Does Ipsilateral Remapping Following Hand Loss Impact Motor Control of the Intact Hand?

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What happens once a cortical territory becomes functionally redundant? We studied changes in brain function and behavior for the remaining hand in humans (male and female) with either a missing hand from birth (one-handers) or due to amputation. Previous studies reported that amputees, but not one-handers, show increased ipsilateral activity in the somatosensory territory of the missing hand (i.e., remapping). We used a complex finger task to explore whether this observed remapping in amputees involves recruiting more neural resources to support the intact hand to meet greater motor control demands. Using basic fMRI analysis, we found that only amputees had more ipsilateral activity when motor demand increased; however, this did not match any noticeable improvement in their behavioral task performance. More advanced multivariate fMRI analyses showed that amputees had stronger and more typical representation—relative to controls’ contralateral hand representation—compared with one-handers. This suggests that in amputees, both hand areas work together more collaboratively, potentially reflecting the intact hand’s efference copy. One-handers struggled to learn difficult finger configurations, but this did not translate to differences in univariate or multivariate activity relative to controls. Additional white matter analysis provided conclusive evidence that the structural connectivity between the two hand areas did not vary across groups. Together, our results suggest that enhanced activity in the missing hand territory may not reflect intact hand function. Instead, we suggest that plasticity is more restricted than generally assumed and may depend on the availability of homologous pathways acquired early in life.

Key words: amputees; plasticity; fingers; fMRI; hand; motor control; primary motor cortex; primary somatosensory cortex

Significance Statement

People with congenital hand absence (one-handers) and amputees rely on their intact hand for everyday actions. This extensive daily training could result in increased motor ability, supported by neural resources within the missing-hand territory (i.e., ipsilateral to the intact hand). However, using a demanding multi-finger configuration task, we observed reduced sensorimotor learning in one-handers in the most difficult configuration. Furthermore, despite increased ipsilateral activity, amputees did not show superior intact hand motor ability. Multivariate fMRI analyses suggested a collaborative relationship between the contralateral and ipsilateral hand territories of amputees, which was unique compared with one-handers and controls. These results suggest that brain plasticity is limited and may depend on the availability of homologous pathways acquired early in life.

The authors declare no competing financial interests.

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Introduction
Specific functions of mature cortical areas are determined by their molecular properties, histological organization, and con- nectional fingerprints. The unique identity of a given area is determined by genetic expression and is moderated by electrical activity over the course of early development (see Sur and Rubenstein, 2005 for review). This phase of increased susceptibility to input in shaping the neural circuit is called a critical period (Levelt and Hubener, 2012). The critical period might be enabled because plasticity “brakes,” such as inhibitory circuits, neural over-growth, and synaptic pruning, normally affording homeo- static balance, have not yet been finalized (Takesian and Hensch, 2013). Yet, even in these earliest stages of development, it seems that the assignment of brain function to a given cortical structure is largely fixed. For example, fMRI studies of children with left hemisphere perinatal stroke found that the (typically left dominant) language areas in the inferior frontal cortex were located in anatomically homologous areas in the right hemisphere (Tillema et al., 2008; Raja Beharelle et al., 2010; Tuckute et al., 2022). In this context, it is interesting to consider how a redundant cortical area’s function changes after hand loss, either because of congenital hand malformation (hereafter—one-handers) or acquired arm amputation later in life (hereafter—amputees).

Extensive research has mainly explored neighboring body part homunculus “remapping” following hand loss (see Muret and Makin, 2021 for review), with a focus on the face (Pons et al., 1991; Flor et al., 1995; Raffin et al., 2016). However, from a functional perspective, the intact hand is expected to gain the most from redundant resources in the missing hand territory to adapt to life with only one hand. If this reallocation of resources from the missing hand territory toward the intact hand is functional, and if brain plasticity is more potent in early development, we should find improved motor abilities and learning in one-handers relative to controls. Surprisingly, while the deprived hand area of one-handers was demonstrated to be activated by multiple body parts (e.g., arm, face, feet, torso), it does not appear to be activated by the intact hand (Hahamy et al., 2017; Hahamy and Makin, 2019). Instead, previous research reported increased activity in the missing hand territory from the (ipsilateral) intact hand in amputees (Kew et al., 1994; Hamzei et al., 2001; Bogdanov et al., 2012; Makin et al., 2013a; Philip and Frey, 2014; Valyear et al., 2020). Thus, bimanual experience might be required to establish a functional connection between the two hands to enable ipsilateral functionality (Philip et al., 2015), but direct evidence supporting this is limited. Moreover, previous studies used basic motor tasks (e.g., opening and closing the hand, moving a single digit). If ipsilateral processing due to plasticity processes provides additional resources to aid motor control of the intact hand, it may require difficult tasks to activate it. Similarly, prior results were primarily inferred from net activity changes in the hand territory, under the (potentially naive) assumption that greater activity reveals greater functional involvement. However, as demonstrated in recent learning studies (Berlot et al., 2020), multivariate approaches provide a more detailed opportunity to establish changes in functional brain representation.

Here we aimed to address the relationship between brain and behavioral adaptations for the intact hand in individuals with a missing hand. To better gauge whether activity changes in one-handers and amputees are functional, we varied task difficulty. Participants had to press three fingers simultaneously on a piano-like keyboard, while keeping the other two fingers relaxed. Some combination of fingers are known to be more difficult than others (Waters-Metenier et al., 2014), and we therefore selected two sets of five configurations that ranged from easy to difficult. Motor difficulty is known to increase activity level, particularly in the ipsilateral hemisphere (Verstynen et al., 2005). Therefore, we used fMRI scans while participants performed the same task to compare the net activity (remapping) and multivariate representational similarity (information content and representational structure) of the missing and intact hand territories. While we provide results from both primary somatosensory and motor territories (S1 and M1), we focus our research on S1, which has been the historical primary culprit for remapping following amputation-related deprivation (Merzenich et al., 1984; Pons et al., 1991). S1 is also known to contain greater information content relating to finger movements (Ejaz et al., 2015; Berlot et al., 2019; Arbuckle et al., 2022) and plays a crucial role for dexterous hand control (Brochier et al., 1999) and motor planning (Ariani et al., 2022). To explore the structural underpinnings of these functional changes, we also used diffusion MRI to examine potential white matter microstructural changes in transcallosal fiber connections linking the two hand areas.

We predicted that if one-handers rely on ipsilateral processing for difficult tasks, we should see increased activity and information content in the missing hand cortex compared with controls, leading to improved performance compared with controls. Alternatively, if the functional availability of homologous resources depends on bimanual experience, we should expect to find greater activity and information content in the missing hand territory of amputees relative to controls. Moreover, this information should be organized in a homologous representa- tional structure relative to the intact hand territory.

Materials and Methods
The experimental procedures described in this manuscript were run as part of a larger study (the full study protocol can be found at https://osf.io/gmvua/). Here we focus on procedures related to the finger coordination task. The motor task was similar to previous studies (Waters-Metenier et al., 2014; Ejaz et al., 2015). Participants took part in the training session outside the scanner first and then an fMRI session.

Participants
Amputees (N = 19; 4 females; mean age = 49.05 ± 12.05), one-handers (N = 16; 9 females; mean age = 43.44 ± 11.40), and two-handed controls (N = 16; 7 females; mean age = 45.37 ± 10.67) were invited to take part in a motor control task. The current experiment was composed of a training session outside the scanner, followed by a scanning session. Not all participants were able to take part in (or complete) the scanning session. Thus, the final sample in the scanning session was N = 16 amputees (4 females; mean age = 48.40 ± 12.6), N = 13 one-handers (7 females; mean age = 45.80 ± 11.30), and N = 14 two-handed controls (5 females; mean age = 44.20 ± 12.20). Furthermore, due to technical reasons, we could not register the responses of two amputee participants in the scanner; therefore, the analyses of the finger coordination task during the MRI session were based on N = 14 amputees (2 females; mean age = 49.5 ± 13.1). For the DTI, we were able to collect data for N = 18 amputees, N = 13 one-handers, and N = 13 controls. Three of the amputee participants lost their dominant hand. However, considering they were amputated for at least 26 years, their nondominant hand had effectively taken on the role of dominance. Table 1 displays the demographic information of the participants. The mean age was not significantly different between groups (motor training, \( F_{(2, 48)} = 1.10, \ p = 0.341; \ \eta_p = 0.034 \); scanner session, \( F_{(2, 39)} = 1.11, \ p = 0.348; \ \eta_p = 0.054 \).
Nevertheless, to take into account any potential inter-individual impact of age, we included participants’ age as a covariate in the analyses. In all analyses, outliers were defined as values exceeding the metrics of interest of three standard deviations from the mean. We made a predetermined decision to retain all data points, including those considered outliers, as long as their removal would not have caused any qualitative changes to the results. For the behavioral data during the training session, we identified one potential outlier, but since this outlier did not impact the results qualitatively, we opted to include the outlier in the final analysis. For the multivariate data, we identified an outlier that we decided to remove because its removal qualitatively changed the significance level.

Recruitment was carried out in accordance with the University of Oxford’s Medical Sciences inter-divisional research ethics committee (MS-IDREC-C2-2015-012). Informed consent and consent to publish was obtained in accordance with ethical standards set out by the Declaration of Helsinki. All participants were compatible with local magnetic resonance imaging (MRI) safety guidelines.

### Apparatus

Responses were recorded using a custom-built five-finger MRI-compatible piano-like device (Wiestler and Diedrichsen, 2013; Ejaz et al., 2015; Wesselink et al., 2019). Each key was equipped with a sensor that could continuously measure isometric force during a finger press. The sensors

### Table 1. All participants

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were connected to a laptop and the applied forces were monitored online. Participants received real-time visual feedback on how much force each finger exerted by means of moving horizontal white cursors corresponding to each key. In the training task outside the scanner, the apparatus was placed on a desk in front of the seated participant, who rested the five fingers of their intact hand (or dominant hand in controls) on the keys that were immobile but able to measure the applied pressure. Participants could choose to keep their fingers extended or flexed, based on comfort. Inside the scanner, the device was placed on their lap or belly, depending on their preference. Ensuring participants’ comfort was paramount in this experiment because we wanted them to be able to control their fingers during a complex task (i.e., the finger configuration task). This is standard practice both inside and outside the scanner.

General procedure

Instructions. The top of the screen showed five vertical gray bars, each corresponding to one of the keys. At rest, participants were required to apply and maintain a minimal force (0.5 N) on the keys, as indicated by a horizontal bar at the bottom of the screen (hereafter baseline area). In a typical trial, three of the vertical bars turned green indicating which of the keys to press. Participants were instructed to wait until the appearance of a go cue that was provided as a green horizontal bar similar in dimension and right above the baseline area. At this point, participants had to use the sensory information provided by all fingers which is fundamental in dexterous manipulation (Pruszynski et al., 2016). Participants received a positive feedback (i.e., a point) as soon as they configured the instructed fingers as required. Once the finger cursors were successfully stabilized in the target area, the area disappeared indicating the participants to go back to the baseline position (this was a requirement to obtaining a point in the next trial) by releasing the pressure on the instructed fingers. At this point, a new trial started. Note that the training session was self-paced, whereas the scanning session was timed (see below for details).

Training session. First, the experimenter explained the task and showed the participants how to use the device. Then, the participants performed a few familiarization trials with a set of configurations not used in the study. This was followed by a single-finger movement block, where the participants had to press only one of the five fingers. This block was repeated one more time at the end of the training (as detailed in Wesselinik et al., 2019). Then, the actual training session started, and it lasted 25 min. Within this time window, participants were encouraged to complete as many blocks as possible. Each block was about 3 min long, depending on the performance, leading to a variable number of blocks across participants. On average, participants completed Md = 5.69 blocks (SDd = 1.42; MAmputees = 5.58; SDAmputees = 1.54; MHb = 5.62; SDhb = 1.36; MC = 5.87, SDC = 1.41), and the three groups did not differ for the number of blocks completed (F(2, 48) = 0.20; p = 0.846; N = 48). Visual aspects of the required chord were presented for 3 s, followed by the go cue. Within each block, instructions for the same finger configuration were repeated four times.

fMRI session. After the training session, participants were invited to take part in a similar motor task as part of the fMRI study. In the scanner, there was no minimal pressure requirement at baseline because the applying the same force (2.5 N) was tiring while lying supine. In addition, the task was timed. Visual instructions were presented for 1.3 s, and participants had to execute the chord (i.e., press and release the keys) within 2.3 s from the onset of the instructions. The same instruction was repeated three times resulting in blocks of 6.9 s. Each finger configuration block was repeated three times within a run, resulting in 45 trials per run (9 trials by 5 finger configurations). Participant took part in four runs and each run lasted around 3.5 min (141 volumes).

Behavioral performance. Behavioral performance was measured as the deviation from the required finger configuration, by taking into account two sources of error: (1) any deviation of the noninstructed fingers from the baseline (0.5 N) and (2) any deviation of the each instructed fingers from the average force as all the instructed fingers were expected to exert a similar force (2.5 N). These two forms of residuals were computed within the response and release time, summed up and averaged across time to obtain a unique measure of performance per trial (see Waters-Metenier et al., 2014). In line with previous studies (Eizaj et al., 2015; Waters et al., 2017), the beginning of the response was defined as the point in time in which at least one of the fingers exceeded the threshold of 1.5 N when pressed.

Finger configuration and difficulty levels

Figure 1 displays the configurations used in the training session (A, easy to difficult from the bottom to the top) and in the scanner (D, easy to difficult from the bottom to the top). In the training session, we used the following finger configurations (1, thumb; 2, index; 3, middle; 4, ring; 5, little finger): 345, 123, 124, 245, and 135. The aim of the training session was twofold: (1) measure the sensorimotor learning of participants and (2) familiarize participants with the task before entering the scanner. In the scanner, we used different finger configurations (145, 234, 134, 125, and 235) in order to minimize any differences across groups that were hypothesized to arise due to different training capacity.

To independently confirm that the previously estimated difficulty levels of finger configurations, defined from a pilot session of a previous study (Waters-Metenier et al., 2014), were appropriately labeled, we also utilized a model-based approach. To this aim, we used the amount of enslavement (percentage of maximal voluntary contraction – MVC) between fingers in a single-finger task (Yu et al., 2010). In particular, considering the task’s complexity that involves simultaneous control of multiple digits, we reasoned that difficulty is influenced by at least three components: easy configurations would be characterized by high amount of enslavement between instructed fingers (component 1) and noninstructed fingers (component 2) and low amount of enslavement between the instructed and noninstructed fingers (component 3). In other words, it is easier to move in parallel fingers with high amount of enslavement, as it is to keep relaxed fingers with high amount of enslavement. Furthermore, it is easier to control a finger configuration where the instructed and noninstructed fingers have low amount of enslavement. More specifically, for each chord, we estimated the three components of enslavement as follows: the total (i.e., sum) of enslavement for the instructed fingers (E1), the enslavement for the noninstructed fingers (E2), and the enslavement between the instructed and noninstructed fingers (E3). Then, we combined the three components (i.e., E1 + E2 – E3) to obtain a unique measure of enslavement such that the configurations with a high score were categorized as easier than the ones with a low score. Using this measurement, we sorted the configurations from easy to difficult and divided them into two groups: easy (345, 145, 234, 123, and 124) and difficult (134, 125, 245, 235, and 135). In our analysis, for the training session, the easy averaged configurations were 345, 123, and 124, and the difficult averaged configurations were 245 and 135; for the scanning session, the easy averaged configurations were 145 and 234, and the difficult averaged configurations were 134, 125, and 235. We also used this scoring to establish the easiest and most difficult configurations for specific fMRI analysis.

MRI data acquisition

MRI images were acquired using a 3 T MAGNETON Prisma MRI scanner (Siemens) with a 32-channel head coil. Functional images were collected using a multiband T2*-weighted pulse sequence with a between-slice acceleration factor of 4 and no in-slice acceleration (2 mm isotropic; TR, 1,500 ms), covering the entire brain. The following acquisition parameters were used: TE, 32.40 ms, and flip angle, 75°, 72 transversal slices. Field maps were acquired for field unwarping. A T1-weighted sequence was used to acquire an anatomical image (TR, 1,900 ms; TE, 3.97 ms; flip angle, 8°; spatial resolution, 1 mm isotropic). Diffusion-weighted fMRI data were acquired using 10 directions: 1,900 ms; TE, 2,951 ms; TE, 79.80 ms; flip angle, 80°; spatial resolution, 1.5 mm isotropic; and 84 transversal slices. Gradients were applied along 60 uniformly distributed directions with a b value of 1,000 s/mm².
non-diffusion-weighted images with $b = 0$ s/mm$^2$ were also acquired. No task was given to the participants during the structural and DTI acquisition. They viewed a calm nature video to prevent them from falling asleep and making large head movements.

fMRI preprocessing and first-level analysis

MRI data were preprocessed using a standard pipeline as implemented in FSL 6.0 (Smith et al., 2004; Jenkinson et al., 2012). The following steps were applied to each functional run: motion correction using MCFLIRT (Jenkinson et al., 2002); B0 fieldmap correction to account from distortions due to magnetic field inhomogeneity; brain extraction using BET (Jenkinson et al., 2002); high-pass temporal filtering of 90 s; and spatial smoothing with a Gaussian kernel of full-width at half-maximum of 5 mm for the univariate analyses and 3 mm for the multivariate analyses.

In order to estimate brain activity related to our configuration task, we employed a voxel-based general linear model (GLM) as implemented in FEAT. For each functional run, time series were predicted using five regressors of interest corresponding to the five configurations that participants had to do in the scanner. These regressors were convolved with a double-gamma function, and their temporal regressors were also added to the design matrix to account for temporal variability of the BOLD response. We also included the motion parameters resulting from the MCFLIRT step and columns indicating outlier volumes as returned from the FSL function *fsl_motion_outliers* with default and recommended parameters (root mean square intensity difference of each volume to the reference volume as metric; as a threshold, metrics that were larger than 75th percentile + 1.5*interquartile rage were considered outliers). The number of volume outliers was small for all groups (amplitude to the reference volume as metric; as a threshold, metrics that were larger than 75th percentile + 1.5*interquartile rage were considered outliers). The cross-validated Mahalanobis (or cross-validated Mahalanobis) distance was used to increase reliability of distance estimates (noisy voxels are down-weighted), based on the voxel’s covariance matrix calculated from the GLM residuals. The advantage of using the cross-validated Mahalanobis distance is twofold: (1) spatially correlated noise is removed using multivariate noise normalization and this improves the estimate of the dissimilarities (Walther et al., 2016); (2) cross-validation ensures that if two patterns only differ by noise, their mean dissimilarity estimate will be zero. As a consequence, the dissimilarity between two patterns can also be negative (Diedrichsen et al., 2016) and thus dissimilarities significantly larger than zero can be taken as evidence that the two patterns are distinct and that the ROI contain task-related information (e.g., distinct measurement of configurations). The cross-validated dissimilarity was computed using the python library for RSA *rsatoolbox* version 0.0.4 (https://github.com/rsagroup/ rsatoolbox).

Diffusion MRI preprocessing

Diffusion data were preprocessed using a custom pipeline that combined tools from MRtrix 3.0 (Tourin et al., 2019), ExploreDTI 4.8.6 (Leemans et al., 2009), and FSL 5.0.9 (Smith et al., 2004; Jenkinson et al., 2012). These included the following: (1) denoising using the MP-PCA (principal component analysis of Marchenko–Pastur) method in MRtrix (Veraart et al., 2016); (2) Gibbs ringing correction using “mrdgibbs” in MRtrix (partial Fourier; Kellner et al., 2016); (3) global signal drift correction using ExploreDTI (Vos et al., 2017); and (4) motion EPI distortion correction using Eddy and Topup within FSL (Andersson et al., 2003). Data were visually checked as part of quality assurance procedures. Whole-brain voxel-wise maps of fractional anisotropy (FA) and mean diffusivity (MD) maps were then derived from the preprocessed data by fitting the diffusion tensor model. FA represents the degree to which diffusion is constrained in a particular direction and ranges from 0 (isotropic diffusion) to 1 (anisotropic diffusion). MD (10−3 mm2 s−1) represents the average diffusivity rate. The diffusion tensor was estimated and fitted using the nonlinear least squares method with Robust Estimation of Tensors by Outlier Rejection (RESTORE) applied (Chang et al., 2005).

Tractography

A multiple-ROI tractography approach enabled specific transcallosal pathways to be constructed in each participant between their left and right S1 hand areas (see also Postans et al., 2020). Initially, each participant’s ROIs in T1 space (see ROI definition above) were registered to their native space diffusion MRI image using the following steps: (1) the T1-to-diffusion transformation matrix was generated using FLIRT with 6 degrees of freedom and the correlation ratio cost function. The fractional anisotropy (FA) map was used as the reference image (rather than the b0 image) as it provided better image contrast; (2) the transformation matrix was then applied to the individual subject ROIs in T1 space using FLIRT. As tractography can be challenging from gray matter ROIs (due to low anisotropy), the diffusion space ROIs were dilated by 1.5 mm to include some white matter voxels (Gschwind et al., 2012).
Tractography was initially performed from all voxels in the left hemisphere ROI in each participant’s native diffusion MRI space in ExploreDTI (v4.8.3; Lemans et al. 2009) using a deterministic tractography algorithm based on constrained spherical deconvolution (Tournier et al., 2008; Jeurissen et al., 2011). Spherical deconvolution approaches enable multiple peaks to be extracted in the fiber orientation density function within a given voxel, allowing complex fiber arrangements, such as crossing/kissing fibers, to be modeled more accurately (Dell’Acqua and Tournier, 2019). The contralateral S1 ROI was then used as an “AND” gate to capture any streamlines that arose from the seed ROI and terminated in the contralateral ROI. Next, the same procedure was repeated, this time starting with the right hemisphere ROI as seed and gating with the right hemisphere. This process was conducted for each participant and then inspected visually by the research team (C.J.H., R.T.). A step size of 0.1 mm and an angle threshold of 60° were applied to prevent the reconstruction of anatomically implausible streamlines. Tracking was performed with a supersampling factor of 4×4×4 (i.e., streamlines were initiated from 64 grid points, uniformly distributed within each voxel). The resulting interhemispheric pathways were then intersected with the whole-brain voxel-wise FA and MD maps (see above) to derive four tract-specific measures of microstructure in each participant (S1-to-S1 and M1-to-M1, in both directions). As in Postans et al. (2020), the FA and MD values for the left-to-right and right-to-left segments were combined into a streamline-weighted mean using the following equation:

\[
\text{Vertex-Weighted Mean FA} = \frac{(N_{L-R} \times \text{FA}_{L-R}) + (N_{R-L} \times \text{FA}_{R-L})}{(N_{L-R} + N_{R-L})}
\]

\[
\text{Vertex-Weighted Mean MD} = \frac{(N_{L-R} \times \text{MD}_{L-R}) + (N_{R-L} \times \text{MD}_{R-L})}{(N_{L-R} + N_{R-L})}
\]

**Tract-based spatial statistics**

We also conducted a complementary voxel-wise statistical analysis of the FA and MD data using tract-based spatial statistics (TBSS; Smith et al., 2006). First, each participant’s FA and MD maps were aligned to the standard MNI template using nonlinear registration (Andersen et al., 2010). Second, the mean FA image was created and subsequently thinned (using the default FA threshold = 0.2) to generate the mean FA skeleton, which represents the center of all tracts common to the group. Third, each participant’s FA and MD data were projected onto the skeleton for voxel-wise analyses using tract-based spatial statistics (TBSS; Smith et al., 2006). First, each participant’s FA and MD maps were aligned to the skeleton for voxel-wise analyses using randomization in FSL (Winkler et al., 2014). For both analyses, correlation values were standardized using the Fisher’s r-to-z transformation. Independent t tests were used to test for group differences. The experimental groups (amputees and one-handers, with the contralateral RDM averaged across the control participants), we used two-tailed one-sample t test for each group and hemisphere. We also used a two-way rMANCOVA with age (de-meaned) as a covariate and group as a between-subject factor, and hemisphere as a within-subject factor. To test for existing functional homotopy (i.e., correlation between finger configuration pairs across hemispheres), we used two-tailed one-sample t test for each group and hemisphere. We also used a one-way rANCOVA with age (de-meaned) as a covariate and group as a between-subject factor to investigate differences in functional homotopy between groups. To investigate similarity to typical contralateral representation in the experimental groups (i.e., correlation between the representation dissimilarity matrices (RDMs) of the experimental participants, amputees, and one-handers, with the contralateral RDM averaged across the control participants), we used two-tailed one-sample t test for each group and hemisphere. We also used a two-way rMANCOVA with age (de-meaned) as a covariate, group (one-handers, amputees) as a between-subject factor, and hemisphere as a within-subject factor to investigate differences in typical contralateral representation between the experimental groups. Prior to these analyses, correlation values were standardized using the Fisher’s r-to-z transformation. Independent t tests were used to test for group differences. The experimental groups (amputees and one-handers) were compared against the control group unless differently specified. To control for age while performing an independent t test, we first ran an ANCOVA and then computed the contrasts of interest using the R package emmeans 1.8.2. For post hoc comparisons that were exploratory (i.e., not a priori and not confirmatory), we adjusted our significance α-level for multiple comparisons using Bonferroni’s approach. In the Results section, we report the uncorrected p values with a note of the adjusted α-level. For nonsignificant results of interest, we reported the corresponding Bayes factor (BF10), defined as the relative support for the alternative hypothesis. While it is generally agreed that it is difficult to establish a cutoff for what consists sufficient evidence, we used the threshold of BF<1/3 as sufficient evidence in support of the null, consistent with others in the field (Wetzels et al., 2011; Dienes, 2014). For Bayesian ANCOVAs, we used a uniform model as a prior, and for Bayesian t tests, we used the Cauchy model with a width of 0.707, which are the default settings in JASP. For all analyses, whenever the normality assumptions were not met, we adopted a permutation approach using the function aovperm of the R package permuco 1.1.2 with default settings (permutation method for fixed effects models, freedman_lanc for mixed effects models, Rd_kheradPajouh_renaud), and we report these results with a note only when they are qualitatively different from the parametric approach.

**Data code and accessibility**

The preprocessed data and the scripts necessary to reproduce the analyses can be found at https://osf.io/hsvkc/.

**Results**

**One-handers show reduced benefits from brief training of difficult finger configurations**

We first explored whether individuals with a missing hand, either due to congenital malformedness (one-handers) or amputation in adulthood (amputees), differ from controls in their ability to learn to perform a finger configuration task with varying levels of difficulty. Mean deviations from the instructed hand configuration for each of the five configurations across the first seven blocks are shown for the three groups in Figure 1B, with more difficult configurations displayed in cooler colors. At the first attempt (block 1), there was no difference in performance between the experimental and control groups, except for a trend...
for the most difficult level, in which one-handers showed worse performance compared with the control group (10 comparisons, no corrections for multiple comparisons). To quantify training effects across groups, we averaged deviation means between easy (configurations 1–2) and difficult levels (configurations 3–5) for each participant and compared performance between the first and last blocks completed during training (Fig. 1C). The resulting 3 (group) × 2 (block) × 2 (difficulty) ANCOVA (controlling for age) resulted in a significant three-way interaction ($F_{(2, 47)} = 5.28; \ p = 0.009; \ \eta_p = 0.18$), indicating that participants across the three groups benefited differently from the practice, with respect to difficulty levels. In addition, a main effect of difficulty ($F_{(1, 47)} = 137.22; \ p \leq 0.001; \ \eta_p = 0.74$) and block number ($F_{(1, 47)} = 17.48; \ p \leq 0.001; \ \eta_p = 0.27$) was found, with no significant main effect of group ($F_{(2, 47)} = 1.31; \ p = 0.280; \ \eta_p = 0.05$). As apparent from the figures, this was driven by a lack of learning effect in the one-handed group, specifically for the difficult configurations. To better quantify this, we ran a separate repeated-measures ANCOVA for each group and observed a significant interaction between block number and difficulty for the one-handers only ($F_{(1, 14)} = 12.61; \ p = 0.003; \ \eta_p = 0.47$). To further explore the differential learning effect observed in the one-handed group, we compared differences in performance between the last and the first block (Fig. 1C) and found a significant learning effect in the easy condition only (easy, $t_{(14)} = -3.58; \ p = 0.003$; difficult, $t_{(15)} = 0.14; \ p = 0.889; \ BF_{10} = 0.258; \text{Bonferroni’s adjusted } a, \ 0.05/2 = 0.025$), suggesting that the impairment in learning was specific for the difficult configurations. This was also confirmed by significant differences in the last block of training between one-handers and controls for the difficult configurations only ($t_{(47)} = 2.32; \ p = 0.024$).

We next examined whether these group differences in performance were replicated in the fMRI task, where five different configurations were used Figure 1D. Figure 1E shows performance across the four runs. To test for differential learning effects, we repeated the analysis mentioned above while comparing performance across groups and difficulty levels between the first and the last runs (Fig. 1F). The three-way interaction in the ANCOVA was not significant ($F_{(2, 38)} = 0.48; \ p = 0.622; \ \eta_p = 0.02$), indicating that the groups did not show different learning effects —indeed as shown in the figure, performance had already plateaued. However, we did observe a significant interaction between group and difficulty ($F_{(2, 38)} = 3.93; \ p = 0.028; \ \eta_p = 0.17$), indicating that participants in different groups responded differently to task difficulty. We also observed a trend toward a main effect of group ($F_{(2, 38)} = 2.71; \ p = 0.080; \ \eta_p = 0.12$), in addition to a main effect of difficulty ($F_{(1, 38)} = 73.61; \ p < 0.001; \ \eta_p = 0.66$) and block number ($F_{(1, 38)} = 9.89; \ p = 0.003; \ \eta_p = 0.21$). The interaction between group and difficulty only showed a trending result and was driven by the one-handers performing worse on the difficult configurations relative to controls ($t_{(38)} = 2.38; \ p = 0.022$; Bonferroni’s adjusted $a, \ 0.05/3 = 0.0167$). This is reflective of the behavioral results found outside the scanner, where the one-handers showed worst performance on the difficult configurations at the end of the training session. Here, we also found a trending result suggesting performance deficits in the amputee group relative to controls in the
difficult configurations ($t_{(38)} = 2.42; p = 0.021$; Bonferroni’s adjusted $\alpha, 0.05/3 = 0.0167$). However, the one-hander and amputees groups did not differ relative to each other in performance ($t_{(38)} = 0.06; p = 0.955$; Bonferroni’s adjusted $\alpha, 0.05/3 = 0.0167$). It is important to note that previous tests comparing the two experimental groups against the control group only showed a trend (i.e., did not survive the multiple comparisons correction as Bonferroni’s corrected $p$ values were both below 0.067), and as such, these findings should be interpreted with caution.

**Amputees show increased averaged ipsilateral activity that scales with difficulty**

Next, we examined univariate activity levels across the bilateral S1 hand ROIs (Fig. 2A). To estimate whether difficulty modulated brain activity differently for the different groups and hemispheres, we first conducted a three-level ANCOVA, including 3 (group) x 2 (hemisphere) x 2 (difficulty) and age (as a covariate). We observed a significant three-way interaction ($F_{(2, 39)} = 5.23; p = 0.010; \eta_p = 0.21$), confirming that difficulty modulates activity differently across hemispheres and groups, as shown in Figure 2B. The analysis also revealed a significant interaction between group and hemisphere ($F_{(2, 39)} = 6.46; p = 0.004; \eta_p = 0.25$), difficulty and hemisphere ($F_{(1, 39)} = 6.16; p = 0.018; \eta_p = 0.14$), and main effects of hemisphere ($F_{(1, 39)} = 71.02; p < 0.001; \eta_p = 0.65$) and difficulty ($F_{(1, 39)} = 11.33; p = 0.002; \eta_p = 0.23$). To further explore the three-way interaction, we conducted two separate two-level ANCOVAs for each hemisphere. As hypothesized, we observed group differences within the ipsilateral cortex only, where we found a significant interaction between group and difficulty ($F_{(2, 39)} = 3.39; p = 0.044; \eta_p = 0.15$) and a main effect of group ($F_{(2, 39)} = 5.95; p = 0.006; \eta_p = 0.23$), while no main effect or interaction involving group was observed in the contralateral hemisphere (all $p > 0.6$). This suggests that activity scales with task difficulty differently across groups in the ipsilateral cortex (which is the missing hand cortex in the experimental groups). The main effect of difficulty was significant in both hemispheres (contralateral, $F_{(1, 39)} = 15.68; p < 0.001; \eta_p = 0.287$; ipsilateral, $F_{(1, 39)} = 5.57; p = 0.023; \eta_p = 0.125$). The ipsilateral interaction between group and difficulty was driven by an increase of activity with difficulty in the amputees ($t_{(39)} = 3.55; p < 0.001$) but not in one-handers or controls ($t_{(39)} = -0.18; p = 0.857; t_{(39)} = 0.90; p = 0.373$). Furthermore, amputees showed significantly larger activity than controls in the ipsilateral cortex for both the difficult ($t_{(39)} = 3.37; p = 0.002$) and the easy conditions ($t_{(39)} = 3.22; p = 0.003$). Together, these findings confirm and extend previous studies—ipsilateral activity for the intact hand was heightened in amputees, particularly with increased task difficulty. Conversely, the one-handed group did not show any significant benefit or disadvantage in activating the missing hand cortex relative to controls.

We repeated the same analysis in bilateral M1 hand ROI (Fig. 4B). The three-way interaction was not significant in this case ($F_{(2, 39)} = 1.50; p = 0.236; \eta_p = 0.07$). We observed a significant interaction between difficulty and hemisphere ($F_{(1, 39)} = 11.48; p = 0.002; \eta_p = 0.23$), driven by activity increase with difficulty in the contralateral hemisphere only (contralateral, $t_{(39)} = 4.07; p = 0.0002$; ipsilateral, $t_{(39)} = 2.02; p = 0.05$; Bonferroni’s adjusted $\alpha, 0.05/2 = 0.025$). We also observed a significant interaction between group and hemisphere ($F_{(2, 39)} = 4.83; p = 0.013; \eta_p = 0.20$), due to the fact that the difference in activity between the two hemispheres was reduced in the amputees relative to the control groups ($t_{(39)} = -3.05; p = 0.004$; Bonferroni’s adjusted $\alpha, 0.05/3 = 0.0167$).

**Figure 2.** BA3b ROI. Amputees showed significantly larger activity than the control and one-hander groups in the ipsilateral cortex for the difficulty condition (B) but this did not result in an increased information content (C). A, Bilateral hand BA3b ROIs used in the analyses (one example participant). B, Brain activity (z scores) in the contralateral and ipsilateral hemispheres averaged across runs and across easy and difficult configurations. C, Information content (dissimilarities between configuration pairs) in the contralateral and ipsilateral hemispheres averaged across runs. Only the dissimilarities between the easiest (e.g., the green square in Fig. 3A) and most difficult (e.g., the blue square in Fig. 3A) finger configurations were selected. The unshaded dots with different colors represent individual participants. Color filled dots with asterisks at the top of plots B and C indicate significant difference (Bonferroni’s corrected) between the groups specified by the colors in a specific or averaged condition. Lines with asterisks refer to significant difference (Bonferroni’s corrected) between conditions within a group. We are reporting here only the relevant comparisons, for the complete analysis, please refer to the Results section.
This is in line with the observation that the amputees showed higher activity in the ipsilateral hemisphere than the control group. We did not find main effects or interaction between group and difficulty (all $p > 0.3$). Overall, these results suggest that contrary to ipsilateral S1, ipsilateral M1 does not scale with difficulty in the amputees. Finally, to confirm that our effects reflect increased difficulty relating to motor performance per se, rather than more general task demands, for example, relating to attentional or arousal effects, we repeated the same analysis in a control visual region (left and right hMT+; Fig. 4E) and observed no significant main effects or interactions (all $p > 0.11$).

**Amputees show bilateral increase in information content relative to one-handers**

We next assessed whether the selective increase in unilateral activity observed in amputees, previously interpreted as functional remapping, translated to a gain in information content. Average distances (across all configuration pairs as shown in Fig. 3A) were significantly greater than zero (all $p$s < 0.05, not corrected for multiple comparisons), confirming that task-relevant information was encoded in both hemispheres. We first examined distances by specifically comparing the easy and difficult configurations separately across hemispheres and groups. To allow us to specifically account for difficulty, this analysis was restricted to the easiest configuration pair (C234–C145) and the most difficult configuration pair (C235–C125) in our representational dissimilarity matrix (highlighted in the contralateral RDM of the Amputees in Fig. 3A, green, easiest; blue, most difficult). If increase in activity translates to information content gain, we should see larger distances between configuration pairs for the amputees, especially across the most difficult conditions. However, we did not find a significant three-way interaction ($F_{(2, 38)} = 0.41; p = 0.668; \eta_p = 0.15$) or a resulting two-way interaction involving group (Fig. 2C). Instead, we found a main effect of group ($F_{(2, 38)} = 3.29; p = 0.048; \eta_p = 0.15$) driven by increased information content in amputees relative to one-handers ($t_{(38)} = 2.55; p = 0.015$; Bonferroni’s adjusted $\alpha = 0.05/3 = 0.0167$). We also observed a main effect of hemisphere ($F_{(1, 38)} = 15.57; p \leq 0.001; \eta_p = 0.29$) and difficulty ($F_{(1, 38)} = 11.91; p = 0.001; \eta_p = 0.24$). Interestingly, we found that information scales down with difficulty, regardless of group, suggesting that the overall increase in information observed in amputees is not linked to their reduced performance.

To take best advantage of our information content analysis, we repeated the analysis while comparing the average distances...
across the entire RDM (10 cells) across groups and hemispheres in a two-way ANCOVA. Again, if increase in activity translate to information content gain, we should see larger averaged distance between configuration pairs for the amputees. Here again, we found no significant interaction ($F_{(2,38)} = 0.65; p = 0.525; \eta_p = 0.03$), suggesting that information content was not modulated differently across group and hemisphere. Instead, again, we found a main effect of group ($F_{(2,38)} = 3.83; p = 0.030; \eta_p = 0.17$) and hemisphere ($F_{(1,38)} = 25.84; p < 0.001; \eta_p = 0.40$). The main effect of group was again driven by increased distances across both hemispheres in amputees relative to one-handers ($t_{(38)} = 2.62; p = 0.013$). Similar to the previous analysis, these effects were not specific to the ipsilateral cortex but were instead generalized. Do these group differences reflect increased information in amputees or decreased information in one-handers? When comparing against controls, the results are ambiguous (amputees vs controls, $t_{(38)} = 1.99; p = 0.054; BF_{10} = 0.90$; one-handers vs controls, $t_{(38)} = -0.57; p = 0.573; BF_{10} = 0.46$; Bonferroni’s adjusted $\alpha$, 0.05/2 = 0.025). Together, it appears that the increased activity found in the ipsilateral hemisphere of amputees for the difficult configurations does not neatly translate to a selective increased information content.

To further confirm the specificity of our effects, we repeated the same analyses in M1 and hMT+ and verified that the increased distances were also significantly larger than zero (all $p < 0.007$, not corrected for multiple comparisons). In M1 (Fig. 4C), when focusing on difficulty as a factor, we observed a main effect of difficulty ($F_{(1,38)} = 5.60; p = 0.023; \eta_p = 0.13$), suggesting larger distances between the easiest pairs than the most difficult ones, and hemisphere ($F_{(1,38)} = 15.83; p < 0.01; \eta_p = 0.29$), suggesting larger distances in the contralateral hemisphere relative to the ipsilateral hemisphere. No main effect of group ($F_{(2,38)} = 2.05; p = 0.143; \eta_p = 0.10$) and no group interactions (group × hemisphere, $F_{(2,38)} = 0.44; p = 0.648; \eta_p = 0.023$; group × difficulty, $F_{(2,38)} = 0.50; p = 0.610; \eta_p = 0.026$) were observed. Similarly, when averaging across all configurations, we observed a main effect of hemisphere ($F_{(1,38)} = 14.79; p < 0.001; \eta_p = 0.28$), suggesting larger distances in the contralateral relative to the ipsilateral hemisphere, no main effect of group ($F_{(2,38)} = 1.94; p = 0.158; \eta_p = 0.09$), and no group interactions ($F_{(2,38)} = 0.18; p = 0.836; \eta_p = 0.000$). Despite higher distances in hMT+ (presumably due to the visual information provided throughout the motor task; Fig. 4F), we did not observe any main effects or interactions (all $p > 0.2$).

**Figure 4.** M1 and hMT+ ROIs. We extended our analyses from BA3b (Fig. 2) to explore (A) M1 and (D) hMT+. In the case of univariate analysis, unlike BA3b, we did not find a three-way interaction in either (B) M1 or (E) hMT+. Specifically, the increase in averaged activity related to task difficulty, which we observed in the ipsilateral hemisphere of the amputees (Fig. 2B), was not evident in these control ROIs. In M1, there was a general increase in averaged activity related to difficulty across all groups, but this effect was seen only in the contralateral hemisphere (B). Additionally, we noted a significant interaction between group and hemisphere in M1, indicating that the difference in activity between the two hemispheres was reduced in amputees compared to the control groups. We did not identify any significant main effects or interactions in hMT+ (E). For the multivariate analysis, in (C) M1, we observed a similar decrease in distance with increasing difficulty across groups and hemispheres, mirroring what we found in BA3b (Fig. 2C). However, in contrast to BA3b, there was no main effect of group in M1. Lastly, in (F) hMT+, our analyses did not reveal any significant main effects or interactions.

**Amputees show increased functional homotopy in representational structure across hemispheres**

Functional homotopy refers to brain regions in opposite hemispheres exhibiting correlated activity patterns during a task or at rest and suggests that two brain regions are functionally associated and working in concert to perform a certain function (e.g., a motor task). We explored the degree of functional homotopy (defined here as the correlation between the representational dissimilarity matrices shown in Fig. 3A) in the hand region across the two hemispheres. We first correlated the 10 configuration pairs of the RDM across the two SI hand areas of each participant. The homotopy correlation values were significantly larger than zero for the amputee and one-hander groups,
but not for the controls (amputees, $t_{(15)} = 2.83; p = 0.013$; one-handers, $t_{(12)} = 3.28; p = 0.006$; controls, $t_{(12)} = 0.59; p = 0.563$; BF = 0.32; Bonferroni’s adjusted $\alpha$, 0.05/3 = 0.016). When comparing across groups (using a one-way ANCOVA, accounting for age), we found a trend toward significance ($F_{(2, 38)} = 2.99; p = 0.062$; $\eta_p = 0.14$; BF$_{10} = 2.05$), which is also reflected in greater homotopy in amputees relative to controls ($t_{(38)} = 2.41; p = 0.021$) but not for one-handers relative to controls ($t_{(38)} = 1.63; p = 0.112$; BF$_{10} = 1.44$; Bonferroni’s adjusted $\alpha$, 0.05/2 = 0.025 for the last two comparisons).

To determine whether the increased homotopy found in amputees reflects typical contralateral representation of the ipsilateral (missing hand) cortex, we next compared the ipsilateral representational structure of amputees and one-handers to the average DCM of controls’ contralateral averaged DCM (Fig. 3C). As shown in Figure 3C, for amputees we found a significant (above zero) correlation between both contralateral and ipsilateral ROIs relative to the typical contralateral representational structure in controls (amputees contralateral, $t_{(15)} = 6.82; p < 0.001$; amputees ipsilateral, $t_{(15)} = 4.38; p < 0.001$; one-sample $t$ test; Bonferroni’s adjusted $\alpha$, 0.05/2 = 0.025), whereas the correlation between one-handers and controls was approaching significance for the contralateral ROI only (one-handers contralateral, $t_{(12)} = 2.51; p = 0.027$; one-handers ipsilateral, $t_{(12)} = 1.70; p = 0.114$; Bonferroni’s adjusted $\alpha$, 0.05/2 = 0.025). The two-way ANCOVA comparing group and hemisphere showed an expected effect of hemisphere ($F_{(1, 26)} = 9.44; p = 0.005$; $\eta_p = 0.26$), reflecting the greater correlation of the contralateral hemisphere, and a significant main effect of group ($F_{(1, 26)} = 4.64; p = 0.041$; $\eta_p = 0.15$). The interaction was not significant ($F_{(1, 26)} = 1.73; p = 0.20$; $\eta_p = 0.06$). This demonstrates that amputees represented the different finger configurations bilaterally in a way that was similar to the typical representation in the contralateral hemisphere in neurotypical controls.

When repeating the same set of analyses in M1, amputees only showed a significant correlation between the contralateral ROI relative to the typical contralateral structure in controls (amputees contralateral, $t_{(15)} = 2.81; p = 0.013$; amputees ipsilateral, $t_{(15)} = 0.45; p = 0.659$; Bonferroni’s adjusted $\alpha$, 0.05/2 = 0.025; one-handers contralateral, $t_{(12)} = 1.62; p = 0.131$; one-handers ipsilateral, $t_{(12)} = 0.57; p = 0.575$; Bonferroni’s adjusted $\alpha$, 0.05/2 = 0.025). Furthermore, the two-way ANCOVA revealed a significant main effect of hemisphere ($F_{(1, 26)} = 5.55; p = 0.026$; $\eta_p = 0.17$), no main effect of group ($F_{(1, 26)} = 0.04; p = 0.840$; $\eta_p = 0.02$), and no interaction ($F_{(1, 26)} = 0.61; p = 0.443$; $\eta_p = 0.02$).

No differences in white matter tracts between the three groups
Finally, we analyzed diffusion MRI data, collected in the same cohort, to explore whether the group differences observed in the functional analysis are also reflected by alterations in structural connectivity. As noted in the Introduction, it is possible that ipsilateral functionality depends on the brain establishing (through bimanual experience) a functional interaction between the two hand territories. One possibility is that this is mediated, at least in part, via transcallosal pathways that connect the two hand areas (Filign, et al., 2013). To address this, we conducted deterministic tractography to examine potential differences in the tissue microstructural properties of the transcallosal fibers connecting the two hand areas. We first compared the vertex-weighted mean FA and MD, derived from tractography-based interhemispheric connections, using two separate ANCOVAs (controlling for age). For both metrics, the main effect of group was not significant (FA, $F_{(2, 35)} = 0.05; p \leq 0.950$; $\eta_p = 0.003$; BF$_{10} = 0.19$; MD, $F_{(2, 35)} = 0.08; p \leq 0.922$; $\eta_p = 0.005$; BF$_{10} = 0.20$). The Bayes factors in both analyses provided evidence in favor of the null hypothesis being no group structural differences in FA and MD.

To explore potential differences between amputees/one-handers and controls beyond these transcallosal interhemispheric connections, we conducted a complementary voxel-wise TBSS analyses at the whole-brain level, as well as within a corticospinal tract ROI (see Methods; Halamy et al., 2015). At the whole-brain level, we found no FA or MD differences between either group (amputees and one-handers) or controls (TFCE- corrected; $p = 0.05$). We also saw no significant clusters when contrasting amputees with one-handers. We did, however, find a negative effect of age, confirming the quality of the data. For the corticospinal tract, we similarly found no significant differences between each experimental group and the controls (both FA and MD), and this was also the case when comparing amputees with one-handers. Together, these findings do not support substantial structural changes in white matter architecture most relevant for interhemispheric coordination for motor control in our experimental groups.

**Discussion**
In this study, we investigated the impact of hand loss, whether congenital or through amputation, on intact hand motor ability and representation. Given the profound behavioral pressure of growing up and/or living with only one hand, perceptual learning combined with practice effects are likely to enhance motor skills of the intact hand in both groups. Critical development periods may be more favorable for training effects to occur (Sur and Rubenstein, 2005; Levelt and Hubener, 2012), favoring plasticity in one-handers and thus improving motor control and learning of the intact hand. Instead, we found that one-handers showed poorer performance in a finger configuration task, particularly when learning more difficult configurations, whereas amputees did not show any clear deviations from controls during task training outside the scanner. This aligns with prior research indicating motor deficits in one-handers but not amputees. For example, one-handers (Philip et al., 2015) but not amputees (Philip and Frey, 2011) exhibited accuracy and speed deficits while planning a grasp with their intact hand. Based on this, it has been postulated that sensorimotor experience of both hands is necessary for the refinement of accurate unilateral motor prediction and performance (Philip et al., 2015). Relatedly, we previously found that one-handers made more errors during visually guided reaching with their artificial arm, relative to amputees, as well as two-handed controls using their nondominant arm (Maimon-Mor et al., 2021; though it is worth noting that in this study, intact hand reaching performance was not significantly different from the other groups). Interestingly, one-handers who started using an artificial arm earlier in life as toddlers showed less motor deficit, hinting at a critical period for integrating a visual-motor representation of a limb. These findings imply that early-life disabilities may impede motor development, even for body parts not directly affected by the malfunction. This does not necessarily contradict the prediction that motor control and learning would be superior in one-handers due to early-life behavioral pressure. It is possible that critical periods trigger both long-term deficits and improved skill that would counterbalance each other. Without early-life over-practice, one-handers might have shown more severe motor impairments in their daily life.

Although the missing hand territory showed increased activity during the finger configuration task (as discussed below), amputees did not display superior motor performance with their...
intact hand. The idea of amputees gaining enhanced abilities following their amputation due to reallocation of central resources in the missing hand cortex has been a topic of much fascination for the past century. Originally, hypotheses (and reports) focused on heightened tactile sensitivity on the residual limb (stump) of human amputees (Katz, 1920; Teuber et al., 1949; Haber, 1955; see Makin et al., 2021 for a critical review of maladaptive consequences of reorganization in amputees). Merzenich et al. (1984) proposed that remapping following finger amputation should improve tactile acuity of the neighboring fingers. Other studies, using short-term and reversible deafferentation, suggested increased acuity for the nondeafferented ("intact") hand due to increased excitability of the deafferented hemisphere (Björkman et al., 2004; Lissek et al., 2009; Dempsey-Jones et al., 2019). More recently, we and other suggested that increased activity for the intact hand in the missing hand territory is a potential neural correlate of adaptive plasticity for motor abilities (Makin et al., 2013a; Philip and Frey, 2014). According to these ideas, the brain can correctly interpret missing hand territory signals as related to the intact hand, thereby providing greater (or better) information about the new representation. This is consistent with physiological studies showing that, while hand and finger movements are mostly controlled through contralateral (crossed) corticospinal projections (Brinkman and Kuypers, 1973), there are also known ipsilateral (uncrossed) motor projections (Soteropoulos et al., 2011). Given that amputees rely heavily on their remaining hand, one might expect improved read-out of neural signals originating from the ipsilateral cortex, which typically has limited functionality in two-handed individuals. This improvement should lead to recruitment of the missing hand hemisphere in the brain’s ipsilateral hemisphere. However, much of the original evidence for perceptual gains in amputees have been since challenged (O’Boyle et al., 2001; Vega-Bermúdez and Johnson, 2002). In our brief training paradigm, we found no evidence for motor behavioral benefits in amputees.

What is then the functional relevance of the increased ipsilateral activity observed in sensorimotor cortex of amputees here, as well as in many previous studies (Kew et al., 1994; Hamzei et al., 2001; Bogdanov et al., 2012; Makin et al., 2013a; Philip and Frey, 2014; Valyear et al., 2020)? One difficulty in interpreting the functional meaning of net changes in activity levels is that they could result from multiple dissociated mechanisms, such as aberrant processing (Makin et al., 2013b), disinhibition (Hahamy et al., 2017), or merely reflect gain changes due to upstream processing (Kambi et al., 2014). Common to these alternative processes is that increased activity does not necessarily entail a change of the underlying information being processed (Arbuckle et al., 2019). In other words, activity changes that underlie remapping do not necessarily entail information content changes. In the present study, we found that while difficulty increases contralateral activity across all groups, in the ipsilateral cortex, difficulty increases activity significantly only in amputees. This is interesting, because it goes against the idea that the increased ipsilateral activity is a simple passive consequence of interhemispheric disinhibition (Werhahn et al., 2002; Ramachandran and Altschuler, 2009; Simões et al., 2012). Instead, amputees selectively recruited the ipsilateral cortex for more difficult configurations. However, this does not necessarily indicate functional recruitment of the ipsilateral cortex. RSA is a multivariate technique designed to determine how separate or distinct one activity pattern is to another. RSA allows us to ask not only if more information is available in a given brain area (dissimilarity distances) but also whether this new information is structured consistently with known representational principles, for example, related to the contralateral hemisphere. By quantifying and characterizing brain function beyond the spatial attributes of activity maps, while providing a more precise model for how information content varies across configurations, we believe RSA provides an arguably better tool for assessing the functional characteristics of the ipsilateral cortex (as explained in Box 1 in Makin and Krakauer, 2023). Furthermore, both in our previous study (conducted on the same set of participants; Wesselinik et al., 2019) and here, we found that the increased activity in amputees did not translate to differentially increased ipsilateral information.

Several previous studies using multivariate pattern analysis have demonstrated that despite activity suppression, ipsilateral sensory and motor cortex contains information pertaining to individual fingers (Diedrichsen et al., 2013b, 2018; Berlot et al., 2019; Wesselinik et al., 2019). These ipsilateral activity patterns appear to be weaker, but otherwise similar in representational structure to those elicited by movement of the mirror-symmetric finger in the opposing hand, at least for single-finger movements (Diedrichsen et al., 2013a, 2018). The ipsilateral representation is not a simple "spill-over" or passive copy from the homologous (contralateral) hand area, as it has been shown to be differentially modulated by behavioral task context (Berlot et al., 2019). Yet, the functional significance of these ipsilateral representations and independence from the contralateral representation is still unknown. Ipsilateral activity in M1 has also been observed in monkey studies during proximal (i.e., shoulders and elbows) motor tasks (Ames and Churchland, 2019; Heming et al., 2019; Cross et al., 2020). These studies seem to suggest that, even if the same population of neurons encodes both ipsilateral and contralateral movements, the two limb representations are distributed differently across neurons (i.e., arm-related activity occupies distinct subspaces), which is proposed to be the mechanism that avoid impacting (i.e., moving) the wrong arm. Furthermore, it has been suggested that the ipsilateral representation is an efference copy resulting from the contralateral activity to inform the ipsilateral cortex about the contralateral arm movement and help with bimanual coordination (Ames and Churchland, 2019). The efference copy would be sent by default, even in the absence of bimanual movements, and ignored if not needed. In other words, the ipsilateral representation could be a consequence of the fact that the two homologous areas inform each other about their respective current state. Although the relationship between level of task complexity and the functional role of the efference copy has not been explored yet, it is interesting to speculate that the relevance of the efference copy will be greater for tasks requiring coordination across hands.

This latter interpretation provides an interesting conceptual framework for our reported findings: in one-handers, the lack of bimanual experience will dampen the mechanistic development of bimanual hand representation, including cross-hemisphere efference copy, reducing bilateral information content. Whereas in amputees, the ipsilateral efference copy from the intact hand will be more prominent in the missing hand cortex due to the reduced utilization of the missing hand, resulting in increased homotopy for the intact hand across the two hemispheres. Importantly, under this conceptual framework, these representational changes do not necessarily impact behavior because the efference copy is presumably involved in improving bimanual coordination, which is impossible for amputees to implement. This interpretation is consistent with a recent study which did not find a functional relevance for increased SI ipsilateral activity (Valyear et al., 2020), in line with our observation
that amputees did not show any behavioral improvement outside the scanner and, if anything, they showed a performance reduction inside the scanner. Our white matter findings also indirectly support the idea that activity changes are functionally irrelevant, as they reveal stable anatomy despite increased activity and better interhemispheric collaboration. In this context, it is interesting to consider previous evidence for persistent representation of the missing hand in amputees (Raffin et al., 2012; Makin et al., 2013b). The intact hand’s homotopic representation in the missing hand cortex might help maintain the missing hand representation. While we and others previously showed that the phantom hand map is activated by phantom hand movements independently of the intact hand (Kikkert et al., 2016; Bruurmijn et al., 2017; Wesselinck et al., 2019), it is still possible that structured inputs from the intact hand (via ipsilateral pathways) sustains the missing hand map, despite the loss of the original peripheral inputs.

As a potential limitation, we did not control for variations in finger postures, device locations, and hand sizes, which could influence task complexity. For example, since we used a single-size keyboard, participants with smaller hands might have found the task more complicated than participants with larger hands. While we acknowledge the potential impact of these factors, we believe they did not substantially impact on our results. Participants were allowed to choose their preferred device position and finger postures based on comfort, aiming to optimize their performance. Furthermore, the keyboard was designed with elongated keys, similar to a piano, to accommodate a broad range of hand sizes. Although we did not collect explicit hand size data, we used gender as a proxy, assuming women generally have smaller hands. Even after including gender in our models, the results remained consistent.

To conclude, our findings reveal a collaborative relationship between contralateral and ipsilateral cortices during task performance in amputees. By focusing on information content and its representational structure above and beyond the salient effects of remapping (defined as increased mean activity), our findings highlight a different aspect of the critical period than normally emphasized, which is based on experience rather than deprivation. Specifically, representations of both hands and some bimanual experience in the early developmental stage is necessary to develop a bilateral motor representation and a typical contralateral representation. Interestingly, while the ipsilateral efference copy interpretation is functionally irrelevant for the unimanual tasks studied here and in previous research, it may provide a useful consideration, and perhaps even new opportunities, for combining novel restorative brain–computer interfaces (Fouda et al., 2015) and augmentation technologies (Dominijanni et al., 2021) for bimanual interactions.


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