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Longitudinal connections and the organization of the temporal cortex in macaques, great apes, and humans

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Abstract

The temporal association cortex is considered a primate specialization and is involved in complex behaviors, with some, such as language, particularly characteristic of humans. The emergence of these behaviors has been linked to major differences in temporal lobe white matter in humans compared with monkeys. It is unknown, however, how the organization of the temporal lobe differs across several anthropoid primates. Therefore, we systematically compared the organization of the major temporal lobe white matter tracts in the human, gorilla, and chimpanzee, great apes and in the macaque monkey. We show that humans and great apes, in particular the chimpanzee, exhibit an expanded and more complex occipital–temporal white matter system; additionally, in humans, the invasion of dorsal tracts into the temporal lobe provides a further specialization. We demonstrate the reorganization of different tracts along the primate evolutionary tree, including distinctive connectivity of human temporal gray matter.

Introduction

Understanding the organization and function of the brain requires an appreciation of its phylogenetic context. In the study of primate brain evolution, the temporal lobe is of particular interest. Indeed, the temporal cortex as found in Primates is thought to be unique to this order and distinct from expansions of the lateral brain surface seen in Proboscidea, Cetacea, and Carnivora [1]. Within the Primate order, the external morphology of the temporal lobe differs substantially, ranging from nearly lissencephalic, possessing only the Sylvian fissure, to highly gyrified with multiple longitudinal sulci in great apes [1]. Functionally, the temporal association cortex is involved in some of the most high-level behaviors of primates, including categorization and conceptual processing [2,3], social cognition [4,5], and in the human semantic
Longitudinal connections in primate temporal lobe

species and blueprint analysis are openly available at [https://git.fmrrib.ox.ac.uk/reia/project_temporal_tracts.git](https://git.fmrrib.ox.ac.uk/reia/project_temporal_tracts.git).

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One way to compare temporal lobe architecture between species is by comparing the longitudinal white matter connections that can be reliably detected using neuroimaging, both in vivo and ex vivo [13]. Earlier work has identified a major difference between humans and other primates in the extension of the arcuate fascicle (AF), a major longitudinal tract running mostly dorsally to the Sylvian fissure between the frontal and posterior temporal cortex [14,15]. However, it has been suggested that other aspects of temporal cortex also differ across primates. One example concerns the relocation of areas due to differential cortical expansion between macaques and humans [16,17]. More recently, it has also been suggested that connectivity of temporal association cortex from tracts originating from primary visual and auditory cortices increased in humans and chimpanzees compared with macaques [18]. Recent data from both Klingler dissections and tractography indicate that the inferior longitudinal fascicle (ILF), which connects inferior temporal cortex with occipital regions, is composed of multiple subcomponents in humans [19]. This could suggest that this fiber system has undergone further specialization in humans in comparison with macaques, in which only one component has been described. Therefore, to understand temporal lobe organization, it is necessary to systematically compare the architecture of temporal tracts across a range of primate species.

To understand how the expansion of the temporal cortex in the ape and human lineages is reflected in changes in its macroscale connectivity, we here compared the longitudinal white matter tracts running along the temporal lobe and the dorsal AF between the human, chimpanzee and gorilla great apes, and macaque monkey brains using diffusion-weighted magnetic resonance imaging (MRI). We start by demonstrating that diffusion MRI tractography in macaques is able to reconstruct white matter tracts close to their definition by other methods such as tracers and dissection. Then, we use the same procedure to reconstruct the same tracts in the other species so that we can reliably compare their organization.

Importantly, any differences in the longitudinal tracts between species are likely to result in differences in the connectivity profile of parts of the temporal cortex gray matter. It has been suggested that each cortical area has a unique set of connections with the rest of the brain, the so-called connectivity fingerprint, and that this fingerprint is a crucial determinant of an area’s function [20,21]. Therefore, we also use the reconstructed white matter tracts to undertake a systematic comparison of the temporal lobe gray matter between the macaque and the human, with the aim to investigate how observed white matter differences between species affect their connectivity fingerprints.
Results

Longitudinal tracts in macaque and human

We used connectivity-based parcellations of coronal regions of interest (ROIs) through the temporal white matter to identify the bodies of the main longitudinal tracts of the temporal lobe, which subsequently informed the position of the masks used to perform probabilistic tractography in order to reconstruct the course of the tracts.

In macaques, we were able to reconstruct tracts similar to earlier studies [13,20,22] (Fig 1A, S1 Fig). The superior temporal gyrus contained a single cluster, from which we reconstructed a tract that can be identified as the middle longitudinal fascicle (MdLF) because of its course through the superior temporal gyrus and termination points mainly in the inferior parietal lobe but also branching toward occipital regions. The inferior temporal gyrus contained a single cluster, from which we reconstructed a tract similar to the ILF, terminating in inferior occipital and inferior parietal regions. Medial to these tracts, we identified a tract that is reminiscent of the inferior fronto-occipital fascicle (IFOF), running from the occipital cortex medially through the temporal cortex at the level of the inferior temporal gyrus. Among the tracts we identified, it was the only one to have very clear projections to the frontal lobe. The IFOF as identified here is likely to be a multisynaptic pathway because we are using the method of tractography and not tract tracing. Indeed, tract tracing identifies a monosynaptic section of the IFOF termed the extreme capsule [22]. Importantly, a recent dissection study identified an IFOF in the macaque similar to our tractography results, demonstrating a biological basis of our finding [23]. We provide a more in-depth comparison of these tracts as obtained using different methodologies in the Supporting information section (S1 Fig, S1 Text).

The same procedure in humans identified a tract through the superior temporal gyrus similar to the MdLF (Fig 1B). In the inferior temporal gyrus, the clustering could reliably identify two clusters with differential connectivity to the rest of the brain where we would expect the territory of the ILF to be. We suggest that the higher number of clusters found in humans reflects a higher complexity in the temporal lobe white matter organization of this species. We interpret this in the light of previous studies supporting a subdivision of the ILF into two subcomponents [19,25]. Accordingly, the tractography identified different tractograms for the two subcomponents. One of them runs more laterally in the inferior temporal gyrus and slightly invades the middle temporal gyrus, whereas the other runs more medially and ventrally, mainly in the fusiform gyrus, and reaches more inferior occipital regions. We refer to these tracts as the ILF lateral (ILFlat) and ILF medial (ILFmed) subcomponents of the ILF. We identified a tract very similar to the IFOF that runs predominantly through the middle temporal gyrus, although it is located more medially than the other two tracts. Indeed, from the surface projections (Fig 2A and 2B, S2 Fig), it can be seen that the middle temporal gyrus in humans is mainly reached by the ILFlat and the IFOF.

In summary, compared with the macaque, the human IFOF is located more laterally and runs predominantly along the middle temporal gyrus (which is not present in the macaque). Also, unlike macaques, human ILF can be subdivided into two subcomponents. The MdLF has a similar path in both species, which indicates that its main connectivity pattern across species should not be fundamentally different.

Longitudinal tracts in great apes

Using the same procedure of connectivity-based clustering followed by tractography as for humans and macaques, we identified longitudinal temporal tracts in the chimpanzee and gorilla. We used both in vivo and ex vivo data in the chimpanzee and obtained comparable
results (Fig 3A and 3B), indicating that our results generalize across the different imaging protocols. However, because the left hemisphere in the postmortem sample was slightly flattened in the extraction protocols, it is hard to interpret the results obtained in that hemisphere.

We identified the MdLF running through the superior temporal lobe following a similar course as human and macaque MdLF, reinforcing the argument for a conserved tract across all studied anthropoid primates (Fig 3A, 3B and 3C). In the inferior temporal lobe, the territory occupied by the ILF was similar to that in humans and could similarly be divided into two subcomponents. We therefore will refer to them as ILFlat and ILFmed as well. Although the differentiation is strong in the chimpanzee, specifically in the right hemisphere, the separation of the ILF into two subcomponents in gorilla, although present, is less pronounced. Different methods to assess the number of clusters in the temporal white matter did not reach a consistent conclusion in the gorilla (S2 Text, S8 Fig), suggesting that the gorilla temporal lobe white matter might not be as complex as in chimpanzees and humans. We suggest that this is related to the anatomy of its temporal lobe, with a less prominent fusiform gyrus and a less distinct difference between the inferior and middle temporal gyrus in the gorilla, which is associated with a shallower inferior temporal gyrus, compared with the chimpanzee. The IFOF was also present in great apes, and its course was similar to the human IFOF, running more medially than other tracts as well as in the middle temporal gyrus. In chimpanzee in vivo, it seems that the IFOF does not reach as lateral as in the other great ape data. In gorilla, it can be observed that the IFOF is most prominent in its posterior section, whereas it is quite limited in its frontal projections. However, having only one sample for the gorilla because of the presence of a frontal lesion on the left hemisphere, we cannot draw too-strong conclusions from this observation.

Fig 1. A 3D representation and coronal sections of longitudinal temporal tracts in macaques (A) and humans (B). Tractograms were log transformed and normalized for display. The coronal sections from left to right are taken from posterior to anterior. Blue, MdLF; green, IFOF; yellow, ILF (macaque) or ILFmed (human); red, ILFlat. Thresholds for the tracts are as follows: 0.7 for MdLF; 0.75 for IFOF; and 0.7 for ILF, ILFmed, and ILFlat. Human data are available from the Human Connectome Project (www.humanconnectome.org). Macaque postmortem data are available from the PRIME-DE repository (http://fcon_1000.projects.nitrc.org/indi/PRIME/oxford2.html) [24]. IFOF, inferior fronto-occipital fascicle; ILF, inferior longitudinal fascicle; ILFlat, inferior longitudinal fascicle lateral; ILFmed, inferior longitudinal fascicle medial; MdLF, middle longitudinal fascicle; PRIME-DE, Primate Data Exchange.
To further illustrate these tracts in the great apes, we represented their tractograms on the cortical surface (Fig 4, S3 Fig). In other words, we examined where the tractogram approached the gray/white matter border. Although one should be cautious in interpreting such results because of the presence of superficial white matter systems [26] and possible gyral bias of tractography toward the surface, it provides an illustration of these fiber tracts in the great ape, in which they are currently underexplored. It allows one to represent not only the differences in tract organization with other species but also the sulcal and gyral patterns of these brains, which are not well known. This showed that for both species, the MdLF reaches mostly superior temporal gyrus areas as it does in the macaque and human. The ILF subcomponents are similar to those in the human, with the majority of the lateral part of the inferior temporal cortex reached predominantly by the ILFlat and the medial part reached predominantly by the ILFmed. The IFOF is the only one of these tracts that has projections to the frontal cortex.

Thus, great apes (including humans) seem to display a similar organization regarding the subcomponents of the ILF running in inferior and fusiform gyri, respectively. The ILFmed reaches an area reminiscent of the fusiform gyrus in all species, which is absent in the macaque, in which we did not detect the subdivision of the ILF. In line with this argument, we notice that these ILF subcomponents are less evident in the gorilla, in which the fusiform gyrus is less prominent than in chimpanzees and humans.
Dorsal longitudinal tract

We have concentrated thus far on the major ventral longitudinal tracts along the temporal cortex. However, a major dorsal longitudinal tract, the AF, runs between the frontal and temporal cortex. Previous studies have developed tractography protocols for this tract in the human, chimpanzee, and macaque, although the details differ between studies and groups [15,27]. Here, we used a consistent protocol to track the AF in all four species, creating a comparable result for the three species in which the AF has previously been defined and in the gorilla (Fig 5, S4 Fig).

In macaques, the AF consisted of only a dorsal component, whereas in humans, its course exhibited a sharp curve to end up in the temporal lobe running mainly in the middle temporal gyrus, although quite medially. In chimpanzees, we observed a small curve as well, with AF reaching the superior temporal gyrus but not the middle temporal gyrus, as in humans. An arcuate path similar to that of the chimpanzee could be observed in gorilla. Thus, the AF expansion to the temporal lobe seems to be unique to humans and not present in other great apes studied here.

Differential in macaque and human temporal cortex connectivity profiles driven by longitudinal tracts

The goal of the present work is to establish similarities and differences between the long-range connections of the temporal lobe across primate species, with the macaque and the human as the two most studied samples. If differences in these long-range connections exist, they should
result in different connectivity profiles of temporal cortex gray matter between the two species. To establish whether this is the case, we compared the connectivity profiles of each gray matter vertex of the temporal lobe—in other words, the temporal lobe connectivity blueprint—across species. The logic of the approach is that for each vertex in each species, we quantify how well its connectivity fingerprint with the temporal cortex white matter tracts resembles that of any vertex in the other species. Thus, vertices with a low divergence score have a connectivity fingerprint also present in the other species, whereas vertices with a high divergence score are more likely to have a connectivity that is unique to that species. The connectivity fingerprint profile of a given vertex in this analysis is represented by the probability that the vertex is reached by each tract.

We first restricted the blueprint to four temporal tracts that have been well established in both humans and macaques—namely, the temporal part of the cingulum bundle, the fornix, the optic radiation, and the uncinate fascicle [27–30]. These tracts are known to be conserved across species, and hence we expect all vertices to have low divergence scores. Indeed, in this blueprint, we can observe that both human and macaque brain organization are predictable from one another (Fig 6A, S6A Fig). With those same tracts, but also adding the MdLF, the combined ILF, and the IFOF in the blueprint, we notice a small increase of the divergence score in some areas of the temporal lobe in both species (Fig 6B, S6B Fig). The example connectivity fingerprints in Fig 6A and 6B illustrate how the increase in divergence scores (computed as Kullback–Liebler [KL] divergence) is due to differences in the connections of these parts of the cortical surface with the underlying white matter. When only using the nonlongitudinal tracts (Fig 6A), all vertices have a low value because the pattern of connections of any
part of the surface can be found in the other species. When adding the ventral longitudinal fibers, some differences are noticeable, and indeed a vertex with a high divergence score has a connectivity profile that differs even from its best matches in the other species (Fig 6B). Finally, when adding the AF, the profiles of connections of parts of the cortex are very different even from their best matches in the other species, indicating that this tract contributes most to the distinctive organization of the human temporal lobe (Fig 6C, S6C Fig).

These results are confirmed by the distribution of the divergence scores obtained with each blueprint (Fig 6D, S6D Fig). We can observe a shift toward higher minimum divergence scores in the distributions when adding the AF, which is confirmed by the distributions being statistically different (as assessed by the two-sample Kolmogorov–Smirnov test with \( p < 0.01 \)). This shift is even more dramatic when focusing the human blueprint on the AF territory, confirming its influence on the increase of KL divergence.

One difference we found between the macaques and all the great apes was the dissociability of the two components of the ILF. Using the blueprint method, we next investigated the different impact of the two subcomponents of the ILF on similarity of macaque and human temporal lobe organization. When we ran the same analysis comparing macaque temporal lobe with a human temporal lobe in which the ILF either consists solely of the ILFmed or consists solely of the ILFmed, we observed that macaque temporal lobe contains substantially more vertices with a high KL divergence in the territory of the ILF when we use ILFmed. In other words, macaque and human temporal lobes are more similar if we assume equivalence of macaque
ILF and human ILFmed than if we assume equivalence of macaque ILF and human ILFlat (Fig 7A and 7B, S7A Fig, S7B Fig). Computing the difference between the two divergence maps analyzed with the two different subcomponents thus allows one to investigate the putative
homology (Fig 7C, S7C Fig). On this difference map, we can also notice that when using the ILFmed, the human brain better predicts the areas where the macaque ILF runs, but the macaque brain does not predict the part where the ILFlat runs anymore. When using the ILFlat in humans, it better predicts an area situated more in the domain of the IFOF in macaques. The distribution of the divergence scores in the territory of the macaque ILF, obtained when using one or the other human ILF, also shows a shift toward higher value when using the human ILFlat compared with the human ILFmed (Fig 7D, S7D Fig). The two distributions are statistically different (as assessed by the two-sample Kolmogorov–Smirnov test with $p < 0.01$).
Discussion

Association cortex volume scales differently in monkeys and apes [31–33], which can be taken to indicate a grade change with respect to this aspect of brain organization because a grade designates a group of species based on similar morphological traits. Here, we investigate how these differences in volume are accompanied by differences in long-range connectivity. We show that the main longitudinal tracts connecting the temporal lobe are present in the most often studied monkey “model” (the macaque), in two separate great ape species (chimpanzee and gorilla), and in the human brain. Importantly, we identify modifications of these bundles across the different lineages (Fig 8). First, the ILF in chimpanzees and humans is reliably separated into subcomponents that were not observed in the macaque. This result is less strong in...
the gorilla. Second, the middle temporal gyrus, which is present in all species except the macaque, seems to be most heavily connected by the IFOF, which is located more laterally than in the macaque. Finally, the AF’s extension into the temporal cortex seems to be a uniquely human specialization, with the lack of expansion previously identified in the chimpanzee replicated here in the gorilla. Although consistent with previous reports of differences in the organization of macaque and human temporal cortex [15,20,34], our report crucially demonstrates that differences between the two species are not due to a single change at a single moment in time but rather to a series of changes in white matter architecture.

We studied examples of two nonhuman great apes, the gorilla and the chimpanzee. Although their temporal lobe organization seems generally similar, some differences were apparent. Because it seems that the sulcal anatomy of the gorilla and chimpanzee differs in how clearly defined their fusiform gyrus is, we could expect differences in their long-range connectivity as well, but because of the scarcity of the gorilla’s data consisting of one hemisphere, we should be careful in interpreting them. One observation concerns the gorilla’s ILF, which seems less clearly subdivided than in the chimpanzee. We can also notice that the IFOF reaches the frontal lobe to a lesser extent and is more biased toward the posterior projections, as well as not invading the lateral middle temporal gyrus as much, as seen in chimpanzee. Finally, we can also raise the point that the AF shows even less invasion of the temporal lobe than in chimpanzee, looking more similar to that of the macaque. Overall, this study would suggest that the limited expansion of the temporal lobe in the gorilla leads to a less complex pattern of white matter organization than that observed in the chimpanzee.

The IFOF carries fibers from occipital cortex through the length of the temporal cortex into the frontal cortex. This tract is frequently discussed in the human literature, but its existence in the macaque monkey has been controversial [35]. It should be noted, however, that most macaque studies are based on tracer injections, which identified monosynaptic pathway, whereas the human results are based on tractography, which identified multisynaptic pathway and therefore usually results in larger white matter bundle. Recent studies using tractography for both species suggested the presence of an IFOF-like bundle extending all the way posteriorly to occipital cortex in both species [13], but it has been argued that this reflects an artifact of the tractography method [36]. This issue now seems resolved by recent Klingler dissections in the macaque monkey that provide evidence for a posterior extension of IFOF even in the macaque [23] and by similar reports in another Old World monkey species [37].

However, differences in IFOF across species are notable, with human IFOF showing more extended prefrontal projection than macaque IFOF. We also noticed some differences between the two species regarding which frontal areas are mostly reached by the IFOF. The macaques’ IFOF projects more to medial frontal regions than the human IFOF, which reaches mainly lateral ones. Concerning the path taken by the IFOF in the temporal lobe, a recent exploratory study provided evidence of IFOF in the chimpanzee and suggested that most connections seeded from the middle temporal gyrus reach this fiber bundle [38]. The present results confirm this finding, showing that in all species that have a middle temporal gyrus, the IFOF is the main longitudinal tract reaching it and that it is placed more laterally than in the macaque. This is interesting, given that IFOF has been suggested to carry information between language-related areas in the human brain [39], especially those involved in semantic processing [40]. A previous study subdividing the human IFOF has also shown that it was the more lateral component of the IFOF that was actually connected to language-related areas as opposed to its medial components, which seem to carry more visual information [41]. The present results, showing similarity between human and great ape in possessing a lateral IFOF, suggest that an anatomical substrate enabling cognitive functions related to language existed in the common ancestor of humans and great apes but not in macaques. This demonstrates that cortical
features supporting behavior commonly associated only or especially with humans cannot be assumed to be unique to the human brain.

Evolution of linguistic abilities has often been associated with another longitudinal tract, the AF. Our results confirmed previous studies that have shown differences between primate species [14,15]. In humans, chimpanzees, and gorilla, we observed that the arcuate is innervating the temporal gyrus, whereas it targets only dorsal regions in macaques. Nevertheless, the extent of fibers reaching the middle temporal gyrus is substantially more elaborate in humans than in any other primate species studied to date, accounting for most of the distinctive organization of the human temporal lobe. However, because we are focusing only on the right hemisphere for the gorilla, we should be careful in generalizing this result, particularly because of the left asymmetry of the arcuate observed in humans [14].

We have reported that the great ape ILF can be subdivided into two subcomponents, the medial and the lateral. The medial ILF of humans seems to be most similar to the macaque ILF as classically defined in tracer data [29]. We can try to interpret the relevance of these subcomponents for brain function, although of course, we do not claim to ascribe a function to any white matter bundle as one can for a gray matter area. A recent review of neuropsychological studies [42] sheds light on some possible contribution of the ILF to functional networks, with one of the best described being face recognition. Damage to the ILF has indeed been associated with impairment in face tasks in humans [43]. Although face-responsive areas have been identified in both humans and macaques [44], their locations in the temporal lobe differ between the two species. In humans, the most often described face area is the fusiform face area (FFA) [45]. This is located in the fusiform gyrus—a gyrus that monkeys do not possess but which is reached by the ILFmed in the present study. The main face areas in monkeys are scattered throughout the middle superior temporal sulcus and more ventral areas such as temporal area TE [46]. Therefore, the finding of ILFmed being the most similar to the macaque ILF is consistent with the observation that ILFmed reaches the fusiform gyrus as macaque ILF reaches homologs of the FFA [29,46]. This argument is also in line with the knowledge of the chimpanzee face-processing system. Indeed, it is located in a similar location to that in humans (in the fusiform gyrus) [47], and the current study shows that the chimpanzee ILF could also be subdivided with the ILFmed reaching the fusiform gyrus.

Neuropsychological work also associates the medial part of ILF to other networks, such as the visual memory, scene perception, and the emotion recognition and valuation networks [42], whereas lateral parts of ILF seem to be more associated with reading, object recognition, and semantic networks. Although we need to be careful with interpreting these linkages, we can observe that the reading and semantic networks, which are more associated with uniquely human cognition, are associated to the ILFlat; the one that we showed here was the most dissimilar to the macaques’ ILF.

We have used diffusion MRI data in combination with probabilistic tractography to reconstruct white matter tracts. It should be noted that there have been critiques on certain aspects of the tractography method in recent years. Mostly, these critiques pointed out that tractography can lead to the identification of false positives and that tracking toward the cortical surface is difficult because of the presence of superficial white matter systems near the gray/white matter border [26,48]. It is important to point out that the manner with which the tractography was applied here mitigates these concerns. Rather than tracking from a point on the cortical surface to another point at the cortical surface, which indeed can be problematic for tractography [49], we focus here predominantly on the body of the tracts, which we define in a data-driven manner using anatomical priors. Reconstructing the bodies of cortical fiber bundles is generally quite reliable [27]. Furthermore, in the one analysis in which we rely on the tracts’ connections to the gray matter—namely, the diversion score analyses at the end of the results.
section—we do not track toward the surface but use a procedure that combines tracking from
the body of the tract and tracking from all of the gray matter toward the white matter, which is
less susceptible to the problems outlined above [20].

In this study, we have used a greater range of species than is common in comparative diffu-
sion MRI studies [13,36,50,51], which allowed us to build confidence in our conclusions. How-
ever, we acknowledge that studies using a wider range of species are still needed. We have
studied two nonhuman great ape species, the chimpanzee and gorilla, but our study neglected
the Asian great apes and lesser apes, as well as taking the macaque monkey as the sole repre-
sentative of the Old World monkeys. The scarcity of data in this field means that other apes
and especially other Old World monkey as well as other primate species remain to be studied,
but the accelerating pace of comparative neuroimaging, of which this study is an example,
means that these studies can now be considered imminent. Studies on gray matter architecture
in a large number of primate species present promising examples [52].

This study does not address the evolutionary pressures that gave rise to the cortical speciali-
izations described here. It has been suggested that cortical specializations in the prefrontal cor-
text along the anthropoid lineage arose from increasingly complex foraging challenges that
came with new ecological niches [53]. In this framework, humans have further reduced forag-
ing errors by relying more than other surviving great apes on mental trial and error based on
mental time travel. A similar argument for temporal cortex is advanced in another monograph
[54], which suggests temporal lobe modifications might have occurred in great apes and
humans when they adapted to their environment by changing their social behavior. Although
it is quite difficult to relate these findings to our study, we are nevertheless able to illustrate
here that any structural specialization is derived from a modification of existing machinery
present in the ancestral state. Hence, to understand any one brain, understanding the organi-
ization of closely related species provides essential information. We must be cautious, however,
when inferring evolutionary pressure from brains that come from distant species because they
have all continued to evolve since their common ancestor.

When observing differences in white matter anatomy across species, such as those reported
here, it is important to ask whether these are differences related to more general changes in
brain structure, such as increased cortical folding due to a larger brain size, or whether they
represent specific adaptations to a particular niche. These different types of changes are often
difficult to disentangle [8,55]. It is known that expansion of temporoparietal cortex has
occurred many times in the primate lineage [56], and this might underlie some of the so-called
uniquely human organization of the temporal cortex, such as the ventral location of the middle
temporal area [57]. However, such general mechanisms cannot describe the phenomenon of
“mosaic evolution,” which demonstrates that expansion can be selective to certain brain struc-
tures in certain lineages [58]. Similarly, these general mechanisms cannot predict the results
observed here, i.e., increased complexity of the ILF in great apes and extension of the AF only
in humans. Indeed, the AF extension in humans has been generally interpreted as a unique
adaptation rather than a result of general mechanisms in the language literature [59–61]. Of
course, this does not mean that the developmental mechanisms that give rise to these adapta-
tions need not be the same generic mechanisms at work in all primates, but it does imply that
the specific tracts can be a unique adaptation [62].

In sum, we show a more elaborate picture of temporal cortex white matter organization
across selected species of the order Primates than previously demonstrated. The human expan-
sion of the AF has become one of the most established results of early comparative diffu-
sion MRI studies, but here we show the beginning of a more complete picture, focusing in particu-
lar on occipital–temporal systems subserved by the IFOF and ILF. The changes in these tracts
result in unique connectivity profiles of the temporal cortex gray matter in the humans and other great apes, which has important implications for differences in their behavioral abilities.

**Materials and methods**

**Ethics statement**

All in vivo data (human and chimpanzee) are from publicly available databases. The human data were obtained from the Human Connectome Project (HCP), and the approved consent procedure is fully described in the core HCP literature referenced here (www.humanconnectome.org). The chimpanzee data were obtained from chimpanzeeframe.org and collected at the Yerkes Primate Center under procedures that were carried out in accordance with protocols approved by YNPRC and the Emory University Institutional Animal Care and Use Committee (approval no. YER-2001206). All in vivo chimpanzee scans were done prior to 2012 and thus prior to the 2015 implementation of United States Fish and Wildlife Service and National Institutes of Health regulations governing research with chimpanzees. The ex vivo data were acquired from deceased animals that died of causes unrelated to this research project. As such, the research did not require a Home Office License as defined by the Animals (Scientific Procedures) Act 1986 of the United Kingdom.

**Data acquisition and preprocessing**

The acquisition and preprocessing of the diffusion MRI, structural, and surface data are described in detail in the Supporting information (S2 Text). Key parameters and information on the diffusion data are summarized in Table 1.

**Reconstruction of ILF, MdLF, and IFOF**

Our first aim was to reconstruct the three major longitudinal tracts running through the temporal cortex in all four species: the ILF, MdLF, and IFOF (sometimes referred to as extreme capsule fiber complex [13]). Given that these tracts have not been previously described in chimpanzee and gorilla, we developed an unbiased connectivity-based clustering approach to define tractography seed and waypoint masks in the temporal lobe white matter of all species. We established and validated the approach using human and macaque data because the tracts have been previously identified in these two species [19,22,63]. Briefly, our approach was divided into two steps. We first used a connectivity-based parcellation of three coronal ROIs in the temporal lobe white matter of each individual to identify the bodies of each of the longitudinal tracts based on their connectivity profile to the rest of the brain (Fig 9A). The clusters

| Number of samples | Type of sample | Protocol of acquisition | Resolution (mm$^3$) | b-values (s/mm$^2$) or q-values (cm$^{-1}$) |
|-------------------|---------------|-------------------------|---------------------|--------------------------------------------|
| Macaque           | 4             | Postmortem              | 2D diffusion-weighted spin-echo multislice
|                   |               |                        | 0.6 × 0.6 × 0.6     | b = 4,000                                  |
| Gorilla           | 1             | Postmortem              | 3D DW-SSFP          | 0.6 × 0.6 × 0.6                           | q = 300                                   |
| Chimpanzee        | 1             | Postmortem              | 3D DW-SSFP          | 0.6 × 0.6 × 0.6                           | q = 300                                   |
| Chimpanzee        | 3             | In vivo                 | Single-shot spin-echo echo planar
|                   |               |                        | 1.8 × 1.8 × 1.8     | b = 1,000                                  |
| Human             | 10            | In vivo                 | Slice-accelerated gradient echo EPI (multishell) | 1.25 × 1.25 × 1.25 | b = 1,000, 2,000, 3,000 |

Abbreviations: DW-SSFP, diffusion-weighted steady-state free precession; EPI, echo planar imaging

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**A. Connectivity-based clustering**

(a) ROI definition:
In each hemisphere:
3 ROIs in white matter made of
3 coronal sections each

(b) Connectivity to whole brain
Voxels in ROI
Voxels in rest of the brain

(c) Similarity matrix
Voxels in ROI
Voxels in ROI

(d) K-means clustering
Voxels in ROI
1 2 3 Clusters

(e) Clustering results
Posterior
Middle
Anterior
Clusters for each tract added across individuals

Green clusters ➤ IFOF masks
Blue clusters ➤ MdLF masks
Yellow clusters ➤ ILF masks

**B. Reconstruction of the tracts**
Based on clustering results
Waypoints posterior
Seeds
Waypoints anterior

Streamlines from seed to waypoints
Probabilistic tractography

Average across subjects
Averaged tractograms

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Fig 9. Methods overview. The macaque is shown as the example; the same methods were used for the other species, except that postmortem analyses were fully performed in subject space. (A) Connectivity-based clustering: (a) In each hemisphere, we defined three coronal ROIs in the white matter of the temporal lobe (excluding the area of the cingulum bundle). We transformed these ROIs to the diffusion space of each individual. (b) We then performed tractography from all the voxels in the ROI to the rest of the voxels in the whole brain. (c) We then computed the similarity matrix,
obtained, belonging to our tracts of interests, were subsequently used in the second step of tract reconstruction as seed and waypoint masks (Fig 9B).

The clustering step was first applied in the macaque and human to validate the approach and was performed as follows. For both species, we defined three ROI masks (anterior, middle, and posterior) in standard space (MNI152 1 mm for humans and F99 0.5 mm for macaques), each composed of three consecutive coronal sections and containing only the temporal lobe white matter (Fig 9Aa). The area corresponding to the body of the cingulum bundle was excluded. These ROI masks were transformed to each subject’s diffusion space prior to performing probabilistic tractography using FSL’s probtrackx from all the voxels in the ROI to all the voxels in the brain. This completely unconstrained tractography is part of the clustering step and is not intended as a method to define the whole tracts but to set up constraints for the actual tractography performed in the later tract reconstruction step. We thus obtained for each ROI a matrix of dimensions “number of voxels in ROI” × “number of voxels in the brain,” representing the connectivity of each voxel in the white matter ROI to the rest of the brain, and transformed it back to standard space (Fig 9Ab). From this matrix, we derived a similarity matrix of dimensions “number of voxels in ROI” × “number of voxels in ROI,” representing the similarity between each white matter ROI voxels regarding their connectivity to the rest of the brain (Fig 9Ac). The three similarity matrices (one for each ROI: anterior, middle, and posterior) were each submitted to k-means clustering to group together those voxels that have a similar profile of connections to the rest of the brain (Fig 9Ad) [64,65]. The logic of this approach is that all voxels belonging to the same fiber tract will show a similar connectivity profile, which enables us to infer each tract’s most likely location in the temporal white matter ROIs.

From the current knowledge of tract anatomy in the temporal lobe [19,22,27], we could then determine which cluster belonged to each tract for each subject. In macaques, we set three clusters in the middle and posterior ROI, which was presumed to reconstruct MdLF, ILF, and IFOF, except for the most anterior slice, in which we set it to four clusters to account for the presence of the uncinate fascicle. These assumptions are based on previous tract tracing literature [66]. In humans, we set one more cluster in each ROI because it has been shown that the ILF could be subdivided [19,25]. We justify a lower number of clusters in macaques by hypothesizing a more complex organization of temporal lobe white matter in humans than in macaque. This hypothesis has been verified by various measures that we detail in S2 Text and illustrate in S8 Fig.

After identifying which cluster belonged to each tract for all individuals in one species, we then added the clusters belonging to the same tract in each coronal ROI to obtain a probability map of cluster position for each tract and ROI. Thus, we obtained probability maps for anterior, middle, and posterior positions of each tract. We refer to these probability maps as “clustering results” (Figs 9Ac and 10, top row). Based on these maps, we defined the seed (from middle position) and waypoint masks (from anterior and posterior positions) for the
subsequent tract reconstruction (Fig 10, bottom row). All the masks for a given species were made of the same number of voxels across tracts, and masks from different tracts did not overlap.

Following the clustering step to identify the bodies of all tracts in three different positions along the anteroposterior axis, we reconstructed, in a second step, each tract by performing probabilistic tractography informed by the clustering step. For each tract, the tractography started from the middle seed mask defined in the previous step and was constrained by two waypoints that were anterior and posterior to the seed and also defined in the clustering step. Probabilistic tractography, as implemented in FSL’s probtrackx algorithm, started streamlines in the seed, and each streamline then followed local orientations sampled from the posterior distribution given by bedpostX [67]; only streamlines that passed through one or the other waypoint were retained. To further constrain the tractography, we also defined exclusion masks based on the anatomical description of the tracts in previous literature. Streamlines hitting these exclusion masks were deleted (Fig 9B). The exclusion masks were drawn as follows: for all tracts, we used sagittal midline sections to avoid tract crossing to the other hemisphere; a dorsal exclusion mask to avoid self–back projection from the longitudinal tracts; and exclusion masks for the fornix, cingulum bundle, and AF seed (defined as described below), as well as the medial section of the optic radiation and the temporal section of the uncinate fascicle. For each tract, waypoints and seeds of the other tracts were also used as exclusion masks. Subcomponents of ILF, however, were not used as exclusion masks for each other. To avoid tracts erroneously crossing gray matter, we also used a termination mask consisting of the entire gray matter defined in subject space (using FAST from FSL [68] for macaques and using the individual surface for humans). When reaching these termination masks, the streamlines were terminated but not deleted. Because the macaques’ gray matter masks were less precise, we added an exclusion mask in insular cortex and putamen. The resulting tractogram was normalized for each subject by dividing its path distribution by the total number of generated streamlines not rejected by waypoint or exclusion mask criteria. We then averaged the tract

![Fig 10. Tractography masks (bottom row) derived from clustering results (top row) in macaques and humans. (A) In macaques, the clustering results are thresholded to show the overlap between at least two subjects out of four. (B) In humans, the clustering results are thresholded to show the overlap between at least four subjects out of 10. Blue, MdLF; green, IFOF; yellow, ILF (macaque) or ILFmed (humans); red, ILFlat. R denotes right hemisphere. Human data are available from the Human Connectome Project (www.humanconnectome.org). Macaque postmortem data are available from the PRIME-DE repository (http://fon_1000.projects.nitrc.org/indi/PRIME/oxford2.html). IFOF, inferior fronto-occipital fascicle; ILF, inferior longitudinal fascicle; ILFlat, inferior longitudinal fascicle lateral; ILFmed, inferior longitudinal fascicle medial; MdLF, middle longitudinal fascicle; PRIME-DE, Primate Data Exchange.](https://doi.org/10.1371/journal.pbio.3000810.g010)
distribution over all subjects and down-sampled the average tractogram to 1-mm resolution (isotropic voxels) in F99 space for macaques and 2 mm in MNI152 space for humans. For visualization, 3D renderings of the log-transformed and thresholded average tractograms were generated using in-house Matlab code.

Having performed these analyses for the human and the macaque, we then applied the same procedure to the chimpanzee and gorilla data. The in vivo chimpanzee data were treated exactly as the human and macaque data. We performed a clustering procedure by setting the number of clusters to four because of the similar anatomy of the temporal lobe gyri and sulci between humans, gorillas, and chimpanzees. Indeed, although previous studies have shown differences in the gyral pattern between hemisphere, including notably an asymmetry of the Sylvian fissure and occipital bending between chimpanzees and humans [69], their overall sulci pattern remains more similar than compared with the macaque, which lacks defined middle temporal and fusiform gyri [1]. We performed a similar analysis as in humans and macaques to confirm that this was an optimal number of clusters (S2 Text, S8 Fig). For the chimpanzees, the measures used pointed toward the solution using a high number of clusters. However, for the gorilla, the different measures do not uniformly point toward solutions using a high or low number of clusters, and having only one hemisphere available for this species, we cannot strongly conclude on this issue (S2 Text, S8 Fig).

In this manner, we were able to define four tracts in the temporal lobe: the IFOF, MdLF, ILFlat, and ILFmed. From the clustering results (Fig 11, top rows), we defined the masks used in the tract reconstruction (Fig 11, bottom rows). The probabilistic tractography algorithm used the same exclusion masks as described for the macaques as well as the gray matter termination masks. The tract distributions obtained were averaged over three subjects and down-sampled to 1-mm resolution in Yerkes29 space. For the two postmortem ape samples (gorilla and chimpanzee), we used the same procedure as described above, except that all the analyses were only performed in the diffusion space of each subject and not transformed to a standard space because only one sample per species was available. Again, for visualization, 3D renderings of the log-transformed and thresholded average tractograms were generated using Matlab code made in-house. We also generated a surface representation of the tracts in great apes using tools from Connectome Workbench [70]. For the in vivo chimpanzees, we projected the averaged tracts to the Yerkes29 template (previously created from 29 chimpanzee scans) and we used the respective individual surfaces in T1 space for the postmortem gorilla and chimpanzee.

For the tract reconstruction, we used the same tractography settings in all species (10,000 samples, 2,000 steps, curvature threshold 0.2), except for the step length to account for differences in brain size and relative resolution (0.5 mm in humans and chimpanzees in vivo, 0.3 in gorilla and chimpanzee postmortem, and 0.25 in macaque postmortem).

**Reconstruction of AF**

The AF is a dorsal longitudinal tract that runs from the inferior frontal cortex in a posterior direction and, in the human brain, curves ventrally to terminate in the temporal cortex. Because of its variability between species [15], we could not use the temporal cortex coronal ROI sections to identify it in all species with the clustering approach outlined above. We therefore adapted the method used previously by Eichert and colleagues [14] to reconstruct the tract from a manually defined seed outside the temporal cortex, just posterior to the level of the central sulcus (Fig 5 top rows). We also placed an axial mask as a posterior waypoint in the parietal–temporal white matter posterior to the caudal end of the Sylvian fissure. The mask was placed in a comparable location in all species to not bias the tract’s posterior course (Fig 5,
To make the tractography comparable to that of the other longitudinal tracts outlined above, we added an anterior waypoint to the protocol of Eichert and colleagues [14], allowing us to track from a middle seed to anterior and posterior waypoints as well. We used the same exclusion masks for all species as in Eichert and colleagues [14]: a midline mask, two subcortical white matter and extreme/external capsule masks, and one mask through the superior parietal cortex.

In the great apes, we examined where the tractogram approached the gray/white matter border to represent their tractograms on the cortical surface (Fig 4, S3 Fig). We normalized, log transformed, smoothed with a kernel of 2 mm full-width half maximum (FWHM), and thresholded these surface tractograms.

The clustering and tract reconstruction steps have also been performed with alternative tractography techniques to validate our approach (S2 Text, S10–S14 Figs).

Comparison of macaque and human temporal lobe connectivity blueprints

To establish the effects of any differences in the longitudinal temporal tracts across species on the connectivity profiles of temporal cortex gray matter, we established the white matter connectivity of each part of the temporal cortex gray matter, the so-called “connectivity blueprint,”
using the method established by Mars and colleagues (2018) [20]. The connectivity blueprint is in effect a matrix that describes how well each vertex of the temporal lobe gray/white matter border, as defined in the surface, is reached by each of the white matter tracts of the temporal lobe. Each row of this matrix describes the “connectivity fingerprint” of each vertex of the temporal lobe with each white matter tract (Fig 12A). Therefore, the connectivity fingerprint is a subset of the connectivity blueprint focusing on a particular vertex instead of the whole brain. Each column of the blueprint matrix corresponds to one tract. Because the tracts are defined in a similar way in the different species, they present a common space that allows us to compare connectivity fingerprints across the two brains, testing whether there are unique connectivity fingerprints in one brain compared with the other.

We directly compared these blueprints between the macaque and the human brain (Fig 12). We focused on these two species because they are the most studied species among the ones presented in the current study. Furthermore, the course and connectivity of other nonlongitudinal tracts reaching the temporal lobe, such as the uncinate fascicle, the cingulum bundle, and the fornix, have been well established in these species [20,22].

We established the connectivity blueprint using the tractograms of ILF, MdLF, IFOF, and AF as established above and supplemented them with tractograms of the other major white matter tracts in the temporal lobe—namely, the temporal part of the cingulum bundle, the fornix, the optic radiation, and uncinate fascicle—using established tractography protocols [20]. A tractogram can be represented as a column vector describing how well each voxel is reached by the tract (Fig 12A). By concatenating the vectors corresponding to each tract, we obtained a matrix representing how each voxel (row) is reached by each tract (column). Second, we also calculated whole-hemisphere vertexwise connectivity matrices obtained by tracking from all vertices in the gray/white matter border of the surface file to all voxels in the brain in volumetric space (Fig 12A). These matrices were computed for both hemispheres and for each subject individually before creating an average for each hemisphere and species. Then, we multiplied the tractogram’s matrix with this averaged matrix in both species, obtaining the blueprint matrix (Fig 12A). We also individually extracted each column of this matrix to obtain a surface representation of the different tracts, which we will refer to as “surface projections.”

The connectivity blueprints of different species can be compared by calculating the KL divergence between the connectivity profiles of each vertex in one brain with that of each vertex in the other brain (Fig 12B). The KL divergence measures how the probability distribution of likelihoods of one vertex to be reached by each of the temporal lobe tracts is different in one brain from the probability distribution of each vertex in the other brain. For each vertex in one brain, we could then determine another vertex in the other brain that had the minimum KL divergence—in other words, the most similar connectivity profile. If a vertex in one species has a very similar connectivity profile to that of a vertex in the other brain, meaning that they are reached by a similar combination of tracts, the minimum KL divergence will be low. If, on the other hand, a vertex’s fingerprint connectivity profile is unique in one brain, its minimum KL divergence will be high (Fig 12C).

We first defined the blueprints based on the well-established tracts: the cingulum bundle, the fornix, the optical radiation, and the uncinate fascicle. Because we did not expect these tracts to be massively different between the two species, we expected the KL to be very low and, therefore, this comparison to serve as a reference point for the following blueprint comparisons. Then, we added the tracts of interest: MdLF, ILF, IFOF, and AF. To assess the reorganization of the human ILF subcomponents, we generated blueprints with both ILF subcomponents combined or either one or the other while keeping the macaque ILF constant. This manipulation enabled us to compare the ILF subcomponents’ distinct prediction of the macaque brain organization. We compared these maps visually by subtracting the KL
Fig 12. Blueprint method to assess connectivity divergence between macaque and human temporal cortex. (A) To compute the blueprint, we multiplied a matrix representing the connectivity of each cortical vertex (surface space) to each brain voxel (volume space) with a matrix representing how each brain voxel (volume space) is reached by each tract. Because we reconstructed the same tracts in both macaques and humans, the columns of the blueprint define a common space to compare the connectivity of the temporal cortex. The rows of the blueprint correspond to the connectivity fingerprints of each vertex. The columns of the matrix correspond to the surface tractograms. (B) Using the KL divergence measure to compare every vertex in one brain to every vertex in the other brain, we can identify
divergence scores spatial map established with human ILFmed from the one established with human ILFlat. Furthermore, we masked the blueprint to only focus on the temporal, occipital, and inferior parietal lobes. To assess the divergence between species more quantitatively, we obtained the distribution of the minimum KL divergence scores obtained with each blueprint: with nonlongitudinal tracts; adding MdLF, ILF, and IFOF; adding the AF; with the human ILFlat; and finally, with the human ILFmed (Fig 6D). We included only nonzero values, which are the vertices with differences in their connectivity fingerprints. Each distribution of minimum KL divergence scores indicates how similar the connectivity fingerprints of this brain are with that of another brain. All distributions were then compared using the two-sample Kolmogorov–Smirnov test (with \( p < 0.01 \)).

For illustration, but not for establishing the blueprint, we smoothed the surface tracts with a kernel of 2 mm FWHM for macaques and 4 mm for humans. We then log transformed and normalized the surface tract distributions to facilitate the threshold determination.

Supporting information

S1 Fig. Comparison of the macaque tractography results with the tract tracing and dissection literature. (A) Comparison with tract tracing results redrawn from Schmahmann and Pandya (2006) [29]. Tracts to be compared are highlighted in similar colors. Blue, MdLF; green, IFOF; yellow, ILF; pink, AF. (B) Comparison with the dissection results redrawn from Decramer and colleagues (2018) [23]; the green crosses illustrate the IFOF pathway. Macaque postmortem data are available from the PRIME-DE repository (http://fcon_1000.projects.nitrc.org/indi/PRIME/oxford2.html). AF, arcuate fascicle; IFOF, inferior fronto-occipital fascicle; ILF, inferior longitudinal fascicle; MdLF, middle longitudinal fascicle; PRIME-DE, Primate Data Exchange.

S2 Fig. Surface projection of longitudinal temporal tracts in macaques (A) and humans (B). Shown are the group averages of the normalized, log-transformed, and thresholded tracts in the left hemisphere only (for space-saving purpose). Blue, MdLF; green, IFOF; yellow, ILF (macaque) or ILFmed (humans); red, ILFlat. Thresholds for the tracts are as follows: 0.7 for MDLF; 0.75 for IFOF; and 0.7 for ILF, ILFmed, and ILFlat. Human data are available from the Human Connectome Project (www.humanconnectome.org). Macaque postmortem data are available from the PRIME-DE repository (http://fcon_1000.projects.nitrc.org/indi/PRIME/oxford2.html). IFOF, inferior fronto-occipital fascicle; ILF, inferior longitudinal fascicle; ILFlat, inferior longitudinal fascicle lateral; ILFmed, inferior longitudinal fascicle medial; MdLF, middle longitudinal fascicle; PRIME-DE, Primate Data Exchange.

S3 Fig. Surface representation of longitudinal temporal tracts in chimpanzees. Shown are the group averages of the normalized, log-transformed, smoothed and thresholded left tracts. Blue, MdLF; green, IFOF; yellow, ILFmed; red, ILFlat. Thresholds for the tracts are as follows:
0.7 for MDLF; 0.75 for IFOF; and 0.7 for ILFmed and ILFlat. In vivo chimpanzee data are available from the National Chimpanzee Brain Resource (www.chimpanzeebrain.org). IFOF, inferior fronto-occipital fascicle; ILFlat, inferior longitudinal fascicle lateral; ILFmed, inferior longitudinal fascicle medial; MdLF, middle longitudinal fascicle.

**S4 Fig. AF tractography protocols and results.** Top panel: AF tractography masks example for one individual macaque, human, and chimpanzee, represented on the principal eigenvector (V1) map weighted by the fractional anisotropy map. The light-pink mask represents the seed, and the white masks represent the anterior and posterior waypoints. Bottom panel: 3D and surface representation of the left tractogram obtained for AF. Threshold of 0.75. R denotes right hemisphere. Human data are available from the Human Connectome Project (www.humanconnectome.org). Macaque postmortem data are available from the PRIME-DE repository (http://fcon_1000.projects.nitrc.org/indi/PRIME/oxford2.html). In vivo chimpanzee data are available from the National Chimpanzee Brain Resource (www.chimpanzeebrain.org). AF, arcuate fascicle; PRIME-DE, Primate Data Exchange.

**S5 Fig. MRtrix results showing tractography streamlines for temporal lobe tracts in macaques (A) humans (B).** Tractograms were log transformed and normalized for display. Blue, MdLF; green, IFOF; yellow, ILF (macaque) or ILFmed (human); red, ILFlat; pink, AF. Thresholds for the tracts are as follows: 0.82 for MDLF; 0.82 for IFOF; 0.8 for ILF, ILFmed, and ILFlat; and 0.8 for AF. Human data are available from the Human Connectome Project (www.humanconnectome.org). Macaque postmortem data are available from the PRIME-DE repository (http://fcon_1000.projects.nitrc.org/indi/PRIME/oxford2.html). AF, arcuate fascicle; IFOF, inferior fronto-occipital fascicle; ILF, inferior longitudinal fascicle; ILFlat, inferior longitudinal fascicle lateral; ILFmed, inferior longitudinal fascicle medial; MdLF, middle longitudinal fascicle; PRIME-DE, Primate Data Exchange.

**S6 Fig. Results of the blueprint analysis.** The figure reports results on the left hemisphere. Shown are the tracts used (top row), the resulting KL divergence (middle row) for human predicting macaque (left) and macaque predicting human (right), and the connectivity fingerprints (bottom row). The greener the vertices of the KL divergence map in one brain, the more their connectivity profile is similar to that of vertices in the other brain. The connectivity fingerprints show the probability of the vertex, highlighted by a white sphere in the brain above, to be reached by each tract (dotted line). The solid line represents the probability of being reached by each tract in the other species, calculated as the mean over the 10 vertices with the smallest KL divergence with the initial species’ vertex of interest. (A) Blueprints established using the cbt, the fx, the or, and the unc. (B) Blueprints established with the tracts in (A) and adding the MdLF, ILF (both subcomponents combined in humans), and IFOF. (C) Blueprint established with the tracts in (B) and adding the AF. Blue, MdLF; green, IFOF; yellow, ILF (macaque) or ILFmed (human); red, ILFlat; pink, AF. (D) Distribution of minimum KL values obtained for each blueprint. From left to right for the macaque, the human, and masked with the human AF. Light gray, nonlongitudinal tracts; gray, adding MdLF, ILFs, and IFOF; black, adding AF. Human data are available from the Human Connectome Project (www.humanconnectome.org). Macaque postmortem data are available from the PRIME-DE repository (http://fcon_1000.projects.nitrc.org/indi/PRIME/oxford2.html). AF, arcuate fascicle; cbt, temporal part of the cingulum bundle; fx, fornix; IFOF, inferior fronto-occipital fascicle; ILF, inferior longitudinal fascicle; ILFlat, inferior longitudinal fascicle lateral; ILFmed, inferior longitudinal fascicle medial; MdLF, middle longitudinal fascicle.
longitudinal fascicle medial; KL, Kullback–Liebler; MdLF, middle longitudinal fascicle; or, optic radiation; PRIME-DE, Primate Data Exchange; unc, uncinate fascicle.

**S7 Fig. Blueprint method to determine differences between the human ILFs.** (A) Blueprint established as in S6C Fig but with the ILFlat for humans (no ILF med). (B) Blueprint established as in (A) but with the ILFmed for humans (no ILFlat). (C) The different ILF tractograms are represented on the top row. On the bottom row is shown the KL difference between the two maps established in (A) and (B). More yellow vertices means that using ILFmed in humans as macaques’ ILF homologous results in lower KL divergence between the two species at these vertices, whereas more red applies to ILFlat. Blue, MdLF; green, IFOF; yellow, ILF (macaque) or ILFmed (human); red, ILFlat; pink, AF. (D) Distribution of minimum KL values in the macaque’s ILF territory obtained for the blueprints with the different human ILF subcomponents. Red, with human ILFlat; yellow, with human ILFmed. Human data are available from the Human Connectome Project (www.humanconnectome.org). Macaque postmortem data are available from the PRIME-DE repository (http://fcon_1000.projects.nitrc.org/indi/PRIME/oxford2.html). IFOF, inferior fronto-occipital fascicle; ILF, inferior longitudinal fascicle; ILFlat, inferior longitudinal fascicle lateral; ILFmed, inferior longitudinal fascicle medial; KL, Kullback–Liebler; MdLF, middle longitudinal fascicle; PRIME-DE, Primate Data Exchange.

**S8 Fig. Verification of cluster number reliability.** (A) Illustration of the correspondence between ILF clusters in the middle (seed) ROI when defined as one ILF (three-cluster solution, top row) or the sum of the two subcomponents (four-cluster solution, bottom row), shown from the clustering results obtained across species (thresholded at two individuals’ overlap for macaques, two for chimpanzee, and four for humans). (The IFOF is not visible on the macaque left hemisphere four-cluster solution because the hypothetical ILF subcomponents are invading that space.) R denotes right hemisphere. (B) Correspondence between ILF as defined as a single cluster in the three-cluster solution and as defined by the sum of two clusters in the four-cluster solution, shown as their percentage overlap. If the ILF is reliably split into subclusters, the “correspondence” should be high. (C) Hierarchy index between the three-cluster and four-cluster solutions is represented in black squares for all species. The black lines represent the range of hierarchy index values obtained with 1,000 random permutations of cluster labeling. (D) Results from all measures investigating cluster number solutions. The y-axis represents the results of the subtraction of macaques’ ideal cluster number from the great apes’ cluster number. Each symbol corresponds to a different measure. Human data are available from the Human Connectome Project (www.humanconnectome.org). Macaque postmortem data are available from the PRIME-DE repository (http://fcon_1000.projects.nitrc.org/indi/PRIME/oxford2.html). In vivo chimpanzee data are available from the National Chimpanzee Brain Resource (www.chimpanzebrain.org). Gorilla postmortem data are available from https://doi.org/10.5281/zenodo.3901205. IFOF, inferior fronto-occipital fascicle; ILF, inferior longitudinal fascicle; ILFlat, inferior longitudinal fascicle lateral; ILFmed, inferior longitudinal fascicle medial; KL, Kullback–Liebler; MdLF, middle longitudinal fascicle; PRIME-DE, Primate Data Exchange.

**S9 Fig. Voxel distributions in the three- to four-cluster solutions in the middle ROI.** The lines link the clusters between the two clustering solutions according to whether they have voxels in common; the thickness of the line represents the percentage of the three-cluster voxels that have been allocated to the four clusters. In humans, the clear division of the ILF in the
three-cluster solution into ILFmed and ILFlat in the four-cluster solution can be observed. This division is not apparent in macaques. Human data are available from the Human Connectome Project (www.humanconnectome.org). Macaque postmortem data are available from the PRIME-DE repository (http://fcon_1000.projects.nitrc.org/indi/PRIME/oxford2.html). IFOF, inferior fronto-occipital fascicle; ILF, inferior longitudinal fascicle; ILFlat, inferior longitudinal fascicle lateral; ILFmed, inferior longitudinal fascicle medial; MdLF, middle longitudinal fascicle; PRIME-DE, Primate Data Exchange.

S10 Fig. FMRIB Software Library probabilistic and deterministic clustering results. The top panel shows the Dice coefficients in black dots between the clustering results obtained with probabilistic and deterministic tractography for each ROI. The light-gray lines represent the range of Dice coefficients obtained between 1,000 random permutations of cluster solutions. The bottom panels show the clustering results obtained with probabilistic (top) and deterministic (bottom) tractography for each ROI. As in Fig 10 and Fig 11, the clustering results are thresholded to show the overlap between at least two subjects out of four in macaques and four out of 10 in humans; no thresholding is applied in chimpanzees and gorilla. R denotes right hemisphere. Human data are available from the Human Connectome Project (www.humanconnectome.org). Macaque postmortem data are available from the PRIME-DE repository (http://fcon_1000.projects.nitrc.org/indi/PRIME/oxford2.html). In vivo chimpanzee data are available from the National Chimpanzee Brain Resource (www.chimpanzeebrain.org). Gorilla postmortem data are available from https://doi.org/10.5281/zenodo.3901205. ant, anterior; post, posterior; PRIME-DE, Primate Data Exchange; ROI, region of interest.

S11 Fig. Comparison of tract reconstruction using different tractography techniques in macaques. (A) Comparison of tract anatomy obtained using FSL probabilistic, FSL deterministic, and MRtrix probabilistic techniques, as well as overlapping tracts between the three. The top panels show the 3D representation of the tracts, and the bottom ones show their organization on coronal sections. Thresholds for the tracts with FSL probabilistic and deterministic techniques are as follows: 0.7 for MDLF; 0.75 for IFOF; 0.7 for ILF; and 0.75 for AF. Thresholds for the tracts with MRtrix are as follows: 0.82 for MDLF; 0.82 for IFOF; 0.8 for ILF; and 0.8 for AF. The overlap is binarized. (B) Comparison of tract overlap when reconstructed with different techniques. We calculated the number of voxels overlapping between a tract of interest and each of the tracts obtained with FSL probabilistic divided by the total number of voxels of the tract of interest. The darker color represents the overlap of the tract of interest obtained with FSL probabilistic, and the lighter color represents the overlap of the tract of interest obtained with MRtrix and the color in between the tract of interest obtained with FSL deterministic. Macaque postmortem data are available from the PRIME-DE repository (http://fcon_1000.projects.nitrc.org/indi/PRIME/oxford2.html). AF, arcuate fascicle; FSL, FMRIB Software Library; IFOF, inferior fronto-occipital fascicle; ILF, inferior longitudinal fascicle; ILFmed, inferior longitudinal fascicle medial; MdLF, middle longitudinal fascicle; PRIME-DE, Primate Data Exchange.

S12 Fig. Comparison of tract reconstruction using different tractography techniques in humans. (A) Comparison of tract anatomy obtained using FSL probabilistic, FSL deterministic, and MRtrix probabilistic techniques, as well as overlapping tracts between the three. The top panels show the 3D representation of the tracts, and the bottom ones show their organization on coronal sections. Thresholds for the tracts with FSL probabilistic and deterministic
techniques are as follows: 0.7 for MDLF; 0.75 for IFOF; 0.7 for ILFlat and ILFmed; and 0.75 for AF. Thresholds for the tracts with MRtrix are as follows: 0.82 for MDLF; 0.82 for IFOF; 0.8 for ILFlat and ILFmed; and 0.8 for AF. The overlap is binarized. (B) Comparison of tract overlap when reconstructed with different techniques. We calculated the number of voxels overlapping between a tract of interest and each of the tracts obtained with FSL probabilistic divided by the total number of voxels of the tract of interest. The darker color represents the overlap of the tract of interest obtained with FSL probabilistic, and the lighter color represents the overlap of the tract of interest obtained with MRtrix and the color in between the tract of interest obtained with FSL deterministic. Macaque postmortem data are available from the PRIME-DE repository (http://fcon_1000.projects.nitrc.org/indi/PRIME/oxford2.html). AF, arcuate fascicle; FSL, FMRIB Software Library; IFOF, inferior fronto-occipital fascicle; ILFlat, inferior longitudinal fascicle lateral; ILFmed, inferior longitudinal fascicle medial; MdLF, middle longitudinal fascicle; PRIME-DE, Primate Data Exchange.

S13 Fig. Comparison of tract reconstruction using different tractography techniques in chimpanzees in vivo. (A) Comparison of tract anatomy obtained using FSL probabilistic and FSL deterministic techniques, as well as overlapping tracts between the two. The top panels show the 3D representation of the tracts, and the bottom ones show their organization on coronal sections. Thresholds for the tracts with FSL probabilistic and deterministic techniques are as follows: 0.7 for MDLF; 0.75 for IFOF; 0.7 for ILFlat and ILFmed; and 0.75 for AF. The overlap is binarized. (B) Comparison of tract overlap when reconstructed with different techniques. We calculated the number of voxels overlapping between a tract of interest and each of the tracts obtained with FSL probabilistic divided by the total number of voxels of the tract of interest. The darker color represents the overlap of the tract of interest obtained with FSL probabilistic, and the lighter color represents the overlap of the tract of interest obtained with FSL deterministic. In vivo chimpanzee data are available from the National Chimpanzee Brain Resource (www.chimpanzeebrain.org). AF, arcuate fascicle; FSL, FMRIB Software Library; IFOF, inferior fronto-occipital fascicle; ILFlat, inferior longitudinal fascicle lateral; ILFmed, inferior longitudinal fascicle medial; MdLF, middle longitudinal fascicle.

(TIF)

S14 Fig. Comparison of tract reconstruction using different tractography techniques in gorilla postmortem. (A) Comparison of tract anatomy obtained using FSL probabilistic and FSL deterministic techniques, as well as overlapping tracts between the two. The top panels show the 3D representation of the tracts, and the bottom ones show their organization on coronal sections. Thresholds for the tracts with FSL probabilistic and deterministic techniques are as follows: 0.7 for MDLF; 0.75 for IFOF; 0.7 for ILFlat and ILFmed; and 0.75 for AF. The overlap is binarized. (B) Comparison of tract overlap when reconstructed with different techniques. We calculated the number of voxels overlapping between a tract of interest and each of the tracts obtained with FSL probabilistic divided by the total number of voxels of the tract of interest. The darker color represents the overlap of the tract of interest obtained with FSL probabilistic, and the lighter color represents the overlap of the tract of interest obtained with FSL deterministic. Gorilla postmortem data are available from https://doi.org/10.5281/zenodo.3901205. AF, arcuate fascicle; FSL, FMRIB Software Library; IFOF, inferior fronto-occipital fascicle; ILFlat, inferior longitudinal fascicle lateral; ILFmed, inferior longitudinal fascicle medial; MdLF, middle longitudinal fascicle.
S1 Text. Tractography as the best method to study white matter in a large range of species.

(DOCX)

S2 Text. Additional methods.

(DOCX)

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References

1. Bryant KL, Preuss TM. A Comparative Perspective on the Human Temporal Lobe. In: Bruner E, Ogi-hara N, Tanabe HC, editors. Digital Endocasts. Tokyo: Springer Japan; 2018. pp. 239–258.

2. Lambon Ralph MA, Sage K, Jones RW, Mayberry EJ. Coherent concepts are computed in the anterior temporal lobes. Proc Natl Acad Sci. 2010; 107: 2717–2722. https://doi.org/10.1073/pnas.0907307107

PMID: 20133780
3. Miller EK, Nieder A, Freedman DJ, Wallis JD. Neural correlates of categories and concepts. Curr Opin Neurobiol. 2003; 13: 198–203. https://doi.org/10.1016/s0959-4388(03)00037-0 PMID: 12744974

4. Schurz M, Radua J, Aichhorn M, Richlan F, Perner J. Fractionating theory of mind: A meta-analysis of functional brain imaging studies. Neurosci Biobehav Rev. Elsevier Ltd; 2014; 42: 9–34. https://doi.org/10.1016/j.neubiorev.2014.01.009 PMID: 24486722

5. Sallet J, Mars RB, Noonan MP, Andersson JL, O'Reilly JX, Jbabdi S, et al. Social network size affects neural circuits in macaques. Science (80- ). 2011; 334: 697–700. https://doi.org/10.1126/science.1210027 PMID: 22053054

6. Price CJ. The anatomy of language: contributions from functional neuroimaging. J Anat. 2000; 197: 332–359.

7. Mars RB, Eichert N, Jbabdi S, Verhagen L, Rushworth MFS. Connectivity and the search for specializations in the language-capable brain. Curr Opin Behav Sci. 2018; 21: 19–26. https://doi.org/10.1016/j.cobeha.2017.11.001

8. Eichert N, Robinson EC, Bryant KL, Jbabdi S, Jenkinson M, Li L, et al. Cross-species cortical alignment identifies different types of anatomical reorganization in the primate temporal lobe. Elife. 2020; 9. https://doi.org/10.7554/eLife.53232 PMID: 32202497

9. Smaers JB, Vanier DR. Brain size expansion in primates and humans is explained by a selective modular expansion of the cortico-cerebellar system. Cortex. Elsevier Ltd; 2019; 118: 292–305. https://doi.org/10.1016/j.cortex.2019.04.023 PMID: 31213287

10. Striedter GF, Belgard TG, Chen C-C, Davis FP, Finlay BL, Güntürkün O, et al. NSF Workshop Report: Discovering General Principles of Nervous System Organization by Comparing Brain Maps across Species. Brain Behav Evol. 2014; 83: 1–8. https://doi.org/10.1159/000360152 PMID: 24603302

11. Lerch JP, Kouve AJW van der, Raznahan A, Paus T, Johansen-Berg H, Miller KL, et al. Studying neuroanatomy using MRI. Nat Neurosci. 2017; 20: 314–326. https://doi.org/10.1038/nn.4501 PMID: 28230838

12. Mars RB, Neubert F-X, Verhagen L, Sallet J, Miller KL, Dunbar RIM, et al. Primate comparative neuroscience using magnetic resonance imaging: promises and challenges. Front Neurosci. 2014; 8. https://doi.org/10.3389/fnins.2014.00298 PMID: 25339857

13. Mars RB, Foxley S, Verhagen L, Jbabdi S, Sallet J, Noonan MP, et al. The extreme capsule fiber complex in humans and macaque monkeys: a comparative diffusion MRI tractography study. Brain Struct Funct. 2016; 221: 4059–4071. https://doi.org/10.1007/s00429-015-1146-0 PMID: 26627483

14. Eichert N, Verhagen L, Folloni D, Jbabdi S, Khrapitchev AA, Sibson NR, et al. What is special about the human arcuate fasciculus? Lateralization, projections, and expansion. Cortex. Elsevier Ltd; 2018; 1–9. https://doi.org/10.1016/J.CORT EX.2018.05.005 PMID: 29937266

15. Rilling JK, Glasser MF, Preuss TM, Ma X, Zhao T, Hu X, et al. The evolution of the arcuate fasciculus revealed with comparative DTI. Nat Neurosci. 2008; 11: 426–428. https://doi.org/10.1038/nn2072 PMID: 18344993

16. Mars RB, Sallet J, Neubert F-X, Rushworth MFS. Connectivity profiles reveal the relationship between brain areas for social cognition in human and monkey temporoparietal cortex. Proc Natl Acad Sci. 2013; 110: 10806–10811. https://doi.org/10.1073/pnas.1302956110 PMID: 23754406

17. Orban GA, Van Essen DC, Vanduffel W. Comparative mapping of higher visual areas in monkeys and humans. Trends Cogn Sci. 2004; 8: 315–324. https://doi.org/10.1016/j.tics.2004.05.009 PMID: 15242691

18. Bryant KL, Glasser MF, Li L, Jae-Cheol Bae J, Jacquez NJ, Alarcón L, et al. Organization of extrastriate and temporal cortex in chimpanzees compared to humans and macaques. Cortex. 2019; 118: 223–243. https://doi.org/10.1016/j.cortex.2019.02.010 PMID: 30910223

19. Latini F, Mårtensson J, Larsson E, Fredrikson M, Åhs F. Segmentation of the inferior longitudinal fasciculus in the human brain: A white matter dissection and diffusion tensor tractography study. Brain Res. Elsevier B.V.; 2017; 1675: 102–115. https://doi.org/10.1016/j.brainres.2017.09.005 PMID: 28899757

20. Mars RB, Sotropoulos SN, Passingham RE, Sallet J, Verhagen L, Khrapitchev AA, et al. Whole brain comparative anatomy using connectivity blueprints. Elife. 2018; 7: 245209. https://doi.org/10.7554/eLife.35237 PMID: 29749930

21. Passingham RE, Stephan KE, Kötter R. The anatomical basis of functional localization in the cortex. Nat Rev Neurosci. 2002; 3: 606–616. https://doi.org/10.1038/nn93 PMID: 12154362

22. Schmahmann JD, Pandya DN, Wang R, Dai G, D’Arceuil HE, De Crespigny AJ, et al. Association fibre pathways of the brain: Parallel observations from diffusion spectrum imaging and autoradiography. Brain. 2007; 130: 630–653. https://doi.org/10.1093/brain/aw359 PMID: 17293361
23. Decramer T, Swinnen S, van Loon J, Janssen P, Theys T. White matter tract anatomy in the rhesus monkey: a fiber dissection study. Brain Struct Funct. Springer Berlin Heidelberg; 2018; 223: 3681–3688. https://doi.org/10.1007/s00429-018-1718-x PMID: 30022520

24. Milham MP, Ai L, Koo B, Xu T, Amiez C, Balezeau F, et al. An Open Resource for Non-human Primate Imaging. Neuron. 2018; 100: 61–74.e2. https://doi.org/10.1016/j.neuron.2018.08.039 PMID: 30269990

25. Panesar SS, Yeh F-C, Jacquesson T, Hula W, Fernandez-Miranda JC. A Quantitative Tractography Study Into the Connectivity, Segmentation and Laterality of the Human Inferior Longitudinal Fasciculus. Front Neuroanat. 2018; 12: 1–13. https://doi.org/10.3389/fnana.2018.00001 PMID: 29440997

26. Reveley C, Seth AK, Pierpaoli C, Silva AC, Yu D, Saunders RC, et al. Superficial white matter fiber systems impede detection of long-range cortical connections in diffusion MR tractography. Proc Natl Acad Sci. 2015; 112: E2820–E2828. https://doi.org/10.1073/pnas.1418198112 PMID: 25964365

27. Catani M, Thiebaut De Schotten M. A diffusion tensor imaging tractography atlas for virtual in vivo dissections. Cortex. 2008; 44: 1105–1132. https://doi.org/10.1016/j.cortex.2008.05.004 PMID: 18619589

28. Catani M, Jones DK, Donato R, Ffytche DH. Occipito-temporal connections in the human brain. Brain. 2004; 127 (Pt 8): 1577–1596. https://doi.org/10.1093/brain/awh164 PMID: 15269090

29. Schmahmann JD, Pandya DN. Fiber pathways of the brain. Oxford: Oxford University Press; 2006.

30. Foloni D, Sallet J, Khrapitchev AA, Sibson N, Verhagen L, Mars RB. Dichotomous organization of amygdala/temporal-prefrontal bundles in both humans and monkeys. Elife. 2018; 8: 1–23. https://doi.org/10.7554/eLife.41775 PMID: 31689177

31. Passingham RE, Smaers JB. Is the prefrontal cortex especially enlarged in the human brain? Allometric relations and remapping factors. Brain Behav Evol. 2014; 84: 156–166. https://doi.org/10.1159/000365183 PMID: 25248097

32. Glasser MF, Goyal MS, Preuss TM, Raichle ME, Van Essen DC. Trends and properties of human cerebral cortex: Correlations with cortical myelin content. Neuroimage. Elsevier Inc.; 2014; 93: 165–175. https://doi.org/10.1016/j.neuroimage.2013.03.060 PMID: 23567887

33. Smaers JB, Gómez-Robles A, Parks AN, Sherwood CC. Exceptional Evolutionary Expansion of Prefrontal Cortex in Great Apes and Humans. Curr Biol. 2017; 27: 714–720. https://doi.org/10.1016/j.cub.2017.01.020 PMID: 28162899

34. Preuss TM. The human brain: rewired and running hot. Ann N Y Acad Sci. 2011; 1225: E182–E191. https://doi.org/10.1111/j.1749-6632.2011.06001.x PMID: 21599696

35. Forkel SJ, Thiebaut de Schotten M, Kawadler JM, Dell’Acqua F, Danek A, Catani M. The anatomy of fronto-occipital connections from early blunt dissections to contemporary tractography. Cortex. 2014; 56: 73–84. https://doi.org/10.1016/j.cortex.2012.09.005 PMID: 23137651

36. Takemura H, Pestilli F, Weiner KS, Keliris GA, Landi SM, Sliwa J, et al. Occipital White Matter Tracts in Human and Macaque. Cereb Cortex. 2017; 27: 3346–3359. https://doi.org/10.1093/cercor/bhx070 PMID: 28369290

37. Sarubbo S, Petit L, De Benedictis A, Chioffi F, Pito M, Dyrbj TB. Uncovering the inferior fronto-occipital fascicle and its topological organization in non-human primates: the missing connection for language evolution. Brain Struct Funct. Springer Berlin Heidelberg; 2019; 0: 0. https://doi.org/10.1007/s00429-019-01856-2 PMID: 30847641

38. Mars RB, O’Muircheartaigh J, Foloni D, Li L, Glasser MF, Jbabdi S, et al. Concurrent analysis of white matter bundles and grey matter networks in the chimpanzee. Brain Struct Funct. Springer Berlin Heidelberg; 2018; 0: 0. https://doi.org/10.1007/s00429-018-1817-8 PMID: 30569281

39. Makris N, Pandya DN. The extreme capsule in humans and rethinking of the language circuitry. Brain Struct Funct. 2009; 213: 343–358. https://doi.org/10.1007/s00429-008-0199-8 PMID: 19104833

40. Almairac F, Herbet G, Moritz-Gasser S, de Champfleur NM, Duffau H. The left inferior fronto-occipital fasciculus subserves language semantics: a multilevel lesion study. Brain Struct Funct. Springer Berlin Heidelberg; 2015; 220: 1983–1995. https://doi.org/10.1007/s00429-014-0773-1 PMID: 24744151

41. Panesar SS, Yeh FC, Deibert CP, Fernandes-Cabral D, Rowthu V, Celtikci P, et al. A diffusion spectrum imaging-based tractographic study into the anatomical subdivision and cortical connectivity of the ventral external capsule: uncinate and inferior fronto-occipital fascicles. Neuroradiology. Neuroradiology; 2017; 59: 971–987. https://doi.org/10.1007/s00234-017-1874-3 PMID: 28721443

42. Herbet G, Zemmoura I, Duffau H. Functionality and Inferior Anatomy of the Inferior Longitudinal Fasciculus: From Historical Reports to Current Hypotheses. Front Neuroanat. 2018; 12: 1–15. https://doi.org/10.3389/fnana.2018.00001 PMID: 29440997

43. Wang Y, Olson IR. The Original Social Network: White Matter and Social Cognition. Trends Cogn Sci. Elsevier Ltd; 2018; 22: 504–516. https://doi.org/10.1016/j.tics.2018.03.005 PMID: 29628441

44. Tsao DY, Moeller S, Freiwald WA. Comparing face patch systems in macaques and humans. Proc Natl Acad Sci. 2008; 105: 19514–19519. https://doi.org/10.1073/pnas.0809662105 PMID: 19033466
45. Kanwisher N, McDermott J, Chun MM. The Fusiform Face Area: A Module in Human Extrastriate Cortex Specialized for Face Perception. J Neurosci. 1997; 17: 4302–4311. https://doi.org/10.1523/JNEUROSCI.17-11-04302.1997 PMID: 9151747

46. Ku SP, Tolas AS, Logothetis NK, Goene J. FMRI of the Face-Processing Network in the Ventral Temporal Lobe of Awake and Anesthetized Macaques. Neuron. Elsevier Inc.; 2011; 70: 352–362. https://doi.org/10.1016/j.neuron.2011.02.048 PMID: 21521619

47. Porr LA, Hecht E, Barks SK, Preuss TM, Votaw JR. Face Processing in the Chimpanzee Brain. Curr Biol. Elsevier Ltd; 2009; 19: 50–53. https://doi.org/10.1016/j.cub.2008.11.048 PMID: 19097899

48. Maier-Hein KH, Neher PF, Houde J-C, Côté M-A, Garyfallidis E, Zhong J, et al. The challenge of mapping the human connectome based on diffusion tractography. Nat Commun. 2017; 8: 1349. https://doi.org/10.1038/s41467-017-01285-x PMID: 29116093

49. Donahue CJ, Sotopoulos SN, Jbabdi S, Hernandez-Fernandez M, Behrens TEJ, Dyrby TB, et al. Using Diffusion Tractography to Predict Cortical Connection Strength and Distance: A Quantitative Comparison with Tracers in the Monkey. J Neurosci. 2016; 36: 6758–6770. https://doi.org/10.1523/JNEUROSCI.0493-16.2016 PMID: 27335406

50. Hecht EE, Gutman DA, Preuss TM, Sanchez MM, Porr LA, Rilling JK. Process Versus Product in Social Learning: Comparative Diffusion Tensor Imaging of Neural Systems for Action Execution—Observation Matching in Macaques, Chimpanzees, and Humans. Cereb Cortex. 2013; 23: 1014–1024. https://doi.org/10.1093/cercor/bhs097 PMID: 22539611

51. Rilling JK, Glasser M, Jbabdi S, Andersson J, Preuss T. Continuity, divergence, and the evolution of brain language pathways. Front Evol Neurosci. 2012; 3: 1–6. https://doi.org/10.3389/fnevo.2011.00011 PMID: 22319495

52. Heuer K, Gulban OF, Bazin P-L, Osoianu A, Valabregue R, Santin M, et al. Evolution of neocortical folding: A phylogenetic comparative analysis of MRI from 33 primate species. bioRxiv. 2018; 379750. https://doi.org/10.1101/379750

53. Passingham RE, Smaers JB, Sherwood CC. Evolutionary Specializations of the Human Prefrontal Cortex. In: Kass J., Preuss TM, editors. Evolution of Nervous Systems. 2nd ed. Elsevier Press; 2017. pp. 207–226. https://doi.org/10.1016/B978-0-12-804042-3.00114-7

54. Murray EA, Wise SP, Graham KS. The Evolution of Memory Systems. Oxford University Press; 2016. https://doi.org/10.1093/acprof:oso/9780199686438.001.0001

55. Krubitzer L, Kaas J. The evolution of the neocortex in mammals: How is phenotypic diversity generated? Curr Opin Neurobiol. 2005; 15: 444–453. https://doi.org/10.1016/j.conb.2005.07.003 PMID: 16026978

56. Chaplin TA, Yu HH, Soares JGM, Gattass R, Rosa MGP. A conserved pattern of differential expansion of cortical areas in simian primates. J Neurosci. 2013; 33: 15120–15125. https://doi.org/10.1523/JNEUROSCI.2909-13.2013 PMID: 24048842

57. Large I, Bridge H, Ahmed B, Clare S, Kolasinski J, Lam WW, et al. Individual Differences in the Alignment of Structural and Functional Markers of the V5/MT Complex in Primates. Cereb Cortex. 2016; 26: 3928–3944. https://doi.org/10.1093/cercor/bhw180 PMID: 27371764

58. Barton RA, Harvey PH. Mosaic evolution of brain structure in mammals. Nature. Elsevier; 2000; 405: 1055–1058. https://doi.org/10.1038/35016580 PMID: 10890446

59. Catani M, Bambini V. A model for Social Communication And Language Evolution and Development (SCALED). Curr Opin Neurobiol. Elsevier Ltd; 2014; 28: 165–171. https://doi.org/10.1016/j.conb.2014.07.018 PMID: 25156623

60. Friederici AD. Language development and the ontogeny of the dorsal pathway. Front Evol Neurosci. 2012; 4: 1–7. https://doi.org/10.3389/fnevo.2012.00001 PMID: 22319496

61. Xiang H, Van Leeuwen TM, Dediu D, Roberts L, Norris DG, Hagoort P. L2-Proficiency-Dependent Laterality Shift in Structural Connectivity of Brain Language Pathways. Brain Connect. 2015; 5: 349–361. https://doi.org/10.1089/brain.2013.0199 PMID: 25594261

62. Ardesch D, Scholtens L, Li L, Preuss TM, Rilling JK, van den Heuvel MP. Evolutionary expansion of connectivity between multimodal association areas in the human brain compared to chimpanzees. Pnas. 2019; https://doi.org/10.1073/pnas.1818512116 PMID: 30886094

63. Makris N, Papadimitriou GM, Kaiser JR, Sorg S, Kennedy DN, Pandya DN. Delineation of the middle longitudinal fascicle in humans: A quantitative, in vivo, DT-MRI study. Cereb Cortex. 2009; 19: 777–785. https://doi.org/10.1093/cercor/bhn124 PMID: 18669591

64. Eickhoff SB, Thirion B, Varoquaux G, Bzdok D. Connectivity-based parcellation: Critique and implications. Hum Brain Mapp. 2015; 36: 4771–4792. https://doi.org/10.1002/hbm.22933 PMID: 26409749

65. Mars RB, Jbabdi S, Sallet J, O’Reilly JX, Croxson PL, Olivier E, et al. Diffusion-Weighted Imaging Tractography-Based Parcellation of the Human Parietal Cortex and Comparison with Human and Macaque
Resting-State Functional Connectivity. J Neurosci. 2011; 31: 4087–4100. https://doi.org/10.1523/JNEUROSCI.5102-10.2011 PMID: 21411650

66. Schmahmann JD, Pandya DN. Composite Summary of Cerebral White Matter Fiber Pathways in the Rhesus Monkey. Fiber Pathways of the Brain. Oxford University Press; 2006. pp. 533–554. https://doi.org/10.1093/acprof:oso/9780195104233.003.0027

67. Behrens TEJ, Berg HJ, Jbabdi S, Rushworth MFS, Woolrich MW. Probabilistic diffusion tractography with multiple fibre orientations: What can we gain? Neuroimage. 2007; 34: 144–155. https://doi.org/10.1016/j.neuroimage.2006.09.018 PMID: 17070705

68. Zhang Y, Brady M, Smith S. Segmentation of brain MR images through a hidden Markov random field model and the expectation-maximization algorithm. IEEE Trans Med Imaging. 2001; 20: 45–57. https://doi.org/10.1109/42.906424 PMID: 11293691

69. Hou L, Xiang L, Crow TJ, Leroy F, Riviere D, Mangin JF, et al. Measurement of Sylvian Fissure asymmetry and occipital bending in humans and Pan troglodytes. Neuroimage. 2019; 184: 855–870. https://doi.org/10.1016/j.neuroimage.2018.08.045 PMID: 30170149

70. Marcus DS, Harwell J, Olsen T, Hodge M, Glasser MF, Prior F, et al. Informatics and Data Mining Tools and Strategies for the Human Connectome Project. Front Neuroinform. 2011; 5: 1–12. https://doi.org/10.3389/fninf.2011.00001 PMID: 21472085