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Does pericentral mu-rhythm “power” corticomotor excitability? — A matter of EEG perspective

Anke Ninija Karabanov a,b, *, 1, Kristoffer Hougaard Madsen a,c, 1, Lærke Gebser Krohne a,c, Hartwig Roman Siebner a,d, e

a Danish Research Centre for Magnetic Resonance, Centre for Functional and Diagnostic Imaging and Research, Copenhagen University Hospital, Amager and Hvidovre, Hvidovre, Denmark
b Department of Nutrition, Exercise and Sports, University of Copenhagen, Copenhagen, Denmark
c Department of Applied Mathematics and Computer Science, Technical University of Denmark, Denmark
d Department of Neurology, Copenhagen University Hospital Bispebjerg, Copenhagen, Denmark
e Institute for Clinical Medicine, Faculty of Medical and Health Sciences, University of Copenhagen, Copenhagen, Denmark

ABSTRACT

Background: Electroencephalography (EEG) and single-pulse transcranial magnetic stimulation (spTMS) of the primary motor hand area (M1-HAND) have been combined to explore whether the instantaneous expression of pericentral mu-rhythm drives fluctuations in corticomotor excitability, but this line of research has yielded diverging results.

Objectives: To re-assess the relationship between the mu-rhythm power expressed in left pericentral cortex and the amplitude of motor potentials (MEP) evoked with spTMS in left M1-HAND.

Methods: 15 non-preselected healthy young participants received spTMS to the motor hot spot of left M1-HAND. Regional expression of mu-rhythm was estimated online based on a radial source at motor hotspot and informed the timing of spTMS which was applied either during epochs belonging to the highest or lowest quartile of regionally expressed mu-power. Using MEP amplitude as dependent variable, we computed a linear mixed-effects model, which included mu-power and mu-phase at the time of stimulation and the inter-stimulus interval (ISI) as fixed effects and subject as a random effect. Mu-phase was estimated by post-hoc sorting of trials into four discrete phase bins. We performed a follow-up analysis on the same EEG-triggered MEP data set in which we isolated mu-power at the sensor level using a Laplacian montage centered on the electrode above the M1-HAND.

Results: Pericentral mu-power traced as radial source at motor hot spot did not significantly modulate the MEP, but mu-power determined by the surface Laplacian did, showing a positive relation between mu-power and MEP amplitude. In neither case, there was an effect of mu-phase on MEP amplitude.

Conclusion: The relationship between cortical oscillatory activity and cortical excitability is complex and minor differences in the methodological choices may critically affect sensitivity.

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Introduction

Alpha oscillations (8–12 Hz) are a distinct feature of human cortical activity. Initially interpreted to reflect “cortical idling” [1] they are today assumed to have an active role in neural processing:

According to the “gating-through-inhibition” hypothesis, alpha oscillations gate information by active inhibition of task irrelevant areas [2,3]. This hypothesis is supported by studies that demonstrated an increase in alpha power over task-irrelevant cortical areas with simultaneous decrease of alpha power in task-relevant areas [2–5]. Instantaneous fluctuations also actively influence task performance and several studies have shown that performance on perceptual tasks increases, when stimuli are presented during periods of low alpha power [5–7].

While alpha oscillations are most prominent in the occipito-parietal cortex, they are also expressed in pericentral,
sensormotor cortex where they are called mu-rhythm [8–11]. Invasive recordings in monkeys provided evidence that mu-oscillations in the sensormotor cortex follow the gating-through-inhibition hypothesis: Low sensormotor mu power predicted an increase in neuronal spiking and increased performance on a tactile perception task [12]. Motor Evoked Potentials (MEPs) induced by single-pulse Transcranial Magnetic Stimulation (spTMS) offer a unique possibility to probe the transsynaptic excitability of fast-conducting corticospinal pathways noninvasively in humans. Several studies used spTMS to examine if the pre-TMS mu-power modulates corticospinal excitability (i.e. the MEP amplitude) using post-hoc grouping of MEPs according to pre-stimulus power. Early studies seem to support the gating-through-inhibition hypothesis and report associations between low mu-power and higher MEP amplitudes [13–15], but these findings could not be replicated by a number of often better-powered studies [16–21] (See Ref. [22] for a detailed, tabular report of studies grouping MEPs according to pre-stimulus power). Real-time EEG-triggered stimulation systems makes it possible to assess the relationship between mu-oscillatory activity and MEP amplitude more effectively through online targeting of specific oscillation states [13–15]. In contrast to the studies using post-hoc grouping, several real-time mu-triggered experiments report a positive relationship between mu-power and MEP amplitude, suggesting mu-facilitation rather than a mu-inhibition in the corticospinal system [23–25].

Considering the contradictory evidence from spTMS studies, it is paramount to understand the physiological effects of basic stimulation parameters before the findings of these studies can be used to support or challenge theoretical ideas like the gating-through inhibition hypothesis. Recently, a real-time EEG-triggered TMS study by Ogata and coworkers found a positive relationship between mu-power and MEP amplitude, but only at higher TMS intensities and when participants had their eyes open. No such relationship emerged at low stimulus intensity or when participants had their eyes closed [25]. This suggests that the relationship between mu-power and corticospinal excitability is not fixed but critically depends on extrinsic (i.e. experimental stimulation and recording set-up) and intrinsic (i.e., brain state) factors.

One important experimental factor is the way how the oscillatory cortical activity of interest is “distilled” from the EEG activity recorded at the scalp level [22,26,27]. In this regard, it is noteworthy that all mu-triggered studies, that report a positive linear relationship between mu-power and MEP amplitude used a sensor-level Laplacian montage centered above the M1 [23,25]. Conversely, post hoc-studies that have shown a negative linear relationship or no relationship at all between mu-power and MEP amplitude have predominantly used source projections or the averaged activity over several surface electrodes [13,14,16,18,19,22].

To re-examine the relation between mu-power and corticospinal excitability, as reflected by MEP amplitude, we employed mu-triggered spTMS using an individual source projection to trace pericentral mu-activity. We applied brain-state informed EEG-TMS for real-time power estimation and targeted the highest and lowest 25% of mu-power in each individual. In addition, we also referenced our data using a sensor level Laplacian montage centered over M1-HAND and used post-hoc trial sorting, to test whether the EEG montage used to extract power influenced the detected relationship between MEP amplitude and mu power. For both methods, we also defined the pre-stimulus phase and the inter-stimulus interval between two pulses and used these as covariates to explore their ability to predict MEP amplitudes.

**Methods**

**Subjects**

15 right-handed healthy participants took part in this study (7 female, average age = 24.1 ± 2.8 years). The sample size was based on previous studies using state-informed brain stimulation to investigate cortico-spinal excitability [22,28]. We did not perform any preselection based on individual TMS or EEG characteristics (e.g. size of the single alpha-band peak or resting Motor Threshold (rMT)). Subjects were allowed to enroll in the study if they were right-handed and between 18 and 40 years of age and did meet the criteria specified in the TMS safety screening questionnaire [29]. All subjects gave informed written consent. The study was approved by the Regional Committee on Health Research and Ethics of the Capitol Region in Denmark and was in accordance with the Helsinki declaration (Protocol H-16017716).

**Experimental setup**

Participants were sitting in a relaxed position in a commercially available TMS-chair (MagVenture, Farum, Denmark). Cushioning provided additional arm and neck support and the participant was instructed to keep the hands and arms relaxed and the eyes open throughout the experiment. A real-time EEG-TMS setup was used for the online analysis of the EEG-signal and triggered single TMS pulses when individual mu-power met the predefined criteria of the current stimulation condition (see Fig. 1).

**Electrophysiological recordings:** Both EEG and EMG were recorded using a NeuOne Tesla system (NeurOne Tesla, Bittium, Oulu, Finland). The amplifiers had a sampling rate of 5 kHz and we used a 2.5 kHz antialiasing low-pass filter with 24-bits resolution per channel, across a range of ±430 mV. To ensure that the latency of the data-delivery for the real-time processing stayed below 5 ms, the data was sent directly from the EEG amplifiers Field-Programmable-Gate-Arya via a user-datagram protocol (1 kHz update rate over a 1 Gb/s Ethernet link). The scalp EEG was recorded using a TMS compatible EEG cap (EasyCap M10, sintered Ag/AgCl multielectrodes Easy Cap, Woerthsee-ETterschlag, Germany), with equidistant spacing between the 63 surface electrodes. All electrodes were prepared using high-chloride, abrasive electrolyte gel (EasyCap, Herrsching, Germany), until the impedance was below 5 kΩ.

The EMG was recorded using self-adhesive, disposable surface electrodes (NeuroLine, Ambu A/S, Denmark), and the EMG signal was transferred in the same way as the EEG signal described above. The electrodes were applied in a belly-tendon montage on first dorsal interosseous (FDI) muscle of the right hand (the ground was placed on the right wrist), making sure that a clear muscular response was measured and that 50 Hz noise was below 20μV.

**TMS:** Single TMS pulses were applied using a figure-of-eight shaped MC-B70 coil connected to a MagPro 100 stimulator (Magventure, Farum, Denmark). The stimulator was set to a monophasic stimulation waveform inducing a P-A current direction in the brain. The motor hotspot (M1-HAND) was defined as the coil position and orientation that resulted in the largest and most reliable MEP amplitude, recorded from the fully relaxed FDI muscle. Stimulation intensity was individually adjusted to elicit a mean MEP amplitude of 1 mV, using an in-house threshold hunting algorithm (implemented in Python) based on a previously described adaptive thresholding method [30]. Threshold hunting was initiated at 47% of maximum stimulator output (MSO), and the relative standard deviation of the true threshold was assumed to be 7% [31] and 20.
stimuli were given to reach the threshold. The same procedure was used to estimate the Resting Motor Threshold (RMT) (MEP amplitude > 50 μV) and the intensity required to elicit MEPs of 1 mV, this intensity was used during experimental blocks. The precise positioning of the TMS coil at the target site (FDI motor hotspot) was continuously monitored using stereotactic neuronavigation (Localite GmbH, Sankt Augustin, Germany).

Source projection: A radial source projection was used to extract the pericentral mu-rhythm for each subject. The source projection matrix was calculated using (i) an individual head model based on a structural T1-MRI scan, (ii) the position of the EEG channels, and (iii) the functionally determined FDI motor hotspot. Both (ii) and (iii) were registered on each individual head model using the neuronavigation. The forward model was estimated using a boundary-element modelling (BEM) approach consisting of three surfaces (brain, skull and scalp) extracted from the T1-weighted structural MRI scan of each participant [32]. Conductivities in the BEM forward model were set to 0.33 S/m for skin and brain and 0.0041 S/m for skull. The nasion, as well as right and left tragus were manually marked on the MRI images and electrode positions were set according to positions digitized using stereotactic neuronavigation. The forward modelling approach resulted in a linear system of equations which were solved through LU-decomposition following the standard procedures in the Fieldtrip toolbox (Version date: 2016-01-26). This procedure decouples the dependence on the properties of the volume conductor model and the source position allowing fast estimation of the potential at all electrodes resulting from a dipole at any source position. Finally, a projection to source space (weighted sum of all EEG electrodes) was estimated using the “dipoli” method described in Ref. [32] implemented in Fieldtrip considering a single dipole with radial orientation at a position 20 mm (also in radial direction) underneath the FDI hotspot.

Real time estimation of pericentral mu power: During acquisition a ring-buffer containing the latest 500 ms of data was updated upon the arrival of each sample. To limit the computational cost of the subsequent processing steps the sampling rate in this buffer was reduced to 1 kHz by averaging the 5 samples received in each UDP package. A processing loop running in a separate process with real-time priority performed source projection and a discrete Fourier transform of the 500 ms of data [33]. As an indicator of mu-band power we considered the fraction of the sum of the absolute squared coefficients within the defined mu-band and the total signal power, thereby obtaining a mu-band power fraction. The average update time for the processing loop was below 0.5 ms, which ensured that only very occasionally the estimate was not updated for each sample, meaning that the additional latency due to signal processing was negligible.

Re-extraction of the pericentral my-rhythm using a Laplacian montage: To test the influence of differences in electrode montages for extracting the mu-power we re-extracted the pericentral mu-rhythm from the raw scalp EEG for each subjects using a sensor level five-channel Laplacian montage centered on the electrode above the M1-HAND area [23]. This enabled the post-hoc determination of pre-stimulus mu-power and mu phase according to a C3-centered Laplacian montage and allowed the direct comparison of source-extracted and surface extracted mu-oscillations.

Experimental sessions

Structural MRI scans: On the day before the experiment each participant was scanned using a structural T1-weighted MRI sequence (T1-w MRI). T1-w scans were done using a magnetization prepared rapid gradient echo (MPRAGE) MRI sequence with 0.85 mm isotropic spatial resolution, TR = 6 ms, TE = 2.7 ms and flip-angle = 8° on a Philips 3 T Achieva scanner (Philips, Best, Netherlands).
The field of view was 245 × 245 × 208mm, such that the scan covered the whole brain. The T1-w scan was required for obtaining the head model for the source projection, as well as for neuronavigation during the state-informed experiment.

**TMS-EEG Experiment:** Each experiment started with a range of pre-measurements to determine the subject specific stimulation criteria. First, the subjects were co-registered with their T1-w scan, then the individual EEG electrode positions were registered to the T1-w scan. The EEG and EMG electrodes were mounted as described above, and the signal quality of both measures were visually monitored throughout the whole experiment, and EEG channels were re-prepared in case the signal quality dropped. The next steps included the determination of the M1-HAND hotspot, the RMT and the stimulation intensity required to evoke MEPs of 1 mV (1 MV-MEP) amplitude for the right FDI. Both RMT and 1 mV-MEP were estimated using the threshold hunting algorithm described in the experimental setup section above. Finally, the source projection matrix was calculated.

To identify the individual mu-rhythm, we recorded 5 min of EEG while the subject was resting with open eyes. The individual peak frequency was determined as the peak of the mu-power spectral density (PSD) within 7–13 Hz, based on the resting EEG data and the estimated source matrix. The mu-frequency band was defined as ±2 Hz around the peak frequency. However, in cases where the lower frequency limit would be below 7 Hz, the limit was set to 7 Hz, resulting in a narrower frequency band for these subjects. Furthermore, the PSD maps were used to determine the highest (q75) and lowest (q25) quantiles of the mu-power, which later were used for thresholding of the power-informed stimulation.

**Power-informed stimulation:** During the experiment TMS stimulation was triggered by instances of high (q75) and low (q25) power of the endogenous pericentral mu-rhythm, using the thresholds estimated during the pre-measurements. As TMS gives rise to large artefacts in the EEG signal just after the stimulation, a non-stimulation trigger was set in 50% of the trials. This enabled post-hoc evaluation of power estimation performance in the non-stimulated trials, while power-dependent effects of spTMS on corticospinal excitability were evaluated in the 50% of trials with TMS. 60 trials were collected for each of the four conditions (q75-TMS, q75-trigger-only, q25-TMS, q25-trigger-only) resulting in a total of 240 trials, split into 4 blocks. Each block contained 15 trials of each condition, resulting in a total of 60 triggers per block. Stimulation was given in blocks of either high or low power and the order between stimulations and non-stimulations triggers was pseudo-randomized, such that no more than 3 of the same kind could appear in a row.

Between each block, a short break of 1–5 min was allowed. For a few subjects, we adjusted the stimulation intensity between blocks, such that the stimulation always elicited an average MEP of around 1 mV, however the total adjustment never resulted in a change of more than 4% stimulator output. Average stimulator output across participants was 70% ± 13%. To avoid any systematic interaction between TMS pulses, the minimum inter-trial-interval (ITI) set by the algorithm was 2s. Due to the constraints set in the power-detection algorithm the actual ITI was considerably longer and had a mean of 10.3 s across all individuals. On a few occasions (less than 5% of trials) the ITI either exceeded 60 s or was undefined (for the first trial for each block) these trials were removed from further analysis.

**Data analysis**

**MEP analysis:** The peak-to-peak amplitude of the MEP was determined trial-by-trial by an in-house developed python script. Similar to a previously published study [22] all trials with an EMG activity > 50 µV during the 100 ms prior to stimulation, or with raw MEP amplitudes more than 2.5 standard derivations away from the mean were excluded from further analysis. On average 6.3% of trials were excluded from further analysis due to these criteria.

**Phase at Stimulation:** The phase at the time point of stimulation was estimated from the recorded data using in-house developed software using a continuous Morlet wavelet transform within 500 ms windows. The transform was done for 51 frequency scales across the mu-band and the phase of the frequency scale which yielded the highest wavelet coefficient 100 ms prior to stimulation was projected to the stimulation time. A more detailed description of the algorithmic procedure can be found in our previous work on phase-triggered TMS [22]. Depending on the estimated phase, all TMS trials were sorted in a post-hoc analysis into four distinct phase bins (0°, 90°, 180°, 270°).

**Statistics:** To test the hypothesis that the cortico-spinal excitability is modulated by mu power at the time of stimulation while simultaneously assessing the relationship between power, phase and interstimulus interval (ISI) on the individual trial basis we performed a mixed-effect analysis that included mu-power fraction (categorical – high, low), the mu-phase (categorical – 0°, 90°, 180°, 270°) and the inter-stimulus-interval between two trials (continuous – ISI) as fixed effects and the participant number as a random effect. Statistical analysis was performed using the statistical software package R (https://www.r-project.org). Mixed effects analysis was performed using the lme 4 package (Team RC 2018) and all continuous variables were log-transformed before they were entered in the model. The significance threshold for null hypothesis testing was set to p < 0.05.

To directly test evidence for the null hypothesis we additionally used Bayesian analysis of covariance as implemented in Jasp v. 0.11.1 with the MEP as dependent variable, high/low power and phase discretized into four bins as a fixed-factors, subject as a random factor and included the logarithm of the ISI as a covariate.

**Analysis of the data extracted by Laplacian Surface Montage re-referencing:** The analysis performed for source-extracted mu-power were also run for the post-hoc re-referenced Laplacian data. Generally, the power in mu-band in the Laplacian montage differed compared to the to the power in the original source projection as indicated by an average correlation value of 0.34 across participants. This meant that the power fraction could no longer be reasonably divided into two separate bins. Therefore, for the mixed model, we only included the highest and lowest quartile of the Laplacian-power fraction in order to keep the data as similar as possible to the source-triggered analysis, even though this meant biasing data. However, in the analyses that considered a continuous representation of the power fraction (Bayesian ANCOVA) we utilized all the available data.

In order to roughly compare the ability of each montage to capture the EEG signal, we calculated the fraction of variance explained by the source projection, both across all electrodes and confining the analysis to fronto-parietal electrodes of the stimulated side (EasyCap M10; Electrodes: 1, 2, 8, 19, 33, 32, 31, 29, 15, 14, 5, 7, 18, 17, 16, 15, 29). For the Laplacian, we only considered the channels that were included in the montage, because Laplacian montages only consider signal in the surrounding electrodes thereby effectively predicting zero signal everywhere else.

**Results**

**Online power-triggered EEG-TMS**

We verified that the online power triggering worked by assessing the power fraction estimate for the non-stimulated trials in a 500 ms window centered at the intended stimulation
timepoint. The mean power fraction was $0.15 \pm 0.07$ in the low-power triggered condition and $0.30 \pm 0.12$ in the high-power condition, indicating that the pericentral mu-power was successfully targeted by brain-state informed spTMS (Fig. 2A). The ISI during the low-power was $13.9 \pm 4.2$ s and $10.5 \pm 2.0$ s during the high-triggered condition. A paired t-test (unequal variance) testing for differences in the ISI indicated that duration was significantly influenced by the triggering condition ($t_{14} = 2.47; p = 0.02$) (Fig. 2B).

The accuracy of the phase detection algorithm was evaluated by comparing the projected phase to a centered phase estimate for non-stimulated trials as also described in a previous publication [22]. As expected, this revealed that the phase estimation accuracy was lower for the low power condition ($q_{25}$) with a mean absolute error of $61\%$, whereas the mean absolute error in the $q_{75}$ condition was comparable to our previous work with a value of $47\%$ [22].

Mu-rhythm extracted by source projection: The fraction of variance explained by the source projection averaged to 7.3% ranging from 5.8% to 8.7%. When the analysis was confined to the frontal, pericentral and parietal electrodes of the stimulated hemisphere the captured variance increased to 32.4% (range 19.2%–47.7%). This was substantially higher than for the Laplacian montage which averaged to 1.9% (0.9%–4.1%), indicating that the Laplacian montage was not superior to the source montage in explaining the signal. This is not surprising because the Laplacian montage is geared to select oscillatory activity which is the same as for the neighboring electrodes but different from the central electrode. The average MEP amplitude was $1.16 \pm 0.39$ mV across both conditions. The mean MEP amplitude in the high-power triggered condition was $1.14$ mV compared to $1.17$ mV in the low-power triggered condition. The linear mixed-effects model, that treated mu-power and mu-phase as fixed effects and participant as a random effect showed no significant main effect for power ($\chi^2(1) = 0.32; p = 0.57$) (Fig. 3A). Also, the main effect of phase was not significant ($\chi^2(3) = 1.54; p = 0.20$) (Fig. 3B), while ISI significantly modulated the MEP amplitude ($\chi^2(1) = 5.14; p = 0.02$). None of the interaction terms were significant. The main effect of ISI was caused by an increase of MEP amplitude at when pulses were given at longer intervals (Fig. 3C).

Bayesian Analysis: Using Bayesian analysis of covariance (ANCOVA) to assess inclusion probabilities for each of the predictors we found no evidence for inclusion of ISI (Bayes Factor (BF): 0.41), strong evidence against inclusion of power fraction (BF: 0.04) and extreme evidence against inclusion of phase (BF: 0.006).

Mu-rhythm extracted by Laplacian Surface montage: To test whether the EEG montage used to extract pericentral power influenced the detected relationship between extracted power and cortical excitability, we re-referenced the EEG data using a sensor level Laplacian montage centered on M1-HAND, similar to previous power-triggered experiments [23]: Laplacian mu-oscillations for each TMS pulse were extracted post-hoc. The linear mixed-effects model, that treated Laplacian mu-power fraction, mu-phase and ISI as fixed effects reveal a significant power-effect ($\chi^2(1) = 5.91; p = 0.01$) indicating that the MEP amplitude was larger when Laplacian mu-power was high (Fig. 4A). For phase, the Laplacian mu-extraction agreed with the initial source-projected data and showed no significant effect of mu-phase ($\chi^2(3) = 0.56; p = 0.64$) (Fig. 4B). Fig. 4 displays the MEP as a function of phase to further illustrate that no effect of phase could be seen. The ISI effect was significant as in the analysis of the source projected data ($\chi^2(1) = 3.9; p = 0.04$).

Bayesian Analysis: A Bayesian ANCOVA considered all the data of the Laplacian montage by including the continuous power fraction as an independent variable. In this analysis the inclusion probabilities showed very strong evidence for inclusion of the power fraction (BF: 75.9) and no evidence for inclusion of ISI (BF: 0.941) and again extreme evidence against the inclusion of phase (BF: 0.002).

Spatial comparison of radial source and Laplacian montage and pairwise electrode comparisons: The location of the Laplacian montage was more posterior that the radial source at the individual hotspot, as indicated by the mean MNI coordinates (Laplacian

![Fig. 2. Precision of Targeting: A) shows the power fraction in the mu-band during non-stimulated trials, centered at the intended stimulation point in high ($q_{75}$) and low ($q_{25}$) power condition. Grey lines represent individuals, the black line the grand average B) shows the average ISI in high ($q_{75}$) and low ($q_{25}$) power condition in stimulated trials. Grey lines represent individuals average, the black line the grand average.](image)
montage, electrode 17: [-62, -22, 73]; radial source at hotspot [-55, -13, 85]). The mean displacement from center of gravity (COG) of each montage was similar in both montages (electrode 17: 9.9; hotspot: 9.5 (both MNI mm)). To compare if specific electrodes preferentially contributed to the effect observed in the Laplacian montage, we also calculated mu-power based on the center electrode (electrode above M1-HAND) and each of its neighbors individually (see Fig. 5B) by using separate Bayesian ANCOVAs with the logarithm of the MEP as dependent variable, subjects as a random factor, the detected phase-bin for the relevant montage as an independent factor and the logarithm of ISI and the relevant power fraction as covariates. We then quantified the effect of power fraction in each of these six electrode pairs at the Bayes factor for inclusion of the continuous power fraction in the model. The results indicated that the significant effect of power was driven by a signal gradient towards the medial/anterior direction in electrode space (difference between electrodes 7 and 17). In all of these models there were extreme evidence against the inclusion of phase (BF ≤ 0.006).

Fig. 3. Source Projection: MEP as a function of Power and ISI A) shows the average MEP amplitude at stimulation point in high (q75) and low (q25) power condition. Grey lines represent individuals, B) shows the relationship between pre-stimulus phase and MEP amplitude and C) shows the relationship between MEP and ISI.

Fig. 4. Laplacian montage: MEP as a function of Power and ISI A) shows the average MEP amplitude at stimulation point in high (q75) and low (q25) power condition. Grey lines represent individuals, B) shows the relationship between pre-stimulus phase and MEP amplitude and C) shows the relationship between MEP and ISI.
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ever, when re-extracting mu-power according to a Laplacian sur-
power-dependent modulation of corticospinal excitability. How-
more hotspot for the contralateral FDI muscle, failed to show
radial source projection, extracting mu-power from the individual
fl
Discussion
Here, we demonstrate that the results of mu-power triggered
spTMS are dependent on the electrode montage used to extract the
fluctuating expressions of individual pericentral mu-oscillation. A
radial source projection, extracting mu-power from the individual
motor hotspot for the contralateral FDI muscle, failed to show
power-dependent modulation of corticospinal excitability. How-
ever, when re-extracting mu-power according to a Laplacian sur-
face montage over M1-HAND, we found a positive relation between
pre-stimulation mu-power and MEP amplitude. While the effect of
mu-power on MEP amplitude was montage-dependent, both
montages concordantly reported no significant effect of mu-phase.
Indeed, Bayesian analysis, confirmed that there was strong evi-
dence in favor of the null hypothesis, indicating that mu-phase
fluctuations in both montages did not play an important role in
influencing MEP amplitude size, thereby replicating the results of
our previous work [22].

Our results have important general implications, they show that
the way the central EEG signal is extracted from the cortical region
of interest will have profound impact on what kind of state-
dependent effects of the expressed power or phase in the
frequency-band of interest will be revealed by the TMS-EEG
approach.

Which property rendered the Laplacian surface montage sen-
sitive to the fraction of pericentral mu-power that has a positive
relationship with cortico-spinal excitability? Both, the source and
the Laplacian surface montage were sensitive to radial sources of
EEG activity around the central sulcus. The fact that they none-
theless produced different results suggests a complex spatial rela-
tionship between pericentral mu-activity and corticospinal
excitability and it is interesting to note the spatial differences be-
tween the tested montages. The Laplacian montage was more
posterior with an average location over the postcentral gyrus (i.e. the
somatosensory cortex) while the source projection was on
average located in the precentral gyrus (i.e. the primary motor
cortex). These results imply that it may primarily be the post-
central sources of pericentral mu-activity that modulate cortico-
spinal excitability. This observation is in good agreement with
several studies that have used invasive and noninvasive methods to
locate cortical sources of sensorimotor Mu-rhythm humans and
animals. They found multiple cortical sources of the Mu rhythm in
frontoparietal networks but quite consistently reported that its
expression peaks in the postcentral somatosensory cortex [34–38].

A predominantly postcentral origin of the Mu-rhythm would also help to explain the contradictory findings of previous work: Studies using the more posterior Laplacian filter [23,25] more often reported significant correlations between MEP amplitude and po-
terior studies that used other ways of mu-extraction, often
focusing on the more anterior precentral cortex [14,16,18,19,22]. It
is worth mentioning that the sensitivity of the Laplacian surface
montage to capture a modulatory effect of mu-power on MEP
amplitude cannot be attributed to a generally increased sensitivity
for the frequency of interest as the Laplacian montage explained
less of the general variance in the frequency band of interest when
compared to the source projection. Finally, both of our montages
were sensitive to radial sources and we cannot exclude that
tangentially oriented sources may be relevant as well. Pairwise
analysis of the electrodes contributing to the power-amplitude
coupling with the Laplacian montage suggest that a signal
gradient towards the medial/anterior direction in electrode space
may have contributed to the positive relationship between mu-
power and MEP amplitude. However, more systematic compari-
sions of radial and tangential pericentral sources are required to
clarify this question in the future.

How do the present results support the gating-through-
inhibition theory of cortical oscillatory activity in the alpha range
[2]? The gating-through-inhibition theory would predict a negative
relationship between ongoing mu-activity and corticospinal excit-
ability (i.e., MEP amplitude). Apart from a few early reports, all
TMS-EEG studies either found no relation or a positive relation
which is in apparent contrast to the gating-through-inhibition
theory. The MEP amplitude may however not be the right mea-
sure to test the gating-through-inhibition theory as Mu is maxi-
mally expressed in somatosensory cortex and primarily focuses on
cortical processing of sensory inputs and not on generating motor
outputs. Phase and amplitude of the mu-rhythm may exert gating-
through-inhibition in postcentral somatosensory cortex but how
this impacts the precentral cortex that is stimulated with TMS re-
mains unclear. There may be secondary effects on the precentral
motor cortex, possibly opposite in sign, and this could cause
corticospinal disinhibition when cortical inhibition in sensory cortex is

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**Fig. 5.** Visualization of the radial source projected to the scalp (red) and the center of the Laplacian montage (blue) (A), the location of the projection 20 mm into the head model using a Gaussian kernel density estimate with bandwidth set according to Ref. [51] (B), and a visualization of the Laplacian montage (C). (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)
maximal but more work needs to be done to determine the interaction between phase and amplitude in the pre and postcentral cortex. Whatever the relationship between pericentral mu-activity and MEP amplitude may be, any relationship will be strongly influenced by the magnitude of cortex stimulation (i.e., the TMS intensity) and the ongoing level of neuronal activity (i.e. the firing rate) [25]. This notion is corroborated by a modelling study using a simple non-compartmental threshold-crossing motoneuron model without dendrites [39]. In that study, Matthews showed that the response of the model neuron to a stimulus depended upon stimulus strength, synaptic membrane noise, and intrinsic tonic firing rate of the neuron (induced by intrinsic background drive). The study revealed a non-linear relationship between background firing rate and the strength of a brief external stimulus: For weak stimuli, the response increased with increasing intrinsic tonic firing rate but for strong stimuli, the response was maximal at a low firing rate and then decreased for higher firing rates. Matthews concluded that “transferring” these findings to corticospinal neurons makes it unlikely that the magnitude of the descending volley elicited by a given cortical stimulus (‘excitability’) will always increase with the initial level of cortical activity. Transferring these observations to TMS-EEG studies, it is unlikely that the MEP amplitude elicited by a TMS pulse will monotonically scale with the level of cortical oscillatory activity (and the associated changes in neuronal firing rates). Instead, strong TMS pulses can be expected to elicit larger MEPs during low levels of intrinsic neural activity (e.g. low firing rate at high mu-power), while weak TMS pulses will elicit larger MEPs during high levels of intrinsic neuronal activity (e.g. high intrinsic firing rate at low mu-power). This flip in the sign of the relationship between mu-power and MEP amplitude is illustrated in Fig. 6. Accordingly, a recent TMS-EEG study demonstrated that high mu-power (e.g. low endogenous activity) was only associated with larger MEP amplitudes during higher TMS intensities [25]. The non-monotonic interaction between intrinsic firing rate (produced by transsynaptic drive) and stimulus response suggested by Matthews [39] also reconcile our present findings (i.e., no relationship or positive relationship between power and MEP depending on montage) with the negative power MEP relationship reported in our recent study [22] in which we only examined the highest power quartile in that work.

An interesting perspective on the relation between evoked activity and pre-stimulus oscillations is afforded by a recent study in which variability in neural responses was interpreted within a framework of criticality, where system dynamics are tuned toward the phase transition between stability and instability [40]. In that study, Stephani et al. tracked both the amplitude of sensory evoked potentials (SEPs) and pre-stimulus alpha oscillations and showed that the amplitude of the early somatosensory evoked potential (SEP) depended on the amplitude of pre-stimulus alpha activity [40]. They also found that both measures display covarying temporal dependencies decaying according to a common power law, pointing to a dependence on a neural network at near-criticality [40].

Our results have major implications for the use of power-informed TMS-EEG as a tool to better control the cortical “state” at the time of stimulation and hereby, to render the brain response to TMS less variable [41]. Overall, the effect size of the MEP amplitude modulation is modest and sometimes not detectable using conventional average statistics (e.g. ANOVAs) [22,23,25]. This suggests that the contribution of the pericentral “ oscillatory state” to the overall trial-to-trial variability of the MEP is generally low. As long as the interactions between spatial filtering, neural background noise, intrinsic firing rate and stimulus intensity are not better understood, the potential of power-triggered TMS to significantly improve the intra-individual variability of single-pulse MEPs remains limited. Additionally, it has to be mentioned that other oscillatory rhythms also have been suggested to modulate cortico-spinal excitability. Especially the sensorimotor beta rhythm but also gamma oscillations and cortico-muscular coherence have also been proposed to modulate cortico-spinal excitability [13,16,18–20,42,43]. More complex interactions like cross-frequency coupling between cortical oscillations with difference frequency bands are also possible modulators that have not yet been explored.

In this study we have chosen to estimate mu-power based on its relative contribution to total EEG power; an alternative approach would have been to estimate power exclusively based on the expression of alpha power. Both approaches are limited in terms of specificity when being used to trigger TMS based on the oscillatory power of an EEG frequency band of interest. When the mu-rhythm read-out is measured as relative contribution to total EEG power, a drop in other frequency bands will be indexed as increase in mu-power. But if the EEG read-out only considers the alpha-band alone, increases in total EEG power will be recognized as specific

![Fig. 6](image_url)
increase in mu-power. Another methodological consideration is that our amplitude thresholds were defined at rest. It is quite possible that oscillatory activity is modified by TMS pulses beyond the acutely evoked oscillatory activity hence an adaptive procedure allowing to dynamically adjust the amplitude threshold for triggering the TMS pulse may have afforded more stringent power criteria.

Regardless of the montage used for extracting pericentral mu-activity, we found no modulatory effects of mu-phase on MEP amplitude. The use of Bayesian statistics also allowed us to move beyond not rejecting the null hypothesis as Bayesian statistics detected strong evidence against the contribution of mu-phase to fluctuations in corticospinal excitability. This contrasts with several studies that reported significant phase modulations of the MEP amplitude using larger numbers of trials (>100) [28,44,45]. While it is possible that our comparatively low trial number (60) prevented us from detecting a subtle phase-effect, we argue that the use of mu-triggered TMS as a tool to reduce interindividual variance is limited, if more than 100 trials are needed to detect statistically reliable effects.

The interval between subsequent TMS pulses is another underexplored aspect of the TMS-EEG approach. In our study, we ensured that TMS was given at very low repetition frequency with a large jitter. This decision was motivated by several TMS studies showing that continuous quasi-repetitive, or jittered application of supra-threshold TMS in the 0.5–0.3 Hz range can induce changes in the excitatory-inhibitory balance in M1 [46–48]. Studies also showed that long ISIs (>0.2 Hz) significantly improve the reliability and lower the variability of intra-individual MEPs [49].

Our experimental procedure resulted in long and highly jittered ISIs, and replicated our previous finding that MEPs tend to be larger when long ISIs are used [22]. This suggests that the preceding pulse may have some conditioning influence on the MEP response produced by the next TMS pulse. These “recency” effects together with the virtue of TMS to shape corticospinal excitability and cortical oscillatory activity, underscore that TMS-EEG always needs to consider whether or how much the repeated administration of supra-motor threshold TMS pulses “actively” shapes regional cortical excitability and oscillatory activity rather than “passively” probing it.

Taken together, both experimental and theoretical work suggests that the relationship between regional neuronal activity and corticospinal excitability is complex and may interact with a range of different experimental choices such as choice of EEG montage, stimulation intensity, ISI, number of stimuli given and the intrinsic inhibitory/excitatory balance in the stimulated cortex [21,22,39,45,50]. The relationship between cortical excitability, cortical activity and basic experimental parameters has to be better understood and investigated before brain-state triggered TMs can become a standard technique to reduce inter-trial variability and increase the effectiveness of TMS protocols.

CRediT authorship contribution statement

Anke Ninja Karabanov: Conceptualization, Formal analysis, interpretation, Writing — original draft. Kristoffer Hougaard Madsen: Conceptualization, Formal analysis, interpretation, Writing — original draft. Lærke Gebser Krohne: Funding acquisition, Writing — original draft. Hartwig Roman Siebner: Conceptualization, Formal analysis, interpretation, Writing — original draft.

Declaration of competing interest

Hartwig R. Siebner has received honoraria as speaker from Sanofi Genzyme, Denmark and Novartis, Denmark, as consultant from Sanofi Genzyme, Denmark and Lundbeck AS, Denmark, and as editor-in-chief (Neuroimage Clinical) and senior editor (NeuroImage) from Elsevier Publishers, Amsterdam, The Netherlands. He has received royalties as book editor from Springer Publishers, Stuttgart, Germany and from Gyldendal Publishers, Copenhagen, Denmark.

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