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Neuroscience

Updating the sulcal landscape of the human lateral parieto-occipital junction provides anatomical, functional, and cognitive insights

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Abstract

Recent work has uncovered relationships between evolutionarily new small and shallow cerebral indentations, or sulci, and human behavior. Yet, this relationship remains unexplored in the lateral parietal cortex (LPC) and the lateral parieto-occipital junction (LPOJ). After defining thousands of sulci in a young adult cohort, we uncovered four previously unidentified small and shallow LPC/LOPJ sulci—one of which (ventral supralateral occipital sulcus, slocs-v) is present in nearly every hemisphere, and is morphologically, architecturally, and functionally dissociable from neighboring regions. A data-driven, model-based approach relating sulcal depth to behavior revealed that the morphology of only a subset of LPC/LPOJ sulci, including the slocs-v, is related to performance on a spatial orientation, but not a relational reasoning task. Our findings build on classic neuroanatomical theories and identify new neuroanatomical targets for future "precision imaging" studies exploring the relationship among brain structure, brain function, and cognitive abilities in individual participants.

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This is an **important** study that provides detailed insights into the morphology of the lateral parietal cortex. Through this work new shallow sulci in the parietal cortex were identified and linked to function and behaviour. The evidence presented is **convincing**, even though some claims about the definition of highly variable sulci should be tampered down, and some **important** precisions about the labelling process are lacking to secure reproducibility. The present work both advances our understanding of the parietal cortex while also stimulating further debate on precise, detailed manual anatomy vs large scale automated data processing.



Introduction

A fundamental goal in psychology and neuroscience is to understand the complex relationship between brain structure and brain function, as well as how that relationship provides a scaffold for efficient cognition and behavior. Of all the neuroanatomical features to target, recent work shows that morphological features of hominoid-specific, shallow indentations, or sulci, of the cerebral cortex are not only functionally and cognitively meaningful, but also are particularly impacted by multiple brain-related disorders and aging (Amiez et al., 2019 2, 2018 2; Ammons et al., 2021 C; Cachia et al., 2021 C; Fornito et al., 2004 C; Garrison et al., 2015 C; Harper et al., 2022 🖸 ; Hathaway et al., 2023 🗳 ; Lopez-Persem et al., 2019 😋 ; Miller et al., 2021 🗳 , 2020 😋 ; Nakamura et al., 2020 C; Parker et al., 2023 C; Voorhies et al., 2021 C; Weiner, 2019 C; Willbrand et al., 2023b 🖸 , 2023c 🗹 , 2022a 🖸 , 2022b 🗹 ; Yao et al., 2022 🖒). The combination of these findings provides growing support for a classic theory proposing that the late emergence of these structures in gestation within association cortices, as well as their prolonged development, may co-occur with specific functional and microstructural features that could support specific cognitive abilities that also have a protracted development (Sanides, 1964 \square). Nevertheless, despite the developmental, evolutionary, functional, cognitive, and theoretical relevance of these findings, they have mainly been restricted to only a subset of association cortices such as the prefrontal, cingulate, and ventral occipitotemporal cortices (Amiez et al., 2019 2, 2018 2; Ammons et al., 2021 C; Cachia et al., 2021 C; Fornito et al., 2004 C; Garrison et al., 2015 C; Harper et al., 2022 C; Hathaway et al., 2023 C; Lopez-Persem et al., 2019 C; Miller et al., 2021 C, 2020 C; Nakamura et al., 2020 C; Parker et al., 2023 C; Voorhies et al., 2021 C; Weiner, 2019 C; Willbrand et al., 2023b C, 2023c C, 2022a C, 2022b C; Yao et al., 2022 C). Thus, examining the relationship among these structures (also known as tertiary sulci) relative to architectonic and functional features of the cerebral cortex, as well as relative to cognition, remains uncharted in other association cortices such as the lateral parietal cortex (LPC).

As LPC is a cortical extent that has expanded extensively throughout evolution (Van Essen et al., 2018 C; Zilles et al., 2013 C), there is great interest in the structure and function of LPC in development, aging, across species, and in different patient populations. Yet, key gaps in knowledge relating individual differences in the structure of LPC to individual differences in the functional organization of LPC and cognitive performance remain for at least four main reasons. First, one line of recent work shows that LPC displays a much more complex sulcal patterning than previously thought (Drudik et al., 2022 🖒; Petrides, 2019 🖒; Segal and Petrides, 2012 🤅; Zlatkina and Petrides, 2014 2), while a second line of work shows that LPC is tiled with many maps and discrete functional regions spanning modalities and functions such as vision, memory, attention, action, haptics, and multisensory integration in addition to theory of mind, cognitive control, and subdivisions of the default mode network (Goodale and Milner, 1992 C; Harvey et al., 2015 C, 2013 C; Humphreys and Tibon, 2023 C; Konen and Kastner, 2008 C; Mackey et al., 2017 C; Schurz et al., 2017 🔼). Second, a majority of the time, the two lines of work are conducted independently from one another and the majority of human neuroimaging studies of LPC implement group analyses on average brain templates—which causes LPC sulci to disappear (Fig. **1**^C). Third, despite the recently identified complexity of LPC sulcal patterning, recent studies have also uncovered previously overlooked tertiary sulci in association cortices (for example, in the posterior cingulate cortex (Willbrand et al., 2023c 2, 2022a 2)). Thus, fourth, it is unknown if additional LPC tertiary sulci are waiting to be discovered and if so, could improve our understanding of the structural-functional organization of LPC with potential cognitive insights as in other association cortices. Critically, while such findings would have developmental, evolutionary, functional, cognitive, and theoretical implications for addressing novel questions in future studies, they would also have translational applications as sulci serve as biomarkers in neurodevelopmental disorders (Ammons et al., 2021 2; Cachia et al., 2021 2; Garrison et al., 2015 C; Nakamura et al., 2020 C) and "corridors" for neurosurgery (Tomaiuolo et al., 2022 C; Tomaiuolo and Giordano, 2016 2).



Fig. 1.

Four previously undefined small and shallow sulci in the lateral parieto-occipital junction (LPOJ).

a. Four example inflated (top) and pial (bottom) left hemisphere cortical surfaces displaying the 13-17 sulci manually identified in the present study. Each hemisphere contains 1-4 of the previously undefined and variable LOC/LPOJ sulci (slocs and pAngs). Each sulcus is numbered according to the legend. **b.** Criteria for defining slocs and pAngs components. (i) Slocs-v is the cortical indentation between the cSTS3 and ITOS. (ii) Slocs-d is the indentation between cSTS3/cSTS2 and IPS-PO. (iii) pAngs-v is the indentation between the cSTS2 and pips. (iv) pAngs-d is the indentation between cSTS2/cSTS1 and IPS. **c.** The variability of the slocs and pAng components can cause them to disappear when individual surfaces are averaged together. Left to right: (i) 10 HCP participants, (ii) 20 HCP participants, (iii) 100 HCP participants, and iv) 650 HCP participants. The disappearance of these sulci on average surfaces, which are often used for group analyses in neuroimaging research, emphasizes the importance of defining these structures in individual hemispheres.



In the present study, we first manually defined LPC sulci in 144 young adult hemispheres using the most recent definitions of LPC sulci (Petrides, 2019^{CC}). By manually labeling over 2,000 sulci, we uncovered four previously undefined (Supplementary Methods and Supplementary Figs. 1-4 for historical details) sulci in the cortical expanse between the caudal branches of the superior temporal sulcus (cSTS) and two parts of the intraparietal sulcus (IPS)—a cortical expanse recently referenced as containing sensory "bridge" regions of the temporal-parietal-occipital junction (Glasser et al., 2016 C)—which we term the supralateral occipital sulci (ventral: slocs-v; dorsal: slocs-d) and posterior angular sulci (ventral: pAngs-d; dorsal: pAngs-d). We then utilized morphological (depth and surface area), architectural (gray matter thickness and myelination), and functional (resting-state functional connectivity) data available in each participant to assess whether the most common of these structures (slocs-v) was dissociable from surrounding sulci. Finally, we assessed whether the updated view of the LPC/LPOJ sulcal landscape provided cognitive insights using a model-based, data-driven approach (Voorhies et al., 2021 🗹) relating sulcal morphology to behavior on tasks known to activate regions within this cortical expanse (for example, reasoning and spatial orientation (Gur et al., 2000 C2; Karnath, 1997 C2; Vendetti and Bunge, 2014 2; Wendelken, 2014 2)).

Results

Four previously undefined small and shallow sulci in the lateral parieto-occipital junction (LPOJ)

In addition to defining the 13 sulci previously described within the LPC/LPOJ, as well as the posterior superior temporal cortex (**Methods**) (Petrides, 2019^{C2}) in individual participants, we could also identify as many as four small and shallow sulci situated within the LPC/LPOJ that were more variable across individuals and uncharted until now (Supplementary Methods and Supplementary Figs. 1-4). Macroanatomically, we could identify two sulci between the cSTS3 and the IPS-PO/ITOS ventrally and two sulci between the cSTS2 and the pips/IPS dorsally.

Ventrally, we refer to these sulci as ventral (slocs-v; sulcus 5 in **Fig. 1**⁽²⁾) and dorsal (slocs-d; sulcus 6 in **Fig. 1**⁽²⁾) components of the supralateral occipital sulcus. The slocs-v, located between the posterior cSTS3 and ITOS, was present in 98.6% of hemispheres (left hemisphere: N = 71/72; right hemisphere: N = 71/72; **Fig. 1**⁽²⁾). Conversely, the slocs-d, located between the cSTS3 and IPS-PO, was present 68.0% of the time (left hemisphere: N = 50/72; right hemisphere: N = 48/72; **Fig. 1**⁽²⁾). Dorsally, we refer to the other newly identified sulci as the ventral (pAngs-v; sulcus 7 in **Fig. 1**⁽²⁾) and dorsal (pAngs-d; sulcus 8 in **Fig. 1**⁽²⁾) components of the posterior angular sulcus. The pAng components were more rare than the slocs components. Specifically, pAngs-v, located between cSTS2 and pips, was identifiable 31.3% of the time (19 left and 26 right hemispheres; **Fig. 1**⁽²⁾). Located between cSTS2 and the IPS, pAngs-d was identifiable only 13.2% of the time (8 left and 11 right hemispheres; **Fig. 1**⁽²⁾). Though we characterize these sulci in this paper for the first time, the location of these four sulci is consistent with the presence of variable "accessory sulci" in this cortical expanse mentioned in prior modern and classic studies (Supplementary Methods).

These incidence rates were significantly different (GLM, main effect of sulcus: $\chi 2(3) = 166.53$, p < .0001; no hemispheric effects: ps > .68). The slocs-v was more common than the other three sulci (ps < .0001), slocs-d was more common than the pAngs components (ps < .0001), and pAngs-v was more common than pAngs-d (p = .002). We could further identify these sulci in post-mortem hemispheres (Supplementary Figs. 2, 3). Finally, to help guide future research on these newly- and previously-classified LPC/LPOJ sulci, we generated probabilistic maps of each of these 17 sulci and share them with the field with the publication of this paper (Supplementary Fig. 6).



The slocs-v is morphologically, architecturally, and functionally dissociable from nearby sulci

Given that the slocs-v was present in the majority of participants (98.6% across hemispheres), we focused our analyses on this stable sulcal feature of the LPOJ. To do so, we first tested whether the slocs-v was morphologically (depth and surface area) and architecturally (gray matter thickness and myelination) distinct from the two sulci surrounding it: the cSTS3 and ITOS (**Fig. 1**, Supplementary Fig. 7 for these metrics in all 17 sulci examined). An rm-ANOVA (within-participant factors: sulcus, metric, and hemisphere for standardized metric units) revealed a sulcus x metric interaction (F(4, 276.19) = 179.15, $\eta 2 = 0.38$, p < .001). Post hoc tests showed four main differences: (i) the slocs-v was shallower than cSTS3 (p < .001) but not ITOS (p = .60), (ii) the slocs-v was smaller than both the cSTS3 and ITOS (ps < .001), (iii) the slocs-v was thicker than both the cSTS3 and ITOS (ps < .001), (iii) the slocs-v was less myelinated than both the cSTS and ITOS (ps < .001; **Fig. 2a**,). There was also a sulcus x metric x hemisphere interaction (F(4.20, 289.81) = 4.16, $\eta 2 = 0.01$, p = .002; Supplementary Results).

We then tested whether the slocs-v was also functionally distinct from the cSTS3 and ITOS by leveraging resting-state network parcellations for each individual participant to quantify "connectivity fingerprints" for each sulcus in each hemisphere of each participant (**Methods**) (Kong et al., 2019 ^{C2}). An rm-ANOVA (within-participant factors: sulcus, network, and hemisphere for Dice coefficient overlap) revealed a sulcus x network interaction (F(32, 2144) = 80.18, $\eta 2 = 0.55$, p < .001). Post hoc tests showed that this interaction was driven by four effects: (i) the cSTS3 overlapped more with the Default A subnetwork than both the slocs-v and ITOS (ps < .001), (ii) the slocs-v overlapped more with the Default C subnetwork than the ITOS (p < .001) and marginally than the cSTS3 and ITOS (ps < .001), and iv) the ITOS overlapped more with the Visual A and Visual B subnetworks than both the cSTS3 and slocs-v (ps < .004; **Fig. 2b** ^{C2}). There was also a sulcus x network x hemisphere interaction (F(32, 2144) = 3.99, $\eta 2 = 0.06$, p < .001; Supplementary Results). Together, these results indicate that the slocs-v is a morphologically, architecturally, and functionally distinct structure from its sulcal neighbors, and thus, deserves a distinct neuroanatomical definition.

We further found that the three caudal STS rami (Petrides, 2019²; Segal and Petrides, 2012²) and intermediate parietal sulci (aipsJ and pips) (Petrides, 2019²; Zlatkina and Petrides, 2014²) are morphologically, architecturally, and functionally distinct structures for the first time (to our knowledge), which empirically supports their distinctions with separate sulcal labels (Supplementary Results and Supplementary Fig. 8).

The morphology of LPC/LPOJ sulci, including the slocs-v, is related to cognitive performance

Finally, leveraging a data-driven approach of cross-validated LASSO feature selection, we sought to determine whether sulcal depth, a main defining feature of sulci, related to cognitive performance (**Methods**). To do so, we primarily focused on spatial orientation and reasoning given that these abilities recruit multiple subregions of lateral parietal and/or occipital cortices (Gur et al., 2000 C; Karnath, 1997 C; Vendetti and Bunge, 2014 C; Wendelken, 2014 C). As in prior work (Voorhies et al., 2021 C; Willbrand et al., 2023 C; Yao et al., 2022 C), we chose the model at the alpha that minimized MSE_{cv}. Participants with a slocs-v in both hemispheres and all behavioral metrics were included (N = 69). Due to their rarity (being in less than 70% of hemispheres at most), we did not include the slocs-d or pAng components in this analysis.

This method revealed an association between spatial orientation scores and normalized sulcal depth in the left hemisphere (MSE_{cv} = 25.63, alpha = 0.05; **Fig. 3a** ⁽²⁾), but not in the right hemisphere (MSE_{cv} = 26.41, alpha = 0.3). Further, we found that no LPC/LPOJ sulci were selected



Fig. 2.

The slocs-v is morphologically, architecturally, and functionally dissociable from nearby sulci.

a. Radial plot displaying the morphological (upper metrics: depth, surface area) and architectural (lower metrics: cortical thickness, myelination) features of the slocs-v (gray), cSTS3 (blue), and ITOS (green). Each dot and solid line represents the mean. The dashed lines indicate \pm standard error. These features are colored by sulcus (legend). Metrics are in standardized units. **b.** Radial plot displaying the connectivity fingerprints of these three sulci: the Dice Coefficient overlap (values from 0-1) between each component and individual-level functional connectivity parcellations (Kong et al., 2019 $\$).



for reasoning in either hemisphere (right: alpha = 0.3, MSE = 24.01; left: alpha = 0.3, MSE = 24.01). Six left hemisphere LPC/LPOJ sulci were related to spatial orientation task performance **Fig. 3a**, **b** \square). Four of these sulci were positioned ventrally: cSTS3 (β = -9.77), slocs-v (β = -3.36), ITOS (β = -4.91), and mTOS (β = -0.06), whereas two were positioned dorsally: pips (β = 5.02), and SPS (β = 4.30; **Fig. 3a**, **b** \square). Using LooCV to construct models that predict behavior, the LASSO-selected model explained variation in spatial orientation score (\mathbb{R}^2 = 0.06, MSE_{cv} = 23.99) above and beyond a model with all left hemisphere sulci ($\mathbb{R}^2 < 0.01$, MSE_{cv} = 27.12). This model also showed a moderate correspondence (\mathbf{r}_s = 0.29, p = .01; **Fig. 3c** \square) between predicted and actual measured scores. We then tested for anatomical and behavioral specificity using the AIC, which revealed two primary findings. First, we found that the LASSO-selected sulcal depth model outperformed a model using the cortical thickness of the six LASSO-selected sulci ($\mathbb{R}^2 < .01$, MSE = 26.02, AIC – AIC = 2.19). This model also showed task specificity as these sulci outperformed a model with processing speed ($\mathbb{R}^2 < .01$, MSE = 254.65, AIC – AIC = 63.57). Thus, our data-driven model explains a significant amount of variance on a spatial orientation task and shows behavioral and morphological specificity.

Discussion

Overview

In the present study, we examined the relationship between LPC/LPOJ sulcal morphology, functional connectivity fingerprints, and cognition. We report five main findings. First, while manually defining sulci in LPC/LPOJ across 144 hemispheres, we uncovered four new small and shallow sulci that are not included in present or classic neuroanatomy atlases or neuroimaging software packages. Second, we found that the most common of these structures (the slocs-v; identifiable 98.6% of the time) was morphologically, architecturally, and functionally differentiable from nearby sulci. Third, using a model-based, data-driven approach quantifying the relationship between sulcal morphology and cognition, we found a relationship between the depths of six LPC/LPOJ sulci and performance on a spatial orientation processing task. Fourth, the model identified distinct dorsal and ventral sulcal networks in LPC/LPOJ: ventral sulci had negative weights while dorsal sulci had positive weights (**Fig. 3b**). These findings are consistent with previous neuroimaging work from Gur et al. (Gur et al., 2000 ^{CD}) who demonstrated separate functional activations in dorsal parietal and the more ventrally situated occipital-parietal cortices for the judgment of line orientation task used in the present study. Fifth, the model identified that the slocs-v is cognitively relevant, further indicating the importance of this new neuroanatomical structure. In the sections below, we discuss (i) the slocs-v relative to modern functional and cytoarchitectonic parcellations in the LPC/LPOJ, as well as anatomical connectivity to other parts of the brain, (ii) underlying anatomical mechanisms relating sulcal morphology and behavior more broadly, and (iii) limitations of the present study. Implications for future studies are distributed throughout each section.

The slocs-v relative to modern functional and cytoarchitectonic parcellations in the LPC/LPOJ, as well as anatomical connectivity to other parts of the brain

To lay the foundation for future studies relating the newly-identified slocs-v to different anatomical and functional organizational features of LPC/LPOJ, we situate probabilistic predictions of slocs-v relative to probabilistic cortical areas identified using multiple modalities. For example, when examining the correspondence between the slocs-v and modern multimodal (HCP-MMP (Glasser et al., 2016 ^{C2})) and observer-independent cytoarchitectural (Julich-Brain atlas (Amunts et al., 2020 ^{C2})) areas (Methods), the slocs-v is located within distinct areas. In particular, the slocs-v aligns with the multimodally- and cytoarchitecturally-defined area PGp bilaterally and



Fig. 3.

The morphology of LPC/LPOJ sulci, including the slocs-v, is related to cognitive performance.

a. Beta-coefficients for each left hemisphere LPC/LPOJ sulcus at a range of shrinking parameter values [alpha (α)]. Highlighted gray bar indicates coefficients at the chosen α -level. Bottom: Cross-validated mean-squared error (MSE_{CV}) at each α level. By convention, we selected the α that minimized the MSE_{CV} (dotted line). **b.** Inflated left hemisphere cortical surface from an example participant highlighting the two groups of sulci—*dorsal positive* (orange) and *ventral negative* (green)—related to spatial orientation performance. **c.** Spearman's correlation (r_s) between the measured and the predicted spatial orientation scores from the LASSO-selected model is shown in a.



cytoarchitecturally-defined hIP4 in the right hemisphere (**Fig. 4**^{C2}). In classic neuroanatomical terms (<u>Cunningham, 1892</u>^{C2}), this indicates that the slocs-v is a putative "axial sulcus" for these regions, which future work can assess with analyses in individual participants.

Aside from recent multimodal and observer-independent cytoarchitectonic parcellations, an immediate question is: What is the relationship between the slocs-v and other functional regions at this junction between the occipital and parietal lobes, as well as potential anatomical connectivity? For example, there are over a dozen visual field maps in the cortical expanse spanning the TOS, IPS-PO, and the IPS proper (see (i), (ii), and (iii), respectively in **Fig. 5a** ⁽²⁾) (Mackey et al., 2017 ⁽³⁾). When projecting probabilistic locations of retinotopic maps from over 50 individuals from Wang and colleagues (Wang et al., 2015 ⁽³⁾) (Methods), the slocs-v is likely located outside of visual field maps extending into this cortical expanse (**Fig. 5a** ⁽³⁾). Nevertheless, when also projecting the map of the mean R² metric from the HCP retinotopy dataset from 181 participants shared by Benson and colleagues (Benson et al., 2018 ⁽³⁾) (Methods), the slocs-v is in a cortical expanse that explains a significant amount of variance (left hemisphere: R² = 19.29, R² = 41.73; right hemisphere: R² = 21.17, R² = 44.23; **Fig. 5b** ⁽³⁾).

In terms of anatomical connectivity, as the slocs-v co-localizes with cytoarchitectonically defined PGp (**Fig. 4**^C) and previous studies have examined the anatomical connectivity of the probabilistically defined PGp, we can glean insight regarding the anatomical connectivity of slocs-v from these previous studies (Caspers et al., 2011^C; Wang et al., 2012^C). This prior work showed that PGp was anatomically connected to temporo-occipital regions, other regions in the temporal lobe, middle and superior frontal cortex, as well as the inferior frontal cortex and insula (Caspers et al., 2011^C; Wang et al., 2011^C; Wang et al., 2011^C; Wang et al., 2011^C; Wang et al., 2012^C). Furthermore, the slocs-v appears to lie at the junction of scene-perception and place-memory activity (a transition that also consistently co-localizes with the HCP-MMP area PGp) as identified by Steel and colleagues(Steel et al., 2021^C). Of course, the location of the slocs-v relative to multimodal, cytoarchitectonic, and retinotopic areas, as well as the anatomical connectivity of the slocs-v, would need to be examined in individual participants, but the present work makes clear predictions for future studies as fleshed out here. To conclude this section, as the multimodal area PGp (**Fig. 4**^C) was recently proposed as a "transitional area" by Glasser and colleagues (Glasser et al., 2016^C) (Supplementary Table 1), future studies can also further functionally and anatomically test the transitional properties of slocs-v.

Underlying anatomical mechanisms relating sulcal morphology and behavior

In this section, we discuss potential anatomical mechanisms contributing to the relationship between sulcal depth and behavior in two main ways. First, long-range white matter fibers have a gyral bias, while short-range white matter fibers have a sulcal bias in which some fibers project directly from the deepest points of a sulcus (Cottaar et al., 2021 🗳; Reveley et al., 2015 🗳; Schilling et al., 2018 🗳, 2023 🗳; Van Essen et al., 2014 🗳). As such, recent work hypothesized a close link between sulcal depth and short-range white matter properties (Bodin et al., 2021 🗳; Pron et al., 2021 🗳; Voorhies et al., 2021 🗳; Willbrand et al., 2023 b 🗳; Yao et al., 2022 🗳): deeper sulci would reflect even shorter short-range white matter fibers, which would result in faster communication between local, cortical regions and in turn, contribute to improved cognitive performance. This increased neural efficiency could underlie individual differences in cognitive performance. Ongoing work is testing this hypothesis which can be further explored in future studies incorporating anatomical, functional, and behavioral measures, as well as computational modeling.

Second, our model-based approach identified separate dorsal and ventral sulcal networks in which deeper sulci dorsally and shallower sulci ventrally contributed to the most explained variance on the spatial orientation task. A similar finding was identified by our previous work in the lateral prefrontal cortex (Yao et al., 2022 C²). These previous and present findings may be



Fig. 4.

The slocs-v relative to modern functional and cytoarchitectonic parcellations in LPC/LPOJ.

a. Top: Left (LH) and right (RH) hemispheres of the inflated fsaverage surface with two areas from the modern HCP multimodal parcellation (HCP-MMP; blue) (Glasser et al., 2016^{C2}) relative to an MPM of the slocs-v (warm colors indicate areas with at least 20% overlap across participants; Supplementary Fig. 6). Bottom: Same as top, except for two observer-independent cytoarchitectonic regions from the Julich-Brain Atlas (Amunts et al., 2020^{C2}). **b.** Overlap between the slocs-v and each area (Methods). Each dot and solid line represents the mean. The dashed lines indicate ± standard error (left: gray; right: white).



Fig. 5.

The slocs-v relative to retinotopy.

a. Top: Left (LH) and right (RH) hemispheres of the inflated fsaverage surface displaying the probabilistic locations of retinotopic maps from over 50 individuals from Wang and colleagues (Wang et al., 2015⁽²⁾) (black outlines). The predicted slocs-v location from the MPMs is overlaid in orange (as in **Fig. 4**⁽²⁾). (i), (ii), and (iii) point out the retinotopic maps in the cortical expanse spanning the TOS, IPS-PO, and IPS, respectively. **b.** Same format as in a, but with a map of the mean R² metric from the HCP retinotopy dataset (Benson et al., 2018⁽²⁾) overlayed on the fsaverage surfaces (thresholded between R² values of 10% and 90%). This metric measures how well the fMRI time series at each vertex is explained by a population receptive field (pRF) model. The mean and max R² values for the slocs-v MPM in each hemisphere are included below each surface.

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explained by the classic anatomical compensation theory, which proposes that the size and depth of a sulcus counterbalance those of the neighboring sulci (Armstrong et al., 1995 , Connolly, 1950 ; Zilles et al., 2013). Thus, a larger, deeper sulcus would be surrounded by sulci that are smaller and shallower, rendering the overall degree of cortical folding within a given region approximately equal (Armstrong et al., 1995 ; Connolly, 1950 ; Zilles et al., 2013). Future work can incorporate underlying white matter architecture into the compensation theory, as well as a recent modification that proposed to also incorporate local morphological features such as the deepest sulcal point (e.g., sulcal pit or sulcal root (Régis et al., 2005)), which has recently been shown to be related to different functional features of the cerebral cortex (Bodin et al., 2018 ; Leroy et al., 2015 ; Natu et al., 2021). Altogether, these and recent findings begin to build a multimodal mechanistic neuroanatomical understanding underlying the complex relationship between sulcal depth and cognition relative to other anatomical features.

Limitations

The main limitation of our study is that presently, the most accurate methodology to define sulci especially the small, shallow, and variable tertiary sulci—requires researchers to manually trace each structure on the cortical surface reconstructions. This method is arduous and timeconsuming, which, on the one hand, limits the sample size in terms of number of participants, while on the other, results in thousands of precisely defined sulci - a push-pull relationship reflecting a conversation in the broader human brain mapping and cognitive neuroscience fields between a balance of large N studies and "precision imaging" studies in individual participants (Allen et al., 2022 C; Gratton et al., 2022 C; Naselaris et al., 2021 C; Rosenberg and Finn, 2022 C). Though our sample size is comparable to other studies that produced reliable results relating sulcal morphology to brain function and cognition (e.g., (Cachia et al., 2021 C; Garrison et al., 2015 C; Lopez-Persem et al., 2019 C; Miller et al., 2021 C; Roell et al., 2021 C; Voorhies et al., 2021 🖸; Weiner, 2019 🖸; Willbrand et al., 2022a C, 2022b C; Yao et al., 2022 C)), ongoing work that uses deep learning algorithms to automatically define sulci should result in much larger sample sizes in future studies (Borne et al., 2020 C; Lyu et al., 2021 C). Finally, the time-consuming manual definitions of primary, secondary, and tertiary sulci also limit the cortical expanse explored in each study, thus, restricting the present study to LPC/LPOJ.

Conclusion

In conclusion, we uncovered four previously-undefined sulci in LPC/LPOJ and quantitatively showed that the slocs-v is a stable sulcal landmark that is morphologically, architecturally, and functionally differentiable from surrounding sulci. We further used a data-driven, model-based approach relating sulcal morphology to behavior, which identified different relationships of ventral and dorsal LPC/LPOJ sulcal networks contributing to the perception of spatial orientation. The model identified the slocs-v, further indicating the importance of this new neuroanatomical structure. Altogether, this work provides a scaffolding for future "precision imaging" studies interested in understanding how anatomical and functional features of LPC/LPOJ relate to cognitive performance at the individual level.

Methods

Participants

Data for the young adult human cohort analyzed in the present study were from the Human Connectome Project (HCP) database (*https://www.humanconnectome.org/study/hcp-young-adult/overview* ^{C2}). Here, we used 72 participants (50% female, 22-36 years old, and 90% righthanded; there was no effect of handedness on our behavioral tasks; Supplementary materials) that



were also analyzed in several prior studies (Hathaway et al., 2023 C; Miller et al., 2021 C, 2020 C; Willbrand et al., 2023 C, 2023 C, 2022 C). HCP consortium data were previously acquired using protocols approved by the Washington University Institutional Review Board (Mapping the Human Connectome: Structure, Function, and Heritability; IRB # 201204036). Informed consent was obtained from all participants.

Neuroimaging data acquisition

Anatomical T1-weighted (T1-w) MRI scans (0.8 mm voxel resolution) were obtained in native space from the HCP database. Reconstructions of the cortical surfaces of each participant were generated using FreeSurfer (v6.0.0), a software package used for processing and analyzing human brain MRI images (<u>surfer.nmr.mgh.harvard.edu</u>) (Dale et al., 1999 ; Fischl et al., 1999). All subsequent sulcal labeling and extraction of anatomical metrics were calculated from these reconstructions generated through the HCP's version of the FreeSurfer pipeline (<u>Glasser et al.,</u> 2013).

Behavioral data

In addition to structural and functional neuroimaging data, the HCP also includes a wide range of behavioral metrics from the NIH toolbox (Barch et al., 2013 ^{CC}). To relate LPC/LPOJ sulcal morphology to behavior, we leveraged behavioral data related to spatial orientation (Variable Short Penn Line Orientation Test), relational reasoning (Penn Progressive Matrices Test), and processing speed (Pattern Completion Processing Speed Test; Supplementary Methods for task details). We selected these tasks as previous functional neuroimaging studies have shown the crucial role of LPC/LPOJ in relational reasoning and spatial orientation (Gur et al., 2000 ^{CC}; Karnath, 1997 ^{CC}; Vendetti and Bunge, 2014 ^{CC}; Wendelken, 2014 ^{CC}), while our previous work relating sulcal morphology to cognition uses processing speed performance as a control behavioral task (Voorhies et al., 2021 ^{CC}; Willbrand et al., 2022b ^{CC}).

Anatomical analyses

Manual labeling of LPC sulci

Sulci were manually defined in 72 participants (144 hemispheres) guided by the most recent atlas by Petrides (Petrides, 2019 2), as well as recent empirical studies (Drudik et al., 2022 2; Segal and Petrides, 2012 C; Zlatkina and Petrides, 2014 C), which together offer a comprehensive definition of cerebral sulcal patterns, including tertiary sulci. For a historical analysis of sulci in this cortical expanse, please refer to Segal & Petrides (Segal and Petrides, 2012 🖒) and Zlatkina & Petrides (Zlatkina and Petrides, 2014 C). Our cortical expanse of interest was bounded by the following sulci and gyri: (i) the postcentral sulcus (PoCS) served as the anterior boundary, (ii) the superior temporal sulcus (STS) served as the inferior boundary, (iii) the superior parietal lobule (SPL) served as the superior boundary, and (iv) the medial and lateral transverse occipital sulci (mTOS and ITOS) served as the posterior boundary. We also considered the following sulci within this cortical expanse: the three different branches of the caudal superior temporal sulcus (posterior to anterior: cSTS3, 2, 1), the supramarginal sulcus (SmgS), posterior intermediate parietal sulcus (pips), sulcus of Brissaud (sB), anterior intermediate parietal sulcus of Jensen (aips]), paroccipital intraparietal sulcus (IPS-PO), intraparietal sulcus (IPS), and the superior parietal sulcus (SPS). Of note, the IPS-PO is the portion of the IPS extending ventrally into the occipital lobe. The IPS-PO was first identified as the paroccipital sulcus by Wilder (1886). There is often an annectant gyrus separating the horizontal portion of the IPS proper from the IPS-PO (Roell et al., 2021 🖄; Zlatkina and Petrides, 2014⁽¹⁾.

Additionally, we identified as many as four previously uncharted and variable tertiary LPC/LPOJ sulci for the first time: the supralateral occipital sulcus (slocs; composed of ventral (slocs-v) and dorsal (slocs-d) components) and the posterior angular sulcus (pAngs; composed of ventral (pAngs-



v) and dorsal (pAngs-d) components). In the Supplementary Methods and Supplementary Figs. 1-4, we discuss the slocs and pAngs within the context of modern and historical sources.

For each participant in each hemisphere, the location of each sulcus was confirmed by trained independent raters (E.H.W., Y.T., and T.G.) and finalized by a neuroanatomist (K.S.W.). All LPC sulci were then manually defined in FreeSurfer using tksurfer tools, as in previous work (Hathaway et al., 2023 ; Miller et al., 2021 , 2020 ; Parker et al., 2023 ; Voorhies et al., 2021 ; Willbrand et al., 2023 ; 2022a , 2022a , 2022b ; Yao et al., 2022 ; Noorhies et al., 2021 ; Willbrand et anatomical features were extracted. For four example hemispheres with these 13-17 sulci identified, see **Fig. 1a** (Supplementary Fig. 5 for all hemispheres). The specific criteria to identify the slocs and pAngs are outlined in **Fig. 1b** .

To test whether the incidence rates of the slocs and pAngs components were statistically different, we implemented a binomial logistic regression GLM with sulcus (slocs-v, slocs-d, pAngs-v, and pAngs-d) and hemisphere (left and right), as well as their interaction, as predictors for sulcal presence [0 (absent), 1 (present)]. GLMs were carried out with the glm function from the built-in stats R package. ANOVA χ 2 tests were applied to each GLM with the Anova function from the car R package, from which results were reported.

Probability maps

Sulcal probability maps were generated to show the vertices with the highest alignment across participants for a given sulcus. To create these maps, the label file for each sulcus was transformed from the individual to the fsaverage surface with the FreeSurfer mri_label2label command (*https://surfer.nmr.mgh.harvard.edu/fswiki/mri_label2label* 2). Once each label was transformed into this common template space, we calculated the proportion of participants for which each vertex was labeled as the given sulcus with custom Python code (Miller et al., 2021 2); Voorhies et al., 2021 2). For vertices with overlap between sulci, we employed a "winner-take-all" approach such that the sulcus with the highest overlap across participants was assigned to that vertex. Alongside the thresholded maps, we also provide constrained maps [maximum probability maps (MPMs)] at 20% participant overlap to increase interpretability (20% MPMs shown in Supplementary Fig. 6). To aid future studies interested in investigating LPC/LPOJ sulci, we share these maps with the field.

Extracting and comparing the morphological and architectural features from sulcal labels

Morphologically, we compared sulcal depth and surface area across sulci, as these are two of the primary morphological features used to define and characterize sulci (Armstrong et al., 1995); Chi et al., 1977); Leroy et al., 2015); Lopez-Persem et al., 2019); Miller et al., 2021 , 2020 ; Natu et al., 2021); Sanides, 1964 ; Voorhies et al., 2021); Weiner, 2019); Welker, 1990 ; Willbrand et al., 2023), 2022); Yao et al., 2022). As in our prior work (Voorhies et al., 2021); Yao et al., 2022), mean sulcal depth values (in standard FreeSurfer units) were computed in native space from the .sulc file generated in FreeSurfer (Dale et al., 1999)) with custom Python code (Voorhies et al., 2021). Briefly, depth values are calculated based on how far removed a vertex is from what is referred to as a "mid-surface," which is determined computationally so that the mean of the displacements around this "mid-surface" is zero. Thus, generally, gyri have negative values, while sulci have positive values. Each depth value was also normalized by the deepest point in the given hemisphere. Surface area (mm²) was calculated with the FreeSurfer mris_anatomical_stats function

(https://surfer.nmr.mgh.harvard.edu/fswiki/mris_anatomical_stats 🖄). The morphological features of all LPC/LPOJ sulci are documented in Supplementary Fig. 7.

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Architecturally, we compared cortical thickness and myelination, as in our prior work in other cortical expanses (Miller et al., 2021 , Voorhies et al., 2021 , Willbrand et al., 2023 , 2022a). Mean gray matter cortical thickness (mm) was extracted using the FreeSurfer mris_anatomical_stats function. To quantify myelin content, we used the T1-w/T2-w maps for each hemisphere, an in vivo myelination proxy (Glasser and Van Essen, 2011). To generate the T1-w/T2-w maps, two T1-w and T2-w structural MR scans from each participant were registered together and averaged as part of the HCP processing pipeline (Glasser et al., 2013). The averaging helps to reduce motion-related effects or blurring. Additionally, and as described by Glasser and colleagues (Glasser et al., 2013), the T1-w/T2-w images were bias-corrected for distortion effects using field maps. We then extracted the average T1-w/T2-w ratio values across each vertex for each sulcus using custom Python code (Miller et al., 2021). The architectural features of all LPC/LPOI sulci are documented in Supplementary Fig. 7.

To assess whether these four metrics differed between the slocs-v and surrounding sulci (cSTS3 and ITOS), we ran a repeated measure analysis of variance (rm-ANOVA) with the withinparticipant effects of sulcus (slocs-v, cSTS3, and ITOS), metric (surface area, depth, cortical thickness, and myelination), and hemisphere (left and right). Rm-ANOVAs (including sphericity correction) were implemented with the aov_ez function from the afex R package. Effect sizes for the ANOVAs are reported with the partial eta-squared metric (n2). Post-hoc analyses were computed with the emmeans function from the emmeans R package (*p*-values corrected with Tukey's method). We also repeated these analyses for the three cSTS components (Petrides, 2019 ^{c2}; Segal and Petrides, 2012 ^{c2}) and the two intermediate parietal sulcal components (ips: aipsJ and pips (Petrides, 2019 ^{c2}; Zlatkina and Petrides, 2014 ^{c2}); detailed in the Supplementary Results and Supplementary Fig. 8) as these components, to our knowledge, have not been quantitatively compared in previous work.

Functional analyses

To determine if the slocs-v is functionally distinct from surrounding sulci, we generated functional connectivity profiles using recently developed analyses (Miller et al., 2021 🖒; Willbrand et al., 2023a C, 2022a C). First, we used resting-state network parcellations for each individual participant from Kong and colleagues (Kong et al., 2019 C), who generated individual network definitions by applying a hierarchical Bayesian network algorithm to produce maps for each of the 17 networks in individual HCP participants. Importantly, this parcellation was conducted blind to both cortical folding and our sulcal definitions. Next, we resampled the network profiles for each participant onto the fsaverage cortical surface, and then to each native surface using CBIG tools (*https://github.com/ThomasYeoLab/CBIG* C). We then calculated the spatial overlap between a sulcus and each of the 17 individual resting-state networks via the Dice coefficient (*Equation* 1 C):

(1) DICE
$$(X, Y) = \frac{2 |X \cap Y|}{|X| + |Y|}$$

This process of calculating the overlap between each sulcus and the 17-network parcellation generated a "connectivity fingerprint" for each sulcus in each hemisphere of each participant. We then ran an rm-ANOVA with within-participant factors of sulcus (slocs-v, cSTS3, and ITOS), network (17 networks), and hemisphere (left and right) to determine if the network profiles (i.e., the Dice coefficient overlap with each network) of the slocs-v was differentiable from the surrounding sulci (i.e., cSTS3 and ITOS). Here we discuss effects related to networks that at least showed minor overlap with one sulcus (i.e., Dice \geq .10). As in the prior analysis, we also repeated these analyses for the three cSTS components and the two intermediate parietal sulcal components (Supplementary Results and Supplementary Fig. 8).



Behavioral analyses

Model selection

The analysis relating sulcal morphology to spatial orientation and/or reasoning consisted of using a cross-validated least absolute shrinkage and selection operator (LASSO) regression to select the sulci that explained the most variance in the data and determined how much variance is explained by sulcal depth as a predictor of behavior, as implemented in our previous work (Voorhies et al., 2021 2; Willbrand et al., 2023b 2; Yao et al., 2022 2). The depths of all LPC/LPOJ sulci were included as predictors in the LASSO regression model (Supplementary Methods for details on demographic control variables). As the shrinkage parameter (alpha) increases, it decreases the coefficient of each of the sulci to zero except for those with the strongest association. Therefore, this technique highlights the sulci whose morphology was most closely related to behavior. We used cross-validation to optimize the shrinking parameter for the LASSO regression. Conventionally, we selected the model parameters that minimized the cross-validated mean squared error (MSE_{cv}) (Heinze et al., 2018 🖸). The optimization was performed with the GridSearchCV function sklearn in Python, which allowed us to determine the model parameters minimizing the MSE_{cv}. To evaluate the performance of any model selected by the LASSO regression, as in prior work (Voorhies et al., 2021 2; Willbrand et al., 2023b 2; Yao et al., 2022 2), we measured the model performance for the relevant behavioral task using nested model comparison. With leave-one-out cross-validation (LooCV), we compared the LASSO-selected model with the predictors to a model with all left hemisphere sulci as predictors.

Assessing morphological and behavioral specificity

To assess whether our findings generalized to other anatomical features, we considered cortical thickness, which is consistently studied in cognitive neuroscience studies relating morphology to cognition (Dickerson et al., 2008 🖒; Gogtay et al., 2004 🖒; Voorhies et al., 2021 🖒; Willbrand et al., 2023b C; Yao et al., 2022 C). To do so, we replaced sulcal depth with cortical thickness as the predictive metric in our LASSO-selected model. As with depth, the model was fit to the data with LooCV. To compare the thickness model to the depth model, we used the Akaike Information Criterion (AIC), which provides an estimate of in-sample prediction error and is suitable for nonnested model comparison. By comparing AIC scores, we are able to assess the relative performance of the two models. If the Δ AIC is > 2, it suggests an interpretable difference between models. If the Δ AIC is > 10, it suggests a strong difference between models, with the lower AIC value indicating the preferred model (Wagenmakers and Farrell, 2004 🖒). To also ascertain whether the relationship between LPC/LPOJ sulcal depth and cognition is specific to spatial orientation performance, or transferable to other general measures of cognitive processing, we investigated the generalizability of the sulcal-behavior relationship to another widely used measure of cognitive functioning: processing speed (Kail and Salthouse, 1994^{c2}). Specifically, we used LooCV to predict processing speed instead of spatial orientation score. As with thickness, we compared the two models with the AIC.

Situating the slocs-v within modern group-level cortical parcellations

To putatively relate the slocs-v to modern multimodal (HCP multimodal parcellation, HCP-MMP (Glasser et al., 2016^[C3])) and cytoarchitectural (Julich-Brain atlas (Amunts et al., 2020^[C3])) regions of the cerebral cortex located in fsaverage template space, we quantified the Dice coefficient overlap between the slocs-v of each participant (resampled to fsaverage space) and the individual regions of interest comprising the HCP-MMP and Julich-Brain parcellations.



Retinotopic response mapping of LPC/LPOJ sulci

To assess whether any of the LPC/LPOJ sulci related to retinotopic representations, we leveraged population receptive field mapping data (Benson et al., 2018). For each sulcal MPM (as the retinotopic data were only available in this template space), we extracted the mean R^2 values (i.e., the percentage of variance in each vertex explained by the population receptive field model) for vertices that showed meaningful retinotopic responses across participants (thresholded at $R^2 > 10\%$) (Mackey et al., 2017).

Competing interests

The authors declare no competing financial interests.

Data availability

The processed data required to perform all statistical analyses and reproduce all Figures, as well as the probability maps, are available on GitHub (*https://github.com/cnl-berkeley/stable_projects*). Anonymized HCP neuroimaging data are publicly available on ConnectomeDB (*db.humanconnectome.org* (*https://db.humanconnectome.org/*)). Raw data will be made available from the corresponding author upon request.

Code availability

The code is available on GitHub (*https://github.com/cnl-berkeley/stable_projects* ⁽²⁾) and Open Science Framework (*https://osf.io/7fwqk*/⁽²⁾).

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References

Allen EJ, St-Yves G, Wu Y, Breedlove JL, Prince JS, Dowdle LT, Nau M, Caron B, Pestilli F, Charest I, Hutchinson JB, Naselaris T, Kay K. (2022) **A massive 7T fMRI dataset to bridge cognitive neuroscience and artificial intelligence** *Nat Neurosci* **25**:116– 126 https://doi.org/10.1038/s41593-021-00962-x

Amiez C, Sallet J, Hopkins WD, Meguerditchian A, Hadj-Bouziane F, Ben Hamed S, Wilson CRE, Procyk E, Petrides M (2019) **Sulcal organization in the medial frontal cortex provides insights into primate brain evolution** *Nat Commun* **10**:1–14 https://doi.org/10.1038/s41467-019-11347-x

Amiez C, Wilson CRE, Procyk E (2018) **Variations of cingulate sulcal organization and link with cognitive performance** *Sci Rep* **8**:1–13 https://doi.org/10.1038/s41598-018-32088-9

Ammons CJ, Winslett M-E, Bice J, Patel P, May KE, Kana RK (2021) **The mid-fusiform sulcus in autism spectrum disorder: Establishing a novel anatomical landmark related to face processing** *Autism Res* **14**:53–64 https://doi.org/10.1002/aur.2425

Amunts K, Mohlberg H, Bludau S, Zilles K (2020) **Julich-Brain: A 3D probabilistic atlas of the human brain's cytoarchitecture** *Science* **369**:988– 992 https://doi.org/10.1126/science.abb4588

Armstrong E, Schleicher A, Omran H, Curtis M, Zilles K (1995) **The ontogeny of human** gyrification *Cereb Cortex* **5**:56–63 https://doi.org/10.1093/cercor/5.1.56

Barch DM, Burgess GC, Harms MP, Petersen SE, Schlaggar BL, Corbetta M, Glasser MF, Curtiss S, Dixit S, Feldt C, Nolan D, Bryant E, Hartley T, Footer O, Bjork JM, Poldrack R, Smith S, Johansen-Berg H, Snyder AZ, Van Essen DC, WU-Minn HCP Consortium (2013) **Function in the human connectome: task-fMRI and individual differences in behavior** *Neuroimage* **80**:169–189 https://doi.org/10.1016/j.neuroimage.2013.05.033

Benson NC, Jamison KW, Arcaro MJ, Vu AT, Glasser MF, Coalson TS, Van Essen DC, Yacoub E, Ugurbil K, Winawer J, Kay K. (2018) **The Human Connectome Project 7 Tesla retinotopy dataset: Description and population receptive field analysis** *J Vis* **18** https://doi.org/10.1167/18.13.23

Bodin C, Pron A, Le Mao M, Régis J, Belin P, Coulon O. (2021) Plis de passage in the superior temporal sulcus: Morphology and local connectivity *Neuroimage*225 https://doi.org/10.1016/j.neuroimage.2020.117513

Bodin C, Takerkart S, Belin P, Coulon O (2018) **Anatomo-functional correspondence in the superior temporal sulcus** *Brain Struct Funct* **223**:221–232 https://doi.org/10.1007/s00429-017-1483-2

Borne L, Rivière D, Mancip M, Mangin J-F (2020) **Automatic labeling of cortical sulci using patch- or CNN-based segmentation techniques combined with bottom-up geometric constraints** *Med Image Anal* **62** https://doi.org/10.1016/j.media.2020.101651



Cachia A, Borst G, Jardri R, Raznahan A, Murray GK, Mangin J-F, Plaze M (2021) **Towards Deciphering the Fetal Foundation of Normal Cognition and Cognitive Symptoms From Sulcation of the Cortex** *Front Neuroanat* **15** https://doi.org/10.3389/fnana.2021.712862

Caspers S, Eickhoff SB, Rick T, von Kapri A, Kuhlen T, Huang R, Shah NJ, Zilles K. (2011) **Probabilistic fibre tract analysis of cytoarchitectonically defined human inferior parietal lobule areas reveals similarities to macaques** *Neuroimage* **58**:362– 380 https://doi.org/10.1016/j.neuroimage.2011.06.027

Chi JG, Dooling EC, Gilles FH (1977) **Gyral development of the human brain** *Ann Neurol* **1**:86–93 https://doi.org/10.1002/ana.410010109

Connolly CJ. (1950) External morphology of the primate brain

Cottaar M, Bastiani M, Boddu N, Glasser MF, Haber S, van Essen DC, Sotiropoulos SN, Jbabdi S. (2021) **Modelling white matter in gyral blades as a continuous vector field** *Neuroimage* **227** https://doi.org/10.1016/j.neuroimage.2020.117693

Cunningham DJ (1892) Contribution to the Surface Anatomy of the Cerebral Hemispheres

Dale AM, Fischl B, Sereno MI (1999) **Cortical surface-based analysis. I. Segmentation and surface reconstruction** *Neuroimage* **9**:179–194 https://doi.org/10.1006/nimg.1998.0395

Dickerson BC, Fenstermacher E, Salat DH, Wolk DA, Maguire RP, Desikan R, Pacheco J, Quinn BT, Van der Kouwe A, Greve DN, Blacker D, Albert MS, Killiany RJ, Fischl B. (2008) **Detection of cortical thickness correlates of cognitive performance: Reliability across MRI scan sessions, scanners, and field strengths** *Neuroimage* **39**:10–18 https://doi.org/10.1016/j.neuroimage.2007.08.042

Drudik K, Zlatkina V, Petrides M (2022) **Morphological patterns and spatial probability maps** of the superior parietal sulcus in the human brain *Cereb Cortex* https://doi.org/10.1093/cercor/bhac132

Fischl B, Sereno MI, Dale AM (1999) **Cortical surface-based analysis. II: Inflation, flattening, and a surface-based coordinate system** *Neuroimage* **9**:195– 207 https://doi.org/10.1006/nimg.1998.0396

Fornito A, Yücel M, Wood S, Stuart GW, Buchanan J-A, Proffitt T, Anderson V, Velakoulis D, Pantelis C (2004) **Individual differences in anterior cingulate/paracingulate morphology are related to executive functions in healthy males** *Cereb Cortex* **14**:424– 431 https://doi.org/10.1093/cercor/bhh004

Garrison JR, Fernyhough C, McCarthy-Jones S, Haggard M, Australian Schizophrenia Research Bank, Simons JS (2015) **Paracingulate sulcus morphology is associated with hallucinations in the human brain** *Nat Commun* **6** https://doi.org/10.1038/ncomms9956

Glasser MF, Coalson TS, Robinson EC, Hacker CD, Harwell J, Yacoub E, Ugurbil K, Andersson J, Beckmann CF, Jenkinson M, Smith SM, Van Essen DC. (2016) **A multi-modal parcellation of human cerebral cortex** *Nature* **536**:171–178 https://doi.org/10.1038/nature18933

Glasser MF, Sotiropoulos SN, Wilson JA, Coalson TS, Fischl B, Andersson JL, Xu J, Jbabdi S, Webster M, Polimeni JR, Van Essen DC, Jenkinson M, WU-Minn HCP Consortium (2013) **The minimal preprocessing pipelines for the Human Connectome Project** *Neuroimage* **80**:105–124 https://doi.org/10.1016/j.neuroimage.2013.04.127



Glasser MF, Van Essen DC. (2011) **Mapping human cortical areas in vivo based on myelin content as revealed by T1- and T2-weighted MRI** *J Neurosci* **31**:11597– 11616 https://doi.org/10.1523/JNEUROSCI.2180-11.2011

Gogtay N, Giedd JN, Lusk L, Hayashi KM, Greenstein D, Vaituzis AC, Nugent TF, Herman DH, Clasen LS, Toga AW, Rapoport JL, Thompson PM (2004) **Dynamic mapping of human cortical development during childhood through early adulthood** *Proc Natl Acad Sci U S A* **101**:8174–8179 https://doi.org/10.1073/pnas.0402680101

Goodale MA, Milner AD (1992) **Separate visual pathways for perception and action** *Trends Neurosci* **15**:20–25 https://doi.org/10.1016/0166-2236(92)90344-8

Gratton C, Nelson SM, Gordon EM (2022) **Brain-behavior correlations: Two paths toward reliability** *Neuron* https://doi.org/10.1016/j.neuron.2022.04.018

Gur RC, Alsop D, Glahn D, Petty R, Swanson CL, Maldjian JA, Turetsky BI, Detre JA, Gee J, Gur RE (2000) **An fMRI study of sex differences in regional activation to a verbal and a spatial task** *Brain Lang* **74**:157–170 https://doi.org/10.1006/brln.2000.2325

Harper L, Lindberg O, Bocchetta M, Todd EG, Strandberg O, van Westen D, Stomrud E, Landqvist Waldö M, Wahlund L-O, Hansson O, Rohrer JD, Santillo A. (2022) **Prenatal Gyrification Pattern Affects Age at Onset in Frontotemporal Dementia** *Cereb Cortex* https://doi.org/10.1093/cercor/bhab457

Harvey BM, Fracasso A, Petridou N, Dumoulin SO (2015) **Topographic representations of object size and relationships with numerosity reveal generalized quantity processing in human parietal cortex** *Proc Natl Acad Sci U S A* **112**:13525– 13530 https://doi.org/10.1073/pnas.1515414112

Harvey BM, Klein BP, Petridou N, Dumoulin SO (2013) **Topographic representation of numerosity in the human parietal cortex** *Science* **341**:1123– 1126 https://doi.org/10.1126/science.1239052

Hathaway CB, Voorhies WI, Sathishkumar N, Mittal C, Yao JK, Miller JA, Parker BJ, Weiner KS (2023) **Defining putative tertiary sulci in lateral prefrontal cortex in chimpanzees using human predictions** *Brain Struct Funct* https://doi.org/10.1007/s00429-023-02638-7

Heinze G, Wallisch C, Dunkler D (2018) Variable selection - A review and recommendations for the practicing statistician *Biom J* **60**:431–449 https://doi.org/10.1002/bimj.201700067

Humphreys GF, Tibon R (2023) **Dual-axes of functional organisation across lateral parietal cortex: the angular gyrus forms part of a multi-modal buffering system** *Brain Struct Funct* **228**:341–352 https://doi.org/10.1007/s00429-022-02510-0

Kail R, Salthouse TA (1994) **Processing speed as a mental capacity** *Acta Psychol* **86**:199–225 https://doi.org/10.1016/0001-6918(94)90003-5

Karnath HO (1997) **Spatial orientation and the representation of space with parietal lobe lesions** *Philos Trans R Soc Lond B Biol Sci* **352**:1411–1419 https://doi.org/10.1098/rstb.1997.0127

Konen CS, Kastner S (2008) **Representation of eye movements and stimulus motion in topographically organized areas of human posterior parietal cortex** *J Neurosci* **28**:8361– 8375 https://doi.org/10.1523/JNEUROSCI.1930-08.2008



Kong R, Li J, Orban C, Sabuncu MR, Liu H, Schaefer A, Sun N, Zuo X-N, Holmes AJ, Eickhoff SB, Yeo BTT (2019) **Spatial Topography of Individual-Specific Cortical Networks Predicts Human Cognition, Personality, and Emotion** *Cereb Cortex* **29**:2533– 2551 https://doi.org/10.1093/cercor/bhy123

Leroy F, Cai Q, Bogart SL, Dubois J, Coulon O, Monzalvo K, Fischer C, Glasel H, Van der Haegen L, Bénézit A, Lin C-P, Kennedy DN, Ihara AS, Hertz-Pannier L, Moutard M-L, Poupon C, Brysbaert M, Roberts N, Hopkins WD, Mangin J-F, Dehaene-Lambertz G. (2015) **New human-specific brain landmark: the depth asymmetry of superior temporal sulcus** *Proc Natl Acad Sci U S A* **112**:1208–1213 https://doi.org/10.1073/pnas.1412389112

Lopez-Persem A, Verhagen L, Amiez C, Petrides M, Sallet J (2019) **The Human Ventromedial Prefrontal Cortex: Sulcal Morphology and Its Influence on Functional Organization** *J Neurosci* **39**:3627–3639 https://doi.org/10.1523/JNEUROSCI.2060-18.2019

Lyu I, Bao S, Hao L, Yao J, Miller JA, Voorhies W, Taylor WD, Bunge SA, Weiner KS, Landman BA (2021) Labeling lateral prefrontal sulci using spherical data augmentation and contextaware training *NeuroImage* https://doi.org/10.1016/j.neuroimage.2021.117758

Mackey WE, Winawer J, Curtis CE (2017) **Visual field map clusters in human frontoparietal cortex** *Elife* **6** https://doi.org/10.7554/eLife.22974

Miller JA, Voorhies WI, Li X, Raghuram I, Palomero-Gallagher N, Zilles K, Sherwood CC, Hopkins WD, Weiner KS (2020) **Sulcal morphology of ventral temporal cortex is shared between humans and other hominoids** *Sci Rep* **10** https://doi.org/10.1038/s41598-020-73213-x

Miller JA, Voorhies WI, Lurie DJ, D'Esposito M, Weiner KS (2021) **Overlooked Tertiary Sulci** Serve as a Meso-Scale Link between Microstructural and Functional Properties of Human Lateral Prefrontal Cortex *J Neurosci* **41**:2229–2244 https://doi.org/10.1523/JNEUROSCI.2362-20.2021

Nakamura M, Nestor PG, Shenton ME (2020) **Orbitofrontal Sulcogyral Pattern as a Transdiagnostic Trait Marker of Early Neurodevelopment in the Social Brain** *Clin EEG Neurosci* **51**:275–284 https://doi.org/10.1177/1550059420904180

Naselaris T, Allen E, Kay K (2021) **Extensive sampling for complete models of individual brains** *Current Opinion in Behavioral Sciences* **40**:45– 51 https://doi.org/10.1016/j.cobeha.2020.12.008

Natu VS, Arcaro MJ, Barnett MA, Gomez J, Livingstone M, Grill-Spector K, Weiner KS (2021) Sulcal Depth in the Medial Ventral Temporal Cortex Predicts the Location of a Place-Selective Region in Macaques, Children, and Adults *Cereb Cortex* 31:48– 61 https://doi.org/10.1093/cercor/bhaa203

Parker BJ, Voorhies WI, Jiahui G, Miller JA, Willbrand E, Hallock T, Furl N, Garrido L, Duchaine B, Weiner KS (2023) **Hominoid-specific sulcal variability is related to face perception ability** *Brain Struct Funct* https://doi.org/10.1007/s00429-023-02611-4

Petrides M (2019) Atlas of the Morphology of the Human Cerebral Cortex on the Average MNI Brain

Pron A, Deruelle C, Coulon O (2021) **U-shape short-range extrinsic connectivity organisation around the human central sulcus** *Brain Struct Funct* **226**:179– 193 https://doi.org/10.1007/s00429-020-02177-5



Régis J, Mangin J-F, Ochiai T, Frouin V, Riviére D, Cachia A, Tamura M, Samson Y (2005) **"Sulcal Root" Generic Model: a Hypothesis to Overcome the Variability of the Human Cortex Folding Patterns** *Neurol Med Chir* **45**:1–17 https://doi.org/10.2176/nmc.45.1

Reveley C, Seth AK, Pierpaoli C, Silva AC, Yu D, Saunders RC, Leopold DA, Ye FQ (2015) Superficial white matter fiber systems impede detection of long-range cortical connections in diffusion MR tractography *Proc Natl Acad Sci U S A* **112**:E2820– 8 https://doi.org/10.1073/pnas.1418198112

Roell M, Cachia A, Matejko AA, Houdé O, Ansari D, Borst G (2021) **Sulcation of the intraparietal sulcus is related to symbolic but not non-symbolic number skills** *Dev Cogn Neurosci* **51** https://doi.org/10.1016/j.dcn.2021.100998

Rosenberg MD, Finn ES (2022) **How to establish robust brain-behavior relationships without thousands of individuals** *Nat Neurosci* https://doi.org/10.1038/s41593-022-01110-9

Sanides F (1964) **Structure and function of the human frontal lobe** *Neuropsychologia* **2**:209–219 https://doi.org/10.1016/0028-3932(64)90005-3

Schilling K, Gao Y, Janve V, Stepniewska I, Landman BA, Anderson AW (2018) **Confirmation of a** gyral bias in diffusion MRI fiber tractography *Hum Brain Mapp* **39**:1449– 1466 https://doi.org/10.1002/hbm.23936

Schilling KG, Archer D, Rheault F, Lyu I, Huo Y, Cai LY, Bunge SA, Weiner KS, Gore JC, Anderson AW, Landman BA (2023) **Superficial white matter across development, young adulthood, and aging: volume, thickness, and relationship with cortical features** *Brain Struct Funct* **228**:1019–1031 https://doi.org/10.1007/s00429-023-02642-x

Schurz M, Tholen MG, Perner J, Mars RB, Sallet J (2017) **Specifying the brain anatomy underlying temporo-parietal junction activations for theory of mind: A review using probabilistic atlases from different imaging modalities** *Hum Brain Mapp* **38**:4788– 4805 https://doi.org/10.1002/hbm.23675

Segal E, Petrides M (2012) **The morphology and variability of the caudal rami of the superior temporal sulcus** *Eur J Neurosci* **36**:2035–2053 https://doi.org/10.1111/j.1460-9568.2012.08109.x

Steel A, Billings MM, Silson EH, Robertson CE (2021) **A network linking scene perception and** spatial memory systems in posterior cerebral cortex *Nat Commun* 12 https://doi.org/10.1038/s41467-021-22848-z

Tomaiuolo F, Giordano F (2016) **Cerebal sulci and gyri are intrinsic landmarks for brain navigation in individual subjects: an instrument to assist neurosurgeons in preserving cognitive function in brain tumour surgery (Commentary on Zlatkina et al.)** *Eur J Neurosci* https://doi.org/10.1111/ejn.13072

Tomaiuolo F, Raffa G, Morelli A, Rizzo V, Germanó A, Petrides M (2022) **Sulci and gyri are topological cerebral landmarks in individual subjects: a study of brain navigation during tumour resection** *Eur J Neurosci* **55**:2037–2046 https://doi.org/10.1111/ejn.15668

Van Essen DC, Donahue CJ, Glasser MF. (2018) **Development and Evolution of Cerebral and Cerebellar Cortex** *Brain Behav Evol* **91**:158–169 https://doi.org/10.1159/000489943



Van Essen DC, Jbabdi S, Sotiropoulos SN, Chen C, Dikranian K, Coalson T, Harwell J, Behrens TEJ, Glasser MF. (2014) **Mapping connections in humans and non-human primatesDiffusion MRI** *Elsevier* :337–358 https://doi.org/10.1016/b978-0-12-396460-1.00016-0

Vendetti MS, Bunge SA (2014) **Evolutionary and developmental changes in the lateral frontoparietal network: a little goes a long way for higher-level cognition** *Neuron* **84**:906– 917 https://doi.org/10.1016/j.neuron.2014.09.035

Voorhies WI, Miller JA, Yao JK, Bunge SA, Weiner KS (2021) **Cognitive insights from tertiary** sulci in prefrontal cortex *Nat Commun* **12** https://doi.org/10.1038/s41467-021-25162-w

Wagenmakers E-J, Farrell S (2004) **AIC model selection using Akaike weights** *Psychon Bull Rev* **11**:192–196 https://doi.org/10.3758/bf03206482

Wang J, Fan L, Zhang Y, Liu Y, Jiang D, Zhang Y, Yu C, Jiang T (2012) **Tractography-based parcellation of the human left inferior parietal lobule** *Neuroimage* **63**:641– 652 https://doi.org/10.1016/j.neuroimage.2012.07.045

Wang L, Mruczek REB, Arcaro MJ, Kastner S (2015) **Probabilistic Maps of Visual Topography in Human Cortex** *Cereb Cortex* **25**:3911–3931 https://doi.org/10.1093/cercor/bhu277

Weiner KS (2019) **The Mid-Fusiform Sulcus (sulcus sagittalis gyri fusiformis)** *Anat Rec* **302**:1491–1503 https://doi.org/10.1002/ar.24041

Welker W, Jones EG, Peters A (1990) **Why Does Cerebral Cortex Fissure and Fold?** *Cerebral Cortex: Comparative Structure and Evolution of Cerebral Cortex, Part II* :3–136 https://doi.org/10.1007/978-1-4615-3824-0_1

Wendelken C (2014) **Meta-analysis: how does posterior parietal cortex contribute to reasoning?** *Front Hum Neurosci* **8** https://doi.org/10.3389/fnhum.2014.01042

Willbrand EH, Bunge SA, Weiner KS (2023) **Neuroanatomical and functional dissociations between variably present anterior lateral prefrontal sulci** *bioRxiv* https://doi.org/10.1101/2023.05.25.542301

Willbrand EH, Ferrer E, Bunge SA, Weiner KS (2023) **Development of human lateral prefrontal sulcal morphology and its relation to reasoning performance** *J Neurosci* https://doi.org/10.1523/JNEUROSCI.1745-22.2023

Willbrand EH, Maboudian SA, Kelly JP, Parker BJ, Foster BL, Weiner KS (2023) **Sulcal** morphology of posteromedial cortex substantially differs between humans and chimpanzees *Communications Biology* **6**:1–14 https://doi.org/10.1038/s42003-023-04953-5

Willbrand EH, Parker BJ, Voorhies WI, Miller JA, Lyu I, Hallock T, Aponik-Gremillion L, Koslov SR, Null N, Bunge SA, Foster BL, Weiner KS (2022) **Uncovering a tripartite landmark in posterior cingulate cortex** *Science Advances* **8** https://doi.org/10.1126/sciadv.abn9516

Willbrand EH, Voorhies WI, Yao JK, Weiner KS, Bunge SA (2022) **Presence or absence of a prefrontal sulcus is linked to reasoning performance during child development** *Brain Struct Funct* **227**:2543–2551 https://doi.org/10.1007/s00429-022-02539-1

Yao JK, Voorhies WI, Miller JA, Bunge SA, Weiner KS (2022) **Sulcal depth in prefrontal cortex: a novel predictor of working memory performance** *Cereb Cortex bhac* **173** https://doi.org/10.1093/cercor/bhac173



Zilles K, Palomero-Gallagher N, Amunts K (2013) **Development of cortical folding during evolution and ontogeny** *Trends Neurosci* **36**:275– 284 https://doi.org/10.1016/j.tins.2013.01.006

Zlatkina V, Petrides M (2014) **Morphological patterns of the intraparietal sulcus and the anterior intermediate parietal sulcus of Jensen in the human brain** *Proc Biol Sci* **281** https://doi.org/10.1098/rspb.2014.1493

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Editors

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Senior Editor **Michael Frank** Brown University, United States of America

Reviewer #1 (Public Review):

Summary: Ever-improving techniques allow the detailed capture of brain morphology and function to the point where individual brain anatomy becomes an important factor. This study



investigated detailed sulcal morphology in the parieto-occipital junction. Using cutting-edge methods, it provides important insights into local anatomy, individual variability, and local brain function. The presented work advances the field and will stimulate future research into this important area.

Strengths:

Detailed, very thorough methodology. Multiple raters mapped detailed sulci in a large cohort. The identified sulcal features and their functional and behavioural relevance are then studied using various complementary methods. The results provide compelling evidence for the importance of the described sulcal features and their proposed relationship to cortical brain function.

Weaknesses:

A detailed description/depiction of the various sulcal patterns is missing. A possible relationship between sucal morphology and individual demographics might provide more insight into anatomical variability. The unique dataset offers to opportunity to provide insights into laterality effects that should be explored.

Reviewer #2 (Public Review):

Summary: After manually labelling 144 human adult hemispheres in the lateral parietooccipital junction (LPOJ), the authors 1) propose a nomenclature for 4 previously unnamed highly variable sulci located between the temporal and parietal or occipital lobes, 2) focus on one of these newly named sulci, namely the ventral supralateral occipital sulcus (slocs-v) and compare it to neighbouring sulci to demonstrate its specificity (in terms of depth, surface area, gray matter thickness, myelination, and connectivity), 3) relate the morphology of a subgroup of sulci from the region including the slocs-v to the performance in a spatial orientation task, demonstrating behavioural and morphological specificity. In addition to these results, the authors propose an extended reflection on the relationship between these newly named landmarks and previous anatomical studies, a reflection about the slocs-v related to functional and cytoarchitectonic parcellations as well as anatomic connectivity and an insight about potential anatomical mechanisms relating sulcation and behaviour.

Strengths:

- To my knowledge, this is the first study addressing the variable tertiary sulci located between the superior temporal sulcus (STS) and intra-parietal sulcus (IPS).

- This is a very comprehensive study addressing altogether anatomical, architectural, functional and cognitive aspects.

- The definition of highly variable yet highly reproducible sulci such as the slocs-v feeds the community with new anatomo-functional landmarks (which is emphasized by the provision of a probability map in supp. mat., which in my opinion should be proposed in the main body).

- The comparison of different features between the slocs-v and similar sulci is useful to demonstrate their difference.

- The detailed comparison of the present study with state of the art contextualises and strengthens the novel findings.

- The functional study complements the anatomical description and points towards cognitive specificity related to a subset of sulci from the LPOJ

- The discussion offers a proposition of theoretical interpretation of the findings

- The data and code are mostly available online (raw data made available upon request).

Weaknesses:

- While three independent raters labelled all hemispheres, one single expert finalized the decision. Because no information is reported on the inter-rater variability, this somehow equates to a single expert labelling the whole cohort, which could result in biased labellings and therefore affect the reproducibility of the new labels.

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- 3 out of the 4 newly labelled sulci are only described in the very first part and never reused. This should be emphasized as it is far from obvious at first glance of the article.

- The tone of the article suggests a discovery of these 4 sulci when some of them have already been reported (as rightfully highlighted in the article), though not named nor studied specifically. This is slightly misleading as I interpret the first part of the article as a proposition of nomenclature rather than a discovery of sulci.

- The article never mentions the concept of merging of sulcal elements and the potential effect it could have on the labelling of the newly named variable sulci.

- The definition of the new sulci is solely based on their localization relative to other sulci which are themselves variable (e.g. the 3rd branch of the STS can show different locations and different orientation, potentially affecting the definition of the slocs-v). This is not addressed in the discussion.

- The new sulci are only defined in terms of localization relative to other sulci, and no other property is described (general length, depth, orientation, shape...), making it hard for a new observer to take labelling decisions in case of conflict.

- The very assertive tone of the article conveys the idea that these sulci are identifiable certainly in most cases, when by definition these highly variable tertiary sulci are sometimes very difficult to take decisions on.

- I am not absolutely convinced with the labelling proposed of a previously reported sulcus, namely the posterior intermediate parietal sulcus.

Assuming that the labelling of all sulci reported in the article is reproducible, the different results are convincing and in general, this study achieves its aims in defining more precisely the sulcation of the LPOJ and looking into its functional/cognitive value. This work clearly offers a finer understanding of sulcal pattern in this region, and lacks only little for the new markers to be convincingly demonstrated. An overall coherence of the labelling can still be inferred from the supplementary material which support the results and therefore the conclusions, yet, addressing some of the weaknesses listed above would greatly enhance the impact of this work. This work is important to the understanding of sulcal variability and its implications on functional and cognitive aspects.

Reviewer #3 (Public Review):

Summary: 72 subjects, and 144 hemispheres, from the Human Connectome Project had their parietal sulci manually traced. This identified the presence of previously undescribed shallow sulci. One of these sulci, the ventral supralateral occipital sulcus (slocs-v), was then demonstrated to have functional specificity in spatial orientation. The discussion furthermore provides an eloquent overview of our understanding of the anatomy of the parietal cortex, situating their new work into the broader field. Finally, this paper stimulates further debate about the relative value of detailed manual anatomy, inherently limited in participant numbers and areas of the brain covered, against fully automated processing that can cover thousands of participants but easily misses the kinds of anatomical details described here.

Strengths:

- This is the first paper describing the tertiary sulci of the parietal cortex with this level of detail, identifying novel shallow sulci and mapping them to behaviour and function.

- It is a very elegantly written paper, situating the current work into the broader field.
- The combination of detailed anatomy and function and behaviour is superb.

Weaknesses:

- the numbers of subjects are inherently limited both in number as well as in being typically developing young adults.

- while the paper begins by describing four new sulci, only one is explored further in greater detail.



- there is some tension between calling the discovered sulci new vs acknowledging they have already been reported, but not named.

- the anatomy of the sulci, as opposed to their relation to other sulci, could be described in greater detail.

Overall, to summarize, I greatly enjoyed this paper and believe it to be a highly valued contribution to the field.