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1 **Distinct neural representations of hand movement direction between**  
2 **motor imagery and execution in the presupplementary motor area**

3

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## 1 **Abstract**

2 Motor simulation theory proposes a functional equivalence between motor execution  
3 (ME) and its simulation, suggesting that motor imagery (MI) is the self-intentioned  
4 simulation of one's actions. This study used functional magnetic resonance imaging  
5 (fMRI) with multivoxel pattern analysis to test whether the direction of hand movement  
6 is represented with a similar neural code between ME and MI. In our study, participants  
7 used their right hand to move an on-screen cursor in the left–right direction with a  
8 joystick or imagined the same movement without execution. The results indicated that  
9 the left–right direction as well as their modality (ME or MI) could be decoded  
10 significantly above the chance level in the presupplementary motor area (pre-SMA) and  
11 primary visual cortex (V1). Next, we used activation patterns of ME as inputs to the  
12 decoder to predict hand move directions in MI sessions and found a significantly  
13 higher-than-chance accuracy only in V1, not in pre-SMA. Moreover, the  
14 representational similarity analysis showed similar activation patterns for the same  
15 directions between ME and MI in V1 but not in pre-SMA. This study's finding indicates  
16 distinct spatial activation patterns for movement directions between ME and MI in pre-  
17 SMA.

18

## 19 **Keywords**

20 motor imagery; motor execution; fMRI; multivoxel classification analysis;  
21 representational similarity analysis; presupplementary motor area

22

## 1 **Introduction**

2 Motor imagery (MI) is a cognitive ability defined as a “mental simulation” of motor  
3 execution (ME) without actual action (Decety, 1996; Grush, 2004; Hanakawa, 2016). It  
4 has been believed that the neural state of an imagined movement is similar to the state  
5 of execution of that action (Jeannerod, 2001). Early neuroimaging studies showed that  
6 MI and ME activate roughly the same brain regions (Hanakawa et al., 2008; Munzert et  
7 al., 2009). Moreover, a large overlap of regions between MI and ME was found in a  
8 meta-analysis study (Hardwick et al., 2018). However, these studies mainly analyzed a  
9 single voxel activity or averaged activities within the region and not the activation  
10 patterns among multivoxels. Thus, although MI and ME could activate the same brain  
11 regions, it remains unclear whether MI and ME use similar neural codes for the same  
12 action.

13         Recently, a technique called multivoxel pattern analysis (MVPA) was developed  
14 (Weaverdyck et al., 2020). MVPA examines the spatial pattern of brain activations,  
15 whereas univariate analyses only consider the overall magnitude of the responses.  
16 MVPA studies showed that MI for different types of right-hand actions could be  
17 decoded significantly above chance level in M1 and premotor cortices (Pilgramm et al.,  
18 2016). Moreover, these different hand actions could also be decoded between MI and  
19 ME (cross-model) in premotor cortices. However, in representational similarity analysis  
20 (RSA), representational dissimilarity matrices showed that MI and ME represent  
21 separate clusters, although the representational organization of action types within these  
22 clusters was identical (Zabicki et al., 2017). Therefore, premotor cortices use similar  
23 neural codes for different types of hand actions between MI and ME.

24         By contrast, it remains unclear whether the different directions of specific hand

1 action use the same neural code between MI and ME. Ogawa and Inui instructed the  
2 participants to perform visually guided movements using a normal mouse and a left–  
3 right reversed mouse. Their study showed that the direction of hand movement could be  
4 decoded in the hand region of the primary motor area (Mot) (Ogawa & Inui, 2012). Our  
5 study thus attempted to decode ME of hand movement direction and investigate whether  
6 we could decode MI in motor-related regions.

7 Our study also examined whether the same hand movement, but with different  
8 directions in MI and ME used similar neural codes. Our participants first performed the  
9 ME tasks by moving their right hand using a joystick to move an invisible cursor to the  
10 left or right target. They subsequently performed the MI tasks, imagining the same  
11 action as the ME tasks. This experimental design allowed us to compare the activity  
12 pattern between ME and MI using multivoxel classification analysis and RSA.

13

## 14 **Materials and Methods**

### 15 **Participants**

16 Participants were 17 volunteers (12 females, 5 males) from Hokkaido University, with  
17 an average age of 23.18 years (range = 20–26, SD = 1.74). Of these, two female  
18 participants were excluded because of excessive head movement during scanning. All  
19 participants were right-handed, according to the Edinburgh Handedness Inventory. The  
20 sample size was estimated from a prior hand-moving decoding study (Ogawa & Inui,  
21 2012) using G\*Power version 3.1.9 (Erdfelder et al., 2009; Faul et al., 2007). We used  
22 15 participants to get power  $(1 - \beta) = .95$ , with  $\alpha = .05$  and Cohen's  $d = 1.03$ .

23

### 24 **Task procedures**

1 All participants completed two practice sessions containing ten trials before three ME  
2 sessions and three MI sessions (20 trials per ME and MI session) in a functional  
3 magnetic resonance imaging (fMRI) scanner without scanning. Experimental stimuli  
4 were controlled by Psychophysics Toolbox Version 3 (PTB-3) (Brainard, 1997; Kleiner  
5 et al., 2007; Pelli, 1997) in MATLAB (The MathWorks, Inc.).

6

### 7 **Practice session**

8 In practice session 1, for each trial, a white fixation was presented in the center of the  
9 screen, and above the fixation, there was a countdown from “3” to “1” that lasted 3 s. At  
10 the end of the countdown, two squares were presented on the left and right sides of the  
11 screen, and the color of the central fixation changed to green or yellow (target phase),  
12 indicating the target (green: left square; yellow: right square). Half of the ten trials were  
13 green, and the other half was yellow, presented in random order. After 2 s, the color of  
14 central fixation changed to red (execution phase). During the execution phase, a joystick  
15 cursor (a small white “x”) was presented centrally on the screen. The participants then  
16 moved the cursor with their right hand to the target square, which was indicated in the  
17 target phase, and maintained the cursor in the target square until the color of the central  
18 fixation changed from red to white. The execution phase lasted for 2 s, and then, the  
19 cursor was frozen, showing the participants the final position of the cursor in the  
20 execution phase for 3 s (result phase). Participants then allowed the joystick to return to  
21 its original position and let the joystick bring their right hand back (Figure 1).  
22 Participants repeated practice session 1 until getting 100% accuracy. Practice session 2  
23 is almost the same as practice session 1. The difference was that in the execution phase  
24 of practice session 2, the joystick cursor was not presented. Moreover, it showed the

1 cursor's last position in the execution phase to the participants in the result phase.

2

### 3 **Execution session**

4 During the execution sessions, participants completed three execution sessions with  
5 fMRI scanning. The differences between execution sessions and practice session 2 were  
6 that each execution session included 20 trials (half were left, and half were right), and  
7 the time of countdown in the countdown phase was from 3 to 9 s (Figure 2).

8

### 9 **Imagery session**

10 After three execution sessions, participants completed three imagery sessions (20 trials  
11 per session). In the imagery session, the execution phase changed to the imagery phase,  
12 and the result phase changed to the evaluation phase with the same duration. In the  
13 imagery phase, participants imagined that they move the cursor by using the joystick  
14 and put the cursor into the target, which was indicated in the target phase, without actual  
15 movement. Participants were instructed to use both kinesthetic and visual images before  
16 the practice session. After the imagery phase, the participants immediately evaluated the  
17 quality of the MI in this trial using their left hand (from 1: very good to 4: very poor). In  
18 the evaluation phase, participants were instructed only to choose "4" when they failed to  
19 imagine before the practice session, which helped us to exclude the error.

20

### 21 **MRI acquisition**

22 "All scans were performed on a Siemens (Erlangen, Germany) 3-Tesla Prisma scanner  
23 with a 64-channel head coil at Hokkaido University. T2\*-weighted echo-planar imaging  
24 (EPI) was used to acquire a total of 170 scans per session, with a gradient EPI sequence.

1 The first three scans within each session were discarded to allow for T1 equilibration.  
2 The scanning parameters were repetition time (TR), 2000 ms; echo time (TE), 30 ms;  
3 flip angle (FA), 90°; field of view (FOV), 192 × 192 mm; matrix, 94 × 94; 35 axial  
4 slices; and slice thickness, 3.0 mm with a 0.75 mm gap. T1-weighted anatomical  
5 imaging with an MP-RAGE sequence was performed using the following parameters:  
6 TR, 2300 ms; TE, 2.41 ms; FA, 8°; FOV; 256 × 256 mm; matrix, 256 × 256; 224 axial  
7 slices; and slice thickness, 0.8 mm without a gap.

8

### 9 **Exclusion criteria for data**

10 For more accurate data analysis, we excluded some fMRI data based on the behavioral  
11 criteria below.

12 a. The trial in which the participant moved the joystick before the execution phase in  
13 execution sessions.

14 b. The trial in which the participant did not put the cursor in the target square at the end  
15 of the execution phase in execution sessions.

16 c. The trial in which the participant moved the joystick to the wrong direction in  
17 execution sessions, although the cursor was in the correct target square at the end of the  
18 execution phase.

19 d. The trial in which participants chose “4 very poor” in the evaluation phase of imagery  
20 sessions.

21

### 22 **Definition of regions of interest (ROIs)**

23 We defined motor-related regions as bilateral pre-SMA, SMA, and left M1, PMv using  
24 Human Motor Area Template (Mayka et al., 2006), and left V1 was defined as

1 Brodmann Area 17. The activity of the left M1 reflected the movement of the right hand,  
2 whereas the direction of movement of the right hand can also be classified in the left M1  
3 (Ogawa & Inui, 2012). SMA and premotor cortex were associated with MI (Decety,  
4 1996). Furthermore, PMv was related to hand actions (Rizzolatti et al., 2002). Pre-SMA  
5 was also related to MI (Hanakawa et al., 2003), and pre-SMA was activated when the  
6 cursor was unavailable during visual guided movement (Ogawa et al., 2006; Ogawa &  
7 Inui, 2007).

8

### 9 **fMRI mass-univariate analysis**

10 Image preprocessing was performed using the SPM12 software (Wellcome Department  
11 of Cognitive Neurology, <http://www.fil.ion.ucl.ac.uk/spm> ). All functional images were  
12 initially realigned to adjust for motion-related artifacts. Volume-based realignment was  
13 performed by co-registering images using rigid body transformation to minimize the  
14 squared differences between volumes. The realigned images were then spatially  
15 normalized with the Montreal Neurological Institute template based on the affine and  
16 nonlinear registration of coregistered T1-weighted anatomical images (normalization  
17 procedure of SPM). They were resampled into 3-mm-cube voxels with the sinc  
18 interpolation. Images were spatially smoothed using a Gaussian kernel of  $6 \times 6 \times 6$ -mm  
19 full width at half-maximum. However, images used for MVPA were not smoothed to  
20 avoid blurring the fine-grained information contained in the multivoxel activity  
21 (Kamitani & Sawahata, 2010; Mur et al., 2009; Ogawa et al., 2019). We analyzed  
22 significantly activated areas during the ME or MI of right-hand movement compared  
23 with activation during rest with univoxel analysis. Activation was the threshold at  $p$   
24  $< .05$ , corrected for multiple comparisons for a family-wise error, with an extent

1 threshold of 10 voxels.

2

### 3 **Multivoxel pattern analysis**

4 In decoding analysis of MVPA, we classified the direction of hand movement in ME  
5 and MI. The classification was performed based on a linear support vector machine run  
6 by LIBSVM (<http://www.csie.ntu.edu.tw/~cjlin/libsvm>) with a fixed regularization  
7 parameter  $C = 1$ . The beta value for each trial of voxels within ROIs (see Table 1) was  
8 used as inputs to the classifier. ROI size did not affect the linear SVM's decoding  
9 accuracy (Misaki et al., 2010). We attempted to interpret the direction of hand  
10 movement in only ME (ME classification) or MI (MI classification) and between ME  
11 and MI (cross-classification). Each participant attended three ME sessions and three MI  
12 sessions. In ME classification and MI classification, we estimated the average  
13 classification accuracy by a three-fold "leave-one-out" cross-validation, in which two  
14 sessions were used as training and the remaining session was used as test data. In cross-  
15 classification, the averaged classification accuracy was estimated via validation between  
16 three ME sessions and three MI sessions (Table 2). Such cross-classification between  
17 different task sets or stimuli has been used to investigate the activation pattern  
18 similarities (Ogawa & Imai, 2016).

19 To compare spatial activation pattern similarities for different directions across  
20 ME and MI, RSA (Kriegeskorte et al., 2008) was also conducted. Beta values of voxels  
21 within ROIs were used as inputs to estimate the representational dissimilarity matrix  
22 among the different directions of hand movement between ME and MI. There were a  
23 total of 30 trials for each direction and modality (3 sessions  $\times$  10 trials). Dissimilarity  
24 was measured with cross-validated Mahalanobis distance (Ejaz et al., 2015), which

1 presents reliable dissimilarity metrics for RSA (Walther et al., 2016). To ensure  
2 invertibility and stability, the voxel-by-voxel noise covariance matrix was separately  
3 estimated within one dataset using an optimal shrinkage algorithm (Ledoit & Wolf,  
4 2003). We then compared the off-diagonal elements of the representational dissimilarity  
5 matrix, which represent the dissimilarity of activation patterns between different  
6 modalities and directions.

7

## 8 **Results**

### 9 **Behavioral analysis**

10 According to the exclusion criteria, we excluded the error trials based on behavior data.  
11 In a total of 60 trials of ME sessions and 60 trials of MI sessions, the percentage (SD) of  
12 error trials per participant in ME and MI sessions was 2.78% (3.77%) and 2.12%  
13 (3.18%), respectively.

14

### 15 **fMRI mass-univariate analysis**

16 We analyzed the activated regions of the brain using a univariate analysis of single  
17 voxels and the regions that were significantly activated by comparing the modalities  
18 (ME vs. MI) and the direction of movements (left vs. right). Activities between the left–  
19 right directions in ME were compared. No areas significantly differed between the left  
20 and right directions in ME at the corrected threshold of  $p < .05$  and an extent threshold  
21 of 10 voxels. Both left and right directions of the right hand moving revealed the  
22 activations in the left M1 and left insula (Figure 3 and Table 3). Next, we compared the  
23 activity during which the participants imagined the right hand moving between the left  
24 and right directions. This comparison also revealed that no areas were significantly

1 differently activated between the left and right directions in MI. Both left and right  
2 directions of hand-moving imagery revealed the activations in the bilateral insula and  
3 SMA (Figure 3). Next, we compared the activated regions between the ME sessions and  
4 the MI sessions. Activations were found to be majorly in the left M1 and vermis during  
5 ME sessions and right M1 during MI sessions. There were no overlapped areas between  
6 “ME > rest” and “MI > rest” at the corrected threshold of  $p < .05$  and an extent  
7 threshold of 10 voxels (Figure 4).

8

9         Next, ROI analysis was performed to compare the averaged parameter estimates  
10 (beta values) between the ME and MI sessions and the left and right directions.

11 Repeated measures analysis of variance was conducted with the modalities (ME and MI)  
12 and the hand movement directions (left and right) as within the subjects’ factors (Figure  
13 5). In both left M1 and bilateral SMA, a significant main effect was observed between  
14 the ME and MI (left M1,  $F(1, 14) = 43.791, p < .001, \mu_p^2 = .758$ ; bilateral SMA,  $F(1, 14)$   
15  $= 8.569, p = .011, \mu_p^2 = .380$ ) and also a significant interaction between the two factors  
16 (left M1,  $F(1, 14) = 11.274, p = .005, \mu_p^2 = .446$ ; bilateral SMA,  $F(1, 14) = 13.074, p$   
17  $= .003, \mu_p^2 = .483$ ). The beta value of the left M1 and bilateral SMA was significantly  
18 higher when the hand was moving to the left than the right in ME sessions (left M1,  $F =$   
19  $8.132, p = .013$ ; bilateral SMA,  $F = 12.149, p = .004$ ). In bilateral pre-SMA, there was  
20 no main effect but a significant interaction was observed ( $F(1, 14) = 8.375, p = .012, \mu_p^2$   
21  $= .374$ ). In the right direction, the beta value of bilateral pre-SMA in the MI sessions  
22 was significantly higher than in the ME sessions ( $F = 5.134, p = .040$ ). Also, in the ME  
23 sessions, the beta value of bilateral pre-SMA was significantly higher when the hand  
24 was moving to the left than the right ( $F = 11.759, p = .004$ ).

1

## 2 **Multivoxel classification analysis**

3 We first conducted MVPA to classify the direction of ME by the subjects using the  
4 activities of each ROI. In left M1, significantly higher-than-chance classification  
5 accuracy was observed ( $t(14) = 6.49, p < .001$ , Cohen's  $d = 1.68$ ). We also found  
6 significantly higher-than-chance classification accuracies in bilateral pre-SMA ( $t(14) =$   
7  $3.38, p = .005$ , Cohen's  $d = .87$ ), SMA ( $t(14) = 2.61, p = .021$ , Cohen's  $d = .67$ ), and left  
8 PMv ( $t(14) = 2.42, p = .030$ , Cohen's  $d = .62$ ), V1 ( $t(14) = 11.89, p < .001$ , Cohen's  $d =$   
9  $3.07$ ) (Figure 6A). Next, we conducted MVPA to classify the direction of MI. We found  
10 significantly higher-than-chance classification accuracies in bilateral pre-SMA ( $t(14) =$   
11  $2.61, p = .010$ , Cohen's  $d = .77$ ) and left V1 ( $t(14) = 3.29, p = .005$ , Cohen's  $d = .85$ ),  
12 but no significant difference in bilateral SMA ( $t(14) = -.87, p = .398$ , Cohen's  $d = -.23$ )  
13 and left M1 ( $t(14) = 1.13, p = .277$ , Cohen's  $d = .29$ ), PMv ( $t(14) = 1.07, p = .304$ ,  
14 Cohen's  $d = .28$ ) (Figure 6B).

15 We also conducted MVPA to classify the direction across ME and MI. We  
16 found significantly higher-than-chance classification accuracies in left V1 ( $t(14) = 5.87$ ,  
17  $p < .001$ , Cohen's  $d = 1.52$ ). However, no significant difference in bilateral pre-SMA  
18 ( $t(14) = .44, p = .665$ , Cohen's  $d = .11$ ), SMA ( $t(14) = -.15, p = .883$ , Cohen's  $d = -.04$ )  
19 and left M1 ( $t(14) = .95, p = .358$ , Cohen's  $d = .25$ ), PMv ( $t(14) = .31, p = .763$ , Cohen's  
20  $d = .08$ ) (Figure 6C). These results indicated distinct spatial activation patterns for the  
21 movement directions between ME and MI in pre-SMA.

22

## 23 **Representational similarity analysis**

24 The RSA was used to investigate similarity in activation patterns of bilateral pre-SMA

1 and left V1 between different directions (left vs. right) across modalities (ME and MI,  
2 Figure 7A). Because bilateral pre-SMA and left V1 were the only ROIs that were  
3 significantly higher-than-chance in the ME classification and MI classification, a paired  
4 sample t-test was conducted with the dissimilarity between the same direction but  
5 different modalities (e.g., ME left and MI left) and different directions with different  
6 modalities (e.g., ME left and MI right) across ME and MI for bilateral pre-SMA and left  
7 V1 (Figure 7B). The result showed that different direction dissimilarities across ME and  
8 MI was significantly higher than the dissimilarity of the same direction in left V1 with  
9 null hypothesis significance testing, whereas there was no evidence for a difference  
10 between the dissimilarities by Bayes factor ( $t(29) = 2.06, p = .049$ , Cohen's  $d = .376$ ,  
11  $BF_{10} = 1.221$ ). There was moderate evidence for no difference between the dissimilarity  
12 of different directions across modalities and the dissimilarity of same direction across  
13 modalities in bilateral pre-SMA ( $t(29) = .33, p = .745$ , Cohen's  $d = .060, BF_{10} = .204$ ).

14

## 15 **Discussion**

16 This study investigated whether the different hand movement directions shared a similar  
17 neural code in motor-related regions between ME and MI. We first analyzed activated  
18 regions in the whole brain with conventional univoxel analysis within the ME and MI  
19 sessions. During the execution phase of the ME session, significantly increased  
20 activations were found in left M1 and left insula in both left and right directions. The  
21 left M1 is well known for its role in right-hand movements (Sanes et al., 1995).  
22 Contralateral insula activation has also been reported during voluntary limb movement  
23 (Chollet et al., 1991) and finger movements (Fink et al., 1997). During the imagery  
24 phase of the MI session, both left and right directions of hand moving imagery revealed

1 the activations in bilateral insula and SMA. A previous study indicated that SMA  
2 activity was associated with MI (Decety, 1996). The comparison between left and right  
3 direction hand movements in both ME and MI sessions showed no significant activated  
4 areas. This result indicated that conventional univoxel analysis could not distinguish  
5 differences in brain activity between the left and right directions of hand movements in  
6 both ME and MI.

7 ROI analysis of the beta value revealed that left M1, SMA, and pre-SMA  
8 activities were significantly higher during right hand moving to the left direction than  
9 the right direction in ME sessions. The previous study showed that right hands that were  
10 rotated in a clockwise direction were more difficult than when they were rotated in a  
11 counterclockwise direction (de Lange et al., 2006). That might cause stronger activities  
12 in left M1, SMA, and pre-SMA during the movement of the right hand to the left side.

13 Our multivoxel classification analysis in ME classification revealed that the  
14 classification accuracies in bilateral SMA, pre-SMA, and left M1, PMv, and V1 were  
15 significantly higher than the chance level. The left M1 and V1 result is consistent with  
16 the previous study (Ogawa & Inui, 2012), which also classified the hand movement  
17 directions while participants used a mouse. Bilateral SMA is associated with the  
18 preparation and readiness for voluntary movements (Cunnington et al., 1996, 2003),  
19 whereas PMv is related to hand actions (Rizzolatti et al., 2002). Pre-SMA is related to  
20 visuomotor imagery (Deiber et al., 1998; Johnston et al., 2004; Leek & Johnston, 2009).  
21 Previous studies showed that pre-SMA activates when visual feedback is unavailable  
22 during visual guided movements (Ogawa et al., 2006; Ogawa & Inui, 2007). In the  
23 present study, there was no visual feedback during ME sessions, the participants might  
24 visually estimate cursor position during right-hand movement. Our multivoxel

1 classification analysis in MI classification revealed that significantly higher-than-chance  
2 classification accuracy occurred only in bilateral pre-SMA and left V1, which showed  
3 that V1 and pre-SMA were also related to hand movement direction in MI.

4 We conducted cross-classification to investigate whether the different hand  
5 movement directions between ME and MI shared a similar neural code in bilateral pre-  
6 SMA and left V1. We only found significantly higher-than-chance accuracy in left V1,  
7 not in the bilateral pre-SMA. This result suggests that the activation of pre-SMA,  
8 although related to both ME and MI, and the activity patterns in pre-SMA are separate  
9 from each other. We conducted RSA to confirm this result further. The result of RSA  
10 showed that the dissimilarity of different directions across ME and MI was significantly  
11 higher than the dissimilarity of the same direction in left V1, and there was moderate  
12 evidence for no difference between them in bilateral pre-SMA. Thus, the bilateral pre-  
13 SMA did not share a similar neural code in hand movement direction across ME and MI.

14 MI is generally classified into two different types: kinesthetic type and visual  
15 type (Jeannerod, 1995). Our participants might unconsciously imagine the trajectory of  
16 the cursor movement while moving the joystick during the ME sessions. In the MI  
17 sessions, the participants were instructed to imagine both the visual and kinesthetic  
18 aspects of hand movement. Although visuomotor imagery was present in both ME and  
19 MI sessions, the participants might have had difficulty visualizing the hand and cursor  
20 movements simultaneously. This might be the reason why pre-SMA activation patterns  
21 were different between ME sessions and MI sessions. Additionally, whereas pre-SMA  
22 is related particularly to visuomotor imagery (Deiber et al., 1998), we cannot deny the  
23 possibility that our participants primarily used the kinesthetic type of MI during MI  
24 sessions, which might also allow pre-SMA to be decoded with significantly higher

1 accuracy than the chance level.

2

### 3 **Conclusion**

4 In summary, using multivariate analysis of fMRI activities, we found that pre-  
5 SMA was the only motor-related region related to hand movement direction in both ME  
6 and MI, but the activity patterns of ME and MI were distinct from each other.

7

### 8 **Glossary:**

9 motor execution, ME; motor imagery, MI; functional magnetic resonance imaging,  
10 fMRI; presupplementary motor area, pre-SMA; multivoxel pattern analysis, MVPA;  
11 representational similarity analysis, RSA; echo-planar imaging, EPI; repetition time, TR;  
12 echo time, TE; flip angle, FA; field of view, FOV; regions of interest, ROIs.

13

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21

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23

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1 **Tables**

2 Table 1: Number of ROIs' voxels in MVPA

ROI	number of voxels (SD)
left M1	925.73 (54.31)
bilateral pre-SMA	598.80 (21.67)
bilateral SMA	696.93 (10.82)
left PMv	788.20 (30.23)
left V1	457.47 (14.63)

3

4 Table 2: Training sets and test sets in multivoxel pattern classification

Training sets	number of trials	Test sets	number of trials
<i>ME classification</i>			
ME session 1&2	40	ME session 3	20
ME session 1&3	40	ME session 2	20
ME session 2&3	40	ME session 1	20
<b>total</b>	120		60
<i>MI classification</i>			
MI session 1&2	40	MI session 3	20
MI session 1&3	40	MI session 2	20
MI session 2&3	40	MI session 1	20
<b>total</b>	120		60
<i>Cross-classification</i>			
ME session 1,2&3	60	MI session 1,2&3	60
MI session 1,2&3	60	ME session 1,2&3	60
<b>total</b>	120		120

5

6 Table 3: Anatomical regions, peak voxel coordinates, and t-values of observed  
7 activations.

Anatomic region	voxels	MNI coordinates	t-value
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		x	y	z	
<i>Execution</i>					
L Precentral cortex	153	-33	-28	62	12.67
Postcentral cortex		-33	-22	50	12.37
R Lingual gyrus	37	21	-79	2	12.58
Calcarine sulcus		12	-79	11	10.34
Vermis	107	3	-61	-13	11.63
R Cerebellum		21	-46	-19	9.89
L Insula	35	-39	-4	11	11.35
Rolandic operculum		-48	-1	5	9.32
L Middle occipital gyrus	13	-15	-95	-1	10.19
L Rolandic operculum	13	-45	-28	17	9.27
<i>Imagery</i>					
R SMA	185	12	8	65	16.33
L SMA		3	14	50	13.63
L Superior parietal lobule	44	-21	-67	53	15.04
R Inferior frontal gyrus	36	48	11	2	14.50
Insula		42	5	2	11
R Insula	33	33	26	-1	13.53
R Fusiform	79	33	-61	-10	13.05
Lingual gyrus		18	-76	-7	12.02
L Insula	90	-39	17	2	12.65
L Precentral cortex	14	-54	8	23	11.4
R Inferior frontal gyrus	16	54	11	17	10.83
L Supramarginal	21	-60	-28	38	10.62
R Precentral cortex	71	39	-13	56	10.61
L Fusiform	47	-33	-55	-16	10.44
Lingual gyrus		-21	-76	-10	9.74
L Middle frontal gyrus	18	-27	-4	53	9.64
<i>Execution &gt; Imagery</i>					
L Postcentral cortex	93	-36	-31	56	13.47
Precentral cortex		-39	-19	59	8.81
Vermis	14	3	-61	-19	10.84
<i>Imagery &gt; Execution</i>					
R Precentral cortex	41	36	-19	53	13.27

*Execution Left*

L Precentral cortex	128	-33	-28	62	13.93
Postcentral cortex		-48	-22	53	11.04
R Lingual gyrus	25	21	-79	2	11.75
Vermis	95	3	-61	-16	10.72
R Cerebellum		18	-46	-19	10.30
L Insula	15	-42	-4	8	9.65

*Execution Right*

L Postcentral cortex	144	-33	-22	50	13.00
Precentral cortex		-33	-28	59	11.82
L Insula	36	-39	-4	11	12.61
Rolandic operculum		-48	-1	5	10.35
R Cerebellum	38	33	-52	-22	10.62
Vermis	33	3	-64	-10	10.49
L Lingual gyrus	10	-12	-76	-1	9.68

*Imagery Left*

R Insula	27	36	26	-1	13.23
R Inferior frontal gyrus	25	48	11	2	11.71
Insula		42	5	-1	10.49
L SMA	49	3	14	50	11.37
L Superior parietal lobule	26	-21	-67	53	11.35
L Insula	50	-36	17	2	10.52
Inferior frontal gyrus		-51	11	2	9.62
R Lingual gyrus	18	18	-76	-7	10.46
R Precentral cortex	10	36	-7	50	10.28
L Lingual gyrus	30	-21	-76	-7	10.10
Fusiform		-27	-61	-10	9.60
R SMA	10	12	8	65	10.01
R Postcentral cortex	11	48	-19	44	9.67
L Supramarginal	10	-60	-31	41	9.66

*Imagery Right*

R Fusiform	149	33	-61	-10	23.91
Lingual gyrus		18	-79	-7	12.09
L Superior parietal lobule	80	-27	-61	44	15.66
R SMA	181	12	8	65	13.82
L SMA		3	14	50	13.46
L Insula	76	-33	23	8	13.59
R Precentral cortex	141	39	-10	59	13.10

Postcentral cortex		45	-22	53	10.23
L Fusiform	72	-33	-58	-16	11.87
Lingual gyrus		-18	-79	-10	10.21
L Supramarginal	24	-60	-31	38	11.77
R Insula	69	33	26	2	11.73
Inferior frontal gyrus		48	11	2	11.51
L Superior frontal gyrus	19	-21	-1	68	10.32
L Precentral cortex	16	-36	-7	41	10.32
L Inferior frontal gyrus	15	-51	11	20	9.58

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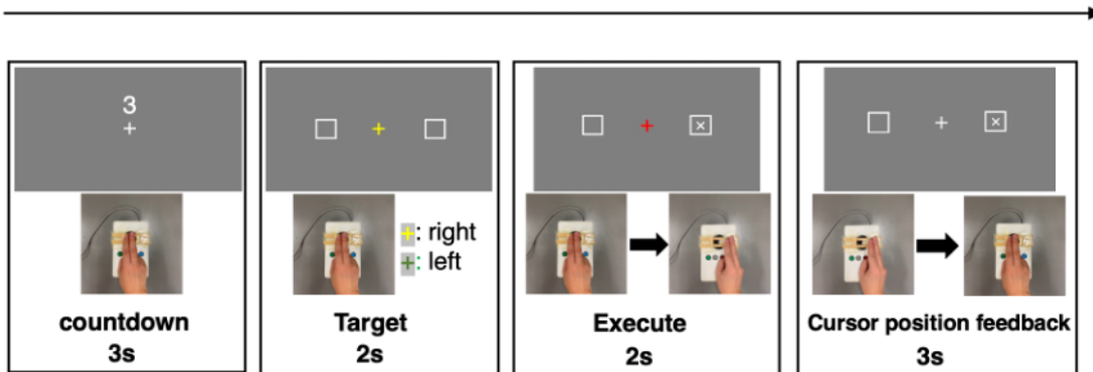
1

2 MNI, Montreal Neurological Institute; L, left hemisphere; R, right hemisphere.

3

4

5

1 **Figures**

2

3 **Figure 1:** Schematic depiction of the time course of a single trial in the practice session.

4 Each practice session included 10 trials. During the target phase, the order in which the

5 colors appear in the central fixation was random, and the same color would not appear

6 three consecutive times. The cursor is available to participants during the execution

7 phase only during the first practice session. The joystick can only be moved to the left

8 and right directions. Participants are instructed to move the joystick after the central

9 fixation turns red.

10

11

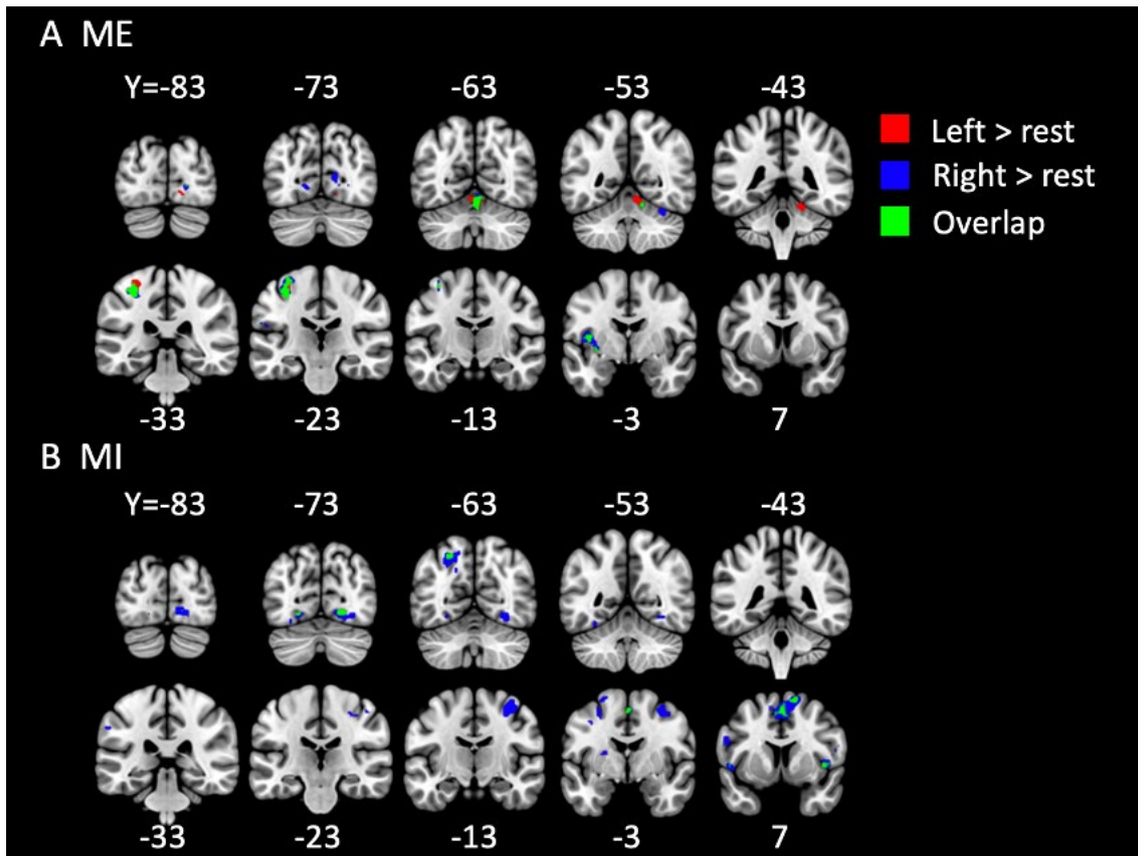


1

2 **Figure 2:** Schematic depiction of the time course of a single trial in ME session and MI  
 3 session. Each session included 20 trials. ME session was almost the same as practice  
 4 session 2, but the countdown phase changed to 9s. Participants were instructed to  
 5 evaluate the quality of the imagery after the picture of the hand appeared on the screen.

6

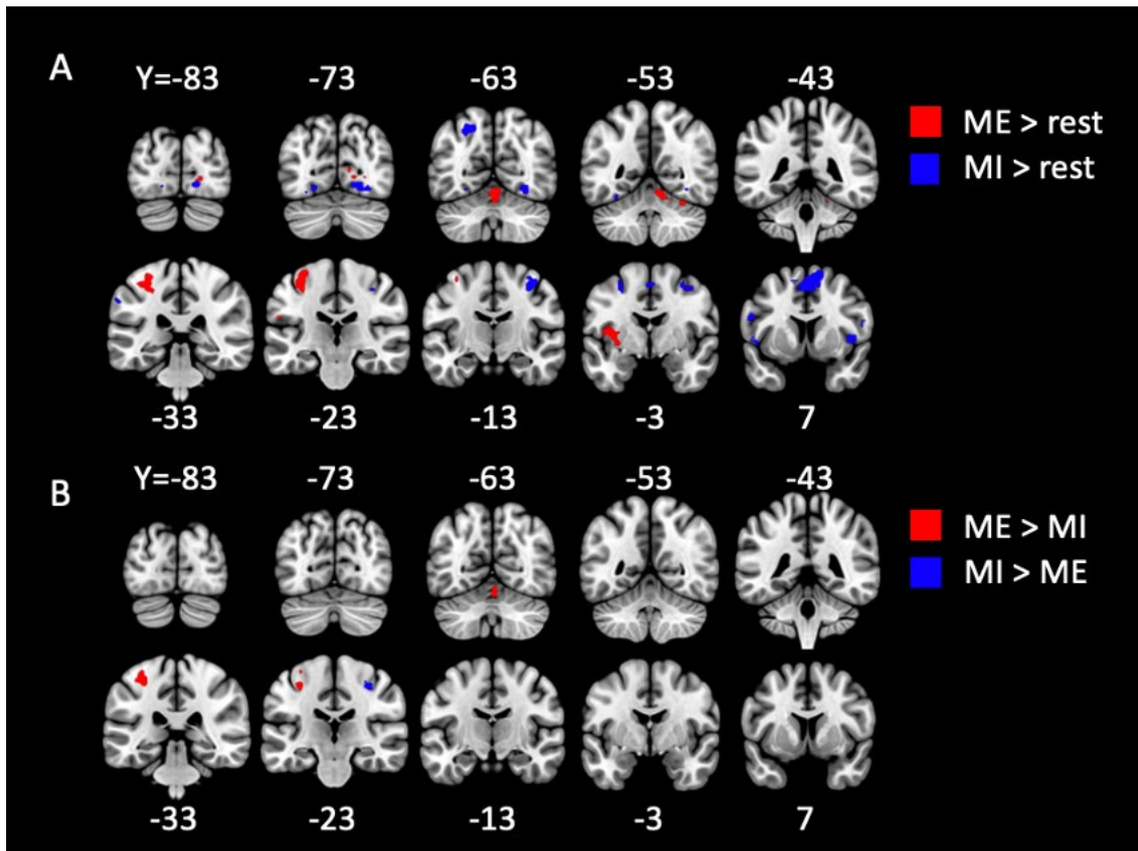
7



1

2 **Figure 3:** Activated regions in the fMRI univariate analysis of ME > rest (A) and MI >  
 3 rest (B). Red showed only the left direction and blue showed only the left. Green  
 4 regions were activated during both left and right directions.

5

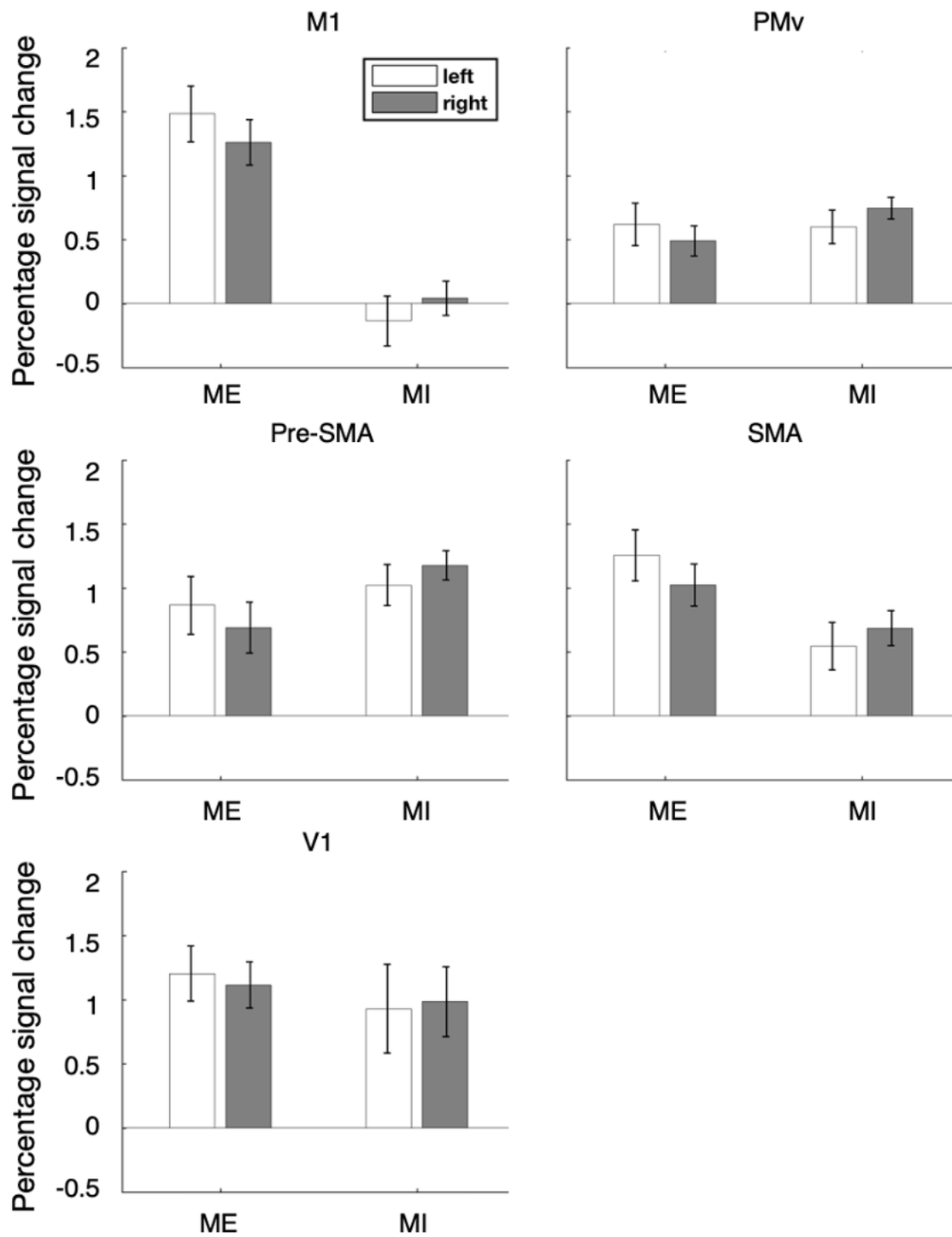


1

2 **Figure 4:** Activated regions in the fMRI univariate analysis of Task vs. Rest (A) and

3 ME sessions vs. MI sessions (B).

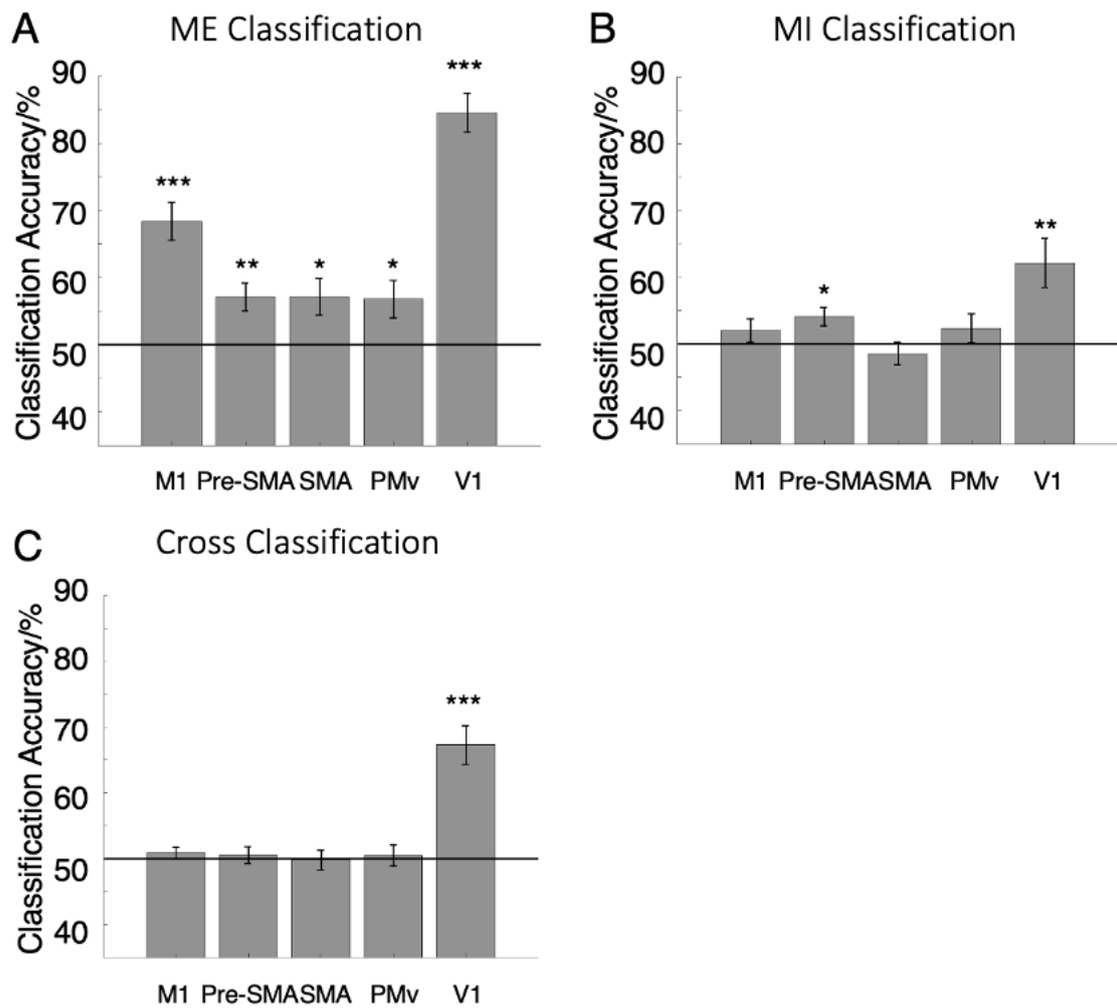
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1

2 **Figure 5:** The averaged activation (beta value) within left M1, PMv, and bilateral pre-  
 3 SMA, SMA, V1; error bars indicate SEMs.

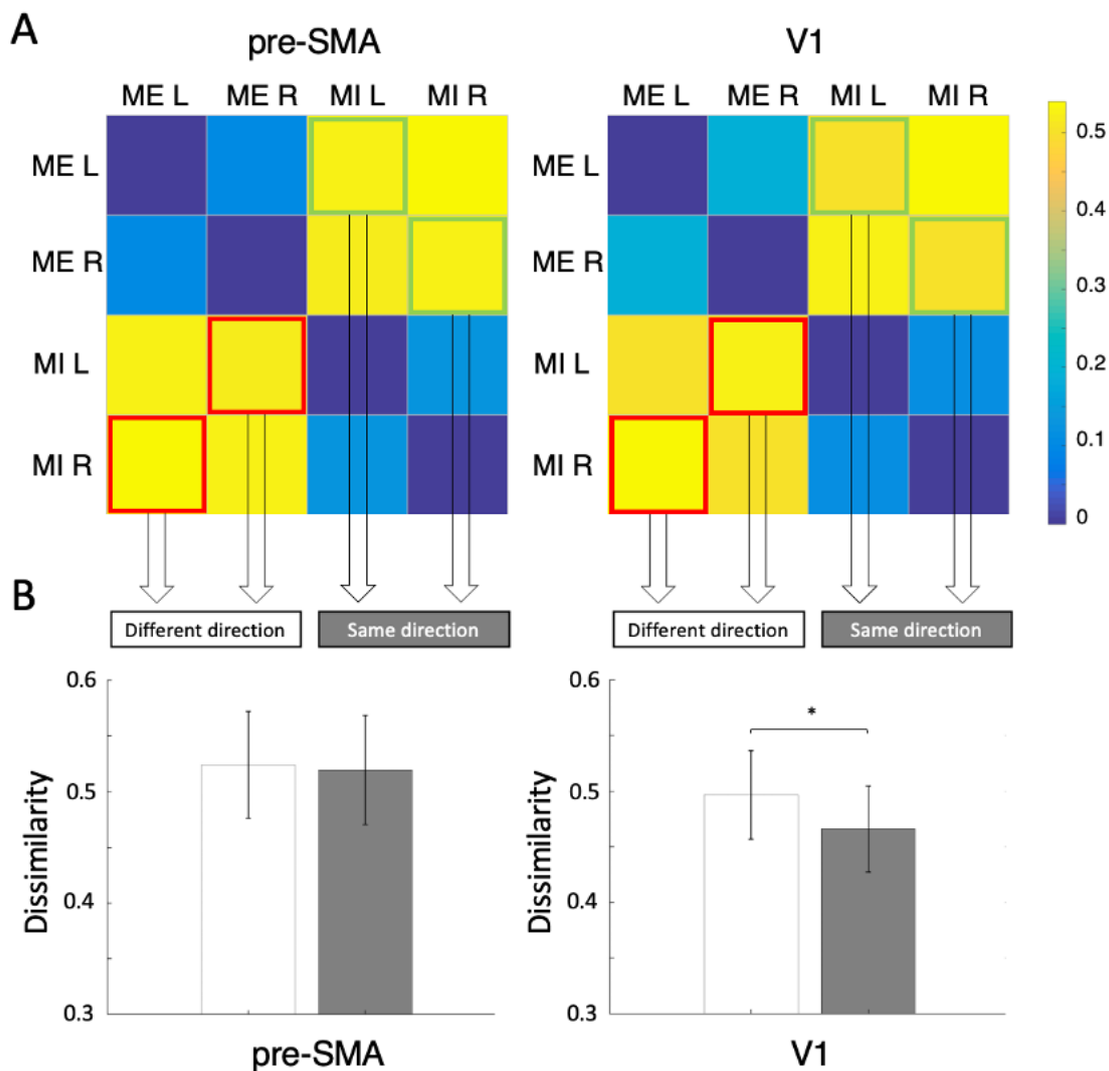
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1

2 **Figure 6:** Classification accuracies with MVPA in ME classification (A), MI  
 3 classification (B), and cross-classification (C) for movement directions in each ROI. (\* $p$   
 4  $< .05$ , \*\* $p < .01$ , and \*\*\* $p < .001$ )

5



1

2 **Figure 7:** Representational similarity analysis (RSA) in bilateral pre-SMA and left V1.

3 Matrix squares show the representational dissimilarity matrix between different

4 directions and modalities, the blue rectangles show the dissimilarity between different

5 modalities but the same direction, and the red rectangles show the dissimilarity between

6 different modalities and different directions (A). The bar plots show the dissimilarities

7 of the same direction and different directions between different modalities; error bars

8 indicate SEMs (B). (\* $p < .05$ , \*\* $p < .01$ , and \*\*\* $p < .001$ ).

9