#### Full title:

# Investigating facial information content in the hand area of individuals with a congenital and acquired missing hand

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# **Abstract**

Cortical remapping after hand loss in the primary somatosensory cortex (S1) is thought to be predominantly dictated by cortical proximity, with adjacent body parts remapping into the deprived area. Traditionally, this remapping has been characterised by changes in the lip representation, which is assumed to be the immediate neighbour of the hand based on electrophysiological research in non-human primates. However, the orientation of facial somatotopy in humans is debated, with contrasting work reporting both an inverted and upright topography. We aimed to fill this gap in the S1 homunculus by investigating the topographic organisation of the face. Using both univariate and multivariate approaches we examined the extent of face-to-hand remapping in individuals with a congenital and acquired missing hand (hereafter one-handers and amputees, respectively), relative to two-handed controls. Participants were asked to move different facial parts (forehead, nose, lips, tongue) during fMRI scanning. We first report evidence for an upright facial organisation in all three groups, with the upper face and not the lips bordering the hand area. We further found little evidence for remapping of all tested facial parts in amputees, with no significant relationship to the chronicity of their PLP. In contrast, we found converging evidence for a complex pattern of face remapping in congenital one-handers across all facial parts, where the location of the

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cortical neighbour – the forehead – is shown to shift away from the deprived hand area, which is subsequently activated by the lips and the tongue. Together, our findings demonstrate that the face representation in humans is highly plastic, but that this plasticity is restricted by the developmental stage of input deprivation, rather than cortical proximity.

# Introduction

Our brains capacity to adapt, known as cortical plasticity, is integral to our successful functioning in daily life, as well as rehabilitation from injury. A key model for exploring the extent, and consequences of, cortical plasticity is upper-limb loss (via amputation or congenital absence). Here, the cortical hand territory in the primary somatosensory cortex (hereafter S1), suffers an extreme loss of sensory input in tandem with dramatic alterations of motor behaviour<sup>1,2</sup>. The functional and perceptual correlates of amputation-related plasticity are currently debated<sup>3,4</sup>. In particular, it is not clear whether functional cortical reorganisation is restricted to early life development or can also occur in adults.

Traditionally, research assessing cortical plasticity after upper-limb loss has followed the tenet that neighbouring body parts of the missing hand, and lower face in particular, shift and encroach into the deprived hand area. This emphasis on the lip representation stems from early electrophysiological work in non-human primates, where numerous studies demonstrated an 'upside-down' facial somatotopy, with the lower face immediately neighbouring the hand<sup>5–13</sup>. Here, the lips and/or lower-chin inputs have been shown to remap into the deprived hand area after sensory loss<sup>14,15</sup>, leading to the well-accepted assumption that remapping is determined by cortical proximity<sup>16,17</sup>. Thereafter, human measurement of topographic shifts has tended to focus on that of the lips, where researchers have reported that shifted lip representation towards and into the deprived hand area is significantly associated with phantom limb pain (PLP) intensity<sup>18–22</sup>. PLP is a neuropathic pain syndrome experienced in the missing, amputated limb by the majority of amputees<sup>23</sup>. This condition is commonly thought to arise from maladaptive cortical plasticity in S1 (although see<sup>24</sup>), specifically from a signal mismatch between the missing hand representation and the remapped inputs of the lips in the deprived hand area<sup>25</sup>.

The research focus on lip cortical remapping in amputees is based on this assumption that the lips neighbour the hand representation, however, it is not well supported. For example, only a handful of neuroimaging studies in humans has supported an inverted (or 'upside-down')

somatotopic organisation of the face, similar to that of non-human primates<sup>26,27</sup>. Contrasting work alternatively suggests an upright orientation of the face in S1<sup>28–32</sup>, with the upper-face (i.e., forehead) bordering the hand area. Indeed, recent work reported that the shift in the lip representation towards the missing hand in amputees was minimal<sup>33,34</sup>, and likely to reside within the face area itself. Surprisingly, there is currently no research that considers the representation of other facial parts, including the upper face in particular (e.g., the forehead), in relation to plasticity or PLP. Detailed mapping of the upper and lower face is therefore needed to assess typical topography of facial sensorimotor organisation, as well as remapping after limb loss.

Remapping after upper-limb loss has also been documented in individuals born without a hand (hereafter one-handers), who do not experience PLP<sup>35</sup>. Here it has been shown that the representation of multiple body parts, including the residual arm, legs and mouth, remapped into the missing hand territory<sup>36,37</sup>. Importantly, cortical remapping in this group does not depend on cortical proximity of the body parts. With regards to the lips, a recent transcranial magnetic stimulation (TMS) study has reported functionally-relevant lip activity in the deprived hand area of one-handers<sup>38</sup>, showcasing that reported remapping may also be functional. It was proposed that the observed remapping of various body parts could have been shaped by compensatory behaviour<sup>36</sup>, as these body parts are all used by one-handers to compensate for their missing hand function, but this hypothesis awaits validation.

Here, we conducted a mapping of face cortical organisation to determine facial orientation (upright versus inverted) in the primary sensorimotor cortex in 22 two-handed controls, 15 amputees and 21 one-handers. We used surface-based comparisons, including cortical (geodesic) distances, to measure the extent of cortical remapping of the upper (forehead) and lower face (lips) in relation to the deprived (or non-dominant) hand area across all groups. We also explored the representation of the tongue, which has not been previously studied in the context of deprivation-triggered brain plasticity. Furthermore, we used multivariate representational similarity analysis (RSA) in order to characterise more subtle alterations in the relationship between facial activity patterns (forehead, nose, lips, tongue) in the deprived hand and face areas, independent of gross spatial somatotopic shifts.

We found that facial topography was arranged in an upright manner, with the forehead (i.e., upper face) bordering the hand area across all groups. Contrary to traditional theories<sup>39</sup>, we did

not find evidence for facial remapping, including the lips, into the deprived hand area of amputees. We did, however, observed significant remapping of all face parts (upper and lower face) in the one-handers' group, validating our methodology as suitable for identifying remapping effects. Interestingly, remapping of the cortical neighbour (upper face) within the one-hander group was away from the missing hand area, while the lips and tongue representations shifted towards the deprived cortex, hinting that the underlying mechanism of remapping is more complex than simple cortical proximity.

# Materials and methods

## **Participants**

Fifteen individuals with acquired unilateral upper-limb amputation (age; M = 55.07, SE = 2.60, women; n = 3, missing right hand; n = 9), twenty-one individuals with unilateral congenital transverse arrest (age; M = 42.67, SE = 3.04, women; n = 13, missing right hand; n = 8) and twenty-two two-handed controls (age; M = 45.55, SE = 2.02, women; n = 10, left-handers; n = 6) were recruited (see Table 1 for full details). Three additional amputees who were recruited for the study did not participate in the scanning session due to MRI safety concerns, and further recruitment was stalled due to Covid-19 restrictions. The proportion of participants with intact/dominant right hand was matched across one-handed and control groups, respectively ( $\chi^2_{(2)} = 4.025$ , p = .134). Statistically significant differences were observed for gender ( $\chi^2_{(2)} = 6.218$ , p = .045) and age ( $H_{(2)} = 9.027$ , p = .011). Importantly, post-hoc comparisons confirmed non-significant differences between amputees and congenital one-handers when compared to controls. Covariates of age and gender were therefore only included in statistical analyses when direct comparisons between amputees and one-handers were carried out. Procedures were in accordance with NHS National Research Ethics Service approval (18/LO/0474), and written informed consent was obtained.

Participants	Age	Gender	Handedness (prior to amputation for amputees)	Affected limb	Level of limb deficiency	Years since amputation	PLS intensity	PLS frequency	Chronic PLS	PLP intensity	PLP frequency	Chronic PLP	Cause of amputation
AA01	60	М	R	R	2	43	100	5	100	60	5	60	Trauma
AA02	34	M	R	R	1	3	50	2.5	14.6	70*	2	17.5*	Trauma
AA03	58	M	R	R	1	33	90	5	90	100	1	20	Trauma
AA04	59	M	R	L	2	16	40	1	8	0	1	0	Trauma
AA05	54	M	Α	L	1	36	100	5	100	80	4	40	Trauma
AA06	47	F	R	L	2	18	80	4	40	0	0	0	Electrocution
AA08	40	F	R	R	1	10	40	3	13.3	0	0	0	Trauma
AA09	47	M	R	R	2	5	70	4	35	10	4	5	Trauma
AA10	53	M	R	L	2	34	20	0	0	0	0	0	Trauma
AA12	66	M	R	R	1	38	60	5	60	0	0	0	Trauma
AA13	65	F	L	L	1	10	90	5	90	80	4	40	Trauma
AA14	66	M	R	L	1	35	80	2	20	100	2	25	Trauma
AA16	64	M	R	R	1	18	75	5	75	65	5	65	Trauma
AA17	65	М	R	R	1	8	70	5	70	0	0	0	Trauma
AA18	48	M	R	R	1	23	85	5	85	65	5	65	Trauma
CA01	32	F	R	L	2								
CA02	32	F	R	L	2								
CA03	35	М	R	L	2								
CA04	48	М	R	L	2								
CA05	22	F	L	R	2								
CA06	54	F	R	L	2								
CA07	56	F	L	R	2								
CA08	53	М	R	L	1								
CA09	54	F	R	L	2								
CA10	58	М	L	R	2								
CA11	22	М	R	L	2								
CA12	30	F	R	L	2								
CA13	24	М	L	R	2								
CA14	33	F	L	R	2								
CA15	39	F	L	R	2								
CA16	55	F	R	L	2								
CA17	67	F	L	R	2								
CA18	30	F	L	R	2								
CA19	43	M	R	L	2								
CA20	63	М	R	L	2								
CA21	46	F	R	L	2								

**Table 1.** Demographic details for one-handed groups. Level of limb deficiency is as follows:  $l = limb \ loss \ above \ elbow \ (transhumeral), \ 2 = limb \ loss \ below \ elbow \ (transradial); \ L = left, \ R$  = right; PLS & PLP frequency:  $0 = no \ sensation \ or \ pain, \ l = once \ or \ less \ per \ month, \ 2 = several \ times \ per \ month, \ 3 = once \ a \ week, \ 4 = daily, \ 5 = all \ the \ time. \ ^*PLP \ intensity \ rating \ was on average. PLS = phantom \ limb \ sensations; PLP = phantom \ limb \ pain.$ 

# Phantom sensations rating

Amputees were asked to rate the frequency of PLP experience within the last year. They also rated the intensity of their worst PLP experience during the last week (or in a typical week involving PLP; 0 = no pain, 100 = worst pain imaginable). A chronic measure of PLP was calculated by dividing the worst PLP intensity in the last week by PLP frequency (1 = all the time, 2 = daily, 3 = once a week, 4 = several times per month, 5 = once or less per month). This approach to categorising PLP, which takes into account the chronic aspect of the condition, has been used successfully before in the literature  $^{33,35,40-44}$ , and has high inter-session reliability  $^{42}$ . We also asked amputees about the vividness and frequency of non-painful phantom sensations (see *Table 1*).

# Functional MRI sensorimotor task

We used an active motor paradigm, where participants were visually instructed to move one of four different face parts (forehead, nose, lips and tongue), as well as moving the left and right thumb (amputees were asked to flex/extend their phantom thumb to the best of their ability; one-handers were asked to imagine such movement), resulting in 6 conditions. Baseline (i.e., rest) was included as a 7<sup>th</sup> condition. Specific instructions involved: raising eyebrows (forehead), flaring nostrils (nose), puckering lips (lips), tapping tongue to the roof of the mouth (tongue), flexing and extending (thumb). The protocol comprised of 8 s blocks, with each condition repeated 4 times per run (5 times for baseline), over 3 functional runs. Before entering the scanner, the participants practised each movement with the experimenter to ensure that the movement could be executed and to standardise each movement across participants (e.g., specificity and pace). Note that multiple participants reported during the experimenter briefing that they could not successfully flare their nostrils, and were therefore instructed to attempt moving their nose in the scanner.

# MRI data acquisition

Functional and anatomical MRI data were obtained using a 3 Tesla Prisma MRI scanner (Siemens, Erlangen, Germany) with a 32-channel head coil. Anatomical data was acquired using a T1-weighted sequence (MPRAGE), with the following parameters: TR = 2530 ms; TE = 3.34 ms; flip angle = 7°; voxel size = 1 mm isotropic resolution. Functional data based on the blood oxygenation level dependant (BOLD) signal were acquired using a multiband T2\*-weighted pulse sequence, with a between-slice acceleration factor of 4 and no in-slice acceleration (TR = 1450 ms; TE = 35 ms; flip angle = 70°; voxel size = 2 mm isotropic resolution; imaging matrix = 106 x 106; FOV = 212 mm). 72 slices were oriented in the transversal plane. A total of 172 whole-brain volumes for each of the three runs were collected per participant. Field-maps were acquired for field unwarping.

# **MRI** pre-processing

Functional data was first pre-processed using FSL-FEAT (version 6.00). Pre-processing included motion correction using MCFLIRT<sup>45</sup>, brain extraction using BET<sup>46</sup>, temporal highpass filtering with a cut-off of 119 s and spatial smoothing using a Gaussian kernel with a FWHM of 3 mm. Field maps were used for distortion correction of functional data.

For each participant, we calculated a midspace between the three functional runs, i.e., the average space in which the images are minimally reorientated. Each functional run was then aligned to the midspace and registered to structural images (within-subject) using FMRIB's Linear Image Registration Tool (FLIRT), and optimised using Boundary-Based Registration<sup>47</sup>. Where specified, functional and structural data were transformed to MNI152 space using FMRIB's Nonlinear Registration Tool (FNRIT<sup>48</sup>).

# Regions of Interest (ROI) definition

Facial topography and remapping were studied using anatomical ROIs for the hand and face areas in S1. Although M1 and S1 are expected to activate during facial movement we primarily focused on S1 remapping due to the traditional focus in the maladaptive plasticity literature on S1 representational shifts<sup>39</sup>. Furthermore, M1 topography tends to be less well-defined<sup>49,50</sup>, and so characterisation of typical facial topography may be more apparent in S1. Nevertheless, we wish to note that due to the proximity of S1 to M1, it is possible that marginal contribution from M1 may have affected our S1 activity profiles.

Firstly, S1 was defined on the average surface using probabilistic cytoarchitectonic maps, by selecting nodes for Brodmann areas (BAs) 1, 2, 3a and 3b<sup>51</sup>. The S1 hand ROI (hereafter hand ROI) was defined by selecting the nodes approximately ~1 cm below and ~2.5 cm above the anatomical hand knob. In contrast to earlier work<sup>52</sup>, we defined a conservative lateral boundary of the hand ROI (~1cm below the hand knob) to ensure there was limited facial activity captured. From the remaining parts of S1, the medial region was discarded and the lateral region was selected as the preliminary approximation for the S1 face ROI (hereafter face ROI). The same procedure was carried out for M1 hand and face ROIs, but with selected nodes in BA 4.

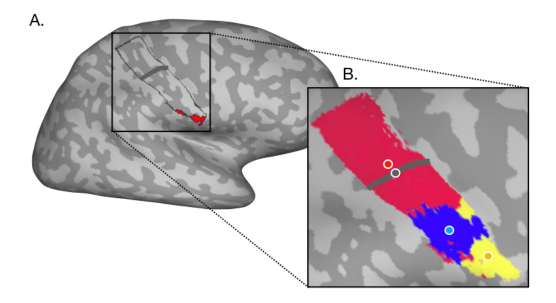


Figure 1. Regions of interest and winner-takes-all analysis in the primary somatosensory cortex for an example participant. A) Hand and face regions of interest in S1 are outlined in black, with the border between the two regions highlighted in grey. ROI overlap with the secondary somatosensory cortex (S2) are highlighted in red, which were removed from the face ROI in order to minimise somatotopic contribution from that region. B) A typical winner-takes-all map from an example participant, with forehead activity in red, lip activity in blue and tongue activity in yellow. The centre-of-gravity for each movement is signified by a coloured dot outlined in white. The hand-face border is outlined in grey, with the midpoint denoted by a grey dot. Cortical geodesic distances were measured from each facial parts CoG to the hand-face border midpoint.

Structural T1-weighted images were then used to reconstruct pial and white-grey matter surfaces using Freesurfer (version 7.1.1) at the individual level. The hand and preliminary face ROIs were then projected into individual brains via the reconstructed individual anatomical surfaces. As the secondary somatosensory cortex (S2) contains a crude somatotopy<sup>53</sup>, the preliminary face ROI was further trimmed in participant's structural space by removing the overlap with S2. S2 was defined in MNI152 space using the Juelich Histological Atlas<sup>54</sup>. The S2 ROI was registered to participants' structural space using an inversion of the nonlinear registration carried about by FNIRT. The remaining preliminary face ROI with the overlap from S2 removed was used as the face ROI for all univariate analyses. We note that due to the

probabilistic nature of these masks, there could be some marginal contribution from S2 in our estimated face area.

# Functional MRI data analysis

Time-series statistical analysis was carried out using FMRIB's Improved Linear Model (FILM). Task-based statistical parametric maps were computed by applying a voxel-based General Linear Model (GLM), as implemented in FEAT. The design was composed of 6 explanatory variables for each movement, convolved with a double-gamma hemodynamic response function<sup>55</sup>, and its temporal derivative. The six motion parameters were included as regressors of no interest. Motion outliers (> 0.9 mm) of large movements between volumes were included as additional regressors of no interest at the individual level (of total *n* volumes per group: amputees: 0.36%; controls: 0.36%; one-handers: 0.42%). For our main comparisons 6 contrasts were set up, corresponding to the facial movements' (forehead, nose, lips, tongue and left/right thumb) relative to rest. Since the nose condition yielded weak activity relative to baseline, we excluded it from the univariate analysis.

The estimates from the three functional runs were then averaged voxel-wise using a fixed effects model in participants structural space, with a cluster forming z-threshold of 2.3 and family-wise error corrected cluster significance threshold of p < 0.05. Each estimates' average was masked prior to cluster formation with a sensorimotor mask, defined as the precentral and postcentral gyrus from the Harvard Cortical Atlas. The sensorimotor mask was registered to the individuals structural scan using an inversion of the nonlinear registration by FNIRT.

# Winner-takes-all approach

In order to characterise facial topography in S1, including possible instances of representational remapping, the hand and face ROIs were combined to produce an overall S1 ROI (minus the medial region), and a winner-takes-all approach was used (Figure 1B). For each participant, thresholded z-statistics averaged across the three functional runs were assigned to one of three face parts (forehead, lips, tongue), dependent on which facial movement relatively showed maximal activity within the S1 ROI. Face-winners (i.e., the output of the winner-takes-all) were then projected to the individual's anatomical surface, and here the centre-of-gravity (CoG) was calculated bilaterally for each movement (weighted by cluster size<sup>33</sup>). Note that we excluded activity for the thumb in order to align our analysis with previous research, and to draw

comparisons of facial somatotopy across all groups (one-handers do not have a phantom limb, and therefore we cannot probe the 'missing' hand representation directly).

#### Cortical distance analysis

To assess possible shifts in facial representations towards the hand area, the cortical distance between each movement's CoG and a predefined cortical anchor was calculated along the individual's anatomical surface using Connectome Workbench (v1.4.2). The cortical anchor was defined as the midpoint of the lateral border of the hand ROI (see Figure 1B). The border was drawn manually and the midpoint calculated for each participant, and were visually confirmed by a second experimenter. The geodesic distance was then calculated from a movement's CoG to the hand border midpoint in each hemisphere. If the movement's CoG was located below the hand border (i.e., laterally), the resulting distance was assigned a negative value. The forehead's CoG location, and therefore assignment of negative distance values, were manually visualised and checked due to its proximity to the hand border.

#### Surface area calculation

To assess possible remapping into the hand area, a secondary winner-takes-all analysis was restricted to the hand ROI only. The surface area coverage (mm²) for each face-winner were computed on the individual anatomical inflated surface using Connectome Workbench (v.1.4.2). We next calculated the proportion of the hand ROI occupied by each face part by dividing each face-winner's surface area by the total hand ROI surface area for each individual. From the resulting percentages, we produced a laterality index for each movement with the following formula:

$$laterality\ index = \frac{(deprived_m - intact_m)}{(\sum deprived_m, intact_m)}$$

whereby  $deprived_m$  and  $intact_m$  represent the percentage of surface area coverage for the facial movement, m, respectively in the deprived and intact hemisphere. A subsequent laterality index of +1 indicates surface area coverage of that movement solely within the deprived hemisphere (or the hemisphere contralateral to the non-dominant hand in controls), whereas a value of -1 relates to surface area coverage solely within the intact hemisphere (or the hemisphere contralateral to the dominant hand in controls). A value of 0 represents an equal balance of surface area coverage across the deprived/non-dominant and intact/dominant hemispheres.

Note that this approach characterises cortical remapping in relation to the intact hemisphere and has been used in numerous previous studies on amputees<sup>18,19,21,22</sup>. It assumes that the intact hemisphere reflects baseline (i.e., that it is truly 'intact'), which may not be the case due to inter-hemisphere plasticity and/or homeostatic mechanisms<sup>2,56,57</sup> and so we compared our results to the control group.

# **Group-level visualisations**

Prior to group-level visualisations, participant information regarding hand dominance (controls) and deprived hemisphere (one-handed participants) were used to sagittal-flip raw pre-processed data, such that the brain activity corresponding to the non-dominant/missing hand is always represented in the left hemisphere (note that a similar proportion of participants were flipped across groups). Group-level statistical parameter maps were then created with a threshold-free cluster enhancement (TFCE) approach using FSL's Randomise tool<sup>58</sup>. TFCE is a nonparametric, permutation-based method for cluster formation, and has been shown to demonstrate improved sensitivity when compared to typical thresholding methods<sup>59</sup>. Group activity mixed-effect maps were calculated for each fixed-effect (i.e., averaged across the three functional runs) parameter estimate of a face movement (forehead, lips, tongue) contrasted to baseline. Prior to permutation (n = 5000), parameter estimates were masked with a sensorimotor mask, defined as the precentral and postcentral gyrus from the Harvard Cortical Atlas. A family-wise error correction of p < 0.05 and variance smoothing of 5 mm (as recommended for datasets with less than 20 participants) were used. Resulting clusters were thresholded at p < .001 and projected to a group cortical surface<sup>60</sup> using Connectome Workbench (v1.4.2).

As well as activity maps, we also visualised the winner-takes-all output at the group-level. Here the 'winners' for each face movement within the S1 ROI (the hand and face region combined) in MNI152 space were concatenated into a single volume per group to produce a consistency map for the individual movements (i.e., how many participants maximally activated the same voxel when moving a given facial part). Resulting consistency maps were then projected to a group cortical surface<sup>61</sup> using Connectome Workbench (v1.4.2).

# Multivariate representational analysis

Representational Similarity Analysis (RSA<sup>62</sup>) was used to assess the multivariate relationship between activity patterns generated by each face part. RSA was conducted in the hand and face ROIs to explore possible remapping across representational features between groups. To ensure the selectivity of the hand and face areas, the ROIs used for univariate analyses were each further trimmed medially by ~1cm, creating a 1cm gap between the hand and face ROIs. For each participant, parameter estimates for the four facial movements (forehead, nose, lips and tongue) and the contralateral thumb (for controls and amputees only) were extracted from all voxels within the chosen ROI, as well as residuals from each runs' first-level analysis (three runs in total). Here we included the nose movement in our analysis due to increased sensitivity to subtle changes in activity patterns that is afforded to us by using a multivariate approach. Multidimensional noise normalisation was used to increase reliability of distance estimates (noisier voxels are down-weighted), based on the voxel's covariance matrix calculated from the GLM residuals. Dissimilarity between resulting facial activity patterns were then measured pairwise using cross-validated Mahalanobis distances<sup>63</sup>. Due to cross-validation, the expected value of the distance is zero if two patterns are not statistically different from each other. Distances significantly different from zero indicate the two representational patterns are different; negative distances indicate noise. Larger distances for movement pairs therefore suggest greater discriminative ability for the chosen ROI. The resulting six unique inter-facial representational distances (10 unique distances when including the thumb for controls and amputees only) were characterised in a representation dissimilarity matrix (RDM). Multidimensional scaling (MDS) was also used to project the higher-dimensional RDM into lower-dimensional space, whilst preserving inter-facial dissimilarity, for visualisation purposes only. Analysis was conducted on an adapted version of the RSA Toolbox in MATLAB<sup>62</sup>, customised for FSL<sup>64</sup>.

## Statistical analyses

All statistical analyses were carried out using JASP (Version 0.14). When appropriate, univariate analyses was compared using parametric statistics. To assess normality for parametric tests, Shapiro-Wilk tests were run on residuals in combination with inspection of Q-Q plots and reporting of Levene's Test for Equality of Variances. Where stated, non-parametric test statistics are reported where the assumption of normality has been violated. Analysis of Variance (ANOVA) was used to explore group differences to controls in cortical distances and surface area coverage. Each mixed ANOVA had a between-subject factor of

Group (Controls x Amputees; Controls x One-handers) and a repeated-measures factor of Hemisphere (Intact/Dominant x Deprived/Non-dominant), and was run separately for each facial movement (forehead, lips and tongue). We controlled for brain size volume when comparing cortical distances between groups. Post-hoc comparisons were conducted with a Bonferroni correction for multiple comparisons (corrected alpha = .025; reported uncorrected *p*-values in text). We reported the corresponding Bayes Factor (BF<sub>10</sub>), defined as the relative support for the alternative hypothesis, for non-significant interactions and post-hoc comparisons. While it is generally agreed that it is difficult to establish a cut-off for what consists sufficient evidence, we used the threshold of BF<1/3 as positive evidence in support of the null, consistent with others in the field <sup>65,66</sup>. Note, however, that this threshold is not considered as providing strong evidence by all accounts<sup>67</sup>. The Cauchy prior width was set at 0.707 (JASP's default). To investigate whether remapping measures were related to PLP, we used Mann-Whitney U tests to compare laterality indices of those with and without PLP for relevant facial parts.

For multivariate analyses, in order to quantify the dissimilarity, or 'information', each ROI holds for the face across groups, a linear mixed model (LMM) analysis was used, allowing us to consider the distinct contribution of face-thumb or face-face pairs. The face-face LMM contained fixed factors of group (controls, amputees and one-handers), hemisphere (intact, deprived) and facial pairs (6 unique representational distances). A random effect of participant, as well as covariates of age and gender, were also included in the model. An additional LMM was used to explore potential remapping in the hand ROI in relation to the non-dominant/phantom thumb in controls and amputees. Here the LMM contained the fixed factors of group (controls and amputees), hemisphere (intact, deprived) and face-to-thumb pairs (4 unique representational distances), as well as a random effect of participant. All LMM's were carried out in jamovi (version 1.6.15) under restricted maximum likelihood (REML) conditions with Satterthwaite adjustment for the degrees of freedom.

# **Results**

# The cortical neighbour of the hand representation is the forehead

When looking at facial organisation at the group-level we found qualitatively similar activity maps across groups (see Fig. 2), highlighting a robust somatotopy of the face with preserved symmetry across the two hemispheres. These facial maps also indicate an upright orientation of the face in S1, with the forehead located closest to the hand area, followed by the lips, and the tongue located laterally, across all groups. The facial somatotopy presented here therefore suggests that the hand's cortical neighbour is the forehead (or upper face), highlighting the need to reassess the often-cited, traditional lip-to-hand marker of cortical remapping in amputees and one-handers. However, conclusions based on group averages may be misleading as they ignore inter-individual differences.

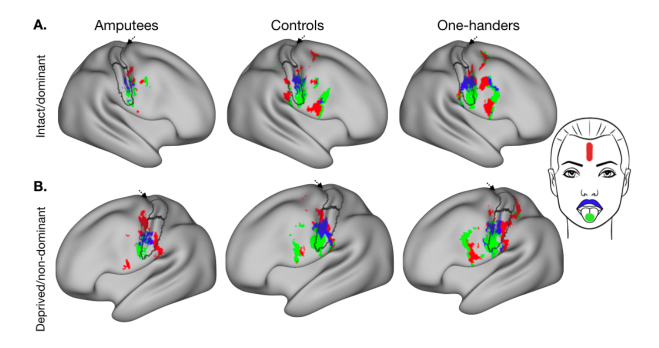


Figure 2. Group-level activity maps for each facial movement. (A) Group average activity for the forehead (red), lips (blue) and tongue (green) movements, contrasted to rest, in the intact/dominant hemisphere for controls (n = 22), amputees (n = 15) and one-handers (n = 21). (B) Group average activity for each facial movement in the deprived/non-dominant hemisphere. All clusters were created using a threshold-free cluster enhancement procedure with a sensorimotor pre-threshold mask (defined using the Harvard Cortical Atlas), and thresholded at p < .001. The hand and face ROIs are outlined in black, and the central sulcus is denoted with a black, dashed arrow.

# One-handers, but not amputees, show lip remapping in the deprived cortex based on univariate topographic mapping

To account for inter-individual differences in functional topography and brain topology, we first explored changes in the cortical (geodesic) distance between the lips and the hand-face border of amputees and controls. Here we found no statistically significant main effects or group x hemisphere interaction ( $F_{(1,34)} = 0.021$ , p = 0.886,  $n^2 = 0.0001$ ,  $BF_{10} = 1.865$ ; controlled for brain size volume; Fig. 3B), indicating that the lip area in amputees is not located differently to that of controls. This is further supported by a lack of evidence for significant remapping of the lips (i.e., greater surface coverage) in the missing hand ROI for amputees when compared to controls (U = 157.000, p = 0.816, d = -0.048,  $BF_{10} = 0.337$ ; Fig. 3C). These results suggest, contrary to popular theories on brain plasticity in amputees<sup>39</sup>, that the lips are likely to not remap into the deprived hand area. We next compared the lips laterality index between those individuals who reported suffering from PLP (n=9) and those who no longer experienced chronic PLP (n=6) and found no significant differences ( $U_{(13)} = 28.000$ , p = 0.955, d = 0.037,  $DF_{10} = 0.461$ ). However, the reported Bayes Factors indicated only anecdotal evidence for the null, and so we cannot be certain that a lack of remapping exists.

When looking at lip plasticity within the one-handers group, however, we did note a slight qualitative shift in the location, and spread, of the lip activity within the deprived hemisphere (Fig. 2B). This is further supported by a visible shift of the one-handers lip group-level consistency map towards, and into, the deprived hand area (Fig. 3A). These changes in the lip representation were statistically significant, with a significant group x hemisphere interaction for the lips cortical distance to the hand-face border in one-handers and controls ( $F_{(1,40)} = 5.419$ , p = 0.025,  $n^2 = 0.032$ ; controlling for brain size; Fig. 3B). Confirmatory comparisons indicated no statistically significant shifts of the lip CoG in the deprived hemisphere when compared to the controls non-dominant hemisphere ( $t_{(41)} = -1.513$ , p = 0.138, d = -0.462,  $t_{(20)} = 0.745$ ; Fig. 3B). However, shorter distances from the lips to the hand area were found in the deprived hemisphere of the one-handers when compared to their intact hemisphere ( $t_{(20)} = -3.073$ ,  $t_{(20)} = -0.006$ ,  $t_{(20)} = -0.671$ ), indicating substantial evidence for lip remapping. These shifts in the deprived hemisphere were also reflected in significantly greater surface area coverage of the lips in the hand ROI when compared to controls ( $t_{(20)} = -0.007$ ,  $t_{(20)} = -0.485$ ; Fig. 3C),

which was significantly different from zero (W = 197.000, p = 0.003, d = 0.706). This evidence of lip remapping is in line with previous work in one-handers<sup>36,38</sup>.

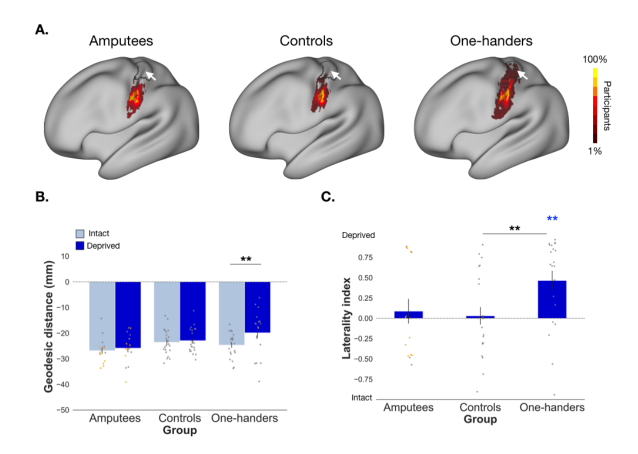


Figure 3. Characterisation of lip (re)mapping in the primary somatosensory cortex. (A) Group-level consistency map for the lips in the S1 ROI (hand and face combined) across groups. The colour gradient represents percentage of participant agreement for maximally activating that particular voxel, relative to other face movements (winner-takes-all approach). For example, yellow reflects all participants (100%) activated that particular voxel when moving their lips, relative to the forehead or tongue. The hand ROI is outlined in black and central sulcus denoted by the white arrow. (B) Cortical geodesic distances from the lip CoG to the hand-face border are plotted for amputees (n = 15), controls (n = 22) and one-handers (n = 21). Distances in the intact/dominant hemisphere are plotted in light blue, and distances in the deprived/non-dominant hemisphere are plotted in darker blue. (C) Laterality indices for the proportion of surface area coverage of the lips in the hand ROI for all groups (amputees, controls and one-handers). Positive values indicate greater surface area coverage in the deprived/non-dominant hemisphere, and negative values reflect greater surface area coverage in the intact/dominant hemisphere. Standard error bars and all individual data-points are plotted in grey and uncorrected for brain size. Amputees with PLP (yes/no) are plotted in

orange. \* p < .05; \*\* p < .01 (please note p-values are uncorrected); coloured asterisk's indicate values are significantly different from zero.

# One-handers, but not amputees, show forehead remapping in the deprived cortex, based on univariate topographic mapping

As we note a qualitative upright orientation of the face (see Fig. 2), the question remains as to whether the neighbour to the hand – the forehead – would reorganise after limb loss in amputees, as hypothesised by traditional theories <sup>39</sup>. Again, we found no significant evidence for cortical remapping of the neighbouring forehead in amputees when assessing changes in cortical distances (group x hemisphere:  $F_{(1,34)} = 1.379$ , p = 0.248,  $n^2 = 0.017$ ,  $BF_{10} = 0.131$ ; controlled for brain size volume; Fig. 4B), with the reported Bayes Factor indicating moderate evidence for the null. A non-significant difference of forehead surface area coverage in the deprived hand ROI when compared to controls was also reported ( $t_{(35)} = 1.795$ , p = 0.081, d = 0.601,  $BF_{10} = 1.105$ ; Fig. 4C; note that here the trend is towards less remapping in amputees). We therefore find little evidence for remapping of the deprived hand's cortical neighbour – the forehead.

However, when looking at the one-handers group we did find significant evidence of forehead remapping with a group x hemisphere interaction ( $F_{(1,40)} = 8.287$ , p = 0.006,  $n^2 = 0.059$ ; Fig. 4B). Confirmatory comparisons indicated a positive trend for shorter distances of the foreheads' CoG to the hand-face border in the deprived hemisphere when compared to their intact hemisphere ( $t_{(20)} = 2.349$ , p = 0.029, d = 0.513,  $BF_{10} = 2.094$ ) and significantly shorter distances when compared to the controls non-dominant hemisphere ( $t_{(41)} = 2.484$ , p = 0.017, d = 0.758). As the forehead's CoG tended to be located above the hand-face border (see Fig. 4A), these results indicate a significant shift of forehead activity away from the deprived hand ROI. This is further supported by a significant decrease of surface area coverage for the forehead in the deprived hand ROI when compared to controls (U = 381.000, p < .001, d = 1.069), which was significantly different from zero (W = 19.000, p < .001, d = -0.835; Fig. 4C). Remapping of the cortical neighbour in one-handers, therefore, manifests in a shifting away of the upper face from the deprived hand area, possibly due to increases in activity of other facial movements, e.g., lips.

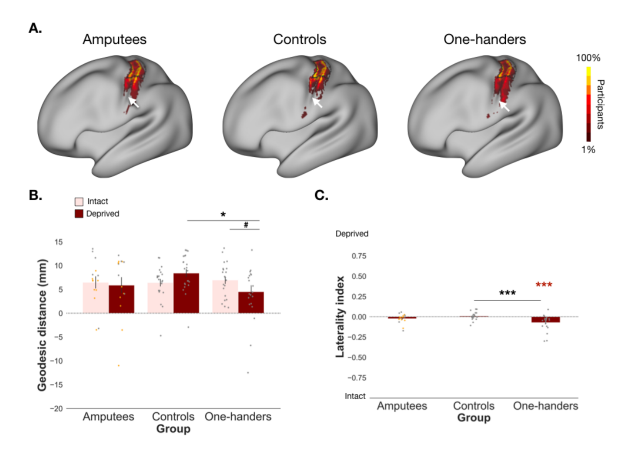


Figure 4. Forehead topography in the primary somatosensory cortex. All annotations are as in Figure 2. Distances in the intact hemisphere are plotted in pink and deprived hemisphere in red. # strong positive trend; \*p < .05; \*\*\* p < .001; coloured asterisk's indicate values are significantly different from zero.

# Tongue movements produce different topographic maps across groups

We also assessed changes in the tongue representation. As tongue somatotopy in amputees and one-handers has not yet been explored (with previous research mainly focusing on the lips<sup>18,20</sup>) we held no prior hypothesises in regards to changes in cortical distances and surface area coverage in the hand ROI. When looking at group-level consistency maps for the tongue (Fig. 5A) we note a lateral 'hotspot' of tongue activity across groups, indicating that the tongue is not an immediate neighbour to the hand. We also did not find significant evidence for shifts in the tongue's CoG towards the hand-face border in amputees when compared to controls (group x hemisphere:  $F_{(1,34)} = 2.572$ , p = 0.118,  $n^2 = 0.007$ ,  $BF_{10} = 0.172$ ; controlled for brain size volume; Fig. 5B). Nevertheless, the tongue did show significantly greater surface area coverage in the deprived hand ROI of amputees when compared to controls ( $U_{(35)} = 82.000$ , p = 0.011, d = -0.503; Fig. 5C), which was also significantly different to zero (W = 77.000, p = 0.030, d = 0.030, d = 0.030, which was also significantly different to zero (W = 0.030, D = 0.030, D

= 0.692). As tongue remapping is not reflected consistently across analyses, and due to the lack of pre-existing hypotheses, this preliminary result should be interpreted with caution. However, it does indicate that some level of cortical remapping may occur in amputees after limb loss.

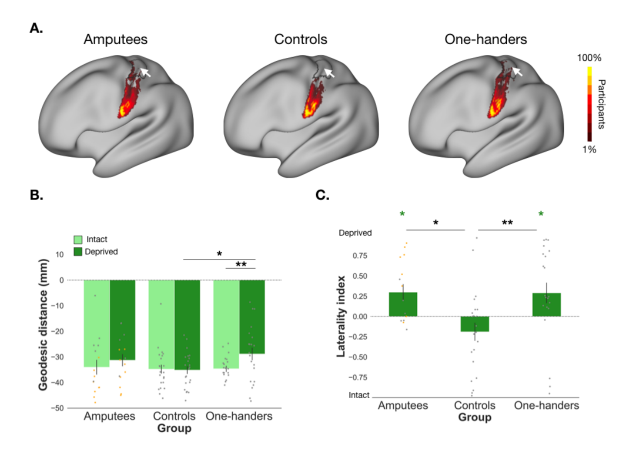


Figure 5. Tongue remapping in amputees and one-handers in the primary somatosensory cortex. Distances in the intact hemisphere are plotted in light green and distances in the deprived hemisphere in dark green. All other annotations are as in Figure 2.

We next explored whether this increase in tongue activity within the deprived hand ROI of amputees was related to PLP (Fig. 5C). To do so, we compared laterality indices for the tongue in those with and without PLP and found a trend towards significance ( $U_{(13)} = 42.000$ , p = 0.087, d = 0.556, BF<sub>10</sub> = 1.071). Since the Bayesian Factor does not indicate support for the hypothesis (i.e., BF < 3), these results provide inconclusive evidence for the relationship between tongue remapping and PLP.

We also found greater surface area coverage of the tongue in the deprived hand ROI of one-handers (see Fig. 5A) that was significantly different from zero (W = 180.000, p = 0.024, d = 0.558) and from controls (U = 109.000, p = 0.003, d = -0.528; Fig. 5C). This increased tongue

activity in the one-handers deprived hand area was also reflected in a significant group x hemisphere interaction ( $F_{(1,40)} = 7.149$ , p = 0.011,  $n^2 = 0.032$ ; controlled for brain size volume; Fig. 5B) of the cortical distance between the tongues' CoG and the hand-face border. Confirmatory comparisons indicated significantly shorter distances to the hand-face border for the tongue in the deprived hemisphere compared to intact ( $t_{(20)} = -2.902$ , p = 0.009, d = -0.633). However, the comparison between one-handers' and controls' non-dominant hemisphere was not significant (U = 150.00, p = 0.050, d = -0.351, BF<sub>10</sub> = 1.164). These results tentatively suggest that cortical remapping in one-handers may extend further to include tongue facial movements.

# Brain decoding in the deprived hand area reveals stable facial representational pattern for amputees, and increased facial information in one-handers

The analyses described above focused on the topographic relationship of the three facial parts, but cortical remapping could potentially manifest subtly, without disrupting the spatial distribution of the face representation. RSA identifies statistical (dis)similarities across activity patterns, providing a much more sensitive measure of representational changes<sup>68</sup>. To further explore any possible remapping in the deprived hand area of the amputees, we first looked at the pattern of dissimilarity for each face part relative to non-dominant/phantom thumb in the hand ROI for controls and amputees. Here we found a non-significant group x hemisphere x face-thumb interaction ( $F_{(3,245.0)} = 0.605$ , p = 0.613), as well as a non-significant group x hemisphere interaction ( $F_{(1,245.0)} = 0.552$ , p = 0.458; Fig. 6A). The pattern of facial activity in relation to the thumb is therefore statistically comparable between amputees and controls, indicating similar representational structure of the face, relative to the hand, between the two hemispheres and groups.

We also found a non-significant group x hemisphere x face-face interaction when looking at face-face pairwise dissimilarity in the hand ROI across all three groups ( $F_{(10,605.0)} = 0.640$ , p = 0.780; controlled for age and gender; Fig. 6B), suggesting a similar representational structure of the face across hemispheres and groups. However, when we looked at the average amount of facial information within the hand ROI, we did find a significant group x hemisphere interaction ( $F_{(2,605.0)} = 13.568$ , p < .001), indicating potential differences in facial information content in the hand area. Post-hoc comparisons (corrected alpha = 0.0125; uncorrected p-values

reported) exploring this effect reported significantly greater dissimilarity between facial-part representations in the deprived hemisphere of the amputees (M = 0.223; SE = 0.0223;  $t_{(605.0)} = -3.399$ , p < .001) and one-handers (M = 0.267; SE = 0.0177;  $t_{(605.0)} = -5.742$ , p < .001), when compared to their respective intact hemisphere (amputees: M = 0.184; SE = 0.0223; one-handers: M = 0.197; SE = 0.0177). When comparing to the controls non-dominant hemisphere, we only found significantly greater facial information in the one-hander's deprived hand area ( $t_{(69.4)} = -3.127$ , p = 0.003) and a non-significant effect for amputees ( $t_{(67.5)} = -1.490$ , p = 0.141). These results are in line with our univariate analyses, which demonstrate significant cortical remapping of facial parts in the one-handers group. It also suggests that there may be an interhemispheric increase in facial information in the deprived hand ROI of amputees.

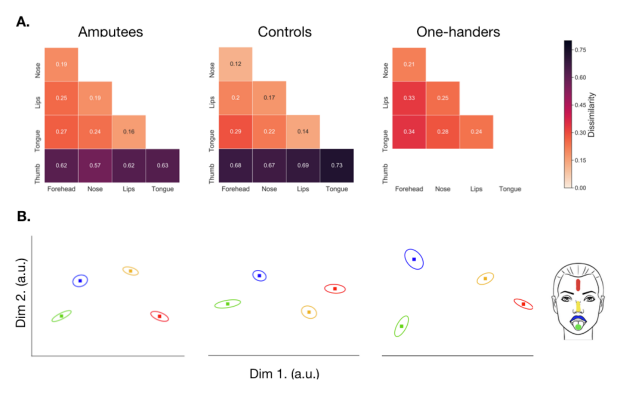


Figure 6. Representational Similarity Analysis (RSA) in the deprived/non-dominant hand area across all groups. A) Representational Dissimilarity Matrices (RDMs) for amputees (n = 15), controls (n = 22) and one-handers (n = 22). Greater dissimilarity between activity patterns for the chosen pairwise comparison indicates more information for that facial part within the hand area. Smaller dissimilarity values of facial (and thumb) activity patterns indicates a reduced ability to discriminate between the chosen movements in the hand area. Face-thumb distance values are only shown for the controls and amputees. B) Multi-dimensional scaling plots for each group, which projects the RDM distances into a lower-dimensional space. Here the distances between each marker reflects the dissimilarity, with more similar activity patterns represented closer together, and more distinct activity patterns positioned further away.

Forehead movements are plotted in red, with the nose in yellow, lips blue and tongue green, and the standard error is plotted around each data point. Please note, a different scale was used compared to the face ROI (Figure. 8).

For completion, we also looked at facial activity patterns (i.e., face-face pairwise dissimilarities) within the face ROI across all three groups. Here we found non-significant differences for a group x hemisphere x face-face interaction ( $F_{(10,605.0)} = 0.113$ , p = 1.000) and group x hemisphere ( $F_{(2,605.0)} = 0.701$ , p = 0.496), suggesting a similar representational pattern of facial activity, i.e., facial information content, across hemispheres and groups (Fig. 7). However, we did note a significant interaction between group x face-face ( $F_{(10,605.0)} = 2.025$ , p = 0.029), indicating that the overall pattern of facial organisation does vary between groups. As our post-hoc comparisons were not significant when correcting for multiple comparisons, we will not interpret this interaction further.

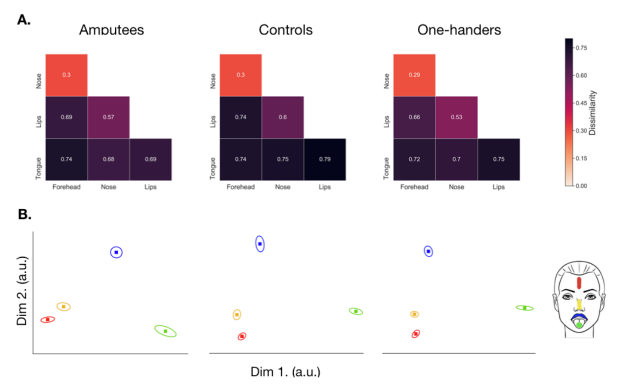


Figure 7. Representational Similarity Analysis (RSA) in the deprived/non-dominant face area across all groups. All annotations are as in Figure 6.

# **Discussion**

It is a well-accepted notion, rooted in non-human primate electrophysiological data, that upperlimb amputation triggers cortical remapping of the assumed neighbour – the lower face – into the missing hand area. This previous work predominantly characterised remapping by investigating shifts of the lip representation <sup>18–22,69,70</sup>. However, by focusing on only one face part, activity elicited by other facial parts (such as the forehead) in areas of interest in S1 (i.e., the hand-face border) are not taken into account (see<sup>2</sup>). Here we explored the relationship of face-to-hand remapping in controls and one-handed groups, and used both univariate (topographic) and multivariate (representational structure) methods to investigate in detail the information content of the face in both the deprived and intact hand and face areas. We found evidence for an upright somatotopy of the face across all groups, suggesting that the cortical neighbour to the hand in humans is the upper, not lower, face. We further found little evidence for remapping of all tested facial parts in amputees, with no significant relationship to the chronicity of their PLP. As a positive control, we also recruited individuals that were born without a hand (one-handers), who have previously shown cortical remapping across multiple body parts<sup>36–38</sup>. Across all facial parts, one-handers showed evidence for a complex pattern of face remapping in the deprived hand area, with consistent and converging evidence across analysis approaches. Together, our findings demonstrate that the face representation in humans is highly plastic, but that this plasticity is restricted by the developmental stage of input deprivation, rather than cortical proximity.

Firstly, our univariate analyses at both group and individual level confirmed an upright orientation of the face in controls, amputees and one-handers. Here we found that the forehead representation borders the hand area, followed by the lips, and tongue located more laterally. These results provide the first neuroimaging evidence of an upright representation of the face in humans. Indeed, while early and recent intracortical recordings and stimulation studies<sup>28,29,31,71</sup> emphasised an upright organisation, in agreement with our results, neuroimaging studies in humans provided contradictory evidence for the past 30 years. For instance, early studies reported an inverted facial somatotopy<sup>27,72</sup>, similar to the topography reported in primates<sup>11,12</sup>. Many subsequent studies then found no clear somatotopic organisation<sup>73–75</sup> or suggested an onion-like segmental topography<sup>76</sup>. These discrepancies and inconsistent neuroimaging findings may arise from the low-resolution of the techniques, small sample sizes and the challenge to find a robust and reliable method to stimulate face parts (and thus elicit detectable cortical activation). Our results provide converging evidence validating the upright orientation of the face, and indicate that the cortical neighbour to the hand is likely to be the upper face, which has important implications for cortical remapping theories based

on cortical proximity<sup>39</sup>. If cortical remapping of neighbours exists, we would therefore expect to see the forehead shifting towards and into the hand area – not the lips.

It may be argued that it is difficult to achieve isolated execution of specific facial muscles when performing gross movements without impacting sensory processing of neighbouring facial parts. For instance, tongue movements in our paradigm (e.g., touching the roof of the mouth with the tongue), may be best considered as a holistic inner mouth movement, and forehead movements may be best considered as engaging the upper face. While this limitation is valid, it may also be relevant (though to a smaller degree) for passive paradigms, as stimulation can induce waves that propagate through the skin<sup>77–79</sup> and Pacinian receptors were found to activate during stimulation of remote sites<sup>80,81</sup>. Despite this caveat, both our univariate and multivariate analyses showed that we were successful in isolating sensorimotor representations of the various movements (forehead, nose, lips and tongue) within our regions of interest, meaning that even if somatosensory information is overlapping across movements, there is still enough distinct information to separate representational patterns. This finding indicates the suitability of our motor paradigm for teasing apart facial somatotopy, allowing us to characterise the face in greater detail than previously attempted.

When looking at face-to-hand remapping in amputees, where the remapping of cortical neighbours has been the prevalent explanation for PLP, we find little evidence of shifts of locality and remapping in the deprived hand area for facial parts, including the neighbour (forehead) and hypothesised neighbour (lips). Our univariate results are further supported by our multivariate analysis, where we find no significant changes in the relationship between face-to-thumb activity in the deprived hand area. These results support previous work reporting a lack of cortical remapping after amputation<sup>35,42,82</sup>, suggesting that in amputees this area might be functionally unipotent – pertaining to hand-related activity alone and lacking the ability to rescope after hand loss. However, due to the inconclusive Bayes Factor in our key analyses, we cannot strongly conclude that remapping does not occur in this group. This could be attributed to our relatively small sample (further recruitment was prevented due to Covid-19 restrictions), and in particular, the small proportion of amputees experiencing PLP (9 out of 15). However, pain is not a necessary condition for deprivation-triggered remapping<sup>83–85</sup> and vice versa, PLP can be experienced in absence of remapping<sup>86</sup>. Moreover, previous studies reporting significant difference between amputees who experienced PLP and those who do not,

often employed a similar sample size<sup>20</sup>, indicating that the expected effect of remapping should be substantial.

We did find anecdotal evidence for remapping for the tongue within the deprived hand area in amputees, which did demonstrate a weak trend towards those with greater chronicity of PLP displaying greater tongue remapping. This was a surprising result, as the tongue is not a cortical neighbour to the hand, and was not specifically hypothesised to remap in amputees. We also found that amputees demonstrated a greater amount of facial information in the deprived hand area when compared to their intact hemisphere. Although this multivariate result was not significantly different to that of controls, it demonstrates the plausibility that cortical remapping in amputees may exist to a certain degree (e.g., an inter-hemisphere imbalance), albeit any relationship to PLP is tenuous. While these latter results require further validation, they support our premise that cortical proximity of representations may not be a necessity for remapping to occur. In this context, as our tongue condition could also be classed as an 'inner mouth' movement, it is important to note that previous work addressing sensorimotor representations of the mouth and the larynx have demonstrated both lateral and more medial 'hotspots' (i.e., a 'double' representation<sup>87</sup>). The potential tongue remapping in amputees, therefore, may reflect changes in the medial mouth representation, but this would need to be investigated further by separating the relative contributions of the tongue and larynx.

We did find converging and conclusive evidence for cortical remapping of all facial parts (neighbours and non-neighbours) in our (congenital) one-hander group. Here the pattern of remapping is strikingly different to that of cortical neighbourhood theories. Specifically, the location of the cortical neighbour – the forehead – is shown to shift away from the deprived hand area, which is subsequently 'taken over' by the lips and the tongue. The increase of facial activity in the deprived hand area is in turn supported by our multivariate results, whereby significantly greater information content for the face was found in the deprived hand area for one-handers when compared to controls. One-handers' deprived hand area, therefore, seems to have increased discriminability between different facial movements. It is difficult to ascertain from our study the drivers of this remapping. It has been suggested previously that remapping within this group may be driven by functionally-relevant behaviour substituting the loss of the limb<sup>36,38</sup>. Alternative explanations relate to an overall and unspecific release of inhibition (i.e., decreased GABA) in the missing hand area, allowing for latent activity of other body parts to be detected<sup>36</sup>. While speculative, our results tend to support the former, as we report remapping

for facial parts which have the ability to compensate for hand function, e.g., using the lips and/or mouth to manipulate an object, and a lack of remapping for those that cannot (the forehead). This increased activity from body parts compensating for hand function may represent a stabilising mechanism, aimed at preserving the integrity of the sensorimotor network and its function<sup>2</sup>. The deprived hand area in one-handers, therefore, may be deemed pluripotent – suitable for adapting to multiple body parts<sup>38</sup>, which may preserve the role of the hand area by sustaining its hand-function related information content.

A limitation that should be acknowledged arises from the potential contribution to S1 from M1 activity. Since these cortical areas are neighbours, it is difficult to separate them with certainty. We tried to minimise the contribution of M1 by taking multiple acquisition and pre-processing steps, including the use of anatomical delineation at the individual level, as well as a comprehensive analytical approach (e.g., both univariate and multivariate techniques). Across the board we find robust evidence for remapping in congenital one-handers and no reliable evidence for similar remapping in amputees. Furthermore, it has been claimed that active movements may produce different cortical maps to those with passive stimulation<sup>88,89</sup>, and previous work demonstrating a relationship between cortical remapping and PLP tended to use passive stimulation<sup>18,19,22,69</sup>. However, we do not think this methodological difference underlies our contrasting results as movement-induced lip activity has been shown to demonstrate lip remapping before<sup>20,70,90</sup>, indicating that an active paradigm is suitable for demonstrating cortical remapping (if it exists). Conversely, a recent study using passive lip stimulation in amputees did not find any evidence for remapping<sup>91</sup>. Moreover, we recently ran a study which found that S1 topography and multivariate representational structure are similar across active and passive paradigms<sup>92</sup>, and so our choice of active paradigm is suitable to identify topographic organisation and remapping (as we clearly find in one-handers). Despite these reservations, the choice of an active paradigm is the most accessible and translatable to a fMRI design, as well as the most reflective of naturalistic tactile inputs in everyday life.

Our findings seem contradictory to the many previous studies reporting lip remapping in amputees<sup>18–22,69,70</sup>. A major difference with regards to these previous studies, which predominantly focused on a single part of the face, lies in the fact that our study was the first to assess the mapping and potential remapping of multiple facial parts at once. By focusing on the lips only, previous designs excluded other facial parts which may have elicited greater activity in certain areas of S1, resulting in a less accurate delineation of the lip-selective

representation. Such down-sampling of body maps, therefore, can lead to biased results and interpretation<sup>2</sup>. While our design is not exempt from this limitation, the fact that we assessed other parts of the face may explain why our results diverged from previous findings.

To conclude, our use of both univariate and multivariate analyses found consistent evidence for a complex pattern of face remapping in congenital one-handers, in line with the theory suggesting remapping in this group reflects compensatory behaviour<sup>2</sup>. This is in contrast to amputees, where we find little evidence for cortical remapping, indicating a stability of both the hand and face representation after limb loss. By and large, remapping measures were not linked to PLP. Our results call for a reassessment of traditional remapping theories based on cortical proximity, and future research into potential remapping of the inner mouth representation after limb loss.

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