**Title** 

Sulcal morphology of posteromedial cortex substantially differs between humans and chimpanzees

# **Authors**

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**Abstract** 

Recent studies identify a surprising coupling between evolutionarily new sulci and the functional organization of human posteromedial cortex (PMC). Yet, no study has compared this modern PMC sulcal patterning between humans and non-human hominoids. To fill this gap in knowledge, we first manually defined 918 sulci in 120 chimpanzee (*Pan Troglodytes*) hemispheres and 1619 sulci in 144 human hemispheres. We uncovered four new PMC sulci, and quantitatively identified species differences in incidence, depth, and surface area. Interestingly, some PMC sulci are more common in humans and others, in chimpanzees. Further, we found that the prominent marginal ramus of the cingulate sulcus differs significantly between species. Contrary to classic observations, the present results reveal that the surface anatomy of PMC substantially differs between humans and chimpanzees — findings which lay a foundation for better understanding the evolution of neuroanatomical-functional and neuroanatomical-behavioral relationships in this highly expanded region of the human cerebral cortex.

## Introduction

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A fundamental question in comparative biology and systems neuroscience is: What features of the brain are unique to humans? Key insights regarding what features of the brain are unique to humans have been gleaned from studies comparing anatomical and functional features of the human brain to features from the brains of our close evolutionary relative, the chimpanzee<sup>1-</sup> <sup>20</sup>. Of all the features to study, researchers particularly focus on the folds of the cerebral cortex, or sulci, as they generally track with evolutionary complexity<sup>21</sup>. For example, while mice and marmosets have rather smooth, lissencephalic cerebral cortices, 60-70% of the cerebral cortex in hominoids is buried within sulci<sup>3,22</sup>. Intriguingly, recent studies have identified "evolutionarily new" shallow sulci in association cortices in hominoid brains that have been linked to functional organization across a broad array of cognitive domains (e.g., 14,19,23-38), several of which reflect cognitive abilities that are arguably unique to humans. Building on this previous work, we compared the sulcal patterning of the posteromedial cortex (PMC) — a region on the medial cortical surface that includes the posterior cingulate, retrosplenial, and precuneal cortices<sup>39</sup> between humans and chimpanzees with a particular emphasis on the smaller, shallower, and relatively overlooked "evolutionarily new" cortical indentations.

The sulcal organization of PMC has been under-documented, even in the most recent neuroanatomical treatises (e.g., 40,41). Nevertheless, PMC is critically important in hominoids as it contains regions implicated in the default mode and cognitive control networks 42-47 with complex structural and functional connections 39,44,47,48. PMC is also implicated in many complex cognitive abilities 43,47,49-52 and is particularly susceptible to neurodegenerative disease 50. Thus, quantifying the similarities and differences in the PMC sulcal patterning between chimpanzees and humans will not only shed light on the comparative neuroanatomy of PMC between species, but also

provide understanding regarding structural-functional relationships between species with potential cognitive insights<sup>53</sup>.

While it is known that the larger (primary) sulci within PMC are present in chimpanzees<sup>54–56</sup> and the inframarginal sulcus — a newly uncovered smaller (tertiary) PMC sulcus — is variably present in chimpanzees<sup>20</sup>, the phylogenetic emergence of a majority of recently clarified PMC sulci<sup>20</sup> has yet to be compared between chimpanzees and humans. Therefore, in the present study, we comprehensively examined the PMC sulcal patterning between humans and chimpanzees using cortical surface reconstructions as in our prior work<sup>15,20,38</sup>. Our analyses were guided by three main questions. First, does the amount of PMC buried in sulci differ between humans and chimpanzees? Second, do the incidence rates of PMC sulci differ between species? Third, do the primary morphological features of these structures (i.e., depth and surface area) differ between species?

### Results

In order to answer these main questions, we examined the PMC of 72 young adult humans [from the Human Connectome Project (HCP; <a href="http://www.humanconnectomeproject.org/">http://www.humanconnectomeproject.org/</a>)] and 60 chimpanzees [from the National Chimpanzee Brain Resource (<a href="https://www.chimpanzeebrain.org/">https://www.chimpanzeebrain.org/</a>)]. These participants were used in prior work to assess the anatomical, functional, and evolutionary significance of a new tripartite landmark in PCC, the inframarginal sulcus (ifrms<sup>20</sup>), but the rest of the PMC sulci were not considered in these previous cross-species analyses until the present study.

To broadly determine how much of the PMC is sulcal vs. gyral in each species, we calculated how much of the regions corresponding to an automated parcellation of PMC in FreeSurfer<sup>57</sup> were buried in sulci (i.e., the percentage of vertices with values above zero in the .sulc

file<sup>58</sup>) via the Dice coefficient (**Fig. 1a**; **Materials and Methods**). Replicating prior postmortem work<sup>3,22,59</sup>, the majority of human PMC was buried in sulci (mean  $\pm$  std = 73.9  $\pm$  1.97%). Chimpanzee PMC was relatively less sulcated (mean  $\pm$  std = 67.4  $\pm$  3.69%; **Fig. 1b**). A linear mixed effects model (LME) with factors of *species* and *hemisphere* (controlling for differences in brain size), confirmed this large difference between species (main effect of *species*: F(1, 130) = 220.57, p < .0001,  $\eta = 0.63$ ; no hemispheric differences: p > .24; **Fig. 1b**).

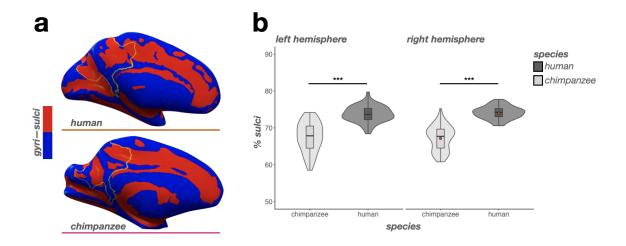
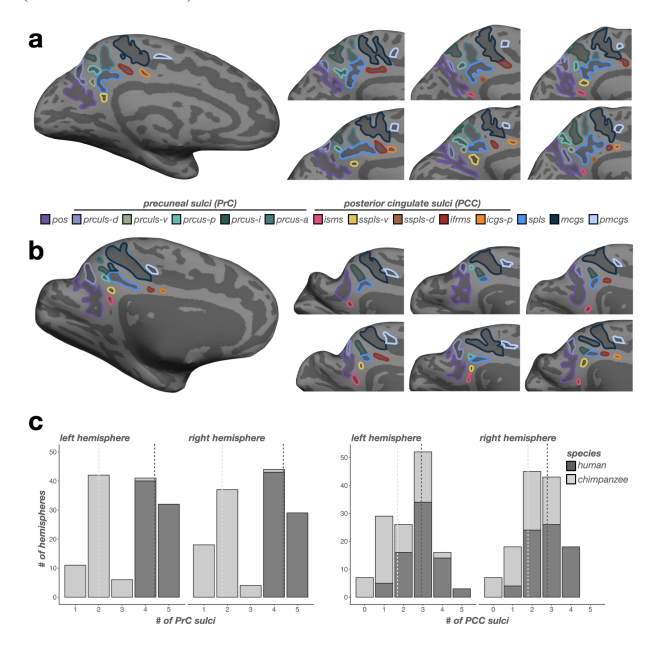


Figure 1. The percentage of PMC buried in sulci differs between humans and chimpanzees. a. Inflated human (top) and chimpanzee (bottom) right hemisphere cortical surface reconstructions (mirrored for visualization purposes). The outline of automatically defined PMC from the Destrieux parcellation<sup>57</sup> is indicated in yellow. The FreeSurfer sulc file<sup>58</sup> is overlaid on each surface (Sulci: red; Gyri: blue). These surfaces present the average PMC sulcation for each species (Human: 73.9%; Chimpanzee: 67.4%). The lines below each surface correspond to the colored individual dots on the plot to the right. b. Violin plots (box plot and kernel density estimate) visualizing the percentage of PMC in sulci (percentage values are out of 100) as a function of species (x-axis) and hemisphere (left: left hemisphere; right: right hemisphere). The significant difference in PMC sulcation between species (as a result of the main effect of species) is indicated with asterisks (\*\*\* p < .001).

Next, we manually defined sulci in precuneal (PrC) and posterior cingulate cortices (PCC) — which are subregions of the PMC<sup>20,39,47</sup> — in all human and chimpanzee brains ( **Materials** and **Methods** for a detailed description of these sulci). All PMC sulci were defined on cortical reconstructions from FreeSurfer (v6.0.0, <u>surfer.nmr.mgh.harvard.edu</u>; **Fig. 2** for example hemispheres; Supplementary Figs. 1-2 for all human and chimpanzee brains). Once all sulci were

defined, we quantified the average sulcal depth (normalized to the max depth in each hemisphere) and surface area (normalized to the total surface area of each hemisphere) of each PMC sulcus (Materials and Methods).



**Figure 2. Humans have more PMC sulci than chimpanzees across hemispheres in both PrC and PCC. a.** Left: An inflated cortical surface reconstruction of an individual human hemisphere. Sulci: dark gray; Gyri: light gray. Individual posteromedial (PMC) sulci are outlined according to the legend at the bottom. Right: Six example hemispheres zoomed in on the PMC depicting variations of sulcal incidence between participants. Right hemisphere images are mirrored so that all images have the same orientation. **b.** Same as a, but for chimpanzee hemispheres. **c.** Left: Incidence rates of precuneal (PrC) sulci (x-axis; see legend in a) across species (colors, see legend) for each

hemisphere (left: left hemisphere; right: right hemisphere). Dashed lines indicate the average number of sulci for each species in each hemisphere. Right: Same as the left, but for posterior cingulate (PCC) sulci.

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Once sulci were defined, we quantified the incidence rates of PMC sulci in three groups: i) sulci that border, or serve as the bounding perimeter of, PMC, ii) PCC sulci, and iii) PrC sulci. Crucially, this procedure revealed four new PMC sulci that were not considered in prior work of PMC sulcal morphology (e.g., <sup>20,44,55,59–63</sup>; **Fig. 2**, Supplementary Figs. 3-4). While we labeled and quantified the incidence rates of these four sulci across species for the first time, some present and modern anatomists often included an unlabeled sulcus in the location of some these sulci in their summary schematics (Supplementary Figs. 3-4). Further, these sulci were identifiable in postmortem chimpanzee hemispheres from a classic neuroanatomical atlas<sup>56</sup>, ensuring that FreeSurfer's computational processes did not artificially create shallow sulci (Supplementary Fig. 4). We described across-species comparisons for each group in turn below using logistic regression GLMs with species (human, chimpanzee) and hemisphere (left, right), as well as their interaction, as factors for sulcal presence. Afterwards, we compared the depth and surface area of PMC sulci between species using LMEs with species (human, chimpanzee), sulcus (PMC sulci), and hemisphere (left, right), as well as their interaction, as factors. Finally, we repeat these analyses on the incidence and morphology of the marginal ramus of the cingulate sulcus — a prominent sulcal landmark in PMC<sup>20,44,55,59–63</sup> that contrary to previous studies, differs substantially between species, which we show here.

Incidence rates of large and deep sulci that border PMC do not differ across species, including the newly identified premarginal branch of the cingulate sulcus (pmcgs)

We identified the following three large and deep sulci serving as borders of PMC: the marginal ramus of the cingulate sulcus (mcgs), splenial sulcus (spls), and parieto-occipital sulcus (pos).

Replicating prior post-mortem work<sup>54–56</sup>, we found that the mcgs, spls, and pos were present in all humans and chimpanzees (**Fig. 3**). We also identified a consistent sulcus just anterior to the mcgs (**Figs. 2-3**). As such, we refer to this sulcus as the premarginal branch of the cingulate sulcus (pmcgs). When present, the pmcgs is located just under the paracentral fossa and serves as the point where the mcgs breaks from the cingulate sulcus (cgs) proper (**Materials and Methods**). The pmcgs was clearly identifiable in 97.22% of left and 94.4% of right hemispheres in humans and in 100% of chimpanzees (**Fig. 3**). The incidence rates for these four sulci were comparable between species (no main effect of *species*:  $\chi 2 = 2.45$ , df = 1, p = 0.12; **Fig. 3**).

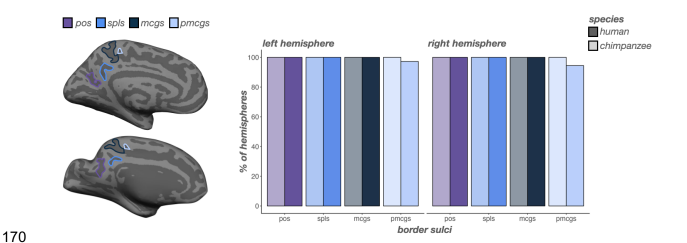


Figure 3. Incidence rates of sulci that border PMC are comparable between humans and chimpanzees. Left: An inflated cortical surface reconstruction of an individual human (top) and chimpanzee (bottom) hemisphere with sulci that border PMC outlined according to the legend at the top of the figure. Right: Bar plots visualizing incidence rates (percent of hemispheres) as a function of sulcus (x-axis), species (darker colors: human; lighter colors: chimpanzee), and hemisphere (left: left hemisphere; right: right hemisphere). Sulci are generally ordered posterior to anterior.

Incidence rates of PrC sulci differ substantially across species, including the newly identified ventral precuneal limiting sulcus (prculs-v)

In human PrC, the posterior (prcus-p), intermediate (prcus-i), and anterior precuneal sulci (prcus-a), as well as the dorsal precuneal limiting sulcus (prculs-d) were present in all hemispheres (**Fig.** 4). Previously, we<sup>20</sup> referred to this latter sulcus as the prculs (mirroring the label from a recent

neuroanatomical atlas<sup>60</sup>). However, here, we also consistently identified a ventral sulcal component in a comparable posterior plane as the dorsal prculs, but more inferiorly situated between the prculs-d and the spls (**Figs. 2, 4**). Consequently, we refer to this sulcus as the ventral prculs (prculs-v), which was identifiable in 44.44% of left and 40.28% of right hemispheres in humans (**Fig. 4**).

In contrast, PrC sulci were far more variable in chimpanzees. Generally, humans contained more sulci than chimpanzees in PrC (F(1, 130) = 1194.13, p < .0001,  $\eta 2 = 0.90$ ; no hemispheric differences: ps > .14; **Fig. 2c**, left). The prculs-d was the only sulcus comparably present between species (*left*: 96.67%; *right*: 96.67%; no main effect of *species*:  $\chi 2 = 3.19$ , df = 1, p = 0.07; **Fig. 4**). Interestingly, among the three recently identified prcus components<sup>20</sup>, prcus-i was the second most present PrC sulcus in chimpanzees, but was still less present than in humans (*left*: 76.67%; *right*: 73.33%; main effect of *species*:  $\chi 2 = 24.09$ , df = 1, p < .0001; **Fig. 4**). Conversely, prcus-p (*left*: 15%; *right*: 5%; main effect of *species*:  $\chi 2 = 125.39$ , df = 1, p < .0001) and prcus-a (*left*: 6.67%; *right*: 5%; main effect of *species*:  $\chi 2 = 150.56$ , df = 1, p < .0001) were quite rare in chimpanzees (**Fig. 4**). Finally, the newly identified prculs-v in humans was not identifiable in any chimpanzee hemispheres examined (main effect of *species*:  $\chi 2 = 47.30$ , df = 1, p < .0001; **Fig. 4**).

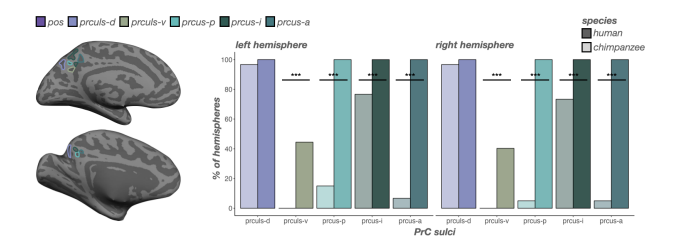


Figure 4. Incidence rates of precuneal (PrC) sulci are generally higher in humans than chimpanzees. Left: An inflated cortical surface reconstruction of an individual human (top) and chimpanzee (bottom) hemisphere with PrC sulci outlined according to the legend at the top of the figure. Right: Bar plots visualizing incidence rates (percent of hemispheres) as a function of sulcus (x-axis), species (darker colors: human; lighter colors: chimpanzee), and hemisphere (left: left hemisphere; right: right hemisphere). Sulci are generally ordered posterior to anterior. Lines and asterisks highlight significant differences in incidence between species (\* p < .05, \*\*\* p < .001). The intermediate precuneal sulcus (prcus-i) is the most common of the three precuneal sulci in chimpanzees. In comparison to the consistency of the prcus-i, prcus-a and prcus-p are extremely rare in chimpanzees.

ventral subsplenial sulcus (sspls-v) and isthmus sulcus (isms) are identifiable as frequently or more frequently in chimpanzees than humans

Sulci in human PCC are more variable than those in human PrC (**Fig. 2**)<sup>20</sup>. Generally, humans contained more sulci in PCC (F(1, 130) = 63.86, p < .0001, η2 = 0.33; no hemispheric differences: ps > .42; **Fig. 2c**, right) than chimpanzees. As shown previously, the inframarginal sulcus (ifrms) is the only PCC sulcus present in 100% of human hemispheres (**Fig. 5**)<sup>20</sup>. The ifrms is identifiable in 50% of chimpanzee hemispheres (**Fig. 5**)<sup>20</sup>. Anterior to the ifrms, the posterior intracingulate sulcus (icgs-p) was present in 65.28% of left and 66.67% right hemispheres in humans, and rarely identifiable in chimpanzees (*left*: 6.67%; *right*: 5%; main effect of *species*: χ2 = 53.74, df = 1, p < .0001; **Fig. 5**). Posterior to the ifrms, the dorsal subsplenial sulcus (sspls-d) was present in 47.22% of left and 50% right hemispheres in humans, and was not identifiable in any chimpanzee hemispheres (main effect of *species*: χ2 = 51.02, df = 1, p < .0001; **Fig. 5**).

While we previously referred to the sspls-d as the sspls<sup>20</sup>, here, we also identified an additional sulcus that was consistently identifiable just ventral and discontinuous with the dorsal component (**Figs. 2, 5**). As such, we refer to this newly-identified sulcus as the ventral sspls (sspls-v), which in humans was present in 66.67% of left hemispheres and 48.61% of right hemispheres (**Fig. 5**). Interestingly, the sspls-v showed no main effect of *species* ( $\chi$ 2 = 1.39, df = 1, p = 0.24), but an interaction between *species* and *hemisphere* ( $\chi$ 2 = 5.34, df = 1, p = 0.02), such that in

chimpanzees it was present in a comparable amount of left hemispheres to humans (56.67%; p = 0.24, Tukey's adjustment), but was present in more chimpanzee right hemispheres than human right hemispheres (66.67%; odds ratio = 0.75, p = 0.03, Tukey's adjustment; **Fig. 5**).

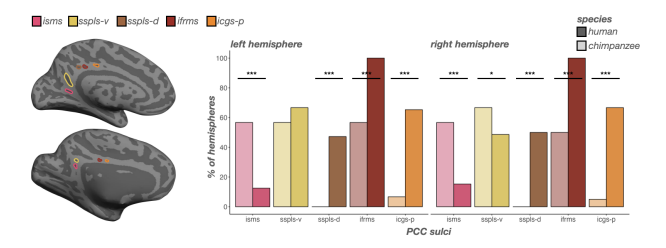
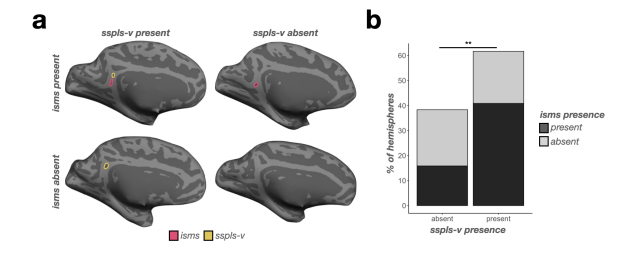


Figure 5. Incidence rates of posterior cingulate (PCC) sulci are variable between humans and chimpanzees. Left: An inflated cortical surface reconstruction of an individual human (top) and chimpanzee (bottom) hemisphere with PCC sulci outlined according to the legend at the top of the figure. Right: Bar plots visualizing incidence rates (percent of hemispheres) as a function of sulcus (x-axis), species (darker colors: human; lighter colors: chimpanzee), and hemisphere (left: left hemisphere; right: right hemisphere). Sulci are generally ordered posterior to anterior. Lines and asterisks highlight significant differences in incidence between species (\* p < .05, \*\*\* p < .001). The isms and sspls-v are more common in chimpanzees than humans. The sspls-d, ifrms, and icgs-p are more common in humans than chimpanzees. ifrms data from  $^{20}$ .

Finally, in a minority of humans (12.50% of left and 15.28% of right hemispheres), we could identify a previously undefined sulcus inferior to the sspls-v within the isthmus of the cingulate gyrus, which we termed the isthmus sulcus (isms; **Figs. 2**, **5**). The isms was present in more chimpanzee hemispheres (56.67% of left and right hemispheres) than humans (main effect of *species*:  $\chi 2 = 30.26$ , df = 1, p < .0001; **Fig. 5**). Interestingly, the incidence of the two more common PCC sulci in chimpanzees (sspls-v and isms) were related in chimpanzees ( $\chi 2 = 7.01$ , df = 1, p = 0.008), such that chimpanzees with an sspls-v were more likely to have an isms (odds ratio = 4.77; **Fig. 6**). No other sulcal incidence rates were related (ps > .10). To further summarize these relationships, there was a *PCC region* (dorsal PCC, ventral PCC) and *species* interaction on

sulcal presence ( $\chi 2 = 74.79$ , df = 1, p < .0001), such that, overall, dorsal PCC sulci (sspls-d, ifrms, and icgs-p) were less common in chimpanzees than humans (odds ratio = -2.33, p < .0001, Tukey's adjustment), whereas ventral PCC sulci (isms and sspls-v) were more common in chimpanzees than humans (odds ratio = 0.96, p < .0001, Tukey's adjustment; **Fig. 5**).



**Figure 6. Incidence of the sspls-v is related to the incidence of the isms in chimpanzees. a.** Four example inflated chimpanzee hemispheres displaying the four combinations of sspls-v (outlined in yellow when present) and isms (outlined in pink when present): both present (top left), sspls-v present (bottom left), isms present (top right), and both absent (bottom right). **b.** Bar plot visualizing the frequency of sspls-v and isms presence (colors, see legend). When the sspls-v is present, the isms is more likely present rather than absent; when the sspls-v is absent, the isms is likely to be absent (\*\* p < .01).

The relative depth and surface area of PMC sulci largely differ between chimpanzees and humans. In terms of depth, a LME model with predictors of sulcus, hemisphere, and species revealed three species-related findings. First, a main effect of species (F(1, 130) = 269.48, p < .0001,  $\eta = 0.67$ ) showed that human PMC sulci were relatively deeper than chimpanzees (**Fig. 7a**). Second, an interaction between species and sulcus (F(7, 1497) = 131.81, p < .0001,  $\eta = 0.38$ ) indicated more complex relationships at the individual-sulcus level. Post hoc analyses revealed three findings: i) the isms, pos, prculs-d, prcus-i, spls, and sspls-v were relatively deeper in humans than chimpanzees (p < .003, Tukey's adjustment), ii) the mcgs was relatively deeper in chimpanzees

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than humans (p = .04, Tukey's adjustment), and iii) the pmcgs was comparably deep between species (p = .45, Tukey's adjustment; Fig. 7a). Third, a three-way interaction among species, sulcus, and hemisphere (F(7, 1497) = 2.43, p = .01,  $\eta 2 = 0.01$ ) showed that the mcgs was deeper in chimpanzees in the left hemisphere (p = .04, Tukey's adjustment), but comparably deep in the right hemisphere (p = .35, Tukey's adjustment; **Fig. 7a**) compared to humans. In terms of surface area, a LME with predictors of sulcus, hemisphere, and species also revealed three species-related findings. First, a main effect of species (F(1, 130) = 6.51, p = .01,  $\eta 2 = 0.05$ ) showed that human PMC sulci were relatively larger than chimpanzees (Fig. 7b). Second, an interaction between species and sulcus (F(7, 1497) = 70.67, p < .0001,  $\eta 2 = 0.25$ ) indicated that the latter main effect was driven by differences at an individual-sulcus level. Post hoc analyses revealed three findings: i) the spls, prculs-d, and prcus-i were relatively larger in humans than chimpanzees (ps < .0001, Tukey's adjustment), ii) the pos, mcgs, and pmcgs were relatively larger in chimpanzees than humans (ps < .02, Tukey's adjustment), and iii) the isms and sspls-v were comparably large between species (ps > .62, Tukey's adjustment; Fig. 7b). Third, a three-way interaction among species, sulcus, and hemisphere (F(7, 1497) = 8.65, p < .0001,  $\eta 2 =$ 0.04) showed that: i) the species difference for prculs-d was larger in the left hemisphere (estimate = -0.0015, p < .0001, Tukey's adjustment) than the right (estimate = -0.0008, p = .01, Tukey's adjustment), ii) the pmcgs was marginally relatively larger in chimpanzees in the left hemisphere (p = .05, Tukey's adjustment) but not the right hemisphere (p = .18, Tukey's adjustment), and iii) the pos is relatively larger in chimpanzees in the left hemisphere (p < .0001, Tukey's adjustment),

but not the right hemisphere (p = .24, Tukey's adjustment; Fig. 7b).

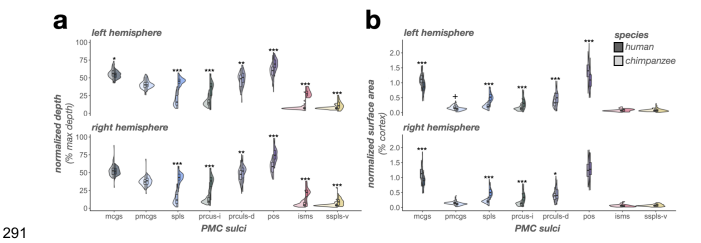


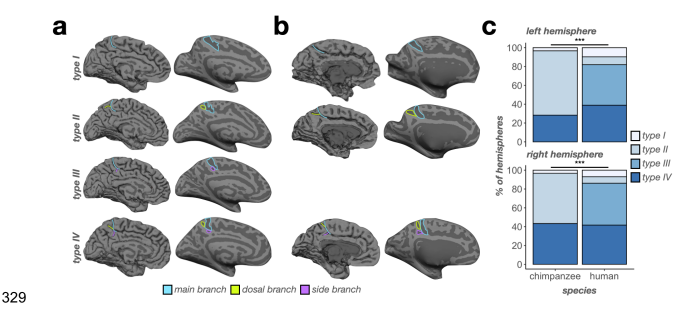
Figure 7. The complex relationship of PMC sulcal morphology in humans versus chimpanzees. a. Split violin plots (box plot and kernel density estimate) visualizing normalized sulcal depth (percent of max depth; percentage values are out of 100) as a function of sulcus (x-axis), species (darker colors, right violin: human; lighter colors, left violin: chimpanzee), and hemisphere (top: left hemisphere; bottom: right hemisphere). Significant differences between species [as a result of the species x sulcus interaction (or the species x sulcus x hemisphere interaction for the mcgs)] are indicated with asterisks (\*p < .05, \*\*p < .01, \*\*\*p < .001). Significant differences between species [as a result of the species x sulcus interaction (or the species x sulcus x hemisphere interaction for the prculs-d, pmcgs, and pos)] are indicated with asterisks (+p = .05; \*p < .05, \*\*p < .01, \*\*\*p < .001).

## Morphological types of the mcgs differ substantially between humans and chimpanzees

Previous work by Bailey and colleagues<sup>55</sup> showed that the chimpanzee mcgs bifurcated into what they termed "vertical" and "horizontal" components. Conversely, Ono and colleagues<sup>61</sup> identified that the human mcgs could variably present with side branches and/or a bifurcated dorsal end. In the present study, we integrated these previous classifications into four patterns based on what branches were present. We could identify up to three different branches of the mcgs: i) the main branch (mb) extending from the cingulate sulcus, ii) a branch extending dorsally from the main branch (db), and iii) a side branch (sb) extending horizontally or ventrally from the main branch (termed cih, as in Bailey *et al.*<sup>55</sup>). In the neuroanatomical literature, it is common to qualitatively describe sulcal "types" based on variation in the shape of a given sulcus and/or patterning of fractionation or intersection with neighboring sulci (e.g., <sup>32,64-66</sup>). Following this terminology, the

combination of these branches fell into four types: I) an mb with no db or sb, II) mb with a db, III) mb with a sb, and IV) mb with both a db and sb (Fig. 8a).

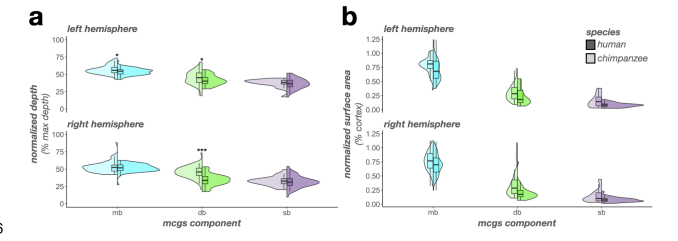
We quantitatively determined whether the incidence rates of the four mcgs types differed by species, as well as between hemispheres for each species with chi-squared ( $\chi$ 2) tests. We observed significant differences in both hemispheres (left:  $\chi$ 2 = 61.95, df = 3, p < .0001; right:  $\chi$ 2 = 52.62, df = 3, p < .0001; **Fig. 8b**). Specifically, type I was comparably present between species in both the left (p = .24; chimpanzee: 3.33%; human: 9.72%) and right hemispheres (p = .34; chimpanzee: 3.33%; human: 6.94%; **Fig. 8b**). Type II was more present in chimpanzees (and the most common type) than humans in both the left (p < .0001; chimpanzee: 68.33%; human: 8.33%) and right hemispheres (p < .0001; chimpanzee: 53.33%; human: 6.94%; **Fig. 8b**). Conversely, type III was only present in humans (and the most common type) in both the left (p < .0001; chimpanzee: 0%; human: 43.06%) and right hemispheres (p < .0001; chimpanzee: 0%; human: 43.06%) and right hemispheres (p < .0001; chimpanzee: 0%; human: 43.06%) and right hemispheres (p < .0001; chimpanzee: 0%; human: 43.67%; human: 38.89%) and right hemispheres (p = .69; human: 43.33%; human: 41.67%; human: 42.21, human: 42.22; human: 43.33%; human: 43.33%; human: 44.44%; human: 45.45%; human: 45.45%; human: 46.45%; human: 47.67%; human: 48.89%) and right hemispheres (human: 49.59%; human: 49.59%; human: 41.67%; human: 41.67%; human: 41.67%; human: 42.50%; human: 42.50%; human: 43.50%) and right hemispheres (human: 43.50%) are species (human: 44.67%; human: 45.50%; human: 45.50%; human: 45.50%; human: 47.50%; human: 48.50%) and right hemispheres (human: 49.50%; human: 41.67%; human: 41.67%; human: 42.50%, human: 43.50%, human: 44.50%, human: 45.50%, human: 45.50%, human: 45.50%, human: 46.50%, human: 47.50%, human: 48.50%, human: 48.50%, human: 49.50%, human: 49.50%, human: 49.50%, human: 49.50%, human: 40.50%, human: 40.50%,



**Figure 8.** Chimpanzees do not have a Type III mcgs. a. Example pial (left) and inflated (right) human hemispheres displaying the four "types" of the mcgs. Type I consists of only a main branch (blue outlines/lines). Type II consists of a main branch and a dorsal branch (green outlines/lines). Type III consists of a main branch and a side branch (purple outlines/lines). Type IV consists of all three branches. **b.** Same as a, but for chimpanzees. Note that no chimpanzees in our sample had an identifiable type III mcgs (empty third row). **c.** Bar plot visualizing the incidence of mcgs types as a function of species (x-axis), type (color, see legend), and hemisphere (top: left hemisphere; bottom: right hemisphere). Lines and asterisks highlight significant species differences in the incidence of mcgs types in both hemispheres (\*\*\* p < .001).

The depth and surface area of mcgs components largely differ between chimpanzees and humans. Finally, we quantitatively tested for species differences in the sulcal depth and surface area of the three mcgs components comprising the different types (mb, db, and sb). In terms of depth, a LME with predictors of component, hemisphere, and species on mcgs component sulcal depth revealed five findings. First, there was a main effect of component (F(2, 341) = 440.90, p < .0001,  $\eta 2 = 0.72$ ), such that the mb was deeper than the db and sb (ps < .0001, Tukey's adjustment) and the db was deeper than the sb (p < .0001, Tukey's adjustment; **Fig. 9a**). Second, there was a main effect of hemisphere (F(1, 130) = 25.25, p < .0001,  $\eta 2 = 0.16$ ), such that components of the mcgs are generally deeper in the left than right hemisphere (**Fig. 9a**). Third, there was a main effect of species (F(1, 130) = 17.29, p < .0001,  $\eta 2 = 0.12$ ) in which chimpanzee mcgs components were

relatively deeper than humans (Fig. 9a). Fourth, there was an interaction between species and component (F(2, 341) = 12.76, p < .0001,  $\eta 2 = 0.07$ ). Post hoc analyses revealed that the db (p < .0001) .0001, Tukey's adjustment) and mb (p = .03, Tukey's adjustment) of the mcgs were relatively deeper in chimpanzees, whereas the sb was comparably deep between species (p = .99, Tukey's adjustment; Fig. 9a). Fifth, there was a three-way interaction among species, component, and hemisphere (F(2, 341) = 5.58, p = .004,  $\eta 2 = 0.03$ ). Post hoc analyses revealed that it was driven by i) the mb of the mcgs being relatively deeper in chimpanzees in the left hemisphere (p = .03, Tukey's adjustment), but not the right (p = .32, Tukey's adjustment; Fig. 9a) and ii) the species difference (i.e., chimpanzee > human) for the db being larger in the right hemisphere (estimate = 0.11, p < .0001, Tukey's adjustment) than the left (estimate = 0.04, p = .02, Tukey's adjustment). In terms of surface area, a LME with component, hemisphere, and species on mcgs component as predictors revealed two findings. First, there was a main effect of *component* (F(2, 341) = 971.27, p < .0001,  $\eta 2 = 0.85$ ), such that the mb was larger than the db and sb (ps < .0001, Tukey's adjustment) and the db was larger than the sb (p < .0001, Tukey's adjustment; Fig. 9b).Second, there was a main effect of species (F(1, 130) = 39.67, p < .0001,  $\eta 2 = 0.23$ ) in which the mcgs components were all relatively larger in chimpanzees compared to humans (Fig. 9b). There were no *species*-related interactions (ps > .16).



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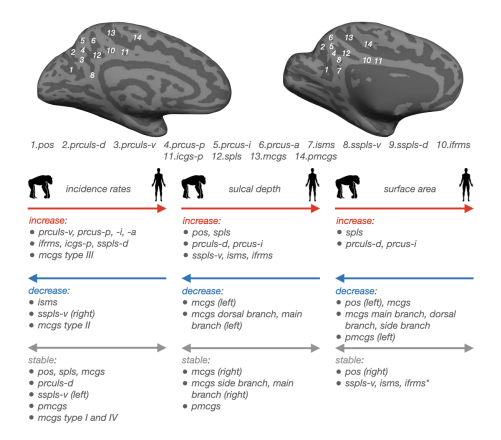
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## Discussion

By manually defining 2,537 sulci spanning the PMC of 144 human and 120 chimpanzee (*Pan Troglodytes*) hemispheres, we show that the surface anatomy of PMC substantially differs between these two hominoid species along three sulcal metrics: i) incidence/patterning, ii) depth, and iii) surface area (**Fig. 10** summarizes the major differences in PMC sulcal morphology between chimpanzees and humans). For sulcal incidence rates, half of PMC sulci are less present in chimpanzees than humans, whereas the other half are either more present in chimpanzees or equally present between species (**Fig. 10**). Further, the prominent megs differs significantly between species (**Fig. 10**). For sulcal depth, the majority of PMC sulci are relatively shallower in chimpanzees compared to humans; however, a minority are relatively deeper in chimpanzees or equally deep in both species (**Fig. 10**). For sulcal surface area, the majority of PMC sulci are relatively smaller in chimpanzees compared to humans; however, a minority are relatively larger in chimpanzees or equally sized across species (**Fig. 10**). This variability is in stark contrast to previous work claiming similarities between PMC between species:

"Overall, the medial aspect of the parietal lobe of the chimpanzee and other apes closely resembles the general appearance of the same structures in the human brain (Bailey et al., 1950)" [Cavanna and Trimble<sup>67</sup>, pg. 565]

In the following sections, we discuss these findings in the context of the evolution of the cerebral cortex and the evolution of complex brain functions and behaviors, as well as discuss limitations and implications for future studies.



**Figure 10. Summary of differences in PMC sulcal morphology between humans and chimpanzees.** Top: Inflated cortical surface reconstructions of the individual human (Left) and chimpanzee (Right) hemispheres shown in Figure 2. Sulci: dark gray; Gyri: light gray. Individual posteromedial (PMC) sulci are numbered according to the key below. Bottom: Overview of differences in PMC sulcal morphology between species. Position (right, left, both) of arrowheads indicates whether sulci increased (right), decreased (left), or remained stable (right and left) in each morphological feature between species. *Left:* incidence rates; *Middle:* sulcal depth; *Right:* surface area. ifrms data from<sup>20</sup>.

The present work adds to the growing literature in comparative neurobiology and paleoneurobiology classifying the presence/absence of sulci across species as a qualitative and quantitative metric to assess the evolution of the cerebral cortex. Such studies have revealed that although the sulcal patterning of primary sensory cortices more or less resembles one another

across species, this relationship is far less consistent in association cortices. For example, while the sulcal organization of visual association cortex was comparable between every human and non-human hominoid hemisphere examined in previous work<sup>15</sup>, the incidence of sulci in medial<sup>14,16,17</sup> and lateral<sup>38</sup> prefrontal cortex, as well as orbitofrontal cortex<sup>64</sup> was substantially different across species. Adding to the complexity, within each of these regions, differences in sulcal incidence rates were greater for some sulci compared to others — elucidating specific areas of cortex that are particularly expanded/more complex in humans. For example, sulcal incidence between humans and chimpanzees in the lateral prefrontal cortex is more consistent across species in the posterior middle frontal gyrus than anterior middle frontal gyrus<sup>38</sup>. Further, some sulci in the human prefrontal cortex are not present in non-human hominoids<sup>2,19</sup>. As shown in the present study, although the PMC (at a regional level) is generally more evolutionarily expanded in humans<sup>68</sup>, the differences in PMC sulcal morphology between humans and chimpanzees was heterogeneous—that is, not all sulci were less present, relatively smaller, and relatively shallower in chimpanzees compared to humans (**Fig. 10**).

Here, we consider two different underlying features that could contribute to this observed heterogeneity: i) differences in the size and depth of border sulci that constrain the macroanatomical definition of PrC and PCC in each species and ii) expansion of PrC, but not PCC, sulci between humans and chimpanzees. First, the main border sulci (pos and mcgs) were relatively smaller and shallower in humans compared to chimpanzees (**Fig. 10**). This finding could be a consequence of the large increase in size, depth, and number of PrC sulci observed in humans compared to chimpanzees (**Fig. 10**). This observation is consistent with the classic compensation theory of cortical folding by Connolly<sup>69,70</sup>, which qualitatively states that the depth and size of sulci are seemingly counterbalanced by those of their neighbors. In terms of the compensation

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theory then, in chimpanzees, the shallow, small (or even absent) precuneal sulci neighbor large and deep pos and mcgs (and the reverse in humans), such that the former "compensate" for the latter and in turn, make the overall degree of cortical folding approximately equal<sup>71</sup>. Second, PrC sulci were relatively larger in humans compared to chimpanzees, whereas PCC sulci were not (Fig. 10). This could be a consequence of the PrC not being topographically constrained along the vertical axis, in contrast to the PCC which is constrained superiorly by the cingulate/splenial sulci and inferiorly by the callosal sulcus. Recent empirical evidence 10 supports this notion, finding that the PrC is the only area of PMC that spatially expands (in the longitudinal direction) between chimpanzees and humans. The majority of sulci in PrC and PCC were also relatively deeper in humans than chimpanzees, which could be due to the fact that both areas are not topographically constrained along this axis. Finally, the decrease in isms presence in humans (Fig. 10) may be a consequence of changes in pos morphology in humans. In nearly all human hemispheres, the pos intersects with the calcarine sulcus (e.g., 61,72-75), which is not necessarily the case in chimpanzees<sup>54–56,72,76,77</sup>. The intersection of these two sulci, which is in the proximity of the isms, may have led to its absence in humans. Considering that the present work only examined the PMC in chimpanzees, future work should seek to also examine the sulcal morphology of PMC in additional species such as macaques, baboons, bonobos, gorillas, orangutans, and gibbons in order to build a larger picture for how the PMC changes along the primate phylogeny.

The present findings also lay the foundation to examine the cognitive and functional role of PMC sulci in species beyond humans. Recent work shows that sulcal morphology relates to the appearance of complex behaviors in non-human hominoids<sup>19,35,78,79</sup>. For example, asymmetries in the depth of multiple sulci<sup>19,35,78</sup>, as well as the presence of the paracingulate sulcus<sup>35</sup> and dorsal fronto-orbital sulcus pattern<sup>19</sup>, relates to the production and use of attention-getting sounds by

chimpanzees. Further, asymmetries in the depth of the inferior arcuate sulcus was related to gestural communication in baboons<sup>79</sup>, as was the presence of the intralimbic sulcus in chimpanzees<sup>35</sup>. Thus, a goal for future work would be to relate the incidence rates and morphological features of PMC sulci to behavioral performance in non-human hominoids.

In conclusion, our findings provide insight regarding how PMC sulcal patterning and morphology differs between humans and our close relative: the chimpanzee. We not only uncover the presence of previously overlooked structures in human and chimpanzee PMC, but also show that the sulcal organization of PMC differs dramatically between chimpanzees and humans along multiple metrics: percent sulci, sulcal presence, surface area, and depth. Future research can seek to further explore how the PMC sulcal patterning differs in humans relative to other non-human hominoids and non-human primates, as well as link the morphology of these structures to the emergence of complex behaviors and functional areas.

## **Materials and Methods**

### Participants:

Humans: Data for the young adult human cohort analyzed in the present study were taken from the Human Connectome Project (HCP) database (https://www.humanconnectome.org/study/hcp-young-adult/overview). Here we used data from 72 randomly selected participants (36 females, 36 males, aged between 22 and 36). HCP consortium data were previously acquired using protocols approved by the Washington University Institutional Review Board. Here, we used the same participants used in our previous work in PMC identifying the ifrms for the first time<sup>20</sup>.

Chimpanzees: 60 (37 female, 23 male, aged between 9 and 51) chimpanzee (Pan Troglodytes)

# **Data Acquisition**

expanses<sup>20,15,38</sup>.

Humans: Anatomical T1-weighted (T1-w) MRI scans (0.8 mm voxel resolution) were obtained in native space from the HCP database. First, the images obtained from the scans were averaged. Then, reconstructions of the cortical surfaces of each participant were generated using FreeSurfer, a software used for processing and analyzing human brain MRI images (v6.0.0, surfer.nmr.mgh.harvard.edu). All subsequent sulcal labeling and extraction of anatomical metrics were calculated from the cortical surface reconstructions of individual participants generated through the HCP's custom-modified version of the FreeSurfer pipeline<sup>80</sup>.

Chimpanzees: Detailed descriptions of the scanning parameters have been described in Keller et al.<sup>5</sup>, but we also describe the methods briefly here. Specifically, T1-weighted magnetization prepared rapid-acquisition gradient echo (MPRAGE) MR images were obtained using a Siemens 3T Trio MR system (TR = 2300 ms, TE = 4.4 ms, TI = 1100 ms, flip angle = 8, FOV = 200 mm) at YNPRC in Atlanta, Georgia. Before reconstructing the cortical surface, the T1 of each

# Manual sulcal labeling: all PMC sulci

*Humans:* For the present study, we re-assessed the 144 human hemispheres analyzed in our prior work<sup>20</sup>. Manual lines were drawn on the FreeSurfer *inflated* cortical surface to define sulci with tools in *tksurfer* based on the most recent schematics of sulcal patterning in PMC by Petrides<sup>60</sup>, as well as by the pial and smoothwm surfaces of each individual as in our prior work<sup>20,27,28,36</sup>. In some cases, the precise start or end point of a sulcus can be difficult to determine on a surface<sup>82</sup>. Thus, using the inflated, pial, and smoothwm surfaces of each individual to inform our labeling allowed us to form a consensus across surfaces and clearly determine each sulcal boundary. For each hemisphere, the location of PMC sulci was identified by trained raters (E.H.W., S.A.M., J.K., B.P., T.H., L.A.G.) and confirmed by a trained neuroanatomist (K.S.W.).

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(prculs-y). The third is a previously uncharted and lone indentation appearing within the isthmus of the cingulate gyrus, which we accordingly term the isthmus sulcus (isms). See Fig. 2A for 7 example human hemispheres with PMC sulci defined, and Supplementary Fig. 1 for every hemisphere with sulcal labels. Chimpanzees: Guided by recent in vivo criteria for defining PMC sulci in humans<sup>20</sup>, we defined PMC sulci in chimpanzees. Prior work leveraging this same chimpanzee sample determined that chimpanzees variably possess an ifrms<sup>20</sup> and it is known that chimpanzees possess an mcgs, pos, and spls<sup>54–56</sup>. Therefore, in the present study, we determined whether or not chimpanzees possessed the pmcgs, as well as the five PrC sulci (prculs-d, prculs-v, prcus-p, prcus-i, prcus-a) and the four other PCC sulci (isms, sspls-v, sspls-d, icgs-p) residing within the bounds of the mcgs, pos, and spls in humans. As with humans, PMC sulci were defined in FreeSurfer using tksurfer tools, and for each hemisphere, the location of PMC sulci was confirmed by the same two-tiered process. See Fig. 2B for 7 example chimpanzee hemispheres with PMC sulci defined, and Supplementary Fig. 2 for every hemisphere with sulcal labels. Manual sulcal labeling: mcgs patterns Linking to prior work by Bailey and colleagues<sup>55</sup> and Ono and colleagues<sup>61</sup>, all 144 human and 120 chimpanzee inflated hemispheres were inspected by authors E.H.W., S.A.M., and K.S.W. to determine which of the four mcgs patterns was present in humans and chimpanzees: I) a main branch (mb) with no dorsal branch (db) or side branch (sb), II) mb with a db, III) mb with a sb, and IV) mb with both a db and sb.

To quantify the amount of cortex buried in PMC across individuals and species, we combined six regions in the Destrieux parcellation<sup>57</sup> corresponding to PMC: *G\_cingul-Post-dorsal*, *G\_cingul-Post-ventral*, *G\_precuneus*, *S\_cingul-Marginalis*, *S\_parieto\_occipital*, and *S\_subparietal* (https://surfer.nmr.mgh.harvard.edu/fswiki/CorticalParcellation). These labels were converted from the Destrieux annotation into individual labels and combined into one "PMC ROI" FreeSurfer label with the *mri\_annot2label* and *mri\_mergelabels* functions in FreeSurfer. To quantify the areas of the cortex defined as sulci, we used the .sulc file<sup>58</sup>. Depth values in the .sulc file 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. To create a "sulci ROI" FreeSurfer label, we thresholded the .sulc file for all vertices with values > 0 with the *mri\_binarize* function in FreeSurfer. To determine the percent of PMC composed of sulci, we calculated the overlap between the PMC ROI and sulci ROI with the Dice coefficient<sup>20,36</sup>:

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

where X and Y are the PMC ROI and sulci ROI, | | represents the number of elements in a set, and  $\cap$  represents the intersection of two sets.

We then ran a linear mixed effects model (LME) with predictors of hemisphere and species, as well as their interaction terms, for percent overlap. Species and hemisphere were considered fixed effects. Hemisphere was nested within subjects. We controlled for differences in brain size in the model (quantified as the total cortical surface area of the given hemisphere). Analysis of variance (ANOVA) F-tests were subsequently applied.

*PMC sulci:* We characterized the frequency of occurrence of each sulcus separately for left and right hemispheres. In line with prior work<sup>14</sup>, for any sulcus that was not present in all hemispheres for either species, we tested the influence of species and hemisphere on the probability of a sulcus to be present with binomial logistic regression GLMs. For each statistical model, species (human, chimpanzee) and hemisphere (left, right), as well as their interaction, were included as factors for presence [0 (absent), 1 (present)] of a sulcus.

To compare whether the incidence of the variable PMC sulci in chimpanzees related to one another, we ran binomial logistic regression GLMs for each variable PMC sulcus [0 (absent), 1 (present)] with the other sulci as factors, while also including an interaction with hemisphere for each sulcus. We iteratively dropped the sulcus that was the dependent variable as a factor from the next model to account for relationships already analyzed. Note that we excluded sulci with an incidence rate of over 90% (prculs-d) and less than 15% (prculs-v, sspls-d, prcus-p, prcus-a, icgs-p) due to the very small sample size.

Marginal ramus of the cingulate sulcus types: We quantitatively determined whether the incidence rates of the four mcgs types differed by species, as well as between hemispheres for each species, with  $\chi 2$  tests.

# Quantification of sulcal morphology

In the present study, we considered depth and surface area as these are two of the most defining morphological features of cortical sulci — especially in PMC<sup>15,20,24–27,29,36,38,65,71,83–89</sup>.

<u>Depth</u>: The depth of each sulcus was calculated in millimeters from each native cortical surface reconstruction. Raw values for sulcal depth were calculated from the sulcal fundus to the smoothed outer pial surface using a modified version of a recent algorithm for robust morphological statistics which builds on the Freesurfer pipeline (Madan, 2019). As the chimpanzee surfaces were scaled prior to reconstruction, we report relative (normalized) depth values for the sulci of interest. For these metrics, within each species, depth was calculated relative to the deepest point in the cortex (i.e., the insula as in previous work<sup>15,20,38</sup>).

Surface area: Surface area (in square millimeters) was generated for each sulcus from the mris\_anatomical\_stats function in FreeSurfer<sup>58,90</sup>. Again, as in prior work<sup>38</sup>, to address scaling concerns between species, we report surface area relative to the total cortical surface area of the given hemisphere.

# Morphological comparisons

To assess whether the depth and surface area of PMC sulci differed between chimpanzees and humans, for both morphological features, we ran a LME with predictors of sulcus, hemisphere, and species, as well as their interaction terms. Species, hemisphere, and sulcus were considered fixed effects. Sulcus was nested within the hemisphere which was nested within subjects. As in our prior analysis, ANOVA F-tests were applied to each model. For brevity, and considering that human PMC sulcal morphology has already been examined in prior work<sup>20</sup>, we only report species-related effects in the main text for this set of analyses. For these analyses we did not include the *ifrms* as our prior work<sup>20</sup> already conducted comparative morphological analyses on this sulcus in

Finally, we repeated the prior analysis, exchanging the factor of PMC sulci for the mcgs branch (main branch, dorsal branch, side branch). As this is the first time these pieces have been quantitatively described, we report all effects in the main text.

# Statistical analyses

All statistical tests were implemented in R (v4.0.1). LMEs were implemented with the *lme* function from *nlme* R package. ANOVA F-tests were run with the *anova* function from the built-in *stats* R package. Effect sizes for the ANOVA effects are reported with the partial eta-squared ( $\eta$ 2) metric and computed with the *eta\_squared* function from the *effectsize* R package. ANOVA chi-squared ( $\chi$ 2) tests were applied to each GLM, from which results were reported. GLMs were carried out with the *glm* function from the built-in *stats* R package and ANOVA  $\chi$ 2 tests were carried out with the *Anova* function from the *car* R package. Relevant post-hoc analyses on ANOVA effects were computed with the *emmeans* and *contrast* functions from the *emmeans* R package (*p*-values adjusted with Tukey's method). Non-ANOVA  $\chi$ 2 tests (for the mcgs type analysis) were carried out with the *chisq.test* function from the built-in *stats* R package. Follow-up post hoc pairwise comparisons on these  $\chi$ 2 tests were implemented with the *chisq.multcomp* function from the *RVAideMemoire* R package.

### Data availability

Data and analysis pipelines used for this project will be made freely available on GitHub upon publication (<a href="https://github.com/cnl-berkeley/stable\_projects">https://github.com/cnl-berkeley/stable\_projects</a>). The colorblind-friendly color

- 663 schemes used in our figures were created using the toolbox available at
- 664 <a href="https://davidmathlogic.com/colorblind/">https://davidmathlogic.com/colorblind/</a>. Requests for further information should be directed to the
- 665 Corresponding Author, Kevin Weiner (<u>kweiner@berkeley.edu</u>).

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