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### **ORIGINAL RESEARCH**



## **Perceptibility and Pain Thresholds in Low‑ and High‑Frequency Alternating Current Stimulation: Implications for tACS and tTIS**

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### **Abstract**

Transcranial electrical stimulation (tES) has emerged as a promising tool for neuromodulation, but its application is often limited by the discomfort associated with higher stimulation intensities. Newer variants like transcranial temporal interference stimulation (tTIS) utilize high-frequency alternating currents ( $\geq$  500 Hz) to penetrate deeper brain regions while mitigating perceptual discomfort. This study sought to examine sensation and pain thresholds across various stimulation frequencies of alternating currents, aiming to explore the boundaries of comfortable intensities. Additionally, we sought to evaluate the efcacy of an anesthetizing topical cream in increasing participant comfort and potentially extending the range of tolerable stimulation levels. We recruited 37 participants and applied alternating current stimulation to the head at various frequencies (10 Hz, 20 Hz, 500 Hz, 1000 Hz, and 2000 Hz) to determine intensity-dependent perception and pain thresholds. Additionally, thresholds were determined under the infuence of a topical anesthetic. Our fndings confrm that as stimulation frequency increases, perceptibility decreases, with higher frequencies allowing a manyfold increase in stimulation intensity before becoming perceptible or causing pain. Additionally, the anesthetizing cream was efficacious in further reducing perceptibility and pain sensations across all frequencies. This study lays the groundwork for future research by establishing comfortable limits for stimulation intensities, particularly in the context of high-frequency stimulation. The reduced perceptibility of high-frequency stimulation, coupled with the efectiveness of anesthetizing creams, enables the administration of higher stimulation intensities for more potent neuromodulatory interventions without causing discomfort.

**Keywords** Transcranial alternating current stimulation (tACS) · Transcranial electrical stimulation (tES) · Transcranial temporal interference stimulation (tTIS) · Somatosensory perception · Nociception · Topical skin anesthetization

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## **Introduction**

Transcranial electrical stimulation (tES) holds signifcant promise for treating various psychological and neurological conditions. Research has explored its potential applications in depression (Arul-Anandam & Loo, [2009](#page-10-0); Vanderhasselt et al., [2015](#page-12-0)), stroke (Convento et al., [2016;](#page-10-1) Khan et al., [2022](#page-11-0); Solomons & Shanmugasundaram, [2019](#page-12-1)), and fatigue (Lin-nhoff et al., [2019;](#page-11-1) Shirvani et al., [2021](#page-12-2)) among others (Cho et al., [2022\)](#page-10-2). Beyond therapeutic applications, there is growing interest in leveraging this technology for neurofacilitation to, for example, enhance motoric performance (Chang, [2022;](#page-10-3) Friehs et al., [2022;](#page-10-4) Perrey, [2023\)](#page-12-3), working memory (Röhner et al., [2018;](#page-12-4) Zaehle et al., [2011\)](#page-13-0), and perception (He et al., [2022;](#page-11-2) Wang et al., [2020](#page-13-1)) in healthy individuals. However, the efficacy and reliability of these interventions in human participants is often constrained by the intensity of stimulation that can be comfortably administered. As

the intensity increases, participants report sensations that evolve from a mere tingling to pronounced discomfort such as prickling sensations and with sufficient intensity even burning or pain sensations (Fertonani et al., [2015](#page-10-5); Hsu et al., [2021;](#page-11-3) Khadka et al., [2020](#page-11-4); Kuhn et al., [2010](#page-11-5); McFadden et al., [2011](#page-11-6); Palmer et al., [1999](#page-12-5); Paneri et al., [2016](#page-12-6); Zeng et al., [2019\)](#page-13-2).

Consequently, tES studies in humans usually stay at or below a stimulation intensity of 2 mA (Antal & Paulus, [2013;](#page-10-6) Bikson et al., [2009\)](#page-10-7), with only very few studies employing higher intensities of up to 4 mA (e.g. Chhatbar et al., [2017](#page-10-8); Hsu et al., [2023\)](#page-11-7), albeit this being still a safe stimulation intensity (Antal et al., [2017;](#page-10-9) Bikson et al., [2016](#page-10-10); Chhatbar et al., [2017;](#page-10-8) Matsumoto & Ugawa, [2017](#page-11-8); Nitsche & Bikson, [2017](#page-12-7)). This may be due to participant compliance issues caused by uncomfortable cutaneous sensations. Sensitivity to stimulation varies based on the method of electrostimulation and specifc parameters of stimulation (Ambrus et al., [2010](#page-10-11); Fertonani et al., [2015](#page-10-5)) such as electrode size (Kuhn et al., [2010](#page-11-5); Turi et al., [2014\)](#page-12-8) or waveform of stimulation (Baker et al., [1988;](#page-10-12) Hsu et al., [2021\)](#page-11-3). There is an ongoing debate about the role that current density plays in perceivability. On one hand, larger electrodes result in a lower current density, which means less current impacts each somatosensory receptor, potentially reducing perceivability (Alon et al., [1994](#page-10-13); Verhoeven & van Dijk, [2006\)](#page-13-3). On the other hand, larger electrodes cover a greater area, leading to a spatial summation efect—namely, the recruitment of more somatosensory receptors to fire, thereby enhancing perceivability (Higashiyama & Tashiro, [1990](#page-11-9); Nielsen & Arendt-Nielsen, [1997\)](#page-12-9). Further, the stimulation duration infuences perceptibility due to adaptation processes attenuating sensations. This was, for example, leveraged by Khadka et al. [\(2020\)](#page-11-4) by gradually increasing intensity over the course of the stimulation in an adaptive procedure, enhancing participant comfort at higher stimulation intensities. When using alternating currents, another key parameter in perceptibility is frequency. Frequencies below 100 Hz, commonly used in transcranial alternating current stimulation (tACS), are more perceptible than higher frequencies (Hsu et al., [2021](#page-11-3); Turi et al., [2013](#page-12-10); Ward & Robertson, [1998](#page-13-4); Zeng et al., [2019](#page-13-2)) due to the spectral specifcity of neurons limiting responsiveness to high frequencies (Anderson & Munson, [1951](#page-10-14); Hawkes & Warm, [1960;](#page-11-10) Hutcheon & Yarom, [2000;](#page-11-11) Palmer et al., [1999](#page-12-5)).

A recent advancement in the area of tES research is transcranial temporal interference stimulation (tTIS). Its efficacy in modulating brain activity has been demonstrated in animal studies, highlighting it as a promising new electrostimulation method (Acerbo et al., [2022](#page-10-15); Carmona-Barrón et al., [2023](#page-10-16); Grossman et al., [2017](#page-10-17); Liu et al., [2023;](#page-11-12) Missey et al., [2021;](#page-11-13) Song et al., [2021a,](#page-12-11) [2021b;](#page-12-12) Sunshine et al., [2021](#page-12-13); Zhang et al., [2022](#page-13-5)). TTIS stands out among other tES methods due to its enhanced stimulation depth. This is achieved through the utilization of two high-frequency alternating currents  $(\geq 500 \text{ Hz})$ , creating an amplitude-modulated signal at the intersection of the two felds (Grossman et al., [2017;](#page-10-17) Karimi et al., [2019](#page-11-14); Mirzakhalili et al., [2020;](#page-11-15) Song, [2019](#page-12-14)). This signal is believed to lead to modulation of neuronal activity via entrainment effects. By carefully configuring electrode placements and adjusting the current ratios of the felds, the point of interference—and consequently, the stimulation focus—can be directed deeper into brain regions. This offers the potential to non-invasively achieve neuronal modulation in deep brain regions. However, evidence supporting tTIS's efficacy in humans remains limited, with some studies even casting doubt on its feasibility (Budde et al., [2023;](#page-10-18) von Conta et al., [2022;](#page-13-6) Iszak et al., [2023](#page-11-16)). A network modeling study by Negahbani et al. [\(2018\)](#page-12-15) as well as single neuron modeling studies (Mirzakhalili et al., [2020](#page-11-15); Wang et al., [2023](#page-13-7)) and an in vitro study by Esmaeilpour et al. ([2021\)](#page-10-19) indicate that higher tTIS intensities compared with tACS are needed to achieve similar neuronal modulation. However, other studies argue that the efficacy of tTIS is largely based on network mechanisms (Cao, [2018](#page-10-20); Martinez et al., [2023](#page-11-17)) as well as a gradual depolarization of neurons over time (Cao et al., [2020\)](#page-10-21). In this context, tTIS offers a significant advantage due to its high-frequency stimulation, which makes the stimulation intensities less perceptible, allowing the application of higher intensities without discomfort. This leads to a growing interest in increasing the stimulation intensity to fully leverage the potential of tTIS, positioning it as a promising method for both treatment and research in non-invasive deep brain stimulation (Grossman et al., [2018\)](#page-10-22).

Our study aimed to establish tolerable stimulation intensity ranges for both low-frequency (10, 20 Hz) and highfrequency (500, 1000, 2000 Hz) alternating currents. We selected low frequencies due to their common use in tACS studies and high frequencies for their relevance in tTIS research. By measuring pain thresholds at these frequencies, we sought to provide a reference for determining safe and tolerable maximum stimulation intensities for future tTIS and tACS research. Similarly, with perception thresholds, our goal was to identify sub-perception intensity levels crucial for ensuring efective blinding.

Studies have advocated for the use of anesthetizing skin creams in tES studies (Antal et al., [2017](#page-10-9); Guleyupoglu et al., [2014](#page-10-23); Liu et al., [2018;](#page-11-18) McFadden et al., [2011\)](#page-11-6), ofering multiple advantages. Foremost, these creams allow for the application of higher stimulation intensities than typically feasible (McFadden et al., [2011](#page-11-6)). The anesthetized skin diminishes pain perceptions, enhancing participant comfort even at higher stimulation levels. Another signifcant beneft is the improved blinding of participants. Many studies compare verum (true) stimulation with sham (false/placebo/ control) stimulation, where sham stimulation involves applying electrical currents that mimic the cutaneous sensations

(such as tingling or itching) of verum stimulation but difer in key aspects. These diferences can include being turned of after a brief period, utilizing a diferent frequency, or targeting a diferent region. The purpose of sham stimulation, as opposed to not applying any stimulation at all, is to create a condition that feels similar to the verum condition for participants, making it harder for them to diferentiate between the two types of stimulation and maintain participant blinding. Using anesthetization as a complementary approach allows to reduce perceptibility of the verum stimulation, thus facilitating participant blinding (Sheffield et al., [2022\)](#page-12-16). A third advantage is the possibility to control for somatosensory perception as a confounding factor in interpretation of stimulation results. Studies have demonstrated that changes in brain activity can be achieved by somatosensory entrainment due to cutaneous sensations of stimulation (Asamoah et al., [2019](#page-10-24); Spooner et al., [2022\)](#page-12-17). Thus, recent studies have begun to control for cutaneous sensations to eliminate somatosensory entrainment as a possible confounding factor (Koganemaru et al., [2020](#page-11-19); Turi et al., [2020](#page-12-18)).

Consequently, the second aim of our study was to investigate the infuence of a topical anesthetic skin cream by quantifying its impact on somatosensory perception. This was measured by observing changes in the perception and pain thresholds resulting from the application of the anesthetization. Specifcally, we aimed to evaluate the anesthetization's efficacy across various stimulation frequencies, to further support it as a future tool in studies to reduce participants' awareness of stimulation conditions and increase the limits of comfortable stimulation intensities.

## **Methods**

#### **Participants**

We recruited 37 participants (12 male, 25 female, mean age=23.6 years,  $SD = 3.93$  years, range=18–36 years) for this study. Participants with a history of epileptic seizures, psychiatric or neurological disorders, metal or electric implants in the head, or those on medication afecting the central nervous system were excluded. Prior to the experiment, all participants were briefed about the procedure, potential risks of electrostimulation, and provided written informed consent. The study received approval from the University Clinic of Magdeburg's local ethics committee and adhered to the Declaration of Helsinki guidelines.

#### **Experimental Design**

We developed the experimental paradigm using MATLAB (Version 2020a, The MathWorks Inc., Natick, MA, USA) and the Psychtoolbox 3 (Kleiner et al., [2007](#page-11-20)).

Prior to the experiment, topical skin anesthetization cream with 25 mg/g lidocaine and 25 mg/g prilocaine (Anesderm, Pierre Fabre Dermo-Kosmetic GmbH) was applied to one side of each participant's head, while the opposite side remained untreated, serving as a non-anesthetized control. The side of anesthetization was counterbalanced among participants. The anesthetizing cream remained in place for 15 min to ensure its full efect. Afterwards, it was removed with a dry tissue to be replaced by an electrically conductive gel for the following stimulation.

The experiment comprised two task blocks: the somatosensory perception threshold block and the pain threshold block, with their order counterbalanced among participants to mitigate efects of task order. In the somatosensory perception block, participants were tasked to indicate if they experienced any cutaneous sensations such as tingling or itching during stimulation. During the pain threshold block, participants had to indicate if the stimulation had induced pain in the form of stinging or burning sensations. We emphasize that the staircase procedure only gradually increased stimulation intensity over trials and decreased if pain was reported. This ensured that participants only ever experienced mild pain sensations.

A block was comprised of 10 conditions: Anesthetization (yes/no) by Frequency (10 Hz, 20 Hz, 500 Hz, 1000 Hz, 2000 Hz). While the frequency variable was operationalized through the frequency of the applied alternating current, anesthetization was operationalized based on the location of stimulation—whether stimulation was administered to the anesthetized side of a participant's head or the untreated side. Each condition had its own staircase, resulting in ten individual staircases during a block. We adopted a random interleaved staircase design, wherein each consecutive stimulation was based on a randomly selected condition (see Fig. [1](#page-4-0)A). This design was implemented specifically to minimize habituation effects, which can occur when sensitivity to stimulation decreases due to the same condition being presented consecutively. By ensuring a varied sequence of frequencies and intensities, we aimed to maintain participant sensitivity and mitigate diminishing responses to the stimulation. The total number of trials needed for a staircase to conclude, depended on how many trials were needed to determine a threshold, i.e., to fulfll one of the conclusion criteria (see "Staircase Procedure" section). After a staircase was concluded, its condition was not presented again. A block ended, when all 10 staircases were concluded.

The starting amplitudes for each condition's staircase varied based on the frequency, aligning with the premise that lower frequencies are generally more perceivable, necessitating it to begin at low intensities. Conversely, higher frequencies are less perceivable, warranting a start at higher intensities to avoid the need for presentation of many unperceivable trials before reaching an intensity level relevant for those



<span id="page-4-0"></span>**Fig. 1 A** Consecutive trials exemplifed. This illustrates how the stimulation intensity for condition 4 (1000 Hz, not anesthetized) changed depending on the subject's answer. After each trial, a question was posed to the subject (Perception block: "Did you feel the stimulation?"; Pain block: "Did you experience pain during the stimulation?"). Indicating a "No" via button press led to an increase in

frequencies. The starting amplitudes for the conditions were as follows: 10 and 20 Hz: 0.2 mA, 500 Hz: 1 mA, 1000 Hz: 1.5 mA, 2000 Hz: 2 mA.

A trial was comprised of a 3-s countdown which was displayed on a screen in front of the participant, followed by 7 s of stimulation and ended with a self-timed period where participants had to indicate via button press if they felt the stimulation (perception threshold block) or felt pain during stimulation (pain threshold block).

Upon concluding the experiment, participants completed a questionnaire regarding potential side efects, such as lasting pain or headaches (Brunoni et al. 2011). They were then debriefed about the study's objectives and compensated with either course credit or monetarily.

## **Staircase Procedure**

We employed an adaptive 1-up-1-down staircase procedure to determine thresholds (Cornsweet, [1962;](#page-10-25) Leek, [2001\)](#page-11-21). This method estimates the stimulation strength at which participants would perceive the stimulation (perception threshold) or experience pain (pain threshold) in 50% of trials, by dynamically adjusting stimulation intensities. For example, at the end of a trial, if participants answered with "yes" to the post-stimulation question ("did you feel the stimulation" or "did you experience pain during the stimulation"), the intensity for the future presentation of that condition was decreased by 20%, based on the last given intensity. Conversely, a "no" lead to a 20% increase of intensity. A signifcant beneft of adaptive staircase procedures is their ability to ensure a high sampling density at and around the most relevant stimulation intensity. This approach prioritizes sampling near the intensity levels where a reversal of answers occurs, while avoiding unnecessary presentation of intensities which are far from the relevant range.

intensity for future presentations of that condition, whereas a "Yes" decreased the intensity. **B** Electrode montage. In this example, blue electrodes represent the side where anesthetization was applied, while grey electrodes indicate the untreated side. The side on which anesthetization was administered varied, being counterbalanced across participants

A condition's staircase could conclude in either of the following ways:

- (1) After a total of fve reversals of "yes/no" responses in a condition. Reversals did not have to be consecutive. The threshold was determined by averaging the last three alternating values in that staircase. To ensure a sufficient number of trials and data collection for accurately pinpointing the thresholds, we chose to require five reversals. We opted to average only the last three reversals because the initial reversals are usually further from the true threshold. In contrast, later reversals tend to be closer, making them more indicative of the actual thresholds (Leek, [2001](#page-11-21)).
- (2) If a condition was presented with a stimulation intensity of 4 mA and received a "no" answer for the third time. Note that "no" answers did not have to be consecutively but were counted over the whole block. This suggests that the actual threshold for inducing sensation or pain in that condition lies above our upper limit of 4 mA. For the purposes of data analysis, we treated these instances as having a threshold of 4 mA, acknowledging that this represents the maximum intensity tested and not the actual somatosensory perception or pain threshold.
- (3) If a condition has been presented for the 20th time during that block. For data analysis, the threshold was assumed as the last three alternating values.

## **Electrical Stimulation**

Stimulation was delivered using two independent batterydriven neuroConn Stimulator systems (Advanced DC-Stimulator Plus for temporal interference stimulation, neuroConn GmbH, Ilmenau, Germany) which were connected via a digital to analog converter (Ni USB-6212, National Instruments, Austin, TX, USA) to a PC and controlled by it using the remote mode of the stimulators. A custom MAT-LAB script (Version 2020a, The MathWorks Inc., Natick, MA, USA) was used to generate and send the stimulation signal. For stimulation, Ag–AgCl electrodes with a 12-mm diameter (Brain Products, Gilching, Germany) were used, which were affixed to an EEG cap (Easycap, Brain Products, Gilching, Germany). Using the international 10–10 system, electrodes were placed at positions F1 and CP1 for the left stimulation site and at F2 and CP2 for the right stimulation site (see Fig. [1](#page-4-0) B). To increase electrode to skin conductivity, we applied a conductive paste (Abralyt 2000 abrasive electrolyte-gel, Brain Products, Gilching, Germany), ensuring impedances remained below 5 kΩ.

Stimulation frequencies were 10 Hz, 20 Hz, 500 Hz, 1000 Hz, and 2000 Hz with a maximum possible intensity of 4 mA. Our rationale for these was as follows: the 2000 Hz and 1000 Hz frequencies were used by Grossman et al. ([2017\)](#page-10-17) in their tTIS study and thus refect proven and efficacious stimulation frequencies. The 500 Hz stimulation frequency is considered to be the lowest frequency feasible for tTIS (Grossman et al., [2017](#page-10-17)). Additionally, we selected 10 and 20 Hz, standard tACS stimulation frequencies prevalent in many tACS studies, to serve as a reference point to benchmark perception diferences between low- and highfrequency stimulation.

#### **Data Analysis**

We performed our statistical analysis using Jamovi version 2.3 (The Jamovi Project 2024). To analyze diferences between thresholds, we conducted repeated-measures analyses of variance (rmANOVAs) separately for perception- and pain thresholds using within-subject factors Anesthetization (Yes, No) and Frequency (10 Hz, 20 Hz, 500 Hz, 1000 Hz, and 2000 Hz). To ensure that factors in our rmANOVAs conformed to the sphericity assumption, we conducted a Mauchly's test.

For rmANOVA results, we report partial eta squared  $(\eta^2$ <sub>p</sub>) to focus on efect size within our chosen design, as well as generalized eta squared  $(\eta^2 G)$  to facilitate comparing effect sizes across studies.

#### **Results**

The descriptive outcomes for perception and pain thresholds are illustrated in Fig. [2](#page-6-0) and detailed in Table [1](#page-7-0). In addition, Table [1](#page-7-0) also presents the intensity values converted to current density, based on the 12 mm electrodes employed in our study.

Statistical analysis revealed violations of the sphericity assumption for perception thresholds [*Frequency*:  $\chi$ 2 (9) = 166.67, *p* < 0.001,  $\varepsilon$  = 0.49; *Frequency* \* *Anesthetization*:  $\chi$ 2(9) = 132.29, *p* < 0.001,  $\varepsilon$  = 0.59] as well as for pain thresholds [*Frequency*:  $\chi$ 2(9) = 105.19, *p* < 0.001,  $\varepsilon = 0.47$ ; *Frequency* \* *Anesthetization*:  $\chi$  2(9) = 47.96,  $p < 0.001$ ,  $\varepsilon = 0.66$ . Given these violations, we adjusted the degrees of freedom using the Greenhouse–Geisser correction to make the test more conservative and control for type I errors.

Results of the rmANOVAs revealed a signifcant main efect of *Frequency* on perception- [*F*(1.95,70.09)=479.16,  $p < 0.001$ ,  $\eta_0 = 0.930$ ,  $\eta_0 = 0.796$ ] and pain thresholds  $[F(1.86,67.11) = 588.70, p < 0.001, \eta_{p} = 0.942,$  $\eta_0 = 0.844$ ]. This indicates that higher frequencies lead to increased thresholds for perception and pain, implying that higher stimulation frequencies induce less cutaneous sensations than lower frequencies. This was confrmed using posthoc analyses: as frequencies increased, so did the thresholds for both perception and pain. This was demonstrated by signifcant increases across all frequency comparisons for perception [all comparisons  $t(36) > 3.22$ ,  $p_{\text{tukev}} < 0.001$ ] and pain thresholds [all comparisons  $t(36) > 5.32$ ,  $p_{\text{ntkey}} < 0.001$ ], with the sole exception being the pain thresholds between 10 and 20 Hz frequencies, which did not differ significantly  $[t(36) = 2.26, p_{\text{tukey}} = 0.181]$ . Additionally, a signifcant main efect of *Anesthetization* on thresholds was revealed, again for both perception  $[F(1,36) = 19.90]$ ,  $p < 0.001$ ,  $\eta_0 = 0.356$ ,  $\eta_0 = 0.086$ ] and pain thresholds  $[F(1,36) = 26.16, p < 0.001, \eta_{p} = 0.421, \eta_{q} = 0.077]$ , indicating that the anesthetization reduced cutaneous sensations to the stimulation in both measures. Additionally, an interaction *Frequency \* Anesthetization* was observed for both perception  $[F(2.36,84.85)=6.08, p=0.002, \eta_0^2=0.144,$  $n_0^2 = 0.031$ ] and pain thresholds  $[F(2.65, 95.31) = 5.96,$  $p=0.001$ ,  $\eta_{p}=0.142$ ,  $\eta_{G}=0.023$ ]. This was due to the efficacy of anesthetization varying based on the stimulation frequency, with higher stimulation frequencies (500 and 1000 Hz) beneftting more from anesthetization than lower frequencies (10 and 20 Hz). At 2000 Hz, however, our analysis found no diference in thresholds between anesthetized and non-anesthetized conditions [perception:  $t(36) = 1.47$ ,  $p_{\text{tukey}} = 0.894$ ; pain:  $t(36) = 1.09$ ,  $p_{\text{tukey}} = 0.983$ ]. It is important to note that this absence of an anesthetization efect at 2000 Hz is a result of many subjects reaching our study's maximum stimulation of 4 mA in both anesthetized and nonanesthetized conditions, refecting a limitation of our study setup.

Exploratory, to mitigate potential confounding efects of task order, we repeated the previous rmANOVAs including *Task Order* (pain task frst, perception task frst) as a between-subject factor. The results indicated that *Task Order* did not significantly affect perception thresholds  $[F(1,35) = 0.225, p = 0.638, \eta_{p}^2 = 0.006, \eta_{q}^2 = 0.002]$  or pain thresholds  $[F(1,35)=0.015, p=0.902, \eta_{p}=0.010,$  $\eta_0 = 0.004$ ]. Furthermore, to analyze a potential difference



<span id="page-6-0"></span>**Fig. 2** Stimulation intensity thresholds. **A** Perception thresholds increase with stimulation frequency, indicating that high-frequency stimulation induces less perception. Conditions with topical anesthetization display higher thresholds. **B** Anesthetization allows for higher stimulation intensities before reaching perception thresholds across all frequencies, as indicated by *Anesthetization Gains*  $(={\rm Threshold}_{\rm WithAnesthetization}~-~{\rm Threshold}_{\rm WithoutAnesthetization}).~~{\rm Notably},$ the reduced anesthetization gain at 2000 Hz is due to some participants not perceiving any stimulation at high frequencies, both with

between sexes, we repeated the rmANOVAs again with *Sex* (male, female) as a between-subject factor. This analysis revealed a signifcant efect of *Sex* on pain thresholds  $[F(1,35)=4.34, p=0.045, \eta_{p}=0.110, \eta_{G}=0.048]$ . Additionally, a signifcant interaction efect *Frequency \* Sex* was observed  $[F(1.99, 69.50) = 5.346, p = 0.007, \eta_0^2 = 0.133,$  $\eta_0^2 = 0.046$ , which is descriptively explained by male participants being able to tolerate higher intensities at high frequencies, though this did not reach signifcance in posthoc analysis [male vs. female; 10 Hz: *Mean Diff*=0.01 mA,  $t(35)=0.085$ ,  $p_{tukey}=1.000$ ; 20 Hz: *Mean Diff*=0.08 mA,  $t(35)=0.665$ ,  $p_{\text{tukey}}=1.000$ ; 500 Hz: *Mean Diff*=0.71 mA, *t*(35)=0.665, *ptukey*=0.250; 1000 Hz: *Mean Dif*=0.42 mA, *t*(35)=2.413, *ptukey*=0.349; 2000 Hz: *Mean Dif*=0.12 mA,  $t(35) = 1.322$ ,  $p_{\text{tukev}} = 0.942$ . No other effects reached

and without anesthetization. For analysis, their threshold was standardized to 4 mA, aligning anesthetized and non-anesthetized conditions and reducing observed anesthetization gains. This refects study constraints more than a decrease in anesthetization efectiveness at higher frequencies. **(C)** Pain thresholds: Higher stimulation frequencies correlate with lower pain thresholds, refecting the pattern seen in perception thresholds. **(D)** Anesthetization gains for pain thresholds: These gains follow a similar trend to perception thresholds, with the same high-frequency constraints previously noted

signifcance. Further, no signifcant efect of *Sex* on perception thresholds  $[F(1,35)=0.097, p=0.758, \eta_0^2=0.003.,$  $\eta_0 = 0.001$ ] could be observed.

#### **Discussion**

Current human applications of tES are limited by the maximum intensity that participants can comfortably tolerate. Even though higher intensities are considered safe (Antal et al., [2017;](#page-10-9) Bikson et al., [2016](#page-10-10); Chhatbar et al., [2017](#page-10-8); Matsumoto & Ugawa, [2017](#page-11-8)), the discomfort from skin sensations or pain often restricts their use. The recent introduction of tTIS (Grossman et al., [2017\)](#page-10-17) leverages the fact that high-frequency alternating currents are

<span id="page-7-0"></span>



 $N_{lim}$ , number of participants reaching the maximum stimulation intensity threshold (4 mA) in that condition;  $N_{Total}$ , total number of participants; *CD*, current density

less perceivable (Hsu et al., [2021](#page-11-3); Hutcheon & Yarom, [2000;](#page-11-11) Turi et al., [2013;](#page-12-10) Zeng et al., [2019\)](#page-13-2), thus potentially allowing for the application of higher stimulation intensities. In our study, we sought to quantify the sensation and pain thresholds for various high-frequency alternating current stimulations used in tTIS, aiming to explore the potential upper limits of intensity for this innovative electrostimulation technique. Furthermore, we explored the possibility to push these limits by employing an anesthetizing topical cream, to reduce cutaneous sensations caused by electrical stimulation. To be able benchmark the perception of these high frequencies against classical methods of electrostimulation, we've also quantified perception of low-frequency alternating currents as commonly used in tACS.

Our fndings are in line with other studies, demonstrating that as stimulation frequency increases, its perceptibility decreases (Hsu et al., [2021](#page-11-3); Imatz-Ojanguren & Keller, [2023](#page-11-22); Turi et al., [2013;](#page-12-10) Ward & Robertson, [1998;](#page-13-4) Zeng et al., [2019](#page-13-2)). Additionally, consistent with fndings from other studies, we verifed that application of an anesthetizing cream reduces perceptibility during transcranial electrostimulation (Guleyupoglu et al., [2014;](#page-10-23) McFadden et al., [2011](#page-11-6)). The reduced perceptibility of high-frequency stimulation offers an added advantage for ensuring participant blinding, given its increased perception thresholds compared to traditional electrostimulation methods. Moreover, the use of an anesthetizing cream appears to be a valuable tool in increasing participant comfort and blinding.

The included lower frequencies of 10 Hz and 20 Hz in our study refect standard frequencies employed in tACS studies (Herrmann & Strüber, [2017;](#page-11-23) Koninck et al., [2023](#page-10-26); Wischnewski et al., [2019;](#page-13-8) Yavari et al., [2018](#page-13-9)). Our fndings demonstrate that these frequencies are already perceptible for most participants at low stimulation intensities, while discomfort or pain became noticeable at slightly higher intensities. In addition, the application of the anesthetizing cream was able to increase these thresholds, proving its efectiveness in reducing somatosensory side efects during stimulation like tingling, itching, or stinging sensations. Furthermore, the higher frequencies of 500 Hz, 1000 Hz, and 2000 Hz selected for this study refect stimulation frequencies used in current tTIS studies (von Conta et al., [2022](#page-13-6); Esmaeilpour et al., [2021;](#page-10-19) Grossman et al., [2017](#page-10-17); Ma et al., [2021;](#page-11-24) Sunshine et al., [2021](#page-12-13); Zhu et al., [2022\)](#page-13-10). Due to reduced sensitivity to higher frequencies in somatosensory perception (Hutcheon & Yarom, [2000](#page-11-11); Palmer et al., [1999](#page-12-5)), we were able to confrm that these frequencies allow for a substantial increase in stimulation intensity before becoming perceptible or inducing pain. In addition, we were able to push the limits of comfortable stimulation intensities even further with the use of anesthetization. Notably, our results indicate that the anesthetic effect seemed to decrease at frequencies of 1000 Hz and above. However, as explained, this is not indicative of reduced anesthetic efficacy, but rather a ceiling efect inherent to our study design. Nonetheless, these results highlight the potential for using high-frequency stimulation in conjunction with topical anesthetization at

Notably, for the higher frequencies, a portion of the participants had their thresholds set to 4 mA for statistical analysis in line with staircase conclusion criterion 2. This indicates that even when the maximum stimulation intensity of 4 mA was reached in a condition, these participants did not report pain or perceivable sensations. This is especially true for stimulation with 2000 Hz, in which 29 out of 37 subjects reached 4 mA. Therefore, it's crucial to recognize that the actual thresholds for these frequencies likely surpass our applied maximum of 4 mA. This limitation is due to the ethical, safety, and hardware constraints within which our study was conducted. Consequently, the recorded thresholds at these frequencies essentially represent the highest stimulation intensities we could safely administer, highlighting that participants might have tolerated even higher intensities during high-frequency stimulation without discomfort. Therefore, our statistical analysis is even on the conservative side and likely underestimates the true tolerable thresholds.

On the other side, some subjects reported pain sensations for high frequencies at considerably lower intensities (e.g., for 2000 Hz, one participant's pain threshold was measured as 2.7 mA). This highlights the interindividual diferences in perception to electrostimulation which should be taken into account. Especially for low frequencies, our results reveal considerable variability in pain and perception thresholds among participants. For instance, in the 10 Hz condition, pain thresholds range from 0.18 to 1.78 mA. This highlights that both high- and low-frequency stimulation vary in individual sensitivity and there is no one-size-fts-all approach. Therefore, our thresholds should be viewed as approximate guidelines rather than absolute values. However, our data indicate that even in common tACS studies, it is highly probable that some subjects will perceive the stimulation and few subjects will even feel pain, even if the stimulation is within a safe range of<2 mA. Thus, for optimal safety and successful blinding, we advise testing individual thresholds before applying electrostimulation at the target intensity.

It is essential to recognize that the efectiveness of stimulation is not solely determined by the applied stimulation intensity. While numerous studies suggest a dose-dependent efect, indicating that higher stimulation intensities often lead to more pronounced effects (Johnson et al., [2020](#page-11-25); Turner et al., [2021;](#page-12-19) Wischnewski et al., [2019](#page-13-8)), this is not an absolute rule. Indeed, some research indicates a complex, non-linear relationship between stimulation intensity and its effects. A study by Moliadze et al.  $(2012)$  $(2012)$  $(2012)$  has shown that while lower intensities might lead to inhibition, increasing the intensity can actually reverse this efect, transforming inhibition into excitation. Moreover, the actual voltage that reaches the target area is not guaranteed by high stimulation intensity alone. Variabilities in individual anatomical factors,

such as skull thickness, and the specifc confguration of the electrode montage, signifcantly infuence the voltage delivered to the target area (Hunold et al., [2023](#page-11-26)). Consequently, it is recommended to utilize current fow modeling tools, like SimNIBS (Puonti et al., [2020](#page-12-21); Thielscher et al., [2015](#page-12-22)) or ROAST (Huang et al., [2019](#page-11-27)). These tools, particularly when modeled on individual anatomical specifcs, can be used to ensure that the target area receives sufficient voltage for effective stimulation (Saturnino et al., [2019\)](#page-12-23).

It is important to emphasize that the stimulation thresholds in our study were determined using round electrodes with a diameter of 12 mm. To ensure that our results are relevant irrespective of electrode size, we have included measures of current density alongside our fndings. However, it's essential to understand the interplay between electrode size, current density, and current intensity. Several studies have posited that larger electrodes generally offer more comfort than smaller ones, attributed to the distribution of currents across a larger area, leading to lower current densities (Alon et al., [1994](#page-10-13); McNeal & Baker, [1988;](#page-11-28) Verhoeven & van Dijk, [2006\)](#page-13-3). Yet, Lyons et al. [\(2004\)](#page-11-29) presented contrasting evidence, showing greater comfort with smaller electrodes. This contradiction may be solved by recent research, which suggests that cutaneous sensation is primarily infuenced by current intensity rather than current density (Fertonani et al., [2015](#page-10-5); Martinsen et al., [2004;](#page-11-30) Turi et al., [2014](#page-12-8)). This is due to a spatial summation efect where larger electrodes engage more cutaneous receptors, increasing sensation (Higashiyama, [1993](#page-11-31); Higashiyama & Tashiro, [1990](#page-11-9); Nielsen & Arendt-Nielsen, [1997](#page-12-9)). Consequently, our results regarding perception and pain thresholds still provide valuable guidance for studies using larger electrodes. However, it's worth noting that cutaneous sensations can be infuenced by other variables, such as the concentration of a saline solution used as a contact medium (Dundas et al., [2007](#page-10-27)), though they are not afected by the shape of the electrode (Ambrus et al., [2011](#page-10-28)).

Certainly, safe stimulation intensities cannot be based solely on the lack of immediate cutaneous pain sensations. Research confrms the safety of classical electrostimulation, as evidenced by rodent model studies. These studies show no changes in several neurotoxicity markers for stimulation intensities commonly used in humans (Jackson et al., [2017](#page-11-32); Liebetanz et al., [2009;](#page-11-33) Zhang et al., [2019\)](#page-13-11). This becomes also evident in human studies, which confrm the absence of neurotoxicity or serious adverse efects of stimulation (Nitsche et al., [2003;](#page-12-24) Nitsche et al., [2004](#page-12-25); 2001; Tadini et al., [2011](#page-12-26)). Currently, stimulation intensities of up to 4 mA are considered safe using electrostimulation methods such as tACS or tDCS (Antal et al., [2017](#page-10-9); Bikson et al., [2016;](#page-10-10) Fertonani et al., [2015](#page-10-5); Matsumoto & Ugawa, [2017;](#page-11-8) Nitsche & Bikson, [2017\)](#page-12-7). Taking into account the novel high-frequency stimulation methods, Grossman et al. ([2017](#page-10-17)) did not fnd tissue damage in rodents,

and Piao et al. ([2022\)](#page-12-27) did not fnd an adverse efect of this stimulation on various tested criteria in humans. This is in line with Cassarà et al. (2022), who explored the safety of tTIS in humans and recommended frequency-based maximum exposure limits, with higher frequencies allowing for greater exposure. Nonetheless, to increase stimulation intensities above 4 mA, a robust body of evidence pointing to its unquestionable safety is needed.

A limitation of our fndings is the brief 5-s duration of stimulation to measure thresholds. Previous research indicates that stimulation sensation decreases over time due to adaptation efects (Hsu et al., [2021](#page-11-3); Khadka et al., [2020](#page-11-4)). This adaptation efect explains why tACS studies can administer intensities of 1 mA or more without causing prolonged discomfort. Consequently, the thresholds identifed in our study are likely conservative, potentially underestimating the maximum tolerable stimulation intensities. Employing a procedure where stimulation is applied for an extended duration with a gradual increase could leverage this adaptation efect to even further increase perception and pain thresholds. This would allow for even more intense, yet still comfortable, electrical stimulation. Future research should explore the limits of this adaptive approach for high-frequency alternating current stimulation.

In addition, in our study, we relied on participants' selfreports to determine thresholds, a method that inherently carries the risk of subjective biases. To enhance the robustness of future studies, incorporating objective indicators, like physiological markers of discomfort including skin conductivity (Storm, [2008;](#page-12-28) Syrjala et al., [2019](#page-12-29)), might offer a more consistent gauge of participant comfort.

Furthermore, is well-documented that pain perception can be infuenced by an individual's physical and psychological state. Factors such as age (Lautenbacher et al., [2017\)](#page-11-34), expectation (Wiech, [2016](#page-13-12); Wiech et al., [2008\)](#page-13-13), fatigue (Lautenbacher et al., [2006](#page-11-35)), or sex (Paller et al., [2009](#page-12-30); Wiesenfeld-Hallin, [2005\)](#page-13-14) infuence how pain is experienced. This is in line with our study's results, where males were able to tolerate higher stimulation intensities compared with females. However, given that the sample in our study had a bias towards female participants (25 female, 12 male), the generalization of our fndings should be done with a degree of caution. A future study systematically comparing sex diferences would be needed to substantiate this result. However, we believe that our results serve as a robust foundation for establishing new limits and possibilities for future stimulation studies.

## **Conclusion**

We demonstrated that the somatosensory perception and pain thresholds for alternating current stimulation are frequency-dependent. Utilizing high-frequency stimulation,

we successfully administered intensities of up to 4 mA without inducing discomfort in participants. This fnding is especially of note for tTIS, whose efficacy has been limited by low-intensity protocols so far. Increasing the stimulation intensity has the potential to enhance the efficacy of tTIS, unlocking the potential for non-invasive stimulation of deeper brain regions. Additionally, the use of topical anesthetic cream further elevates these thresholds, enabling even higher intensities. This fnding also translates to tACS applications in general, allowing for more potent neuromodulatory interventions without compromising participant comfort.

In summary, our fndings reveal signifcant interindividual diferences in perception and pain thresholds, particularly under high-frequency conditions, emphasizing the need for customized stimulation intensities in tTIS/tACS experiments. To ensure participant comfort and efective blinding, we recommend tailoring stimulation based on individual responses. Additionally, our study shows that using a topical anesthetic can raise these thresholds, ofering a viable method to enhance participant tolerance or blinding in future electrostimulation studies.

## **AI and AI‑Assisted Technologies in the Writing Process**

During the preparation of this work, the authors used ChatGPT-4 in order to improve readability and language of the manuscript. After using this tool, the authors reviewed and edited the content as needed and take full responsibility for the content of the publication.

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Cornelius Tamm: Investigation, formal analysis, writing—review and editing.

Philipp Ruhnau: Resources; writing, review and editing; supervision; project administration; funding acquisition.

Tino Zaehle: Conceptualization; resources; writing, review and editing; supervision; project administration; funding acquisition.

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**Data Availability** The datasets generated and analyzed during the current study are available from the corresponding author on reasonable request.

#### **Declarations**

**Conflict of Interest** The authors declare no competing interests.

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