of 0.6 mA (47.8 A/m² electrode current density) (Gellner et al., 2016), suggesting the lesion threshold in rats may be lower than previously reported. Rodent studies evaluating tDCS safety through microglial analysis have shown microglial activation can occur after anodal or cathodal stimulation at 0.5 mA for 15 min (Rueger et al., 2012) (c.f. (Liebetanz et al., 2009)). Microglial changes in morphology associated with neurodegeneration after anodal tDCS have been reported at current intensities as low as 0.4 mA (31.8 A/m² electrode current density) (Gellner et al., 2016).

Considering the available lesion safety data and the variations in polarity, animal size, and electrode area used across studies, the robustness of average electrode current density (or electrode charge density) as a generalized predictor of injury remains unclear, undermining the use of animal data to support clinical safety thresholds. Indeed, computational current models notably show brain current density is not simply a function of the electrode current density, but also anatomy and details of electrode size and position (Datta et al., 2009a, 2012; Miranda et al., 2009; Saturnino et al., 2015). Therefore, animal models of tDCS safety can benefit from being updated in regards to: 1) variation of stimulation polarity/dose (anodal vs. cathodal); 2) alternative indications of injury (Wachter et al., 2011; Rueger et al., 2012; Wong et al., 2014; Gellner et al., 2016); and 3) the suitability of electrode parameters to set a safety threshold given computational current models show brain current density is not a simple, linear function of the applied current or electrode current density (Datta et al., 2009a, 2012; Miranda et al., 2009; Saturnino et al., 2015).

We initially developed an in vivo rodent model of anodal tDCS using a 25.0 mm² electrode and evaluated the effect of various stimulation dosages on tissue damage. We evaluated current intensity (0.15–2.5 mA) which span the range of previously established safety limits (Lieberanz et al., 2009; Rueger et al., 2012; Gellner et al., 2016). Ionized calcium-binding adapter molecule 1 (Iba1) activation was also examined as a more sensitive predictor of brain lesion. Brain tissue histology indicated lesions at a lower electrode current density (20.0 A/m²) than previously reported. Therefore, we systematized our next experiments to critically evaluate this 20.0 A/m² limit while controlling the number and area of electrodes (10.6 mm² and 5.3 mm²). Dissociating current intensity from electrode current density (e.g. same current intensity but different electrode current density), combined with high-resolution FEM computational models of current flow in rat, supported testing the hypothesis that brain current density, rather than simply electrode current intensity or electrode current density, predicts the propensity for lesions. This has important implications for how animal (rodent) models of tDCS, especially aimed at safety, are rationalized and applied to develop clinical guidelines.

2. Materials and methods

2.1. 25.0 mm² electrode placement surgery

Animals were anesthetized with isoflurane (Piramal Critical Care, Shop Med Vet, Mettawa, IL) using 5% induction and 2–3% maintenance. Animals were treated with standard pre- and postsurgical care. The animal was placed into a stereotaxic apparatus and a caudo-rostral incision was made on top of the head, followed by a lateral incision was made at the shoulders. The periosteum was removed, the skull wiped clean, and a head electrode of 0.25 cm² (Valutrode, Axelgaard Manufacturing Co., Fallbrook, CA, 1.25-inch diameter electrode cut to 5 mm × 5 mm) along with SignaGel (Parker Laboratories, Fairfield, NJ) was applied to the skull with the center of the electrode resting on the midline 2.5 mm caudal to Bregma (rostro-caudal: 0.0 mm to −5.0 mm). The insulated electrode was tunneled subcutaneously and exited the lateral incision made at shoulders. The electrode was held in place by a plastic head clamp which caught on the ridges of the skull (AFRL designed and produced) and two types of adhesives: C&B Meta-bond Adhesive Luting Cement (Parkell Inc., Edgewood, NY) was applied to the electrode and skull to create an initial bond, followed by acrylic cement (Stoelting, Co. Fisher Scientific, Pittsburg, PA) to bond the electrode to the clamp. Incisions were sutured closed around cement and wire. Animals recovered from surgery for at least 1 week prior to inclusion in experiments. Prior to stimulation, animals were randomly placed into six anodal tDCS treatment groups: 0.15 mA (n = 4), 0.3 mA (n = 4), 0.5 mA (n = 2), 1.0 mA (n = 4), 2.5 mA (n = 3), and sham stimulation (n = 3).

2.2. 10.6 mm² and 5.3 mm² electrode placement surgery

Animals were prepared as described above and an electrode jacket with a surface area of 5.3 mm² (DIXI Medical, Besançon, France) was placed at −2.5 mm Bregma and 2.5 mm left of sagittal suture. The electrode jacket was secured with FUJI I glass ionomer (Dental Wholesale Direct, FL, USA), and a layer of dental cement was placed on top to further secure the electrode. Prior to stimulation, animals were assigned into 2 groups based on electrode placement: 1) a single 5.3 mm² electrode placed at −2.5 mm Bregma and −2.5 mm left of the sagittal line (n = 6), and 2) two 5.3 mm² electrodes placed at −2.5 mm Bregma and 2.5 mm from the midline bilaterally, for a total electrode surface area of 10.6 mm² (n = 5). Prior to stimulation, the animals were assigned into stimulation groups based on lesion results from the previous 25.0 mm² electrode experiment. The single electrode stimulation group was subdivided into groups based on current intensity: 2.0 mA (n = 1), 1.0 mA (n = 1), 0.75 mA (n = 1), 0.5 mA (n = 2), 0.05 mA (n = 1), and Sham (n = 1). The dual electrode group was also divided into subgroups based on current intensity: 2.0 mA (n = 1), 1.0 mA (n = 1), 0.5 mA (n = 1), 0.25 mA (n = 1), and sham (n = 1).

2.1. tDCS application

2.3.1. 25.0 mm² electrode stimulation

Animals were anesthetized with isoflurane (Piramal Critical Care, Shop Med Vet, Mettawa, IL), using a 5% induction and 2–3% maintenance schedule. The reference electrode (8.04 cm²,
Valutrode, Axelgaard Manufacturing Co., Fallbrook, CA) was placed between the shoulders and SignaGel (Parker Laboratories, Fairfield, NJ) electrode gel was used as a conducting medium. A Petflex cohesive bandage (Andover, Shop Med Vet, Mettawa, IL) was wrapped around the midsection of the animal's torso to hold the reference electrode in place. tDCS was applied using a constant current stimulator (Magstim DC-stimulator, NeuroConn, Ilmenau, Germany) for 60 min in addition to a ramp-up/ramp-down period of 10 s. The sham animal group experienced the same experimental set-up as the treatment group, except the constant current stimulator was off. The highest current the stimulator could provide (based on the current-delivery limitations of the stimulator) is 5.0 mA, which was intended to be the highest stimulation group. Throughout the stimulation at the highest intensity, however, the resistance levels of the circuit would increase and cause the stimulator to shut off. Following the automated shut down, the device was restarted at a current intensity 0.5 mA below the previous current intensity. The total duration remained at 60 min, but the current intensity varied during stimulation from 5.0 mA to 2.0 mA for the highest intensity group. The average intensity over the 60-min duration was approximately 2.5 mA and is referred to as the 2.5 mA condition throughout this manuscript.

2.3.2. 10.2 mm² and 5.3 mm² electrode stimulation

Animals were anesthetized with isoflurane (Piramal Critical Care, Shop Med Vet, Mettawa, IL), using a 5% induction and 2–3% maintenance schedule. The reference electrode (surface area: 13 cm²; DIXI medical, Besançon, France) was placed on the chest and either saline or SignaGel was used as the electrolyte (Table 1). A Petflex bandage was wrapped around the animal’s midsection to hold the reference electrode in place. SignaGel was placed into the electrode jacket, and tDCS was applied for 60 min using either a NeuroConn constant current stimulator (Jali Medical Inc., Waltham, MA, USA) or a Caputron constant current stimulator (Caputron Medical, New York, NY, USA) with a Silver Chloride (AgCl) electrode as the anode (Table 1). Note due to the smaller form factor, a change in material was required – none-the-less, the electrode current densities are below electrochemical injury thresholds (Merrill et al., 2005).

2.2. Histological processing and analysis

Immediately following the cessation of stimulation, animals were euthanized by Euthasol injection followed by exsanguination by cardiac perfusion using Phosphate Buffer Saline Solution (PBS) followed by 4% paraformaldehyde (PFA) to fix and preserve tissue. After perfusion, brains were removed and placed into 4% PFA solution for twenty-four hours and then transferred to 30% sucrose. Brains were kept in solution at 4°C until processed. A sliding microtome with a freezing plate (Leica SM2010R) was used for serial collection of 16-μm thick coronal sections and were placed into a cryoprotectant solution and stored at −20°C until further processing.

2.2.1. Tissue staining: H&E, Iba1

Tissue sections were removed from the cryoprotectant solution, washed for 5 min in 1X Phosphate Buffered Saline (PBS) 5 times and mounted onto slides using a 0.1X Phosphate Buffer Solution (PB) with 5% gelatin. Tissue sections were stained with hematoxylin and eosin (H&E). For staining of the ionizing calcium-binding adaptor molecule 1 (Iba1) protein, a set of six sections from each animal was removed from the cryoprotectant solution and washed 5 times for 5 min each in 1X PBS. Endogenous peroxidase was blocked (10% methanol, 3% H2O2 in PBS) for 15 min and slices were then transferred into blocking buffer (PBS solution with 5% Triton-X100 and 2% goat serum) for one hour. Sections were then incubated with the rabbit polyclonal anti-ionized calcium binding adapter molecule antibody (Iba1, Wako Chemicals USA, Inc., Richmond, VA, USA, 1: 3000 dilution) in blocking buffer solution for 2 h. Sections were washed and incubated for 2 h at room temperature with the biotinylated secondary goat anti-rabbit antibody (Vector Laboratories, Burlingame, CA at 1: 3000 dilution) in blocking buffer solution for 2 h. Sections were washed

Table 1

<table>
<thead>
<tr>
<th>Animal</th>
<th>Stimulation (mA)</th>
<th>Anode Electrode</th>
<th>Size (mm²)</th>
<th>Placement (mm Bregma)</th>
<th>Electrode</th>
<th>Size (cm²)</th>
<th>Placement</th>
<th>Area (mm²)</th>
</tr>
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<tbody>
<tr>
<td>3</td>
<td>0.25</td>
<td>DIXI</td>
<td>10.6</td>
<td>−2.5, 2.5 Bilateral</td>
<td>Ag Ring</td>
<td>0.10</td>
<td>Chest</td>
<td>0.00</td>
</tr>
<tr>
<td>1</td>
<td>1</td>
<td>DIXI</td>
<td>5.3</td>
<td>−2.5, 2.5 Left</td>
<td>Ag Ring</td>
<td>0.10</td>
<td>Chest</td>
<td>0.00</td>
</tr>
<tr>
<td></td>
<td>0.25</td>
<td>DIXI</td>
<td>10.6</td>
<td>−2.5, 2.5 Bilateral</td>
<td>Ag Ring</td>
<td>0.10</td>
<td>Chest</td>
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<td>Chest</td>
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again and incubated for 1 h with Vectastain Elite ABC kit (Vector Laboratories, Burlingame, CA, USA) for enzyme conjugation. Finally, sections were stained using 0.05% dianisobenzidine and 0.5% nickel ammonium sulfate and 0.03% H2O2. Tissue sections were then mounted onto slides and allowed to dry at room temperature overnight. Subsequently, sections were counterstained with H&E as previously described for co-localization of lesion and microglial cells. DPX mountant (Sigma, Saint Louis, MO, USA) was used as a coverslip adhesive.

2.2.2. Image analysis
All images were collected using an Olympus BX63 microscope with a Q100 Blue Camera designed for CellSens Dimension Software. Researchers quantifying lesion were blinded to treatment groups throughout tissue processing, image collection and analysis. CellSens Dimension software was used to determine and quantify the area of lesion damage, and a Rat Brain Atlas (Paxinos and Watson, 2007) was used to order sections rostral to caudal according to their distance from Bregma (mm). Lesion was visually identified by parenchymal discoloration and striation in addition to visible gross cellular alterations such as edema, karyolysis and apparent cell loss in the outermost cortical layer. Percent area of lba-1 positive staining was quantified in cortical samples per electrode placement group using 10X magnification. The software used hue, saturation and intensity (HSI) to segment the image fields. Thresholds for object segmentation were established with images of high and low levels of staining to identify positive staining over any background level. These limits were held constant for the analysis of every section in the study. Cell body area and roundness outcomes were measured using the count and measure component of high and low levels of staining to identify positive staining over any background level. These limits were held constant for the analysis of every section in the study. Cell body area and roundness outcomes were measured using the count and measure component of CellSens Dimension software. Briefly, cell bodies were outlined from a sampled cortical area from each section. Placement of the region of interest analyzed was kept constant at all sections and positioned in areas where current-induced lesions were mostly likely to occur. Cell body area (in μm²) and roundness were measured, where perfectly round objects are scored as 1.

2.3. Computational modeling
To determine the effect of various current densities on brain lesion, a state-of-the-art model was constructed from an MRI (7.0 Tesla 70/30 Bruker Biospec) and micro-CT scan (Siemens Inveon) of a female rat (Song et al., 2015).

2.3.1. MRI data collection and segmentation
MRI resolution was 0.282-mm, as previously mentioned (Song et al., 2015). The scans were segmented into 9 tissues: skin, skull, cerebral spinal fluid (CSF), air, gray matter, white matter, hippocampus, cerebellum, and spinal cord. A Rat Brain Atlas (Paxinos and Watson, 2007) was used to identify the hippocampal region of the brain, and the remaining brain regions were appropriately grouped as either gray or white matter.

Manual segmentation was used to generate an initial segmentation of scalp, skin, CSF, air, gray matter, white matter, hippocampus, cerebellum, and spinal cord. Tissue continuity was verified after segmentation by extensively reviewing the data, and further manual adjustments were made to guarantee continuity and improve the segmentation accuracy to closely match the tissue masks to the real anatomy of the rodent using ScanIP 7.0 (Simpleware Ltd, Exeter, UK).

2.3.2. Modeling of tDCS
The three in vivo electrode placement protocols described above were modeled in SolidWorks (Dassault Systemes Corp, Waltham, MA) and imported into ScanIP for meshing. Three montages were modeled: 1) 5 mm × 5 mm anode (surface area: 23.21 mm²) placed on the skull at 0.0 mm Bregma; 2) a single 5.3 mm² electrode (surface area: 4.84 mm²) placed on the skull at –2.5 mm Bregma and 2.5 mm left of the sagittal suture; and 3) two 5.3 mm² electrodes (total surface area: 9.68 mm²) placed on the skull at –2.5 mm Bregma and 2.5 mm bilaterally from the sagittal suture. The first montage used an 8.04 cm² cathode placed between the shoulder blades, while the last two used a 1.3 cm² cathode placed on the chest. An adaptive tetrahedral meshing algorithm was used in ScanIP to generate meshes of 8 × 10⁶ quadratic elements.

FEM models were created in COMSOL Multiphysics 4.3 (COMSOL, Inc., Burlington, MA) using the meshes mentioned above. Models were created using electrostatic volume conductor physics with material conductivities defined as follows (in S/m): skin, 0.465; skull, 0.01; CSF, 1.63; air, 1e⁻¹⁵; spinal cord, 0.126; gray matter, 0.276; white matter, 0.126; hippocampus, 0.126; cerebellum, 0.276; dental cement, 1e⁻¹⁵; electrode jacket, 1e⁻¹⁵; saline, 1.4; and electrode, 5.99e⁷. These conductivity values were taken from a combination of in vitro and in vivo measurements (Datta et al., 2009b; Minhas et al., 2010). Current boundaries were applied to simulate direct current stimulation, and internal boundaries between tissues were assigned the continuity condition (n * (J₁ − J₂) = 0), and the Laplace equation (∇² * (σ/ε)∇V = 0) was solved. The surface of the cathode was grounded (V = 0) while the surface of the anodes for the three stimulation montages had a current density at which lesion first appeared in vivo: 1) 20.0 A/m²; 2) 23.5 A/m²; and 3) 94.2 A/m². All other exterior surfaces were electrically insulated, and brain current density data from the medial cortex between the most superficial regions of the corpus callosum were collected and averaged for analysis. High-resolution models predicted the concentration and distribution of brain current density for each in vivo stimulation group using the three electrode montages. Brain current density is shown for representative coronal sections in false-color, where brain current density values less than 0.34184 A/m² are represented in blue, and brain current density values greater than 7.04619 A/m² are presented in dark red. The maximum threshold was determined by preliminary analysis corresponding the first appearance of lesion in vivo.

2.4. Statistical analysis
Statistical analysis was performed using SigmaPlot (SigmaPlot 11.0, San Jose, CA) for only the 25.0 mm² electrode group unless otherwise noted. One-way analysis of variance (ANOVA) was used for group comparisons followed by Tukey post hoc test. Statistical significance was established with p < 0.05 for all tests. Two-way ANOVA, three-way ANOVA, and one tailed t-tests were also used and are distinguished in the text.

3. Results
3.1. Brain lesion from anodal tDCS
This study aimed to compare the effects of electrode size (25.0 mm², 10.6 mm², and 5.3 mm²) on brain lesion at various electrode current densities using anodal tDCS (Table 1). Computational modeling was used to further analyze and compare histological brain lesion (Fig. 1). No apparent lesions were present using the 25.0 mm² electrode in the sham, 0.15 mA, or 0.3 mA stimulation groups, whereas the 0.5 mA, 1.0 mA, and 2.5 mA stimulation groups had increasingly larger areas of parenchymal discoloration and cellular alterations. Tissue sections from the 0.5 mA stimulation group showed small lesions at the medial cortex on the outer cortical surface (Fig. 1A). Compared to tissue sections obtained from the 0.5 mA group, morphological changes in the 1.0 mA group
were more extensive, penetrating deeper into the cortex and expanding laterally across the outer cortical layer. Lesion in the 2.5 mA group was the most extensive compared to the other stimulation groups (Fig. 1A) (Table 2).

FEM analysis predicts the greatest brain current density (above 7.04619 A/m²) will occur in the medial cortex and in deeper and more lateral brain regions at higher applied currents, such as the corpus callosum, hippocampus, and thalamus (Fig. 1A). Histologically, however, the alterations in morphology from each stimulation group were only observed in cortical brain regions.

Histological lesion was not present in the Sham group using the dual electrode design (total electrode surface area: 10.6 mm²), but lesion was present on the medial cortex at current intensities at and above 0.25 mA (Fig. 1B). Lesion was more extensive for higher stimulation groups, spreading laterally and penetrating deeper into the tissue. Brain current density analysis of the dual electrode design at 0.25 mA showed brain current density was greatest in the superficial cortical regions. At 0.5 mA, brain current density above 0.34184 A/m² was present in the entire medial cortex as well as the medial corpus callosum at 1.0 mA (Fig. 1B). Histological observation of tissue damage was only noted in the superficial cortical regions.

The single 5.3 mm² electrode design did not produce histological lesions in the Sham or 0.043 mA stimulation groups, but morphological changes were detected at 0.5 mA, 0.75 mA, 1 mA, and 2 mA (Fig. 1C). The morphological changes began in the motor cortical region at 0.5 mA, and spread laterally across the outer cortex at higher stimulation groups; there was bilateral lesion present at

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**Fig. 1.** Micrograph representation (4X) of both H&E stained coronal sections from tDCS-induced brain lesion and current distribution at or near 2.5 mm Bregma. Increasing tDCS current increases brain lesion in the rat as well as computationally regardless of electrode size (A, B, C). The anode (red) was placed on the skull, while the cathode (blue) was placed on either the back or chest. A. Using a 25.0 mm² electrode, a minimum electrode current density of approximately 20.0 A/m² induces cortical lesion beginning in the medial superficial cortex, which spreads laterally and penetrates into the cortex as the electrode current density is doubled. The 2.5 mA stimulation group showed a lesion was greater laterally and penetrated further into the cortex compared to the lower stimulation groups. B. Cortical damage begins to appear in the medial superficial cortex with a 0.25 mA (electrode current density: 23.5 A/m²) stimulation using a 10.6 mm² electrode. The lesion area is also greater when the current intensity is increased. C. Using a 5.3 mm² electrode, lesion began to appear at 0.5 mA (electrode current density: 94.2 A/m²) beginning in the motor cortex and expanded as the current intensity was increased. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
2.0 mA (Fig. 1C; bottom). The greatest brain current densities (above 7.04619 A/m²) for the single 5.3 A/m² electrode occurred unilaterally in the cortex and corpus callosum with an applied current of 0.5 mA (Fig. 1C) and spread bilaterally at greater applied current intensities. Brain current densities above 0.34184 A/m² were bilaterally present in the cortex and unilaterally in the CA1 region of the hippocampus using 1.0 mA. At 2.0 mA, brain current densities above 7.04619 A/m² were seen bilaterally in the corpus callosum, hippocampus, and superficial thalamus (Fig. 1C).

Total brain lesion was quantified by measuring and averaging the area of parenchymal alterations for each stimulation group (Fig. 2). Using a 25.0 mm² electrode, brain damage was present at 20.0 A/m² (average area = 0.50 mm² ± 0.304), 40 A/m² (average area = 15.77 mm² ± 2.365), and 100 A/m² (average area = 40.92 mm² ± 0.937). Animals receiving 40 A/m² and 100 A/m² had a significantly greater lesion area compared to all other groups (One-way ANOVA: F (5) = 178.8, p < 0.001, for both groups). No statistically significant differences occurred between the sham, 0.15 A/m², 0.3 A/m², or 0.5 A/m² groups (Fig. 2A).

Cortical lesion at similar electrode current densities using different electrode sizes was evaluated and appears to be positively correlated with both electrode size and electrode current density (Fig. 2B). Across the three electrode groups, lesion was significantly different at an electrode current density of 100.0 A/m² (One-way ANOVA: F (1) = 519.1, p < 0.001: 25.0 mm² = 360.0 mm² ± 39.2, 40.0 mm² = 100.0 mm² ± 9.0, 100.0 mm² = 40.92 mm² ± 7.4). Across the three stimulating current intensities, lesion was significantly different at an electrode current density of 100.0 A/m² (Two-way ANOVA: F (2) = 18.543, p < 0.001) in tissue sections that were between +2.0 mm and −6.5 mm from Bregma using a 25.0 mm² electrode (black lines above Fig. 3A).

Brain lesion area was quantified in coronal slices based on distance from Bregma (Fig. 3). Lesion area in tissue sections that were between +2.0 mm and −6.5 mm from Bregma (two-way ANOVA; p < 0.02); 2) 2.5 mA and 1.0 mA from +0.5 mm to −6.5 mm Bregma (two-way ANOVA; p < 0.02); and 3) 1.0 mA and 0.5 mA from +2.0 mm to −4.5 mm (two-way ANOVA; p < 0.05). Lesion area differences based on distance from Bregma were also present for the 10.6 mm² (Fig. 3B) and 5.3 mm² (Fig. 3C) electrodes. Using FEM rodent models to predict the average brain current density in coronal sections at 0.5 mm increments from +5 mm Bregma to −6.5 mm Bregma, a larger electrode corresponds to a greater average brain current density across all coronal sections at approximately 100 A/m², (Fig. 3D).

We compared the reliability of electrode current density and average brain current density from models for predicting brain lesion area across the three electrode sizes (Fig. 4). For each electrode, lesion area increased linearly (R² = 0.2442) with electrode current density, but the electrode’s sensitivity (slope) to lesion varied (Fig. 4A). At an electrode current density of 100 A/m², the lesion area for the 5.3 mm², 10.6 mm², and 25.0 mm² electrodes were 6.1 mm², 18.9 mm², and 40.9 mm², respectively. Lesion area also increased linearly with brain current density (R² = 0.8139) and sensitivity to lesion development was similar across the three electrode sizes (Fig. 4B).

3.2. Iba1 immunoreactivity

To investigate whether tDCS affected inflammation in the brain, tissue sections were stained with an antibody against Iba1, a protein that is present in macrophages and microglia and upregulated when in their active state (Hanisch and Kettenmann, 2007). There were no differences in the percentage of positive Iba1 staining in the cortical areas immediately below the electrode (p = 0.50, data not shown). However, microglial morphology tended towards a more activated phenotype, with enlarged cell bodies and thickened processes, in stimulation groups that produced lesion (0.5 mA and higher; Fig. 5). No statistical differences occurred in Iba1 positive cell body surface area (p = 0.61, Fig. 5E), but there was a statistical significance between current intensity and cell body roundness. At 1.0 mA, microglial cell bodies were significantly more rounded compared to all other groups (p = 0.02, Fig. 5F).

4. Discussion

We detected cortical lesions during anodal stimulation using an electrode current density of 20.0 A/m², which is well below the commonly referenced cathodal electrode current density threshold
of 142.9 A/m² (Liebetanz et al., 2009). Histological analysis did not reveal any lesion at or below 0.3 mA using a 25.0 mm² electrode, corresponding to an electrode current density of 12 A/m². This was the greatest applied current intensity that could be administered for 60 min without evidence of tissue damage using the 25.0 mm² electrode.

The lesions from our study expanded laterally and penetrated deeper into the tissue as the electrode current density was increased (Fig. 1). However, the lesion induced from the Liebetanz study appeared to penetrate more deeply and less laterally with increasing electrode current densities, which may reflect differences in electrode montage (including position of the return electrode) or polarity specific mechanisms of injury (anodal vs. cathodal tDCS). Once correcting for average brain current density, our finding of comparable dose response across changes in electrode material, shape, and number are consistent with brain current flow, rather than superficial electrochemistry, causing injury.

Microglial cells have been shown to increase after sub-lesion tDCS when assessed by measuring Iba1 + cells three days after stimulation (Rueger et al., 2012), possibly providing evidence of a more sensitive marker of tDCS related injury than lesion. Our exploratory analysis of microglia and macrophage immunoreactivity showed no significant changes in the percentage of immunoreactive cells present in the medial cortex following a 60-min stimulation (Fig. 5), but higher current densities were likely already shifting the microglial phenotype from its resting to active state (Davalos et al., 2005; Stence et al., 2001). Despite the untimely approach for histological analysis of microglia, our data did show an increase in cell body roundness at 1.0 mA immediately after tDCS (Fig. 5F), suggesting either tDCS or the induction of lesion shifted microglia to its active. This study was not able to make a distinction histologically due to our flawed approach, but it has been shown microglial changes associated with neuroinflammation can occur at approximately 31.8 A/m² using anodal tDCS (Gellner et al., 2016).

Although investigation of multiple tDCS parameters is possible utilizing a rodent model, there are translational limitations. One of those limitations is the ability to easily approximate and translate...
Fig. 3. Averaged lesion area by brain tissue section and electrode size according to its distance from Bregma. The average brain lesion was quantified in 0.5 mm increments, with positive numbers indicating the distance rostral from Bregma, and negative numbers indicating the distance caudal from Bregma. A, B, C. Local differences in cortical lesion for the 25.0 mm² (A), 10.6 mm² (B), and 5.3 mm² (C) electrodes, separated by the stimulation group. FEM images show rostro-caudal changes in brain current density at representative coronal sections. The black line at the bottom of the figure indicates the location of the electrode. Significant differences between current intensity and the average lesion produced at specific locations from Bregma are indicated at the top of (A). D. FEM analysis of brain current density in the medial cortex corresponding with in vivo tissue sections (mm Bregma). The box around ~2.5 mm Bregma indicates the location of the FEM images below the figure.

Fig. 4. Effect of electrode current density and brain current density on total brain lesion area. Electrode size was found to be a relevant variable when measuring tissue lesion obtained from brains stimulated at different current densities. A. Larger electrodes induced more lesion at similar electrode current densities. B. Across electrode sizes, cortical lesion was related to predicted brain current density (R² = 0.8139).
tDCS dosages between small, 300–500 g animals and large 70 kg humans. However, FEM analysis of in vivo rodent experiments has attempted to translate rodent lesion studies to determine a lesion threshold in humans by developing a scaling factor, with the most conservative prediction of injury from rodent FEM models occurring in humans at 6.3 A/m² (Bikson et al., 2016). Understanding and translating minimum lesion thresholds in rodents could allow a greater range of safe stimulation protocols for human applications, potentially enhancing benefits already seen in human tDCS. Our data show damage can occur at a level of 20.0 A/m² (Fig. 4). However, this threshold is 10-fold higher than the typical electrode current density of 0.28–2.0 A/m² utilized in human studies (Poreisz et al., 2007; McKinley et al., 2013). Moreover, our model is hypersensitive compared to the clinical case where electrodes are placed on the skin (not epicranial) and the distance between the electrode and brain is significantly higher. Taken together with our consideration of appropriate safety metrics (above), our threshold for electrode current density in rat models may be conservatively considered (with a safety factor) applicable to the human case.

5. Conclusions

Our anodal animal tDCS study shows tissue damage can occur at 20.0 A/m², which is below the often referenced damage threshold of 142.9 A/m² using cathodal tDCS (Liebetanz et al., 2009). In considering the relevance of animal studies to human safety guidelines, consideration should also be given to the role of electrode size, location, and brain region of interest. Consistent with prior studies, our data does not suggest current clinical tDCS are injurious.

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