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Investigating the stability of fine-grain digit somatotopy in individual human participants

Abbreviated title: Stable fine-grain somatotopy in human SI.

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Abstract

Studies of human somatosensory cortex have placed a strong emphasis on the cortical representation of the hand and the propensity for plasticity therein. Despite many reports of group differences and experience-dependent changes in cortical digit somatotopy, relatively little work has considered the variability of these maps across individuals, and to what extent this detailed functional architecture is dynamic over time. With the advent of 7-tesla fMRI, it is increasingly feasible to map such detailed organisation non-invasively in individual human participants. Here we extend the ability of ultra-high field imaging beyond a technological proof of principle to investigate the inter-subject variability of digit somatotopy across participants, and the stability of this organisation across a range of intervals. Using a well-validated phase-encoding paradigm and an active task, we demonstrate the presence of highly reproducible maps of individual digits in SI, sharply contrasted by a striking degree of inter-subject variability in the shape, extent and relative position of individual digit representations. Our results demonstrate the presence of very stable fine-grain somatotopy of the digits in human SI, and raise the issue of population variability in such detailed functional architecture of the human brain. These findings have implications for the study of detailed sensorimotor plasticity in the context of both learning and pathological dysfunction. The simple task and 10-minute scan required to derive these maps also raises the potential for this paradigm as a tool in the clinical setting.
We apply ultra-high resolution fMRI at 7 tesla to map sensory digit representations in the human cortex at the level of individual participants across multiple time points. The resulting fine-grain maps of individual digits in somatosensory cortex reveal both the stability in this fine-grain functional organization over time, contrasted with the variability in these maps across individuals.
The somatotopic organization of primary somatosensory cortex is well established in the human brain, both at the level of whole-body topography (Penfield and Boldrey, 1937; Walter et al., 1992; Zeharia et al., 2015) and the more fine grain organisation in the representations of the face and the hand (Moulton et al., 2009; Sanchez-Panchuelo et al., 2010). The somatotopic digit map is the subject of continuing interest, with its relative cortical overrepresentation (Mountcastle, 2005). More generally, it is increasingly clear that SI plays a critical role in motor function (Vidoni et al., 2010; Platz et al., 2012; Jacobs et al., 2014).

Studies of human somatotopy have focused considerable effort on attempting to map the representations of digits in the cortex (Baumgartner et al., 1991; Gelnar et al., 1998; Kurth et al., 1998; Francis et al., 2000; Overduin and Servos, 2004; Nelson and Chen, 2008; Schweizer et al., 2008) and the cerebellum (van der Zwaag et al., 2013). Further work has provided evidence for marked topographical differences in the cortical spacing and organisation of SI somatotopy in specific sub-populations, for example in musicians and individuals with focal dystonia (Bara-Jimenez et al., 1998; Elbert et al., 1998; Meunier et al., 2001; Butterworth et al., 2003; Nelson et al., 2009). Other studies, chiefly using MEG, report a propensity for experience-dependent plasticity in SI somatotopy (Braun et al., 2000; Schwenkreis et al., 2001; Candia et al., 2003; Stavrinou et al., 2007; Vidyasagar et al., 2014), building upon seminal studies undertaken in non-human primates (Allard et al., 1991; Zarzecki et al., 1993). However the spatial
resolution of both MRI and MEG is typically insufficient to make a strong
argument about fine-grain digit somatotopy in the somatosensory hand area.

With the increasing prevalence of 7-tesla MRI scanners, it is now possible to
resolve SI representations of all of the digits in the hand at the level of individual
participants. A number of elegant studies at ultra-high field have used tactile
stimulation to demonstrate not only the ability to map digit somatotopy in SI
with a number of paradigms, but also the existence of within-digit somatotopy
and cortical overlap between adjacent digit pairs (Sanchez-Panchuelo et al.,
2010; 2012; Besle et al., 2013a; 2013b)

Reports of SI digit maps to date have showcased the novel capabilities of ultra-
high field fMRI (Sanchez-Panchuelo et al., 2012; Besle et al., 2013a; 2013b;
Martuzzi et al., 2014; Stringer et al., 2014). In light of the considerable interest in
cortical digit maps, and specifically their capacity for plasticity, more thorough
cross-sectional and longitudinal analyses are necessary.

While non-human primate data demonstrates considerable inter-subject
variability in SI digit somatotopy (Merzenich et al., 1987), little evidence exists to
demonstrate stability of the shape and position of SI digit maps over time.
Human studies to date have only considered the reproducibility of isolated
individual digit representations, or used relatively crude measurements (e.g.
centre of mass) (Vidyasagar and Parkes, 2011; Martuzzi et al., 2014). It therefore
remains unclear to what extent reports of inter-subject variance in primates
could actually reflect intra-individual instability in digit representations in SI. In
order to meaningfully interpret previous reports of use-dependent plasticity and
group variability in human cortical digit representations, it is vital to develop a
more thorough understanding of SI digit somatotopy in the healthy population.

Here we address this fundamental gap in the literature, using 7T fMRI mapping
to explore the fine grain functional organization of SI at the level of individual
human participants. More specifically, we apply a phase-encoding paradigm
well-validated for sensory body mapping (Sereno and Huang, 2006; Orlov et al.,
2010; Sanchez-Panchuelo et al., 2010; Mancini et al., 2012; Zeharia et al., 2015)
to investigate whether stable and reproducible maps of individual digits exist in
human SI.

**MATERIALS AND METHODS**

**Participants**

Thirteen healthy control participants [Table 1; Mean age: 28.6 ± 5.66; six female]
were recruited in accordance with local central university research ethics
committee approval (University of Oxford; MSD-IDREC-C2-2013-05). All
participants were right handed according to the Edinburgh Handedness
Inventory (Oldfield, 1971).

**Experimental design**

Participants attended three scan sessions. Two of the sessions were separated by
a period of 24 hours (0 hours and +24 hours). The third session took place four
weeks before or after the other two sessions. During each session participants
underwent a one-hour fMRI scan. One of the sessions also involved an additional scan to acquire a structural image.

**MRI acquisition**

Functional MRI data were acquired using a Siemens 7T Magnetom system with a 32-channel head coil. An initial functional localiser scan was used to identify hand movement-related activity in order to aid slice placement for subsequent task fMRI scans (Multislice gradient echo EPI, TR: 3000ms, TE: 25ms, flip angle: 90°, bandwidth: 1568Hz, 43 axial slices, 2x2x2mm resolution, GRAPPA factor = 2).

Task fMRI data were then acquired using a field of view based on the results of the functional localizer; true axial slices were centred on the hand knob activation in the z-axis of the left hemisphere (Multislice gradient echo EPI, TR: 1500ms, TE: 25ms, flip angle: 90°, bandwidth: 1562Hz, 22 axial slices, 1.2x1.2x1.2mm resolution, GRAPPA factor = 2).

For image registration purposes, single volume high-saturation EPI images were acquired: one whole brain image and one partial field of view (FOV) image with the same slice placement as the task fMRI. T1-weighted Multi-Echo Magnetization Prepared Rapid Acquisition Gradient Echo (MEMPRAGE) structural scans were acquired during one of the three sessions (van der Kouwe et al., 2008) using a 3T Siemens Trio system (TR: 2530ms, TE: 1.69, 3.55, 5.41 and 7.27ms, 1x1x1 mm, GRAPPA factor = 2).

**fMRI tasks**
Participants performed a series of tasks involving visually-cued movements of individual digits in the scanner: digit 2 (D2: index finger), digit 3 (D3: middle finger), digit 4 (D4: ring finger) and digit 5 (D5: little finger). An active motor task was selected to optimally activate a range of inputs to the cortical somatosensory system, analogous to daily use of the hand. Movement recruits a combination of peripheral receptors encoding a range of somaesthetic modalities, from surface mechanoreceptors to deeper cutaneous receptors and proprioceptors, as well as efference information from the motor system.

During the three minute functional localiser scan, participants were instructed to appose their right thumb with each of the digits of their right hand sequentially during 15 second movement blocks, contrasted with equivalent periods of rest. All subsequent task fMRI involved individual movements of D2, D3, D4 and D5 in the form of button presses using an MRI-compatible four-finger button-box (manufactured in-house) resting on the participant’s right thigh during the scan. Participants were presented with four white circles, corresponding to the four digits of the right hand, shown on a visual display projected into the scanner bore. The displayed circles flashed individually at a constant frequency to cue participants to make button presses at the specified rate. Further discussion of the caveats associated with using an active motor task in the study of somatosensory cortex are outlined in Limitations below.

A phase-encoding task was used, which involved continuous button presses with no rest periods (Figure 1A). The task consisted of movement blocks of 8 seconds in duration, during which participants moved one digit (D2-D5) at a rate of 1 Hz.
The phase-encoding forward task cycled through blocks of D2, D3, D4 and D5 in a repeating sequence (8 repetitions of the cycle; Figure 1A). The phase-encoding backwards task cycled through blocks of D5, D4, D3 and D2 in a repeating sequence (8 repetitions of the cycle). The total duration of the phase-encoding task was 8 mins 50 secs. The activation maps derived from the phase-encoding forward and backward tasks were averaged voxel-wise; further details below.

A standard block task was also undertaken, which involved movement blocks and rest blocks, both 12 seconds in duration. A total of four movement blocks were acquired per digit (16 movement blocks total), in a counterbalanced order, randomized across visits. During movement blocks, participants were instructed to perform movements of a specific digit at 1 Hz (e.g. D2, D2, D2, D2 . . . ). Each movement block was separated by a rest block. The total duration of the block task was 6 mins 24 secs.

**MRI analysis**

MRI analysis was undertaken using tools from FSL and Connectome Workbench (Smith et al., 2004; Woolrich et al., 2009; Marcus et al., 2011; Jenkinson et al., 2012). MRI data were projected to cortical surface reconstructions produced with FreeSurfer T1-weighted MEMPRAGE images (Dale et al., 1999; Fischl et al., 2001).

**MRI preprocessing**

All fMRI data were subject to the following preprocessing steps: motion correction using MCFLIRT (Jenkinson et al., 2002); removal of non-brain tissue
using the Brain Extraction Tool (BET) (Smith, 2002), high pass temporal filtering (Gaussian-weighted least squares straight line fitting with sigma = 100 seconds) and spatial smoothing using a Gaussian kernel of FWHM 1.5 mm.

Image registration

Image registration was undertaken within participant using FLIRT (Jenkinson and Smith, 2001; Jenkinson et al., 2002) and Freesurfer’s Freeview. Task fMRI data from the three scan sessions were first registered to a partial-FOV high-saturation EPI image acquired during an additional scan session to avoid biasing any single time point (6 degrees of freedom, normalized correlation cost function). The partial-FOV high-saturation EPI image was then registered to the T1-weighted MEMPRAGE image using boundary-based registration (BBR) (Degrees of freedom: 6, FMRIB’s Automated Segmentation Tool (FAST) white matter segmentation, no search) (Greve and Fischl, 2009), initialised with an affine registration matrix. The results of the BBR were used as a starting point for manual alignment of the single volume partial-FOV high-saturation EPI image to the structural MEMPRAGE white matter and pial surfaces using blink comparison as implemented in Freeview, an approach applied previously in studies of fine-grain topography (Mancini et al., 2012).

Phase-encoding analysis

The phase-encoding task fMRI data were analysed using a cross-correlation approach previously applied in retinotopy, and more recently for body mapping (Engel et al., 1997; Wandell et al., 2007; Orlov et al., 2010); see Limitations for further discussion. This analysis used cross-correlation to find the time point in
the phase-encoding forward (D2-D5) and phase-encoding backwards (D5-D2) tasks at which each cortical voxel responded maximally.

To achieve this, the preprocessed BOLD EPI data were correlated against a series of reference models. The model was composed of a gamma-convolved boxcar: 8 second ‘on’ and 24 second ‘off’, repeated eight times, mirroring the eight 32-second cycles of the phase forward and phase backward tasks (Figure 1B: black). The model was shifted in time iteratively by a number of lags so that activity throughout the cycle could be modeled (Figure 1B). This approach increases sensitivity to track a wave of activation (Engel, 2012), in this case associated with the cycles of movement which progress either from D2-D5, or D5-D2. A correlation was calculated between the raw BOLD signal of each voxel (Figure 1B: red) and the reference model at each lag (Figure 1B: black). Each iteration shifted the model by a given lag (1.5 seconds). With each lag the 8 second ‘on’ of each 32 second cycle was time shifted (e.g. model 1: 8s on/24s off; model 2: 1.5s off/8s on/22.5s off; model 3: 3s off/8s on/21s off ...), with sufficient shifts to cover one 32-second cycle. By plotting for each voxel the cross-correlation at each voxel as a function of the lag, a tuning curve was created for each voxel, demonstrating the optimal model fit for that voxel (Figure 1C). Each lag was assigned to a given digit in the cycle. Voxels responsive to a particular digit demonstrate a peak cross-correlation within the lags corresponding to movement that digit in the cycle.

For each participant and session, the $r$-values resulting from the cross-correlation analysis specified above (Figure 1C) were averaged across the lags.
assigned to the same digit to yield digit maps (D2, D3, D4, D5) for each of the
phase-encoding forward and phase-encoding backwards tasks. For each
participant and each session, the maps for each digit from the phase-encoding
forward and phase-encoding backward tasks were resampled to the single
volume partial FOV high-saturation EPI space and averaged to give a single
voxel-wise r-value map for each session and for each digit (Figure 1D). For each
participant, session and digit, a corresponding voxel-wise z-statistic map was
calculated on the basis of the distribution of values within the brain tissue of the
FOV for which BOLD EPI data were acquired. These soft-edged maps were
further masked using a winner-take-all approach to produce digit maps in which
each voxel was assigned exclusively to one digit.

Block task analysis
The block task fMRI data were analysed using a GLM in the FMRIB Expert
Analysis Tool (FEAT). All analysis was undertaken on the single-participant level,
using FMRIB’s Improved Linear Model (FILM) to estimate timeseries
autocorrelation and pre-whiten each voxel. Each digit was modeled with a
separate boxcar regressor with gamma-HRF convolution and its temporal
derivative, giving a total of eight regressors. FEAT was used to produce
activation maps corresponding to each of the four digits by contrasting a given
digit regressor to the rest blocks (e.g. D2 > rest).

Surface projection
Phase-encoding winner-take-all z-statistic digit maps were projected to two-
dimensional surface space using a cortical ribbon mapping method implemented
in Connectome Workbench. This approach estimates the contribution of multiple voxels to one point on the cortical surface and weights the values therein accordingly.

For visualization on the cortical surface of individual participants, winner-take-all z-statistic maps for each digit and time point were thresholded using FDR to determine a corrected \( p \)-value threshold on the basis of the observed \( p \)-value distribution within the data (\( \alpha=0.001 \)) (Genovese et al., 2002). This resulted in individually defined FDR thresholds for each map under consideration (Table 2).

For inter-subject comparison, volumetric winner-take-all z-statistic maps were resampled into atlas space using Combined Volumetric and Surface (CVS) registration to achieve accurate and robust alignment with the CVS atlas (Postelnicu et al., 2009), and projected to the atlas two-dimensional surface using the cortical-ribbon mapping method.

**Intra-subject reproducibility**

The Dice coefficient (Dice, 1945) was used to assess the reproducibility of phase-encoding digit maps over time, quantifying the spatial similarity of digit map areas. The Dice coefficient varies from 0 (no overlap between digit maps) to 1 (perfect overlap between digit representations). For each digit, the winner-takes-all digit maps for that digit were thresholded (FDR \( \alpha=0.001 \)) and overlap was calculated between each possible pairing across each of the three time intervals. Where A and B are the area of the two digit representations, the Dice Coefficient is expressed as:
Inter-subject variability

The variability in the spatial location of individual digit representations across participants was assessed using the Dice coefficient. Surface-area based thresholding was applied to the winner takes digit maps, such that the maximal 80mm² of activation within SI for a specific digit was considered. Digit representations with this surface area were present within S1 for every participant and time point. The interpretability of a Dice coefficient (Equation 1) calculated across different subjects could be affected by inter-subject differences in the size of digit representations. The use of a fixed surface area for each digit representation excluded any effect of inter-subject variability in the spatial size of digit maps on inter-subject comparisons. Each winner-takes-all digit representation at time point 0 hour was compared with each other winner-takes-all digit representation at time point +/- 4 weeks across all participants.

Dice coefficients were used to construct a large inter-subject Dice comparison matrix (36 x 36 cells), composed of submatrices (9 x 9 cells) for all possible digit pairings. To compare digit representation overlap intra-subject (submatrix diagonal), to digit representation overlap inter-subject (submatrix off-diagonals), measures of matrix dominance ratio (Mdr) were calculated for each submatrix (Greene and Cunningham, 2006). The Mdr of a square matrix \( K \), of width and height \( n \) can be expressed as:

\[
Mdr = \frac{2 \times |A \cap B|}{|A| + |B|}
\]
Values of Mdr greater than one would therefore be observed in cases where the average Dice overlap of two digit representations within the same subjects (the matrix diagonal) is of greater magnitude than the average Dice overlap made across different subjects (the matrix off-diagonals).

A higher-level matrix of the Mdr values for each digit pairing was constructed, from which an overall Mdr value was calculated. A high value of overall Mdr in this matrix would suggest high matrix dominance in comparisons of 'same' digits (e.g. D2-D2, D3-D3...) and low levels of matrix dominance in comparisons on 'different' digits (e.g. D2-D3, D3-D5 etc.). This is turn would support the hypothesis that intra-subject overlap in 'same' digits is greater than inter-subject overlap in 'same' digits.

Bootstrap resampling was applied to the large inter-subject Dice comparison matrix (36 x 36 cells) in order to quantify the likelihood of observing the reported pattern by chance, and therefore the statistical significance of the overall Mdr value.

Additional measures of reproducibility and variability

As well as the primary Dice measures of intra-subject reproducibility and intra-subject variability, additional features of the phase-encoding digit maps were
assessed. At each time point soft-edged phase-encoding maps (FDR $\alpha=0.01$) were used to assess the amount of overlap between different digit representations within S1. These measures sought to characterize the extent of shared cortical territory between different digit representations, which the winner-takes-all maps do not capture, and to assess the consistency in the extent of this overlap at each time point. The extent of this overlapping shared territory was expressed as a Dice coefficient. The pattern of overlap in these soft-edged digit representations was represented in 4x4 matrices for each participant and each time point. The pattern in these matrices were compared using a ranked Mantel test (Mantel, 1967), in order to quantify both the intra-subject consistency in the overlap pattern, and the inter-subject variability therein.

In a complementary analysis, peak z-stat coordinates for each digit were calculated with SI on the inflated cortical surface, allowing for the calculation of geodesic distances between adjacent digits for each time point (D2-D3, D3-D4, D4-D5), which were again assessed for consistency.

**Identifying additional digit maps**

To increase statistical power in order to identify further somatotopic digit maps previously reported in SI (Pons et al., 1985; Huffman and Krubitzer, 2001; Yau et al., 2013), an additional all-session phase-encoding map was produced by co-registering and averaging the forward and backward lags from all three sessions, before processing the resulting maps as outlined above, using a more liberal FDR threshold ($\alpha=0.01$).
Statistics

All statistical analysis and graphing were undertaken using JMP (Version 11.0, SAS Institute, Cary, NC, USA) and Statistics Package for the Social Sciences (SPSS, Version 19.0, IBM Corporation, Armork, NY, USA).

RESULTS

All BOLD EPI data were assessed for excessive motion both visually and using motion estimate outputs from MCFLIRT: data from three participants exhibited visible spin history motion artifact as a result of sharp motion during one or more scan sessions (greater than 1mm of absolute mean displacement in fewer than five volumes; Table 1); these participants were excluded. Further analysis found no significant or systematic correlation between the task design and motion parameters in the remaining participants. One further participant was excluded on the grounds of an incidental finding. Nine participants were considered in further analysis [Table 1: Participants 1-9; Mean age: 28.5 ± 6.54; four female; -4 weeks: four participants; +4 weeks: five participants].

Phase-encoding digit maps in SI

The thresholded activation maps from the phase-encoding analysis displayed a clear and specific pattern in the left post-central gyrus around the anatomically characteristic hand knob (Figure 2A/D) (Yousry et al., 1997). Maps showed a pattern of progression from the lateral-most representation of digit two, to the medial most representation of digit five (Figure 2C/D), a pattern that was consistent across all participants and time points (Figure 3/4). However, the maps showed striking qualitative differences in shape and orientation across
participants, in keeping with reports of inter-subject variability from the non-
human primate literature (Merzenich et al., 1987).

Activation was isolated to the primary somatosensory cortex on the postcentral
gyrus, with minimal extraneous activation within the FOV in which BOLD EPI
data were acquired (Figure 2C); no mask or ROI has been applied to any digit
maps presented herein. The Brodmann areas that constitute the primary
somatosensory cortex cannot be defined accurately on the basis of gross
anatomy alone. However, the location of the observed digit representations is
broadly anatomically consistent with the location of Brodmann area 3b: on the
posterior bank of the central sulcus, posterior to area 3a in the nadir and
anterior to area 1 at the apex of the post-central gyrus. Some subjects displayed
partial additional maps more posteriorly in regions consistent with Brodmann
area 2 or 1 (Figure 3/4: participants 3 and 9).

A post-hoc region of interest (ROI) analysis was used to explore the BOLD signal
underlying the phase-encoding digit maps. The average BOLD signal time course
was extracted within each digit representation (Z >3.5) (Figure 2B). These
showed clear and specific activation patterns in a sequence consistent with the
phase-encoding task digit order.

**Intra-subject digit map reproducibility over 24 hours and four weeks**

Surface-projections of digit maps derived from phase-encoding analysis
qualitatively display a striking degree of reproducibility of these fine-grain maps
at the single-participant level across both the 24 hour and four week map-remap
intervals (Figure 3/4). To quantitatively assess the intra-individual reproducibility of the phase-encoding derived digit maps, Dice similarity coefficients were used to compare the spatial extent of digit representations across sessions.

A Dice coefficient was calculated between every possible digit pairing and every possible time point pairing (Figure 5A) within a FreeSurfer anatomically defined ROI of SI overlapping the fMRI acquisition volume. This analysis demonstrated a very high degree of spatial concordance between ‘same’ digit representations across all time intervals.

The reproducibility matrices were averaged across the two different time intervals and further averaged into three digit pairing categories: homologous digits, first order neighbours and second/third order neighbours (Figure 5A; part iv) in order to assess whether the Dice coefficient for homologous digits was significantly greater than the equivalent value between non-homologous digit pairings. A one-way repeated measures ANOVA was performed with one factor of digit pairing category. There was a significant main effect of digit pairing category on the Dice coefficient: $F(2, 19) = 119.429$, $p < .0005$. Post hoc analysis with a Bonferroni adjustment revealed that the Dice coefficient was significantly greater for homologous digit pairings (Average Dice coefficient: 0.542, 95% CI: 0.380 to 0.584) compared with pairings of first order neighbours (Average Dice coefficient: 0.010, 95% CI: 0.000 to 0.020), and pairings of second and third order neighbours (Average Dice coefficient: 0.001, 95% CI: 0.000 to 0.002); all $p < .0005$ (Bonferroni-adjusted). The same pattern of results was also seen using
an equivalent analysis approach on volumetric data rather than surface
projected data.

**Inter-subject digit map variability**

Consistent with qualitative observations (Figure 3/4), Dice analysis comparing
the spatial location of individual digit representations (Figure 5B; part i)
demonstrates a considerable degree of inter-subject variability when compared
to the consistency seen intra-subject. In comparison of ‘same’ digits over time
(e.g. D2-D2, D3-D3) the degree of overlap observed intra-subject exceeds that
observed inter-subject, resulting in values of Mdr >1. This is consistent with the
notion of variability in the spatial position of individual digit representations. In
contrast, for ‘different’ comparisons (e.g. D2-D4), the degree of overlap observed
inter-subject exceeds the degree of overlap observed intra-subject, yielding
values of Mdr <1. This further strengthens our claim that intra-subject
consistency is driven by reproducibility of the spatial patterns for the same digits
over time, rather than other irrelevant aspects of the map (e.g. geometrical
cluster characteristics).

Values of Mdr for each digit comparison are summarized in Figure 5B(ii). From
this matrix a value of overall Mdr was calculated at 19.67 (Equation 2; Average
Mdr for ‘same’ digit pairings / Average Mdr for ‘different’ digit pairings). In order
to substantiate the observed pattern and value of overall Mdr yielded, we applied
bootstrap resampling to the inter-subject Dice comparison matrix (50,000
iterations) to account for the likelihood of observing this value by chance (Figure
5B; part iii), yielding $p < .0005.$
Overall the observed pattern of inter vs. intra-subject Dice overlap provided strong evidence supporting the presence of considerable variability in the spatial distribution of individual digit representations across participants, contrasted with consistency within participants over time.

Additional features of cortical digit maps

Measures of shared cortical territory between different digit representations were calculated at each time point (Figure 6). The inter-subject average matrix reveals previously established features of SI digit representations, with higher overlap in digit pairs such as D4 and D5 and low overlap between D2 and D3 (Figure 6A), consistent with patterns of daily usage (Ejaz et al., 2015). The similarity of cortical overlap patterns for each participant and time point (Figure 6B) was assessed using a ranked Mantel test. An intra-subject value was derived for each participant from the average of matrix comparisons within subject but over time. An inter-subject value was derived for each participant from the average of matrix comparisons between that subject and all other subjects for a given time point; this was repeated for each time point and the results were averaged. Comparison of the intra- versus inter-subject Mantel test values revealed greater similarity of values within a given subject compared with across different subjects: paired sample t-test, t(8)= -7.17, p<.0005.

Measures of peak-to-peak distance for adjacent digit representations were calculated at each time point. These measurements are provided in full in table 3. The intra-subject consistency in these measured was quantified using Cronbach’s
\[\alpha,\text{ which returned the following values: } D2-D3: 0.9714, D3-D4: 0.8526, \text{ and } D4-D5: 0.8422. \text{ These measures support a high degree of consistency across the observed digit maps over time.}\]

**Multiple digit maps across SI**

To reveal additional maps previously reported in SI with weaker digit selectivity (Kaas et al., 1979; Pons et al., 1985; Huffman and Krubitzer, 2001), an all-session average phase-encoding map was produced for each participant and resampled into a common space (FDR thresholding, \(\alpha=0.01\)). Additional maps were seen in a subset of participants. A more anterior map was observed in some individuals (Figure 7B,C,E,F), and a more posterior map (Figure 7G,H) in others, both within SI (see Discussion for further information regarding SI subdivisions).

**Strong concordance between phase-encoding and block design activation**

In order to validate finger selectivity identified using the phase-encoding task, concordance with independently derived sets of digit map data from a GLM analysis of the block task was assessed. The normalised beta values from the block task GLM contrasts comparing each digit to rest (e.g. D2 > rest) were extracted at the peak voxel of each phase-encoding derived digit representation (Figure 8). These values were averaged for each digit across the three scan sessions for each participant.

A two-way repeated measures ANOVA was performed to assess the agreement of the two mapping methods, with one factor of phase-encoding digit representation (D2-D5) and one factor of block design digit representation (D2-
There was a significant interaction between phase-encoding digit representation and block design digit representation on the normalised beta value: $F(9,72) = 69.15, p < .001$, sphericity assumed. Post-hoc paired samples $t$-tests demonstrated a significantly stronger relationship between the phase-encoding and block design digit representations for ‘same’ digits compared with ‘different’ digits (all $p < .0005$, uncorrected). For example, the peak voxel for phase-encoding D2 has larger beta value for the D2 > rest block task contrast compared with other digit contrasts from the block design (e.g. the D4 > rest GLM contrast). These results indicate an agreement between the two mapping methods: the peak voxel from the phase-encoding-derived map of a given digit shows a maximal normalised beta value for the GLM contrast specific to the same digit.

**DISCUSSION**

In the present study we report highly reproducible maps of fine grain digit somatotopy in SI at the level of individual participants, as demonstrated in all nine participants under study (Figure 3/4). These maps were reproducible across up to a four-week interval (Figure 5A). The consistency across this interval is particularly striking given the minimally supervised and easily implemented motor task used in this study. Unlike previously reported passive sensory stimulation paradigms used in digit mapping (Huang and Sereno, 2007; Sanchez-Panchuelo et al., 2010; Martuzzi et al., 2014; Stringer et al., 2014), the motor task applied here is more akin to everyday use of the hand.
The map reproducibility observed within individuals was sharply contrasted by a high degree of spatial variability in these maps across different participants. Despite a common ordering and progression of digits along the central sulcus (Figure 3/4), the shape and relative position of these representations differed, as has been shown previously in primates (Merzenich et al., 1987). We demonstrate the existence of considerable inter-subject variability in the spatial distribution of individual digit representations (Figure 5B; Table 3). Taken together these results robustly demonstrate the presence of very stable fine-grain somatotopy of the four digits under study in human SI, but also highlight the population variability in such detailed functional architecture in the human brain.

Digit maps in Brodmann area 3b and beyond

Using the FDR threshold applied here (α=0.001), the maps observed at each time point across all participants under study were located in a region anatomically consistent with Brodmann area 3b (Figure 3/4). The presence of well-delineated maps in Brodmann area 3b is well described in microelectrode mapping studies of individual digits in non-human primates (Kaas et al., 1979; Merzenich et al., 1987). The winner-takes-all phase-encoding approach applied herein is well suited to revealing such regions of high digit selectivity. This strong digit selectivity was an important feature in being able to address the question of consistency in such fine-grain cortical organization.

Digit maps in Brodmann area 3a, 1 and 2 show more limited digit selectivity (Kaas et al., 1979; Pons and Kaas, 1986; Huffman and Krubitzer, 2001). Although it was not possible to resolve evidence for these maps at each time point (Figure
3/4), pooling phase-encoding data across the three time points under study to produce an all-session average and using a more liberal FDR threshold provided further insight (Figure 7). Some individual participants displayed very clear smaller maps anterior and posterior to that presumed to be area 3b. These maps are potentially consistent with Brodmann area 3a and Brodmann area 1/2 respectively.

Since our data does not allow us to reliably define the constituent Brodmann areas in S1 at the level of individual participants, we are unable to discuss the position of these additional maps. While atlases do provide Brodmann area boundaries, these vary considerably across individuals and accurate definition would rely on cytoarchitectural information rather than gross anatomy (Zilles and Amunts, 2010). Indeed certain subdivisions (3a/4) are challenging to definitely delineate even on the basis of cytoarchitecture (Mountcastle, 2005).

Digit map reproducibility and variability

In this study we demonstrate, both qualitatively (Figure 3/4) and quantitatively (Figure 5A), a strikingly high degree of reproducibility in digit somatotopy. Previous work at 7-tesla has reported measures of digit map reproducibility either only across different runs within a single scanning session (Stringer et al., 2011) or consistency in the relatively crude measure of centre of mass location of digit representations in subjects scanned on two occasions with variable intervals between them (Martuzzi et al., 2014). Here we were able to provide evidence for very clear reproducibility in digit maps based on the two dimensional area of digit representations on the cortical surface. This was also
supported by additional measures of reproducibility: measures of shared cortical
territory of different digit representations and peak-to-peak distance between
adjacent representations (Figure 6; Table 3). These same measures also
highlight the variability seen across participants.

Somatosensory cortex and motor function

SI acts broadly as both a processing region for afferent sensory inputs, as well as
a more central node in the redirection of incoming sensory information across
the sensorimotor network (Mountcastle, 2005). The region shows highly
organized reciprocal connections with primary motor cortex (M1) (Darian-Smith
et al., 1993; Moore et al., 2000) and is co-activated with M1 during both active
and illusory movement of the hand (Porro et al., 1996; Naito et al., 2005).
Furthermore, it is increasingly clear that SI exerts a strong influence on the
function of M1 (Sakamoto et al., 1987; Widener and Cheney, 1997; Vidoni et al.,
2010; Platz et al., 2012; Jacobs et al., 2014).

In light of the structural and functional interplay between SI and M1, a natural
sensorimotor task such as hand movement will elicit robust activation of SI. The
phase-encoding paradigm applied in this study is targeted as resolving the kind
of ordered smooth somatotopy reported previously in SI rather than M1
(Sanchez-Panchuelo et al., 2010; Martuzzi et al., 2014; Stringer et al., 2014).
However other approaches have provided evidence for representation of specific
movements or digits in different neuronal populations or cortical regions of M1,
though not strictly digit somatotopy (Schieber and Hibbard, 1993; Nudo et al.,
1996). Indeed, work in humans suggests motor representations may be encoded
in a higher dimensionality space rather than as individual body parts (Overduin et al., 2012; Diedrichsen et al., 2013; Wiestler et al., 2013). However, recent work combining fMRI and electrocorticography does provide evidence for some ordered digit topography in M1 (Siero et al., 2014).

Digit mapping: plasticity and disease

The presence of stable but variable somatotopic maps raises the possibility of investigating the factors underlying individual differences in cortical functional architecture. In addition, the observation of stability in even the most fine grain SI somatotopy provides a firm foundation for studies of plasticity, for example using within-subject longitudinal study designs. Such work might consider the potential for remapping in health and disease, building upon previous studies using MEG (Braun et al., 2000; Schwenkreis et al., 2001; Candia et al., 2003; Stavrinou et al., 2007). Furthermore, the reproducibility of these maps combined with the short 10-minute acquisition time and simple motor paradigm, provides encouraging evidence for the clinical utility of single-participant fMRI. Mapping techniques could be of particular interest in presurgical planning or monitoring longitudinal changes in patient populations (Hirsch et al., 2000; Yoo et al., 2005; Bosnell et al., 2008; Gountouna et al., 2010).

Phase-encoding and digit mapping

We provide evidence of concordance between digit maps derived from phase-encoding and more traditional block designs (Figure 8). Previous work in body mapping has also demonstrated agreement between phase-encoding maps and mapping results from other fMRI paradigms, including block designs (Orlov et al.,
2010; Besle et al., 2013a), event related designs (Besle et al., 2013a), and resting state functional connectivity data (Zeharia et al., 2015). The results presented here provide further compelling evidence that the phase-encoding analysis provides a meaningful method of mapping patterns of topography.

Limitations

The use of a motor task in assessing SI topography has a number of limitations. Firstly, it is not possible to isolate the exact somaesthetic sub-modality responsible for these maps, which could be induced by stimulation of cutaneous or subcutaneous receptors, or deeper proprioceptors. However, typical use of the hand recruits a combination of such receptors, as such this task represents a more naturalistic assessment of SI function than somatosensory stimulation in the absence of movement. In light of the active task applied in this study it would also be challenging to make inferences about Brodmann area somaesthetic sub-modality specificity.

Given the anatomical enslavement of certain adjacent digit pairs, it is possible that in moving certain fingers, adjacent fingers will also be moved to a lesser extent. Variability in this enslavement could contribute to the inter-subject differences reported in this study. However, given the relatively universal anatomical basis of enslavement (Yu et al., 2010), it seems unlikely that this could account for the considerable variance observed in the functional architecture of the cortex observed here. Moreover, the phenomenon of enslavement is more marked in extension rather than the flexion involved in button press tasks (Yu et al., 2010).
The coverage limitations of ultra-high resolution fMRI at 7T constrained the region of interest to SI, preventing any assessment of secondary somatosensory cortex or subcortical grey matter structures, where somatotopy has previously been reported (Lenz and Byl, 1999; Ruben et al., 2001; Huang and Sereno, 2007).

**Conclusions**

This study robustly demonstrates the presence of stable digit somatotopy of four digits in human SI, as well as the considerable inter-subject variability in these representations. The use of fMRI to demonstrate this reproducibility at the level of single participants provides a firm foundation for this non-invasive imaging technique to investigate highly detailed functional organization of the human brain. The mapping paradigm validated in this study has potential applications both in the study of sensorimotor plasticity in the context of both learning and pathological dysfunction, as well as in the clinical setting.

**References**


Event-related fMRI at 7T reveals overlapping cortical representations for adjacent fingertips in S1 of individual subjects. Hum Brain Mapp 35:2027–2043.


Figure 1: Overview of phase-encoding digit mapping task and analysis

(A) The phase-encoding paradigm: 8 x 32-second cycles of continuous button presses at 1Hz. Each 32-second cycle consists of four 8-second blocks, with each block cycling through either D2-D5 (forward) or D5-D2 (backward). (B) BOLD timecourses from individual voxels (one timecourse shown) cross-correlated against reference models (8-second 'on', 24-seconds 'off'), shifted iteratively by a number of lags to capture activation throughout the movement cycles. (C) Plotting cross-correlation of each voxel's timecourse as a function of lag reveals peak cross-correlation at a given lag. Four different voxels shown, each with a cross-correlation peaking in lags corresponding to different digits. (D) r-values for each voxel averaged across lags assigned to specific digits. Resulting digit r-value maps for forward and backwards cycled are also averaged to yield voxel-wise r-value maps for each digit for one subject/timepoint (thresholded maps displayed).

Figure 2: Phase-encoding digit maps from a single participant and timepoint

(A) Digit maps in BOLD EPI volumetric space across five adjacent transverse slices (z: 11-15); FDR threshold (α=0.001). Digit 2: pink, digit 3: orange, digit 4: green, digit 5: blue. R: right, L: lateral, M: medial. (B) Post-hoc analysis of BOLD signal from individual digit representations in this participant. (C/D) Surface projection of digit maps shown on the inflated pial surface (black: sulcal pattern). Red highlighted region (C: inset) indicates coverage of BOLD EPI task fMRI data partial field of view. No masking has been applied within the acquisition field of view.

Figure 3: Temporal reproducibility of phase-encoding digit maps within-participant.

Comparison of phase-encoding digit representations at three scan timepoints for three participants. Although there is a high degree of between-subject variability (as shown by the large differences between rows), there is very little within-subject variability over time (demonstrated by the small differences across each row). Cortical maps shown on the inflated pial surface with the sulcal pattern in black (positive curvature). Zoomed panels are centred on the hand knob of the central sulcus. All digit maps are subject to FDR thresholding (α=0.001); full details of thresholds and maxima for each time point provided in table 2; colour bars represent range from zero to maximum.
Comparison of phase-encoding digit representations at three scan timepoints continued from figure 4 for remaining participants. All digit maps are subject to FDR thresholding (α=0.001); full details of thresholds and maxima for each time point provided in table 2.

Figure 5: Quantifying intra-subject reproducibility and inter-subject variability in phase-encoding digit maps

(A i-iii) Dice coefficients demonstrate a clear pattern of reproducibility for maps of homologous digits across the three timepoints under study compared with first order and second/third order neighbours. (A iv) Dice coefficients for homologous digits were greater than the equivalent value between non-homologous digit pairings: one-way repeated measures ANOVA: significant main effect of digit pairing category (homologous, first order neighbour, second/third order neighbour). **: Post hoc analysis (Bonferroni adjusted): p < .0005. (B i)

Dice coefficients comparing all combinations of individual digit representations across different participants (after accounting for differences in digit map size) across 0 hour and +/- 4 week timepoints. (B ii) Patterns in each digit pair sub-matrix were summarised by the matrix dominance ratio (Mdr; Equation 2) Mdr > 1 suggests greater intra-subject overlap in digit representations; Mdr < 1 implies greater inter-subject overlap in digit representations. For ‘same’ pairings (e.g. D2-D2) a pattern of high overlap was seen intra-subject (B i: submatrix diagonals), contrasted lower overlap values for comparisons inter-subject (B i: submatrix off-diagonals). For ‘different’ pairings (e.g. D2-D4) no such pattern was observed, suggesting intra-subject consistency is driven by reproducibility of the spatial patterns for the same digits over time. (B iii) Calculation of the overall Mdr (from B ii) was subjected to bootstrap resampling (50,000 iterations) to account for the likelihood of observing these dominance ratios by chance. Bootstrapping returned p < .0005 for the observed value of overall Mdr, consistent with the notion of inter-subject variability in fine grain digit representations.

Figure 6: Patterns of overlap between different digit representations

Soft-edged phase-encoding digit maps provide information regarding shared cortical territory of different digit representations. (A) Average measures of cortical overlap between different digit representations across all subjects and time points reveal pattern of greater shared territory across functionally coupled digits: the relative independence of D2, with increasing levels of cortical overlap between more synergistic D3/D4 and D4/D5. (B) Cortical overlap matrices for
individual participants and time points; ranked Mantel test statistics were used
to compare matrices. Intra-subject comparisons: average Mantel test statistic for
intra-subject comparisons across the three time points. Inter-subject
comparisons: average of the Mantel test statistic calculated between each
participant at a given time point and all other participants at that time point;
calculated for each time point and averaged. (C) Comparison of the intra- versus
inter-subject Mantel test statistics revealed greater similarity of values within a
given subject compared with across different subjects: paired sample t-test, **: $p$
<.0005.

Figure 7

Figure 7: Resolving additional digit maps within SI.

An all-session average phase encoding map was produced for each participant
and resampled into a common space (FDR thresholding, $\alpha=0.01$). Additional
maps were seen in a subset of participants. A more anterior map was observed
(Arrowhead: B,C,E,F) in some individuals, and a more posterior map
(Arrowheads: G,H) in others, both within SI. In the remaining participants (A, D,
I) no clear evidence for additional maps was found.

Figure 8

Figure 8: Validation of phase-encoding digit maps using block design data

Beta values from the block design task fMRI data were extracted for each GLM
contrast (digit > rest) at the peak voxels of the phase-encoding digit
representations (D2-D5). This process was repeated for each of the three scans
to derive average values for each participant. For each phase-encoding digit
representation the beta value of the homologous GLM contrast (e.g. D2 phase-
coding vs. D2 > rest GLM contrast), was significantly greater than for non-
homologous GLM contrasts (e.g. D2 phase-encoding vs. D4 > rest GLM contrast).
RM-ANOVA: significant interaction between phase-encoding digit representation
and the digit contrast of the block design GLM on the normalised beta value. **
Post-hoc t-test $p<.0005$ (uncorrected).
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Table 1: Participant demographic information

Participant demographics for the thirteen participants recruited to this study. F: female, M: male, R: right-handed. Laterality index calculated using Edinburgh Handedness Score (Oldfield et al. 1971).
Table 2

Table 2: FDR thresholds for single-participant digit maps

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FDR thresholds (maxima in parentheses) for z-statistic phase-encoding derived digit maps for individual participants shown in Figures 3 and 4 across the three timepoints under study. FDR: false discovery rate (α=0.001). A two-way repeated measures ANOVA was performed to assess for systematic differences in FDR-defined thresholds, with one factor of digit (D2-D5) and one factor of session (0 hour, +24 hour, +/− 4 weeks). There was no significant main effect of session on FDR threshold: \( F(2,16) = 0.218, p=0.806 \), sphericity assumed.
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</table>

Table 3: Peak-to-peak distances for single participant digit maps across sessions

Peak-to-peak distances (mm) derived from phase-encoding digit maps.
**A** Phase-encoding task digit order

**B** BOLD timecourse vs. lag-shifted HRF-convolved models: cross-correlation

<table>
<thead>
<tr>
<th>Lag</th>
<th>r</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>-0.3</td>
</tr>
<tr>
<td>2</td>
<td>0.2</td>
</tr>
<tr>
<td>3</td>
<td>0.6</td>
</tr>
</tbody>
</table>

**C** Cross-correlation reveals digit specificity for individual voxels

**D** Voxel-wise averages across lags/orders
FDR thresholds and maxima in table 2

Participant 1

0 hours
+24 hours
+4 weeks

Participant 2

0 hours
+24 hours
-4 weeks

Participant 3

0 hours
+24 hours
+4 weeks
0 hours +24 hours +/-4 weeks

Participant 4

Participant 5

Participant 6

Participant 7

Participant 8

Participant 9

FDR thresholds and maxima in table 2
A  Intra-subject digit map reproducibility: Dice coefficient

i  Dice coefficients: 'same' and 'different' pairings

ii  Quantifying intra- vs. inter-subject patterns in Dice coefficient submatrices

iii  Overall matrix dominance ratio: 19.67

(B) Inter-subject digit variability: Dice coefficient

i  Dice coefficients: 'same' and 'different' pairings

ii  Quantifying intra- vs. inter-subject patterns in Dice coefficient submatrices

iii  Overall matrix dominance ratio: 19.67

iv  Bootstrap resampling overall matrix dominance ratio
A  Overlap of different digit representations: all time point inter-subject average

B  Inter- versus intra-subject digit overlap patterns

C  Intra- versus inter-subject digit overlap similarity
Phase-encoding derived digit representation (peak voxel)

GLM contrast
- Digit 2 > rest
- Digit 3 > rest
- Digit 4 > rest
- Digit 5 > rest