1 Title: Divergent Connectional Asymmetries of the Inferior Parietal Lobule Shape

2 Hemispheric Specialization in Humans, Chimpanzees, and Macaque Monkeys

- 3 Short title: Comparative Study of IPL Connectional Asymmetry
- 4 Luqi Cheng^{1,2,3}, Yuanchao Zhang¹, Gang Li^{2,3,5}, Jiaojian Wang^{1,6}, William D.
- 5 Hopkins⁷, Chet C. Sherwood⁸, Gaolang Gong^{9,10}, Linzhong Fan^{2,3,4,5,*}, Tianzi
- 6 Jiang^{1,2,3,4,5,11,*}
- 7 ¹Key Laboratory for NeuroInformation of Ministry of Education, School of Life Science and
- 8 Technology, University of Electronic Science and Technology of China, Chengdu 610054, China
- 9 ²Brainnetome Center, Institute of Automation, Chinese Academy of Sciences, Beijing 100190,
- 10 China
- ³National Laboratory of Pattern Recognition, Institute of Automation, Chinese Academy of
- 12 Sciences, Beijing 100190, China
- 13 ⁴CAS Center for Excellence in Brain Science and Intelligence Technology, Institute of Automation,
- 14 Chinese Academy of Sciences, Beijing 100190, China
- ⁵School of Artificial Intelligence, University of Chinese Academy of Sciences, Beijing 100190,
- 16 China
- ⁶Center for Language and Brain, Shenzhen Institute of Neuroscience, Shenzhen 518057, China
- ⁷Keeling Center for Comparative Medicine and Research, The University of Texas MD Anderson
- 19 Cancer Center, Bastrop, Texas
- ⁸Department of Anthropology and Center for the Advanced Study of Human Paleobiology, The
- 21 George Washington University, Washington, DC 20052, USA
- ⁹State Key Laboratory of Cognitive Neuroscience and Learning & IDG/McGovern Institute for
- 23 Brain Research, Beijing Normal University, Beijing 100875, China
- ¹⁰Beijing Key Laboratory of Brain Imaging and Connectomics, Beijing Normal University,
- 25 Beijing 100875, China
- ¹¹The Queensland Brain Institute, University of Queensland, Brisbane, QLD 4072, Australia
- 27
- 28 *Corresponding Author: Tianzi Jiang, Institute of Automation, Chinese Academy of Sciences,
- 29 Beijing 100190, China. Email: jiangtz@nlpr.ia.ac.cn, Phone: 010 82544778, Fax: 010 -

- 30 82544777.
- 31 *Co-Corresponding Author: Lingzhong Fan, Institute of Automation, Chinese Academy of
- 32 Sciences, Beijing 100190, China. Email: lingzhong.fan@ia.ac.cn, Phone: 010 82544523.

34 Abstract

35 The inferior parietal lobule (IPL) is one of the most expanded and structurally and 36 functionally asymmetric regions in the human cerebral cortex. Whether the structural 37 and connectional asymmetries of IPL subdivisions differ across primate species and 38 whether this relates to functional asymmetries remain unclear. We identified IPL 39 subregions that exhibited symmetric positive allometric scaling across macaque 40 monkeys, chimpanzees, and humans. Patterns of IPL subregions asymmetry were 41 similar in chimpanzees and humans, whereas no IPL asymmetries were evident in 42 macaques. Among the comparative sample of primates, humans showed the most 43 widespread asymmetric connections in the frontal, parietal, and temporal cortices, 44 constituting leftward asymmetric networks that may provide an anatomical basis for 45 language and tool use. Unique human asymmetric connectivity between the IPL and 46 the primary motor cortex may be related to handedness. These findings suggest that 47 structural and connectional asymmetries may underlie hemispheric specialization of 48 the human brain.

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51 Keywords: inferior parietal lobule; brain asymmetry; brain evolution; anatomical
52 connectivity; parcellation

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55 Introduction

The association cortex has expanded greatly in size and exhibits modified 56 connectivity patterns in human brain evolution ^{1, 2, 3, 4}. Compared to the primary 57 sensory and motor cortical regions, the association cortex displays disproportionate 58 59 expansion in conjunction with overall neocortical volume enlargement across primates ⁵. Accordingly, association areas comprise a large percentage of the 60 neocortex in human brains^{2, 6, 7}. Functional and neuroanatomical asymmetries are 61 pronounced in the human brain, appearing to be more extreme compared with other 62 primate species, especially in the association cortex ⁸. Recent evidence suggests that 63 cerebral asymmetry exists not only in humans but also in nonhuman primates ^{9, 10}. For 64 65 example, olive baboons and chimpanzees showed population-level leftward 66 volumetric asymmetry in the planum temporale, which is thought to be homologous 67 to part of Wernicke's area in humans and may have played a facilitating role in the evolution of spoken language ^{11, 12}. Comparative studies on brain asymmetry are 68 69 crucial for understanding the evolution and function of the modern human brain.

70 Language and complex tool use, which show considerable lateralization in the human brain, are considered to be universal features of humans^{13, 14, 15}. These specialized 71 72 functions all involve the inferior parietal lobule (IPL), an area of the association cortex that represents a zone of topographical convergence in the brain ^{15, 16}. 73 74 Moreover, the IPL is one of the most expanded regions in humans compared with nonhuman primates ^{1, 2, 6, 17}. The functional diversity and expansion of the IPL imply 75 76 that it contains subdivisions that may have been elaborated or developed in the 77 ancestors of modern humans, allowing new abilities such as extensive tool use and communication using gestures ¹⁷. However, due to the scarcity of data, different 78 criteria, and methodological limitations for defining regions or subregions³, whether 79 80 the internal organization of the IPL differs across species and whether this relates to 81 different asymmetric functions remain unclear.

82 A major challenge for neuroscience is to translate results obtained using one method

83 and in one species to other methods and other species. Although the IPL has been 84 subdivided into distinct subregions using cytoarchitecture and this technique has 85 provided invaluable information, cellular microstructure alone is insufficient to completely represent brain organization, especially long-range connections, which are 86 the major determinant of regional specialization ^{18, 19}. Furthermore, histological 87 methods with postmortem brains cannot be readily scaled to large populations. Recent 88 89 advances in diffusion magnetic resonance imaging (MRI), which allow the 90 quantitative mapping of whole-brain neural connectivity in vivo, provide an 91 alternative technique called connectivity-based parcellation to subdivide specific regions of the brain or even the entire cortex ^{20, 21}. In previous studies, this technique 92 93 was successfully used to characterize the subdivisions in different species as well as to perform cross-species comparisons^{22, 23}. 94

Previous studies have assessed asymmetries of the IPL using local characteristics, 95 such as cortical volume, thickness, and surface area ^{24, 25}. However, although such 96 97 regional asymmetries have been identified, additional analyses are need to address the architecture of neural connectivity ²⁶. A recent "connectomic hypothesis for the 98 99 hominization of the brain" suggests neural network organization as an intermediate 100 anatomical and functional phenotype between the genome and cognitive capacities, which are extensively modified in the human brain 27 . Beyond that, the functions and 101 interactions of brain regions are determined by their anatomical connections ¹⁸. 102 103 Therefore, identifying connectional asymmetries may provide new insights into the 104 structural and functional specializations of the human brain.

This study investigated asymmetries of IPL subregions in terms of both structure and anatomical connectivity in macaques, chimpanzees, and humans. We first used connectivity-based parcellations to subdivide the IPL to reveal consistent cross-species topographical organization. We then investigated the volumetric allometric scaling and asymmetries of the IPL subregions across species. Using vertex-, region of interest (ROI)-, and tract-wise analyses, we examined asymmetries of the IPL subregions in terms of their connectivity profiles and subcortical white 112 matter pathways to identify evolutionary changes.

113 **Results**

114 Connectivity-based parcellation

115 For each species, a data-driven connectivity-based parcellation was applied to group 116 the vertices in the IPL into functionally distinct clusters based on anatomical 117 connectivity (Fig. 1). Because spectral clustering does not require a specific number 118 of clusters, we iterated the number of subregions from two to twelve to search for the 119 optimal number of subregions. To accomplish this, we identified the optimal number 120 of subregions of the IPL by choosing the maximum number of subregions that showed 121 a coherent topological organization across all species while balancing that by the minimum number of subregions that could be identified based on their 122 cytoarchitectural definitions in macaques, chimpanzees, and humans ^{28, 29}. The two- to 123 124 five-cluster solutions are shown in Supplementary Fig. 1. The two- to four-cluster 125 solutions showed a consistent rostral-caudal pattern in all three species, but in the 126 five-cluster solution a ventral cluster emerged in chimpanzees and a dorsal cluster 127 emerged in humans. The four-cluster solution revealed a rostral-caudal topological 128 pattern that was consistent with previous parcellations based on cytoarchitecture and anatomical connectivity ^{19, 20, 23, 29}. Also, the cytoarchitectural definition of macaques 129 revealed four subregions in the IPL²⁹, which was fewer than the seven 130 cytoarchitectural subregions of the human IPL¹⁹. Although the four-cluster solution 131 132 was not the finest, especially in humans, it contained potentially valuable information 133 about the differences between species. Furthermore, the aim of our research was not 134 to find the "best" cluster solution for the IPL but to identify an appropriate 135 parcellation that could shed light on the lateralization of the structure and connectivity 136 of the IPL and its subregions in this particular sample of three primate species. As 137 such, we chose four clusters as the optimal solution for the cross-species comparison. 138 It is widely accepted that the IPL contains two cytoarchitecturally distinct areas across

species, the anterior (PF) and posterior (PG) areas ^{30, 31, 32}. Our parcellation results 139 140 were consistent with this two-way parcellation and refined it into four subdivisions, 141 specifically, two anterior clusters (the C1 and C2) in the PF and two posterior clusters 142 (the C3 and C4) in the PG. In macaques and chimpanzees, the IPL was previously parcellated into four distinct areas ^{28, 29} in keeping with our four-cluster solution. In 143 humans, the IPL was cytoarchitecturally parcellated into seven distinct areas. 144 145 Although we proposed a four-cluster solution that has fewer areas than the cytoarchitectural map, it is also consistent with it ¹⁹. Specifically, the rostral anterior 146 147 cluster (C1) is similar to the PFt and part of PFop area defined using cytoarchitecture by Caspers et al.¹⁹, the caudal anterior cluster (C2) corresponds to the PF and PFm 148 149 areas, the rostral posterior cluster (C3) is similar to the PGa area, and the caudal 150 posterior cluster (C4) is similar to the PGp area. Our results did not include the PFcm 151 area because it is located deep in the parietal operculum. Given the limited 152 descriptions of subdivisions and connectivity of the IPL in chimpanzees, our 153 parcellation of the IPL can depict the subregions and connectivity of the IPL in 154 chimpanzees from an evolutionary perspective.

155 To assess which hemisphere was dominant with respect to a given function of the 156 human IPL subregions, we decoded the functions of the human IPL subregions from the Neurosyth database ³³ and calculated differences in the correlation values between 157 158 the left and right corresponding subregions (Supplementary Fig. 2). The term tool 159 showed a much higher correlation with the left C1 than with the right C1, suggesting 160 that the left C1 is more involved in tool use. Terms such as *tool* and *semantics* showed 161 relatively high correlations with the left C2, whereas terms such as nogo and 162 *inhibition* showed relatively high correlations with the right C2, suggesting that the 163 left C2 is more involved in tool use and language whereas the right C2 is more 164 involved in executive function. Terms such as retrieval, episodic, recollection, 165 *memories*, and *coherent* showed relatively high correlations with the left C3, whereas 166 terms such as nogo, inhibition, and beliefs were correlated with the right C3, 167 suggesting that the left C3 is more involved in memory and language whereas the

right C3 is more involved in executive and social cognitive functions. Terms such as *episodic* and *coherent* showed relatively high correlations with the left C4, whereas terms such as *spatial*, *attention*, *mentalizing*, and *relevance* showed relatively high correlations with the right C4, suggesting that the left C4 could be more involved in memory and language whereas the right C4 could be more involved in spatial attention and social functions.

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Figure 1. Framework of the connectivity-based brain parcellation for macaques, chimpanzees, and humans. (a) Defining the seed masks of the inferior parietal lobule (IPL) in surface space according to the gyri and sulci. (b) Connectivity-based parcellation using anatomical connectivity. Probabilistic tractography was applied by sampling 5000 streamlines at each vertex within the seed mask. Whole-brain

connectivity profiles were used to generate a connectivity matrix with each row representing the connectivity profile of each seed vertex. Next, a correlation matrix was calculated as a measure of similarity between the seed vertices. Then, a group similarity matrix was calculated by averaging the correlation matrix across subjects and spectral clustering was applied to it. (c) Parcellation results of the IPL across species. The entire framework was applied independently for each hemisphere and each species.

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189 Allometric scaling and structural asymmetry of IPL subregions

190 When examining the relationship between the volume of each of the IPL subregions 191 and the total grey matter volume, the scaling of all the IPL subregions showed 192 positive allometry (all slopes > 1) (Fig. 2a). A statistical analysis revealed no 193 significant relationships between the slopes of each pair of the bilateral IPL 194 subregions. The asymmetry indices (AIs) for the IPL subregions were calculated and 195 are shown in Fig. 2b. The macaques showed no significant asymmetry after 196 Bonferroni correction for any of the subregions. The chimpanzees and humans 197 showed a similar asymmetry pattern, that is, leftward asymmetry in the rostral IPL 198 (the C1 and C2, all p < .001) and rightward asymmetry in the caudal IPL (the C3 and 199 C4, all *p* < .001).



202 Fig. 2. Structural allometric scaling and asymmetries of the inferior parietal lobule 203 (IPL) subregions across species. (a) Volumes of the IPL subregions plotted against 204 total cortical gray matter volume (GMV). Solid lines represent the best fit using mean 205 macaque, chimpanzee, and human data points; dotted lines represent 95% confidence 206 intervals. (b) Volumetric asymmetries of the IPL subregions. Negative asymmetry 207 index indicates leftward asymmetry and positive index indicates rightward asymmetry. 208 * denotes significance at the Bonferroni corrected level of p < .05. The error bar 209 indicates the standard error of the mean.

210 Connectional asymmetries of IPL subregions

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211 To investigate the connectional asymmetries of the IPL subregions, we first calculated 212 the connectivity profiles of the left and right subregions in macaques, chimpanzees, 213 and humans using probabilistic tracking (Supplementary Fig. 3). Visualization of the 214 connectivity patterns of the IPL did not show significant interhemispheric asymmetry 215 in monkeys or chimpanzees but did in humans, especially in connections with the 216 inferior frontal gyrus (IFG) and lateral temporal cortex. A vertex-wise analysis was 217 then performed to examine the connectional asymmetry of each subregion for each 218 species by calculating the AIs between its connectivity profiles for the two 219 hemispheres (Fig. 3). Additionally, ROI- and tract-wise analyses were used to 220 examine the asymmetry of cortical regions and subcortical white matter pathways

connected to the subregions, respectively (Fig. 4). No significant asymmetries were
found in macaques in any of the statistical analyses after correction for multiple
comparisons.

224 In chimpanzees, the C1 showed significant leftward asymmetry mainly in connections 225 with the anterior middle frontal gyrus (MFG), anterior IFG, planum temporale, and 226 insula. The C2 showed significant leftward asymmetric connections with the insula 227 and rightward asymmetric connections with the superior parietal lobule (SPL) and 228 superior longitudinal fasciculus 2 (SLF2). The C3 showed significant leftward 229 asymmetric connections with the anterior superior temporal gyrus (STG), anterior 230 superior temporal sulcus (aSTS), and occipitotemporal area and rightward asymmetric 231 connections with the SPL and posterior cingulate gyrus (PCC). The C4 showed 232 significant leftward asymmetric connections with the anterior STG (aSTG) and 233 rightward asymmetry with the SPL and PCC.

234 In humans, the C1 showed significant leftward asymmetric connections with the 235 ventral premotor and motor cortices and insula, which was consistent with regional 236 leftward asymmetric connections with the precentral gyrus (PreG) and insula. The C1 237 also showed significant leftward asymmetric connections with the posterior MFG, 238 aSTG, and posterior middle temporal gyrus (MTG) and rightward asymmetric 239 connections with the orbital part of the IFG, posterior STS, and dorsal precuneus. The 240 C2 showed significant leftward asymmetric connections with the posterior MFG, 241 ventral premotor and motor cortices, SPL, anterior temporal lobe, and posterior MTG, 242 which was consistent with regional leftward asymmetric connections with the IFG, 243 PreG, postcentral gyrus (PostG), SPL, and STG and was supported by leftward 244 asymmetric subcortical connections with the SLF2, SLF3, and arcuate fasciculus (AF). 245 The C2 also showed rightward asymmetric connections with the orbital part of the 246 IFG and posterior cingulate sulcus. The C3 showed significant leftward asymmetry 247 mainly in the connections with the anterior IFG, SPL, and almost all the lateral 248 temporal cortex, which was consistent with regional leftward asymmetric connections 249 with the MTG and inferior temporal gyrus (MTG/ITG). The C3 also showed

rightward asymmetric connections with the IFG, which was supported by leftward asymmetric subcortical connections with the SLF3. The C4 showed significant leftward asymmetry mainly in the connections with the IFG and anterior and posterior temporal cortex. The C4 also showed significant regional leftward asymmetric connections with the PreG, PostG, and SPL in the ROI-wise analysis.

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257 Fig. 3. Connectional asymmetries of the IPL subdivisions in the vertex-wise analysis 258 across species. Effect size (Cohen's d) related to asymmetric connections of IPL 259 subdivisions displayed on the left hemisphere of a species-specific standard brain 260 (leftward asymmetry: yellow, rightward asymmetry: blue) for each species for areas 261 showing a significance at the level of p < .05 corrected for multiple comparisons 262 using false discovery rate correction. PreG, precentral gyrus; SPL, superior parietal 263 lobule; aSTG, anterior superior temporal gyrus; aSTS, anterior superior temporal sulci; 264 PT, planum temporale; VPMC, ventral premotor cortex; pMTG, posterior middle 265 temporal gyrus; IFG, inferior frontal gyrus; Ins, insula.

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269 Fig. 4. (a) Connectional asymmetries of IPL subdivisions in the region of interest 270 (ROI)-wise analysis across species. Connectional asymmetry was calculated for the 271 connections between each IPL subregion and eleven ROIs. (b) Connectional 272 asymmetries of the IPL subdivisions in the tract-wise analysis across species. 273 Connectional asymmetry was calculated for the connections between each IPL 274 subregion and the seven tracts. For all plots, the four quadrants of each circle 275 correspond to the four IPL subregions. The outermost circles represent ROIs or tracts. 276 The three inner circles from inside to outside represent macaques, chimpanzees, and 277 humans, respectively. For all plots, only the connectivity showing a significance at a 278 Bonferroni corrected level of p < .05 are displayed. SFG, superior frontal gyrus; IFG, 279 inferior frontal gyrus; CGa, anterior cingulate gyrus; Orb, orbitofrontal cortex; PreG, 280 precentral gyrus; PostG, postcentral gyrus; SPL, superior parietal lobule; STG, 281 superior temporal gyrus; MTG/ITG, middle temporal gyrus and inferior temporal 282 gyrus; Ins, insula; SLF1, SLF2, SLF3, the three branches of the superior longitudinal 283 fasciculus; AF, arcuate fasciculus; MdLF, middle longitudinal fasciculus; ILF, inferior 284 longitudinal fasciculus; IFOF, inferior fronto-occipital fasciculus.

286 **Discussion**

287 In the present study, we investigated asymmetries of the IPL in the structure and 288 connectivity of macaques, chimpanzees, and humans. In the structural analysis, the 289 IPL and its subregions exhibited a similar pattern of positive allometric scaling 290 between hemispheres for all species. In addition, the chimpanzees and humans shared 291 similar asymmetric patterns in the IPL subregions, i.e., left asymmetry in the anterior 292 part and right asymmetry in the posterior part, whereas macaques did not display 293 asymmetry. In the connectivity analysis, the chimpanzees showed some connectional 294 asymmetric regions including the SPL, insula, planum temporale, aSTG, and aSTS. 295 The humans showed widespread connectional asymmetric regions including the 296 primary motor and premotor cortices, SPL, insula, and the entire lateral temporal lobe. 297 These regions are associated with language, tool use, and handedness, suggesting a 298 potential relationship between the connectional asymmetry and the functional 299 hemispheric specialization of the human brain.

300 Positive allometric scaling and structural asymmetry of IPL subregions

301 Brain allometry describes the quantitative scaling relationship between changes in the 302 size of one structure relative another structure, often the whole brain or cerebral 303 cortex^{3, 34}. Previous allometric studies suggested that the association cortex 304 (prefrontal, temporal, and parietal regions) scales with positive allometry (i.e., increases in size disproportionally, or more rapidly) across primates ^{3, 35}. Utilizing 305 306 parcellation-based delineations, a recent study provided evidence that human brains 307 have a greater proportion of prefrontal cortex gray matter volume than other primates ⁷ and other studies demonstrate that human prefrontal expansion is greater than would 308 be expected from allometric scaling in nonhuman primates ^{34, 35}, although there 309 remains some conflicting analyses ³⁶. In the present study, we used macro-anatomical 310 311 boundaries to identify the boundaries of the IPL and a connectivity-based parcellation 312 approach to subdivide the IPL, which helped to reveal its internal organization. We 313 found that the bilateral IPL subregions exhibited consistent, positive allometric 314 scaling, which suggests that allometric scaling of the internal organization of the IPL 315 was similar and was also consistent between homotopic regions during the evolution 316 of the IPL in anthropoid primates. With only three species in the sample, our dataset 317 does not allow us to use phylogenetic comparative statistical methods or determine 318 whether human IPL subregions fall significantly above allometric expectations from 319 nonhuman primates; future research that incorporates a broad phylogenetic sample of 320 diverse primate brains would be necessary.

321 We found that chimpanzees and humans showed a similar dichotomous asymmetric 322 pattern in their IPL subregions, i.e., leftward asymmetry in the anterior portion (the 323 C1 and C2) and rightward asymmetry in the posterior portion (the C3 and C4). The 324 result in humans is consistent with a recent study using data from a large consortium 325 showing leftward asymmetry in the supramarginal gyrus and rightward asymmetry in the angular gyrus in terms of surface area ²⁴. The divergent volumetric asymmetries 326 327 suggest functional heterogeneities of the IPL and emphasize the importance of 328 analyzing subregions within the IPL. The shared asymmetric pattern also suggests that 329 divergences in the internal organization of the IPL evolved prior to the common 330 ancestor of chimpanzees and humans.

331 Connectional asymmetries underlying human language and complex tool use

332 Recent neuroimaging studies have highlighted specific brain regions and pathways that may be necessary for tool use ^{13, 37}. We found that humans showed leftward 333 334 asymmetric connectivity between the IPL (the C2) and the primary motor cortex, 335 ventral premotor cortex, SPL, and posterior MTG, all of which were activated in tasks 336 related to tool use and might constitute a cortical network underlying complex tool use ¹³. In addition, portions of this network appeared to represent part of a system that 337 338 is tightly linked with language systems. The interaction between the tool use system 339 and the language system, though with a clear left hemisphere bias, is responsible for 340 representing semantic knowledge about familiar tools and their uses and for acquiring

the skills necessary to perform these actions ^{3, 13, 16, 37}. Several theories suggest that the 341 342 evolutionary path leading to language and tool use in humans may largely be based on 343 the ability to gesture and imitate, which is partially founded on a "mirror neuron" 344 system ¹³. Macaques are thought to emulate the goals and intentions of others, whereas chimpanzees can also imitate certain specific actions, but humans have an 345 346 even stronger bias for high-fidelity copying of precise sequences of actions, which has been called "overimitation" ³⁸. Our findings provide a potential explanation for these 347 348 phenomena in that the macaques showed no asymmetric network connections, the 349 chimpanzees showed a few asymmetric connections, but the humans showed a large 350 number of asymmetric connections. These species differences in leftward asymmetric 351 connections involving language and tool use may reflect human specializations for 352 language and complex tool use.

353 Unlike the humans, who showed considerable leftward asymmetry connectivity 354 between the IPL and the lateral temporal cortex, the chimpanzees showed few 355 leftward asymmetric connections between the IPL and the temporal cortex, including 356 the planum temporale, aSTG, and aSTS. The planum temporale is considered to include part of Wernicke's area homolog 11, and displays leftward anatomical 357 asymmetry in humans and great apes ^{39, 40}. Our result of increased asymmetric 358 359 connections between the IPL and planum temporale in human brains compared to 360 chimpanzees and macaques reinforces the evidence that the evolutionary origin of human language capacities are related to further left hemispheric specialization of 361 neural substrates for auditory processing that are shared with other primates ⁴¹. Since 362 363 the aSTG and aSTS have been implicated in semantic and phonologic processing in humans ⁴², the leftward asymmetric connections of IPL with the aSTG and aSTS may 364 365 be relevant to the evolution of human language processing.

366 Species-specific differences in asymmetric connectivity in chimpanzees and 367 humans

368 Species-specific differences in asymmetric connectivity between the IPL and SPL

369 were found in chimpanzees and humans, with leftward asymmetry in the former and 370 rightward asymmetry in the latter. These species differences in hemispheric 371 asymmetry may reflect evolutionary changes responsible for adaptations or the 372 production of new abilities in the human brain. Structurally, in chimpanzees, right anatomical asymmetry in the white matter below the SPL ⁴³ may increase the right 373 connectivity between the IPL and SPL compared with the left side. In humans, the 374 leftward volumetric asymmetry in the SPL⁴⁴, together with leftward volumetric 375 376 asymmetry in the IPL (the C2), may support the leftward asymmetric connectivity. 377 Functionally, interaction between the IPL and SPL is crucial for tool use, which is 378 dominant in the left hemisphere, and visuospatial function, which is dominant in in the right ^{13, 45, 46}. As for tool use, in contrast to the relatively simple tools used by 379 380 chimpanzees and other species, humans can create complex artifacts through a 381 sequence of actions that may incorporate multiple parts, reflecting a deep 382 understanding of the physics of our bodies, surrounding objects, and the unique demands of the external environments in which we live ^{16, 47}. In addition, complex 383 384 tool use requires the SPL to code the location of the limbs relative to other body parts during planning and executing tool-use movements or hand gestures ^{13, 14, 48}. Leftward 385 asymmetric connectivity between the IPL and SPL may have provided a connectional 386 387 substrate for complex tool use during human evolution. As for visuospatial functions, 388 the rightward asymmetric connectivity between the IPL and SPL in chimpanzees may 389 indicate that visuospatial functions are dominant in the right hemisphere and had 390 already been lateralized to the right hemisphere. During evolution, these lateralized 391 functions may be retained in the human brain. Meanwhile, the lateralized directional 392 reversal of this connectivity from the right to the left hemisphere may reflect 393 evolutionary adaptations for the emergence of new abilities, such as sophisticated and 394 complex tool making and use.

395 Human unique asymmetric connectivity of IPL subregions

396 Unlike the chimpanzees, humans showed leftward asymmetry in the connection

397 between the rostral IPL (the C1 and C2) and the primary motor cortex, which is 398 consistent with a larger neuropil volume in the left primary motor cortex than in the right side ⁴⁹. Meanwhile, the leftward asymmetric volume of the anterior IPL and the 399 400 primary motor cortex may also increase the neural connectivity between these two 401 regions in the left hemisphere compared with the right side. Such a leftward connection is thought to be related to handedness and hand manual skills ^{49, 50}. In 402 403 contrast to humans, chimpanzees and macaques did not show any asymmetric 404 connectivity between the IPL and the primary motor cortex. Although previous 405 studies have shown that chimpanzees exhibit population-level handedness in the use 406 of tools and a corresponding asymmetry in the primary motor cortex, inferior frontal cortex, and parietal operculum ^{51, 52}, they do not show handedness as a more universal 407 408 trait or exhibit manual dexterity to the same extent as humans. One possible 409 explanation is that humans developed the asymmetric connectivity that became the 410 structural basis for specific behaviors of handedness and hand skills during evolution.

411 An unexpected finding was that in humans the IPL, particularly the C3, showed 412 rightward asymmetric connection with the IFG. Since the IPL and the IFG are interconnected though the SLF3, which is strongly rightward asymmetric ⁴⁶, it may 413 414 also increase the connection between the IPL and IFG in the right hemisphere. 415 Functionally, the left IFG is involved in various aspects of language functions, including speech production and semantic, syntactic, and phonological processing²², 416 417 while the right IFG is associated with various cognitive functions, including attention, motor inhibition, and social cognitive processes ⁵³. Our result of rightward asymmetry 418 419 in this connection seems to be associated with attention and social function, but not 420 language, although language dominance in the left hemisphere is considered to be a 421 common characteristic in humans.

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The widespread asymmetric connections of the IPL in humans compared with the other two primates is in keeping with the inter-hemispheric independence hypothesis, in which, during evolution, brain size expansion led to hemispheric specialization due

426 to decreased inter-hemispheric connectivity, resulting in increased intra-hemispheric connectivity ^{54, 55}. While having more cortical neurons (local characteristics) in one 427 428 hemisphere than the other seems to be a necessary condition for asymmetries of 429 complex and flexible behaviors, it is not a full condition for such behaviors. Given 430 that a function or behavior in an area is determined by its connectivity or networks in which it is involved ¹⁸, the widespread lateralized connections may provide the human 431 432 brain with the increased computational capacity necessary for processing language 433 and complex tool use and may play a facilitating role in human specialization.

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435 In conclusion, we identified similar topographical maps of the IPL to study the 436 structural and connectional asymmetry in macaques, chimpanzees, and humans. The 437 structural analysis revealed that the structural asymmetry of the IPL was independent 438 of the allometric scaling of this region. The connectional analysis revealed that 439 humans had the largest connectional asymmetries of IPL subregions compared to 440 macaques and chimpanzees. The regions showing larger asymmetric connections with 441 the human IPL were associated with language, complex tool use, and handedness, 442 which provided potential anatomical substrates for functional and behavioral 443 asymmetries in humans. The opposite asymmetric connection between the IPL and 444 SPL in chimpanzees and humans may reflect evolutionary adaptation during the 445 course of the human evolution.

446

447 Methods

448 Human data

Data from 40 right-handed healthy adults (age: 22–35, 14 males) were randomly
selected from the S500 subjects release of the Human Connectome Project (HCP)
database ⁵⁶ (http://www.humanconnectome.org/study/hcp-young-adult/). T1-weighted
(T1w) MPRAGE images (resolution: 0.7mm isotropic, slices: 256; field of view: 224

453 \times 320; flip angle: 8°), and diffusion-weighted images (DWI) (resolution: 1.25mm

454 isotropic; slices: 111; field of view: 210×180 ; flip angle: 78° ; b-values: 1000, 2000,

and 3000 s/mm²) were collected on a 3 T Skyra scanner (Siemens, Erlangen, Germany)

456 using a 32-channel head coil.

457 Chimpanzee data

458 Data from 27 adult chimpanzees (*Pan troglodytes*, 14 males) were made available by 459 the National Chimpanzee Brain Resource (http://www.chimpanzeebrain.org, 460 supported by the NIH National Institute of Neurological Disorders and Stroke). Data, 461 including T1w and DWI, were acquired at the Yerkes National Primate Research 462 Center (YNPC) on a 3T MRI scanner under propofol anesthesia (10 mg/kg/h) using previously described procedures ⁵⁷. All procedures were carried out in accordance 463 464 with protocols approved by YNPRC and the Emory University Institutional Animal 465 Care and Use Committee (Approval no. YER-2001206).

466 DWI were acquired using a single-shot spin-echo echo-planar sequence for each of 60 467 diffusion directions ($b = 1000 \text{ s/mm}^2$, repetition time 5900 ms; echo time 86 ms; 41 468 slices; 1.8 mm isotropic resolution). DWI with phase-encoding directions (left–right) 469 of opposite polarity were acquired to correct for susceptibility distortion. For each 470 repeat of a set of DWI, five $b = 0 \text{ s/mm}^2$ images were also acquired with matching 471 imaging parameters. T1w images were also acquired for each subject (218 slices, 472 resolution: 0.7x0.7x1mm).

473 Macaque data

Data from 8 male adult macaque monkeys (*Macaca mulatta*) were obtained from TheVirtualBrain ⁵⁸. All surgical and experimental procedures were approved by the Animal Use Subcommittee of the University of Western Ontario Council on Animal Care and followed the Canadian Council of Animal Care guidelines. Surgical preparation and anesthesia as well as imaging acquisition protocols have been previously described ⁵⁸. Images were acquired using a 7-T Siemens MAGNETOM

487 Image preprocessing

488 The human T1w structural data had been preprocessed following the HCP's minimal preprocessing pipeline ⁵⁹, while the chimpanzee and monkey T1w structural data had 489 490 been preprocessed following the HCP's nonhuman preprocessing pipelines described 491 in previous studies ^{7, 59}. Briefly, the processing pipeline included imaging alignment to 492 standard volume space using FSL, automatic anatomical surface reconstruction using 493 FreeSurfer, and registration to a group average surface template space using the Multimodal Surface Matching (MSM) algorithm 60. Human volume data were 494 495 registered to Montreal Neurological Institute (MNI) standard space and surface data 496 were transformed into surface template space (fs_LR). Chimpanzee volume and surface data were registered to the Yerkes29 chimpanzee template ⁷. Macaque volume 497 498 and surface data were registered to the Yerkes19 macaque template ⁷.

Preprocessing of the diffusion-weighted images was performed in a similar way in the human, chimpanzee, and macaque datasets using FSL. FSL's DTIFIT was used to fit a diffusion tensor model for each of the three datasets. Following preprocessing, voxel-wise estimates of the fiber orientation distribution were calculated using Bedpostx, allowing for three fiber orientations for the human dataset and two fiber orientations for the chimpanzee and macaque datasets due to the b-value in the diffusion data.

506 **Definition of the IPL**

507 The IPL, located at the lateral surface of the ventral posterior parietal lobe, is 508 surrounded by several sulci including the Sylvian fissure, superior temporal sulcus (STS), and intraparietal sulcus (IPS) ^{29, 30, 31, 32}. In the absence of detailed homologous 509 510 definitions, it is necessary to use cytoarchitectonic delineations and macroscopic 511 boundaries, such as gyri and sulci, that can be reliably identified in all species as the 512 boundaries of the IPL. The region of interest (ROI) of the IPL was manually drawn on the standard surface template using Connectome Workbench ⁵⁹. In the present study, 513 514 we restricted the ROI to the lateral surface of the IPL and excluded the cortex buried 515 in the sulci, especially the lateral bank of the IPS and the upper bank of the Sylvian 516 fissure. Rostrally, the IPL borders the vertical line between the Sylvian fissure and the 517 rostral lip of the IPS. Dorsally, the IPL borders the lateral bank of the IPS. Ventrally, 518 the anterior ventral IPL borders the upper bank of the Sylvian fissure. The border of 519 the posterior and ventral IPL is formed by the extension of the Sylvian fissure to the 520 top end of the STS in chimpanzees and macaques but by the extension of the Sylvian 521 fissure to the posterior end of the IPS in humans.

522 Connectivity-based parcellation

523 We used a data-driven connectivity-based parcellation framework modified from Fan et al 20 (Figure 1). All steps in the framework were processed on surface data because 524 the surface-based method has advantages, such as cortical areal localization⁶¹, over 525 526 the traditional approach and because the use of surface meshes is a straightforward 527 way to improve existing tractography processing pipelines, such as the precise locations of streamline seeding and termination ⁶². The surface ROI was first 528 registered to native surface using MSM ⁶⁰. The probabilistic tractography was 529 530 performed on the native mesh representing the gray/white matter interface using 531 Probtrackx. The pial surfaces were used as stop masks to prevent streamlines from 532 crossing sulci. 5000 streamlines were seeded from each of the white matter surface 533 vertices in the seed region to estimate its whole-brain connectivity profile and were

downsampled to 5 mm isotropic voxels to construct the native connectivity M-by-N, a matrix between all the IPL vertices (M) and the brain voxels (N). Based on the native connectivity matrix, a symmetric cross-correlation M-by-M matrix was calculated to quantify the similarity between the connectivity profiles of each IPL vertex. A group cross-correlation matrix was calculated by averaging the cross-correlation matrix across subjects.

540 Data-driven spectral clustering was applied to the group cross-correlation matrix to 541 define the anatomical boundaries of the IPL. Spectral clustering can capture clusters 542 that have complicated shapes, making them suitable for parcellating the structure of 543 complicated brain regions such as the IPL. In addition, the spectral clustering 544 algorithm was successfully used to establish the Brainnetome Atlas ²⁰. However, the 545 number of clusters must be defined by the experimenter when using this method. In 546 the current study, we explored from two to twelve parcellations.

547 Volumetric analysis of the IPL

The cortical gray matter volumetric measurements were calculated using Freesurfer. Total cortical volumes were determined by the space between the white and pial surfaces in native space. Each subregion drawn on standard surface space was registered to native surface space using an existing mapping between the two meshes. The volume of the IPL and its subregions was determined by averaging all the vertices for each subject.

554 Functional decoding of each subregion of the human IPL

Each subregion was first mapped to MNI volume space using a ribbon-constrained method in Connectome Workbench. To decode the functions of each subregion, we used the automated meta-analysis database, Neurosynth ³³ to identify the terms that were the most associated with each subregion. The top five non-anatomical terms with the highest correlation values were kept for all subregions and redundant terms, such as *'semantic'* and *'semantics'*, were only considered once. For simplicity, we

561 only showed the positive correlations found by decoding because negative 562 correlations do not directly inform us about the functions of the subregions. The 563 lateralization for each term was obtained by calculating the difference in the 564 correlation values of the subregions between the left and right hemispheres.

565 Mapping anatomical connectivity profiles

566 To map the whole-brain anatomical connectivity pattern for each cluster, we 567 performed probabilistic tractography by drawing 5000 samples from each vertex in 568 each cluster. The resulting tractograms were log-transformed, normalized by the 569 maximum, and then projected onto surface space using 'surf_proj' command in FSL 570 to obtain tractograms in surface space. The surface tractograms were smoothed using 571 a 4 mm kernel for humans, 3 mm kernel for chimpanzees, and 2 mm kernel for 572 macaques. We subsequently averaged the surface tractograms across subjects for the 573 left and right hemispheres separately to obtain population tractograms, which were 574 thresholded by a value of 0.5 for humans, 0.2 for chimpanzees, and 0.3 for macaques 575 due to data quality. The resultant population tractograms represented approximately 576 twenty percent of the non-zero vertexes in the non-thresholded population tractograms 577 and were used for the vertex-wise and ROI-wise comparisons. The volumetric 578 tractograms were used for the tract-wise comparison.

579 Vertex-wise analysis

For each subregion, we restricted the analysis to the group mask defined by the combination of the left and mirrored right population tractograms described above. We here used the connectivity probabilistic value to quantify the connectivity between the IPL and each vertex of the rest of the brain. A higher value in the vertex means a higher likelihood of being connected to the IPL than other vertices.

585 **ROI-wise analysis**

586 Although previous studies have devoted much effort to establishing homologous

587 regions in primates, these are still limited to a few regions, particularly in 588 chimpanzees. To make comparisons across species possible, here we used the 589 common principle of macroscopic anatomical boundaries based on the gyri and sulci 590 to define ROIs in the cerebral cortex. Specifically, the Desikan-Killiany-Tourville (DKT) atlas was used for humans⁶³, a modified DKT atlas for the chimpanzees, and 591 the Neuomaps atlas for the macaques 64 . Because the Neuomaps atlas is volumetric, 592 593 we first mapped it to surface space for the subsequent calculations. A total of eleven 594 cortical ROIs were chosen for each hemisphere: the superior frontal gyrus (SFG), 595 inferior frontal gyrus (IFG, a combination of the pars triangularis and pars opercularis 596 in humans and chimpanzees), anterior cingulate gyrus (CGa, a combination of the 597 rostral and caudal anterior-cingulate in humans and chimpanzees), orbitofrontal cortex 598 (Orb), precentral gyrus (PreG), postcentral gyrus (PostG), superior parietal lobule 599 (SPL), precuneus, superior temporal gyrus (STG), middle temporal gyrus and inferior 600 temporal gyrus (MTG/ITG), and insula. The MTG/ITG was a combination of the 601 MTG and ITG in humans and chimpanzees due to the absence of the MTG in 602 macaques. The connectional value of each ROI was calculated by averaging all 603 vertices in the ROI on the individual surface tractogram for each subregion.

604 **Tract-wise analysis**

605 To investigate which subcortical fiber tracts are associated with lateralization of 606 cortical areas connected to the IPL, we analyzed the lateralization of the subcortical 607 white matter tracts connected to the IPL across species. A total of seven tracts were 608 chosen: the three branches of the superior longitudinal fasciculus, arcuate fasciculus, 609 middle longitudinal fasciculus, inferior longitudinal fasciculus, inferior 610 fronto-occipital fasciculus. The automated tractographic protocols for tracts for each species were from previous studies ⁶⁵ and these tracts were reconstructed using the 611 Xtract tool ⁶⁶. The mean value of each tract was calculated by averaging all voxels in 612 613 the tract in the individual volumetric tractogram for each subregion.

614 Statistical analysis

To investigate the allometric relationship between the volume of each of the IPL subregions and the total gray matter volume using log-transformed data ⁷, linear regression was performed by pooling the human, chimpanzee, and macaque data for each of the IPL subregions, separately. To test whether the scaling regression slopes differed significantly between the two hemispheres, we performed an ANCOVA for comparisons across the two regression slopes for each plot.

621 In all the analyses of the structural and connectional asymmetries (i.e., volumetric, 622 vertex-wise, ROI-wise, and tract-wise), the asymmetry index (AI) was defined as the 623 difference between values for the left and right hemispheres according to the formula 624 AI = 2*(R-L)/(R+L). For the vertex-wise analysis, a one-sample t test was performed 625 at each vertex on the group mask for each species using PALM, with 5000 permutations with a sign-flip strategy ⁶⁷. The statistically significant level was set at 626 627 false discovery rate corrected p < .05. The effect sizes (Cohen's d) were displayed on 628 the average surface. For the volumetric, ROI-wise, and tract-wise analysis, a 629 two-sided Wilcoxon signed-rank test was performed for each subregion. Bonferroni 630 correction was then used for multiple comparisons for seeds, ROIs or tracts, and 631 species, with statistical significance set at p < .05.

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656 The authors declare no competing interests.

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