Voluntary Inhibition of Physiological Mirror Activity: An EEG-EMG Study

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Abstract
Physiological mirror activity (pMA), observed in healthy human adults, describes the involuntary co-activation of contralateral homologous muscles during unilateral limb movements. Here we provide novel evidence, using neuromuscular measurements (electromyography; EMG), that the amplitude of pMA can be voluntarily inhibited during unilateral isometric contractions of intrinsic hand muscles after informing human participants (10 male, 10 female) about its presence and establishing a basic understanding of pMA mechanisms through a standardized protocol. Importantly, significant suppression of pMA was observed immediately after participants were asked to inhibit it, despite the absence of any online feedback during task execution and without special training. Moreover, we observed that the decrease of pMA was specifically accompanied by an increase in relative frontal $\delta$ power recorded with electroencephalography (EEG). Correlation analysis further revealed an inverse association between the individual amplitude of pMA and frontal $\delta$ power that reached significance once participants started to inhibit. Taken together, these results suggest that $\delta$ power in frontal regions might reflect executive processes exerting inhibitory control over unintentional motor output, in this case pMA. Our results provide an initial reference point for the development of therapeutic applications related to the neurorehabilitation of involuntary movements which could be realized through the suppression of pMA observed in the elderly before it would fully manifest in undesirable overt movement patterns.

Key words: $\delta$ power; EEG; EMG; inhibition; physiological mirror activity

Significance Statement
Unilateral movements evoke unintended activation of contralateral muscles, especially when such movements are performed with high effort. This motor phenomenon, termed mirror activity (MA), is observed not only in pathologic conditions of the CNS but also in healthy individuals, where it is known as physiological MA (pMA). Surprisingly, as we show in our study, participants are capable of suppressing this unintended co-activation after being informed of its presence. Suppression was accompanied by increased $\delta$ power in frontal areas. This suggests that unintentional motor output can be suppressed via executive processes, i.e., inhibitory control. We offer an initial reference point for therapeutic applications regarding neurorehabilitation of involuntary movements and a possible strategy to counteract the age-related increase in MA.
Introduction

Humans rely heavily on the bilateral independence of the limbs. A critical phenomenon that can impair such independence is termed mirror activity (MA) or associated movements (Post et al., 2009). MA involves involuntary muscle activity in the contralateral homolog of the purposefully activated muscle. In severe cases, known as pathologic mirror movements (Galléa et al., 2011), this involuntary co-activation of the contralateral limb leads to overt movement that can impair even simple unilateral tasks like reaching and finger tapping. However, even in neurologically healthy adults, physiological MA (pMA) has been observed in homologous muscles (Cincotta and Ziemann, 2008). This phenomenon is hypothesized to occur because of a spillover of the initially lateralized motor command from the contralateral (controlling the active limb) toward the ipsilateral hemisphere (controlling the inactive limb), a mechanism termed motor overflow (Yensen, 1965; Perez and Cohen, 2008). Crossed facilitation within the human motor cortex, i.e., bilateral variation in the excitability of corticospinal projections, is generally observed during intentionally unilateral actions (Carson, 2005). The resulting pMA depends on the functional requirements of unilateral motor tasks (Cincotta and Ziemann, 2008). This is especially true for the level of neural drive toward the muscles engaged in the voluntary movement (Todor and Lazarus, 1986), i.e., the strength and duration of contractions (van Duinen et al., 2008; Sehm et al., 2010). Furthermore, it has been proposed that the occurrence and extent of pMA follows a U-shaped distribution across the life-span (Koerte et al., 2010). Typically, stronger pMA is observed in young children and rapidly decreases as the child enters adolescence (Mayston et al., 1999). In the third decade, pMA gradually reappears (Koerte et al., 2010). This course of pMA occurrence may reflect the processes of increasing inhibition during adolescence, with the later reappearance suggesting an age-related loss of central inhibition (Carr, 2010). Thus, it has been speculated that increases in pMA might be one of many altered patterns of neural mechanisms that could provide an initial reference point for the development of therapeutic applications aimed at involuntary movement prevention. This approach could further provide a possible strategy to counteract or prevent the increase in pMA documented in the elderly.

Materials and Methods

Ethical approval

This study was supported by the local ethics committee of the University of Leipzig (ref.-nr. 423/18-ek). All participants gave written informed consent to participate in the experiment and were compensated for involvement, according to the Helsinki Declaration.

Participants

Twenty neurologically healthy adults participated in the experiment (10 males, 10 females, age: 25.6 ± 5.4; mean ± SD). All participants were right-handed according to the Oldfield handedness inventory (Oldfield, 1971). Exclusion criteria consisted of neurologic pathologies, pregnancy, centrally active drug use, regular sport, or musical participation exceeding 2 h/week.

Experimental design

Participants were seated comfortably in a chair and operated a custom-made force sensor with their thumb and index finger of one hand. The other hand rested on a
Before starting the experiment, all participants performed a maximum voluntary contraction (MVC) test with both hands in the form of a pinch-contraction of the thumb and index finger. Three 5-s maximum contractions per hand were conducted, with 30 s of rest between each. The trial with the largest peak force was selected as the individual’s MVC. This was done to normalize EMG amplitudes during data processing and scale the required force level of the following experiment (40% MVC) to the individual strength capabilities. Subsequently, participants had to vertically move a cursor on a computer screen (placed in ~1.5-m distance) repetitively into a target field by applying force to the sensor (set to require 40 ± 5% MVC) with their left hand while the right hand was resting in the prone position on the pillow. Continuous force data were recorded and used to (1) control that participants reached the target field and (2) compute the duration until participants reached the target field after it appeared on the computer screen for each contraction, i.e., ramp contraction time (RCT). The first 30 contractions served as a baseline block (BL), during which participants were naive concerning the study aim, i.e., at no time during BL were they aware of, or receive feedback about, ongoing pMA. However, pMA occurrence was confirmed through continuous observation of EMG signals during task execution. Following the BL, participants were thoroughly educated on the pMA phenomenon using standardized verbal instructions where their attention was directed toward involuntary muscle activity of their resting hand (Table 1). Afterward, participants were asked to actively inhibit the occurrence of pMA in subsequent force blocks by a self-chosen mental strategy. The goal was to minimize EMG amplitude of the non-performing hand without changing the motor performance of the left hand. Feedback about participants’ inhibition of pMA was provided verbally based on online monitoring by the researcher after the completion of each block. Online feedback was not provided. That means that participants neither received online EMG feedback nor verbal instructions during contractions. Participants could see both hands at all times but were instructed to fixate their gaze on the computer screen in front of them during all contractions. In addition to the BL, four more blocks of 30 contractions each were performed (I-block, II-block, III-block, and IV-block), resulting in a total of 150 epochs. In between blocks, a resting period of 3 min was granted.

Table 1: Standardized verbal instruction given during the experiment

| Timepoint | Instruction |
|-----------|-------------|
| After baseline (BL) block | The goal of this study is to investigate the so-called mirror activity phenomenon. That means that every time you perform a contraction with one hand, we can also measure a weak activity in the contralateral resting hand. From now on, please try to suppress this involuntary activity as much as possible by a self-chosen strategy. However, keep the performance of the left hand identical. |
| After each consecutive block | Good job. However, we still see some involuntary muscle activity in your resting right hand. In the following block, please try to suppress this involuntary activity again as much as possible. |

Data acquisition

EMG data were recorded using a BrainAmp ExG amplifier system (Brain Products GmbH). Skin preparation consisted of shaving, slight abrasion of superficial keratin layers, and cleaning with alcohol. Sintered Ag/AgCl electrodes (4 mm in diameter) were used in a monopolar setup mounted on bilateral first dorsal interossei muscles of the actively contracting hand (FDIVol) and mirror hand (FDIpMA). Reference electrodes were placed on the right and left processus styloideus radii, respectively. One ground electrode was placed on the right epicondylus lateralis humeri. This bilateral setup allowed us to capture voluntary muscle activity of the actively contracting muscles as well as involuntary occurring pMA of the homologous resting muscles in a time-locked manner. EMG data were recorded with a sampling frequency of 1000 Hz. EEG data were acquired using a wireless 64-channel EEG system (LiveAmp, Brain Products GmbH) with an active electrode setup (actiCAP, Brain Products GmbH). Electrodes were mounted individually on an electrode-cap to densely cover bilateral sensorimotor cortices (Fig. 1A). Ground and reference electrodes were placed on AFz and left processus mastoideus, respectively. Electrode impedance was kept below 5 kΩ throughout the experiment. Data were continuously recorded with a sampling frequency of 500 Hz. EMG and EEG signals were segmented around the trigger at the start of every contraction performed by the participant, resulting in a total of 150 epochs each lasting 5 s.

Data processing

All data analyses were performed in MATLAB (v. R2019b, The MathWorks Inc.) using built-in functions, the BBCI toolbox (Blankertz et al., 2016), and the EEGLAB toolbox (Delorme and Makeig, 2004). EMG signals were first decimated to 500 Hz after low-pass filtering [Chebyshev type I (IIR) eighth-order filter, cutoff 200 Hz]. Furthermore, EMG data were high-pass (fourth-order Butterworth, cutoff 20 Hz) and band-stop filtered at 50 Hz (eight-order Butterworth) to remove power-line noise. Both EMG signals (FDIVol and FDIpMA) were full-wave rectified, overlaid, and time locked to preserve the temporal relationship between voluntary and involuntary muscular activity. Mean EMG amplitudes of FDIVol, as well as FDIpMA, were computed by the estimation of root mean square (RMS) values (window length, 50 ms). Mean EMG amplitudes of the FDIVol were computed over the time window defined by the externally triggered start and end of each contraction. Latency of pMA...
Figure 1. Experimental setup depicting the unilateral motor task combined with EEG and EMG measurements. 

A, The motor task consisted of unilateral isometric contractions of the left hand through a custom-build force sensor while the right hand was resting in the prone position. For EEG measurements, 64 electrodes were mounted individually on a 128-channel-electrode-cap to densely cover frontal and bilateral sensorimotor cortices (blue and gray electrodes). Predefined ROIs (blue electrodes): left ROI (FCC3h, FC1, FC3, C1, C3, C5, CP1, CP3, CP5), right ROI (FCC4h, FC2, FC4, C2, C4, C6, CP2, CP4, CP6), and frontal ROI (AF3, AF4, F1, F2, F3, F4, Fz, FCz).

B, Participants had to vertically move a cursor on a computer screen repetitively (in total 150 times) into a target field by applying pinch force to the sensor (set to require 40 ± 5% of MVC force). The target field was visible for 5 s and was followed by a 5-s rest period during which muscles should be completely relaxed. The experiment was divided into five blocks of 30 contractions each. C, EMG was recorded from the bilateral FDI muscle of the voluntary contracting hand (FDI_{vol}) and the mirroring hand (FDI_{pMA}). Please note the different scaling of the EMG traces demonstrating the subliminal nature of pMA.

Figure Contributions: D. Maudrich designed the figure.
was subsequently defined automatically during the delay between burst-onset of FDIvol and the time point at which muscular activity in the contralateral (resting) FDIpMA exceeded, for a time window of at least 10 ms, a threshold of its mean baseline activity (1000 ms pre-FDIvol burst onset) + 2 SD (Maudrich et al., 2019). Concerning the amplitude of pMA, latencies were taken into account so that mean EMG amplitudes of FDIpMA were computed over the time window from the determined point of significant elevation of neuromuscular activity until the muscle offset of FDIvol. Mean EMG amplitudes were additionally estimated for pre contraction epochs (4-s rest periods between consecutive contractions) for FDIvol, as well as FDIpMA. The absence of overt muscle activity in these epochs was checked and confirmed visually, i.e., the relaxation of participants between each consecutive contraction. This control condition was implemented to test for systematic fatigue-related or attention-related changes in baseline EMG amplitudes across the experiment. All EMG amplitudes were normalized with respect to individual MVC values measured at the beginning of every session for each hand separately. MVC values were estimated by the mean EMG activity over a time window of 500 ms during maximum unilateral force production for FDIvol and FDIpMA, respectively. For each participant mean amplitudes of FDIvol and FDIpMA were averaged block wise across all 30 contractions (30 precontraction epochs) before statistical analysis. The same procedure was performed for latencies of FDIpMA.

EEG data were first bandpass filtered (fourth-order Butterworth, 1–100 Hz). Subsequently, independent component analysis (infomax algorithm; Bell and Sejnowski, 1995) implemented in EEGLAB was used to remove artifact components related to eye movement, eye blinking, and muscle activity (Jung et al., 2000). On average, 5.2 ± 2.5 components were removed per participant. Following Laplacian spatial filtering, power spectral densities (PSDs) were calculated for 30 concatenated epochs of each block (corresponding to 30 voluntary muscle contractions), for each of the 64 EEG channels separately, employing Welch’s method with a Hamming window of 500 samples and an overlap of 50%. Resulting PSDs were subsequently normalized to individual total power (1–100 Hz). Next, relative power within distinct frequency bands of interest (δ: 1–3 Hz, θ: 4–8 Hz, α: 9–12 Hz) was calculated by summation of respective frequency bins and extracted for each participant and electrode separately. Frequency bands of interest were chosen based on previous studies investigating motor inhibitory processes (Kaiser et al., 2019) and executive processes involved in lateralized inhibition of symmetrical contractions (Tisseyre et al., 2019). Subsequently, PSDs were averaged across three regions of interest (ROIs; Fig. 1A): left sensorimotor ROI (FCC3h, FC1, FC3, C1, C3, C5, CP1, CP3, CP5), right sensorimotor ROI (FCC4h, FC2, FC4, C2, C4, C6, CP2, CP4, CP6), and frontal ROI (AF3, AF4, F1, F2, F3, F4, Fz, FC2). Finally, relative power within each frequency band (δ, θ, α), block (BL–IV-block), and ROI (FRONTAL, LEFT, RIGHT) was log-transformed before statistical analyses.

The procedure was repeated for precontraction epochs (5-s rest periods between consecutive contractions) for all blocks. This control condition was implemented to test whether changes in spectral power between naive and informed blocks represent general longitudinal effects or are indeed specific to voluntary inhibitory mechanisms during the task execution.

**Statistical analysis**

Normality of all parameters (i.e., RCT, amplitude and latency of EMGs, log-transformed power) was assessed and confirmed through Lilliefors testing (α = 0.05). All further statistical analyses were performed using JASP (version 0.13.1; JASP Team, 2020). Block-wise differences in RCT were investigated using a repeated-measures ANOVA with the within-subject factor BLOCK (BL, I, II, III, IV). Bonferroni adjusted p values of post hoc comparisons are reported within the results. For FDIvol and FDIpMA (contraction and precontraction epochs), block-wise changes in amplitude were investigated using separate repeated-measures ANOVAs with the within-subject factor BLOCK (BL, I, II, III, IV). Here, the significance level was Bonferroni adjusted for four pairwise post hoc comparisons (BL vs I, BL vs II, BL vs III, BL vs IV; p adj < 0.0125). The same model was implemented to test for block-wise changes in latency of FDIpMA. In the case of a violation of the sphericity assumption, Greenhouse–Geisser correction was implemented.

In order to compare PSD during unilateral contractions performed in a naive versus informed state, two conditions were defined for each frequency band and ROI separately: (1) power of BL-block (NAIVE) and (2) power of I-block (INHIBIT). The comparison of these two conditions reflects immediate power changes accompanying initial behavioral inhibition. Changes in log-transformed spectral power (for contraction epochs as well as precontraction epochs separately) were subsequently analyzed through two-way repeated-measures ANOVAs with the within-subject factors ROI (FRONTAL, LEFT, RIGHT) and CONDITION (NAIVE, INHIBIT) for each frequency band separately (δ, θ, α). In the case that sphericity was violated, Greenhouse–Geisser correction was implemented. Simple main effect and pairwise post hoc analyses were conducted in case of significant interaction effects.

To test our hypothesis that the amplitude of pMA observed during naive (BL) and informed blocks (I, II, III, IV) is inversely associated with the individual δ power in the frontal ROI for each respective block, linear regression analyses were conducted on all data per block. δ Power served as dependent variables and corresponding pMA amplitudes served as predictors. Next, we assessed normality of residuals by way of Lilliefors testing (α = 0.05). All residuals were normally distributed. Thereafter, Pearson correlation coefficients were estimated for each block. To evaluate the significance of our results, δ power and pMA amplitude vectors were randomly permuted, resulting in a distribution of Pearson correlation coefficients from 1000 permutations (Manly, 2006). The Pearson correlation coefficients were deemed significant.
when they eclipsed the 95th percentile of permuted data. Family-wise error corrected p values of Pearson correlation coefficients are reported with the results. Additionally, Fisher’s r to z-transformation was implemented to test for a significant difference in the correlation coefficients of I-block and IV-block (Fisher, 1921).

The same procedure was used to estimate the association between participants’ initial pMA amplitudes (BL-block) and immediate rates of inhibition [pMA amplitude BL-block–I-block (%)].

For all statistical comparisons, p < 0.05 was considered significant. All p values Bonferroni adjusted for multiple comparisons are reported with the results.

An overview of statistical methods is provided in Table 2.

Results

Behavioral parameters

All participants consistently achieved the performance goal of the motor task by reaching the target field (corresponding to 40 ± 5% MVC) in every 5-s contraction during the experiment (i.e., 150 contractions). Repeated-measures ANOVAs indicated a significant effect for the factor BLOCK on RCT (F(2,1,39.9) = 11.975, p = 6.660 × 10⁻⁵, ηp² = 0.387). Post hoc comparison showed that RCT significantly increased from BL to II (MD = 257.098, SE = 48.409, p = 1.049 × 10⁻⁵, d = 1.189) and stayed elevated thereafter compared with BL during III (MD = 262.977, SE = 48.409, p = 6.442 × 10⁻⁴, d = 1.215) and IV (MD = 274.906, SE = 48.409, p = 1.049 × 10⁻⁵, d = 1.270).

EMG parameters

Repeated-measures ANOVAs indicated a significant effect for the factor BLOCK on mean amplitudes of FDVol (F(2,517.47,832) = 10.829, p = 4.092 × 10⁻⁵, ηp² = 0.363) and FDpMA (F(1,460.27,740) = 7.140, p = 0.006, ηp² = 0.273). More specifically, post hoc analyses showed a significant increase in amplitude from BL to II (MD = 0.032, SE = 0.012, p = 0.006, d = 0.630) and BL to IV (MD = 0.070, SE = 0.012, p = 4.748 × 10⁻⁴, d = 0.966) for FDVol (Fig. 2A). In all blocks where participants were informed about the nature of the experiment, the amplitude of FDpMA was significantly reduced, compared with the naive BL (Fig. 2B): BL to I (MD = −0.008, SE = 0.002, p = 2.851 × 10⁻⁵, d = −0.652), BL to II (MD = −0.008, SE = 0.002, p = 4.058 × 10⁻⁵, d = −0.645), BL to III (MD = −0.07, SE = 0.002, p = 5.683 × 10⁻⁴, d = −0.677), and BL to IV (MD = −0.006, SE = 0.002, p = 0.006, d = −0.515). Mean EMG amplitudes of FDVol during precontraction epochs showed no difference between blocks, suggesting the absence of systematic changes in EMG baseline over the course of the experiment (F(2.954,56.120) = 1.615, p = 0.179, ηp² = 0.078).

In case of FDpMA precontraction epochs, a significant effect of BLOCK was found (F(2.284,43.440) = 5.327, p = 0.006, ηp² = 0.219). Corresponding to contraction epochs of FDpMA, mean EMG amplitudes were significantly reduced in all blocks where participants were informed about the nature of the experiment (except for IV-block) compared with BL: BL to I (MD = −0.003, SE = 8.368 × 10⁻⁶, p = 0.006, d = −0.802), BL to II (MD = −0.003, SE = 8.368 × 10⁻⁶, p = 0.002, d = −0.865) and BL to III (MD = −0.003, SE = 8.368 × 10⁻⁶, p = 0.003, d = −0.855). For latency of FDpMA no significant effect for the factor BLOCK was found (F(4.000,76.000) = 1.456, p = 0.224, ηp² = 0.071).

δ Band

With regards to spectral power in the δ band (1–3 Hz) during contraction epochs, a repeated-measures ANOVA revealed a significant interaction effect for the factors ROI × CONDITION (F(2,38) = 3.706, p = 0.034, ηp² = 0.183, Table 3). A simple main effect analysis (simple effect factor: CONDITION; moderator factor: ROI) suggests the interaction was present in the FRONTAL ROI (p = 0.003). Pairwise post hoc comparisons confirmed a significant increase in δ power from the NAIVE to INHIBIT condition in the FRONTAL ROI (MD = 0.159, SE = 0.046, ηrob = 0.020, d = 0.777; Fig. 3). Furthermore, a significant main effect of the factor CONDITION (F(1,19) = 5.864, p = 0.026, ηp² = 0.236) was found, with spectral power being significantly higher during INHIBIT compared with NAIVE (MD = 0.092, SE = 0.038, ηrob = 0.026, d = 0.541). No significant main or interaction effects were found for precontraction epochs (all p > 0.05).

θ Band

In the θ band (4–8 Hz) no significant main or interaction effects were found, neither for contraction nor precontraction epochs (all p > 0.05).

α Band

For the α band (9–12 Hz) in contraction epochs, significant main effects for ROI (F(2,38) = 31.171, p = 9.720 × 10⁻⁹, ηp² = 0.621) and CONDITION (F(1,19) = 5.946, p = 0.025, ηp² = 0.238) were found. Pairwise post hoc analyses indicated significantly lower power in the FRONTAL ROI compared with the LEFT (MD = −0.387, SE = 0.065, ηrob = 2.887 × 10⁻⁵, d = −1.334) and RIGHT ROIs (MD = −0.367, SE = 0.058, ηrob = 1.475 × 10⁻⁵, d = −1.406). Furthermore, a significant increase in power from the NAIVE to INHIBIT condition was revealed (MD = 0.080, SE = 0.033, ηrob = 0.025, d = 0.545). However, no significant interaction effect was found.

For precontraction epochs, a similar result was obtained. Significant main effects for ROI (F(1,458,27.698) = 48.306, p = 1.042 × 10⁻⁸, ηp² = 0.718) and CONDITION (F(1,19) =

| Table 2: Statistical table |
|---------------------------|
| Data structure       | Type of test       | Power             |
| Normal distribution   | Repeated-measures ANOVA | Frequentist 95% confidence interval     |
| Normal distribution   | Pearson correlation | 95% confidence interval (estimated using 1000 permutations) |

September/October 2020, 7(5) ENEURO.0326-20.2020 eNeuro.org
were found. Pairwise post hoc analyses indicated significantly lower power in the FRONTAL ROI compared with the LEFT (MD = 0.447, SE = 0.063, \( p_{\text{bonf}} = 2.782 \)) and RIGHT ROIs (MD = 0.424, SE = 0.052, \( p_{\text{bonf}} = 3.937 \)). Furthermore, a significant increase in power from the NAIVE to INHIBIT condition was revealed (MD = 0.091, SE = 0.028, \( p_{\text{bonf}} = 0.004 \), \( d = 0.726 \)). Again, no significant interaction effect was found.

Correlation analysis

During the naive BL, frontal \( \delta \) power and the amplitude of pMA were not significantly associated (\( r_{(20)} = -0.340, p = 0.084 \)). Once participants started to inhibit pMA, a significant negative correlation emerged during I (\( r_{(20)} = -0.397, p = 0.046 \)), II (\( r_{(20)} = -0.406, p = 0.038 \)), III (\( r_{(20)} = -0.558, p = 0.006 \)) and IV (\( r_{(20)} = -0.670, p = 0.001 \); Fig. 4). Pearson correlation coefficient of I-block and IV-block were not significant different from each other (\( z \) score = 1.139, \( p = 0.127 \)).

Furthermore, we found a significant negative correlation between initial levels of pMA (pMA amplitude in BL-block) and the percentage change in pMA amplitude from BL-block to I-block. Asterisks indicate significant correlation after permutation testing (1000 permutations). BL, baseline block; I, first inhibition block; II, second inhibition block; III, third inhibition block; IV, fourth inhibition block.
showed the highest rates of inhibition from BL-block to I-block.

**Discussion**

Using a multimodal approach by combining EMG and EEG recordings, we provide novel evidence of pMA suppression in healthy adults. Suppression was accompanied by increased $\delta$ power in frontal areas. We show that the amplitude of pMA can be voluntarily inhibited during unilateral isometric contractions of intrinsic hand muscles. This was accomplished by asking participants to do so, after informing them about the presence of pMA and offering a basic understanding of pMA mechanisms through a standardized protocol. Importantly, significant inhibition of pMA was observed despite the absence of any online feedback during task execution and without special training.

A consistent and robust observation is that the amplitude of pMA is modulated by force demands of unilateral movements, i.e., increases with rising force requirements (Todor and Lazarus, 1986; Arányi and Rosler, 2002; Shinohara et al., 2003; van Duinen et al., 2008; Sehm et al., 2010, 2016; Maudrich et al., 2017). To avoid any influence of this relationship, the force level in this study was kept constant (40 ± 5% MVC). In addition, the quality of task performance, i.e., consistently achieving the desired force level, is expected to have an influence on pMA amplitude. However, all subjects reached the target field in each trial. Together, we were, therefore, able to keep two factors that mainly influence pMA amplitude constant. Interestingly enough, a significant reduction of pMA amplitude was achieved in all informed force blocks. As we intentionally interfered with the force dependency of pMA by directing participants’ attention toward the presence of pMA and asking them to inhibit any involuntary co-activation, this observation might suggest executive control, through directed attention, exerts inhibitory drive on involuntary motor output, suppressing pMA. In support of this, it has been shown in Go/No-Go reaction tasks, that primary motor cortex excitability is suppressed (Hoshiyama et al., 1996, 1997; Leocani et al., 2000), and short intracortical inhibition (SICI) is enhanced (Waldvogel et al., 2000; Sohn et al., 2002) during voluntary inhibition of movements, providing a potential neurophysiological basis for voluntary inhibition of pMA. Interestingly, we
observed different rates of inhibition between participants (Fig. 2C). In conjunction with this, we found a strong negative correlation between initial levels of pMA (pMA amplitude in I-block) and the percentage change in pMA amplitude from BL-block to I-block (Fig. 2D). This suggests that participants with higher initial pMA amplitudes showed the highest rates of inhibition. However, this correlation must be interpreted with caution. In some participants, very low initial pMA values and inhibition rates were present, suggesting on the one hand that pMA cannot demonstrate a strong decrease because of the “floor effect.” On the other hand, it might be possible that the signal-to-noise ratio of the surface EMG recordings could be limiting the potential for inhibition of participants with initially low pMA amplitudes in the present experiment. Intramuscular needle EMG might be better suited to detect the most subtle involuntary discharges of motor units and could, therefore, be used in future studies to resolve the question of whether pMA can be completely eliminated.

Investigations of right-handed adults, using similar isometric contractions to our implemented task, reported no side differences in pMA between right and left-hand contractions (Addamo et al., 2009; Koerte et al., 2010; Sehm et al., 2016; Maudrich et al., 2017, 2018). Therefore, it seems reasonable to assume that the choice of the contracting hand in the current study (left), most likely did not influence the obtained result. However, future studies are needed to replicate the voluntary inhibition of pMA during dominant hand contractions in right-handed adults.

In previous investigations regarding pMA, it has been ensured that participants were naïve to the study aim and generally the phenomenon of motor overflow to prevent any voluntary inhibition (Post et al., 2009; Maudrich et al., 2019). In the present experiment it seems remarkable that by shifting attention to the involuntarily contracting hand and establishing a basic understanding of pMA, most of the participants were able to significantly inhibit the amplitude of pMA in successive contraction blocks without any online feedback about their performance or special training. After the experiment, participants were asked to describe their self-chosen strategy used to inhibit pMA. While many different and highly individualized strategies have been mentioned, a main common denominator can be inferred. All participants, in one way or another, mentally focused on non-activation of the non-performing hand during each contraction of the performing hand. Thus, constantly paying attention to muscle relaxation in non-performing limbs might be a prerequisite for successful pMA inhibition besides being aware of its presence. However, no clear relation between individually employed strategies and participants inhibition rates is recognizable so that a decisive conclusion cannot be drawn from this dataset. Another strategy that was not reported but is manifested in increased RCTs during the experiment might be to perform more deliberate, less explosive contractions of the performing hand. Slower contractions of the performing hand, without changing the required target force level, might have allowed for easier suppression of pMA in the non-performing hand.

In line with the behavioral inhibition, we observed an increase in relative $\delta$ power from BL-block to the consecutive I-block in the frontal ROI (sensor space). This task-related increase in $\delta$ power was most likely not because of general longitudinal or non-stationarity effects during the experiment as our control comparison of non-task-related epochs (directly preceding muscle contractions) failed to reach significance. Previously, it has been hypothesized that there is a connection between $\delta$-oscillations and suppression of unwanted neural activity (Vogel et al., 1968). Furthermore, studies have shown that increases in $\delta$ power in frontal regions are associated with behavioral inhibition (Kamarajan et al., 2004; Knyazev, 2007; Putman, 2011), and it has been proposed that midfrontal $\delta$ power might serve as a selective marker for motor inhibition (Kaiser et al., 2019). Other investigations incorporating Go/No-Go tasks have observed an increase in $\delta$ activity during inhibition of movement in frontal electrodes (Harmony et al., 2009). Moreover, another study investigating the neurocognitive effects of alcoholism proposed that decreased $\delta$ and $\theta$ power in frontal regions, associated with No-Go processing, might suggest a deficient inhibitory control and information-processing mechanism in alcoholics (Kamarajan et al., 2004). In addition, we found a significant inverse association between the amplitude of pMA and frontal $\delta$ power that emerged once participants were instructed to inhibit pMA and persisted throughout all successive force blocks (Fig. 4). Overall, these results suggest that $\delta$ power in frontal regions might reflect executive processes relating to inhibitory control (Chambers et al., 2009; Munakata et al., 2011), modulating unintentional motor output, in our case pMA. Importantly, the fact that participants were able to suppress pMA immediately after they were instructed indicates that no special training was required. Thus, a reduction of involuntary motor output had rather a categorical character akin to a termination of a given behavioral output. Such regulation of behavior is a hallmark of executive functions, particularly inhibitory control, which often has a character of a veto, once undesirable consequences of the behavior become known (Kühn et al., 2009). In such a case, the over-riding of pMA is similar to not performing a Go-response in a No-Go paradigm, once the participant is made aware that she/he was committing this response but were not yet aware of such behavior. Bringing the presence of pMA to their attention thus requires them to perform a Go-response with one hand (voluntary contraction) while simultaneously preventing this response with the other hand (involuntary muscle activation).

Contrary to our hypothesis, we did not observe changes in $\theta$ power from BL-block to I-block. This seems surprising, as a previous investigation found a $\theta$ power increase associated with lateralized inhibition of symmetrical movements in the fronto-mesial area, attributed to executive processes (Tisseyre et al., 2019). This contradiction may be explained by differences between our task and those from previous studies showing $\theta$ power increases during motor inhibition. Such increases in $\theta$ power were often demonstrated during voluntary response inhibition paradigms (Cohen et al., 2011; Andreou...
et al., 2017). In this context, voluntary response inhibition is closely related to feedback on task performance, meaning that inhibitory strategies and their success rates can be assessed and optimized during task performance (Yamanaka and Yamamoto, 2010). Thus, this ability to strategically fine-tune inhibitory strategies (Kirmizi-Alsan et al., 2006; Yamanaka and Yamamoto, 2010; Tisseyre et al., 2019) is a major difference compared with the task we employed in this study. Because of the inherent characteristics of the motor task in the present study, where participants were entirely unaware regarding successful inhibition of pMA throughout the experiment, strategic fine-tuning is unlikely to have occurred because of participants’ uncertainty regarding successful inhibition. This mechanistic difference may serve as an explanation for the absence of an increase in $\theta$ power as participants could not have been certain regarding successful pMA inhibition and, therefore, stable $\theta$ dynamics, reflecting voluntary inhibitory response mechanisms did not emerge.

With regards to $\alpha$-oscillations, which among other functions generally reflect inhibitory control mechanisms in the motor domain (Klimesch et al., 2007; Sauseng et al., 2013), we observed overall global power increases from naive to informed contractions regardless of ROI. Furthermore, this global increase in $\alpha$ power was also observed in our control analysis of non-task related epochs. An increase in $\alpha$ power over time has previously been associated with the level of sustained attention and fatigue (Boksem et al., 2005; Craig et al., 2012). Recently, multiple endogenous non-stationarity processes in $\alpha$ band activity, i.e., an increase in power, have been shown to occur over time (Benwell et al., 2019). Therefore, the global $\alpha$ power increase observed in the present study most likely reflects the longitudinal effects of fatigue, sustained attention, or endogenous non-stationarity processes. However, the absence of specific changes in $\alpha$ power further supports our interpretation that inhibition of pMA in this study is mediated by executive control, rather than by purely motor inhibition processes.

As a limitation of the present study, we are not able to disentangle the distinct effects of directed focus on pMA presence and actual inhibitory processes on the observed behavioral suppression of pMA. One previous study observed that pMA remained unchanged after young adults were informed of it but asked to ignore their motor overflow. Only when participants were requested to inhibit involuntary co-activation with and without visual feedback were they able to reduce pMA (Addamo et al., 2010). While this within-subject study protocol seems appropriate to disentangle the effects of directed focus and actual inhibitory processes on pMA suppression in general, we suggest that a between-subject protocol might be an alternative. As part of this between-subject protocol, one intervention group would be informed about pMA presence but instructed to ignore it, while a separate intervention group is additionally instructed to actively inhibit pMA. A control group would be naive regarding pMA throughout the experiment. This experimental design could be repeated over several days to investigate the stability of these mechanisms or even learning effects in pMA inhibition over time. However, as stated before, force measurements are insufficient in capturing neurophysiological aspects of pMA in its entirety, rendering future studies incorporating EMG measurements necessary to answer these questions raised above.

In conclusion, we provide novel evidence for voluntary suppression of pMA in healthy adults. As proposed by others, the knowledge of voluntary pMA modulation via attentional processes might be helpful for therapeutic applications in the context of neurorehabilitation, i.e., for the development of cognitive-behavioral strategies for the treatment of mirror movements (MM) that are not a result of major anatomic abnormalities (e.g., uncrossed corticospinal tract; Addamo et al., 2007, 2009). One possible intervention might be to implement online feedback of involuntary muscular activity to voluntarily decrease the level of EMG and even possibly the occurrence of overt MM, as it has been demonstrated previously in brain-injured young adults (Lazarus, 1992). However, the transfer of pMA-mechanisms to pathologic MM remains elusive and requires further research. Nevertheless, this inhibitory feedback-based approach appears additionally promising in the context of counteracting the age-related increases in pMA, which might help prevent the decline in motor performance documented in the elderly.

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