High Angular Resolution Diffusion MRI Reveals Conserved and Deviant Programs in the Paths that Guide Human Cortical Circuitry

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Abstract

Diffusion magnetic resonance (MR) tractography represents a novel opportunity to investigate conserved and deviant developmental programs between humans and other species such as mice. To that end, we acquired high angular resolution diffusion MR scans of mice [embryonic day (E) 10.5 to postnatal week 4] and human brains [gestational week (GW) 17–30] at successive stages of fetal development to investigate potential evolutionary changes in radial organization and emerging pathways between humans and mice. We compare radial glial development as well as commissural development (e.g., corpus callosum), primarily because our findings can be integrated with previous work. We also compare corpus callosal growth trajectories across primates (i.e., humans and rhesus macaques) and rodents (i.e., mice). One major finding is that the developing cortex of humans is predominated by pathways likely associated with a radial glial organization at GW 17–20, which is not as evident in age-matched mice (E 16.5, 17.5). Another finding is that, early in development, the corpus callosum follows a similar developmental timetable in primates (i.e., macaques and humans) as in mice. However, the corpus callosum grows for an extended period of time in primates compared with rodents. Taken together, these findings highlight deviant developmental programs underlying the emergence of cortical pathways in the human brain.

Key words: brain pathways, callosal, cingulum, diffusion MRI, fetus, human, mouse, tractography
**Introduction**

Diffusion magnetic resonance imaging (MRI) is a high-throughput and noninvasive method that can be used to investigate neural pathways in adulthood, as well as their development (Mori and Zhang 2006; Qiu et al. 2015). Questions focused on the development and evolution of connectivity patterns and cell migration have traditionally been addressed with tracers or carbocyanine tracers. Although these methods have revealed key developmental processes in the brain, they are time-consuming and invasive and suffer from a number of technical limitations (Nudo and Masterton 1989, 1990; Chen et al. 2012; Hynd 2005; Greig et al. 2013), which have undergone a dramatic extension in the duration of cortical neurogenesis should entail an extended period in which fibers are radially organized to allow newly born neurons to migrate from the progenitor pool to the CP. Extending the duration of neurogenesis might also lead to major changes in cortical circuitry by potentially altering the timing of initial axon extension of pioneer neurons and/or extending the duration over which cortical association fibers grow. Many of the callosally projecting neurons are situated in the upper layers (i.e., layer II–IV) (Isseroff et al. 1984; Meissner et al. 1991; Greig et al. 2013), which have undergone a dramatic expansion in primates (Charvet et al. 2017a, 2019), and a concomitant increased expression of select genes encoding ion channels, neurofilament, and synaptic proteins in upper layers (Zeng et al. 2012; Krienen et al. 2016). In addition, callosally projecting neurons express a suite of genes, some of which are differentially expressed in macaques compared with mice (Fame et al. 2011, 2016a). These studies highlight major modifications in callosal organization between primates and rodents. Thus, we can use diffusion MR tractography to investigate deviations in developmental processes giving rise to variation in cortico-cortical association pathways between species.

It is well known that primate isocortical neurogenesis is unusual compared with rodents (Rakic 1995; Zecevic et al. 2005; Kriegstein et al. 2006; Martinez-Cerdeño et al. 2006). After controlling for differences in developmental schedules, isocortical neuronal production extends for an unusually long time in humans and macaques compared with mice (Clancy et al. 2001; Workman et al. 2013; Cahalane et al. 2014; Charvet et al. 2017a). The extension in the duration of isocortical neurogenesis leads to an amplification of upper layer neurons (i.e., layer II–IV) and a concomitant expansion of long-range cortico-cortical association pathways in adult primates (Charvet et al. 2015, 2017a, 2019; Krienen et al. 2016). Given these data, it may be expected that there are differences in radial glial organization between humans and mice during neurogenesis because an extension in the duration of cortical neurogenesis should entail an extended period in which fibers are radially organized to allow newly born neurons to migrate from the progenitor pool to the CP. Extending the duration of neurogenesis might also lead to major changes in cortical circuitry by potentially altering the timing of initial axon extension of pioneer neurons and/or extending the duration over which cortical association fibers grow. Many of the callosally projecting neurons are situated in the upper layers (i.e., layer II–IV) (Isseroff et al. 1984; Meissner et al. 1991; Greig et al. 2013), which have undergone a dramatic expansion in primates (Charvet et al. 2017a, 2019), and a concomitant increased expression of select genes encoding ion channels, neurofilament, and synaptic proteins in upper layers (Zeng et al. 2012; Krienen et al. 2016). In addition, callosally projecting neurons express a suite of genes, some of which are differentially expressed in macaques compared with mice (Fame et al. 2011, 2016a). These studies highlight major modifications in callosal organization between primates and rodents. Thus, we can use diffusion MR tractography to investigate deviations in developmental processes giving rise to the evolution of cortical association pathways in the human lineage.

Considerable work has been conducted on radial glial development, pioneer, as well as callosal development in model organisms such as mice, rats, monkeys, and cats (Wise and Jones 1976; Marin-Padilla 1978; Innocenti 1981; O’Leary and O’Leary 1992; Koester and O’Leary 1994; Rash and Richards 2001; Paul et al. 2007). As more and more cells exit the cell cycle, the progenitor pool, the radial organization of the developing human fetal cortex wane, and the white matter of the developing cortex become increasingly replaced with long-range projecting pathways as assessed from diffusion MRI (Takahashi et al. 2012). The aim of the present study is to compare the timetable of "scaffolds" such as radial glial fibers and pioneer axons between species. Although "pioneer" might imply a guidance function, we define pioneer exclusively temporally, and we focus on the temporal sequence of fibers emerging across the hemisphere of the developing cortex. Another aim of the present study is to compare the timing of cortical association pathway maturation (i.e., corpus callosum) between human and mice to investigate evolutionary changes in developmental processes giving rise to variation in cortico-cortical association pathways between species. It is well known that primate isocortical neurogenesis is unusual compared with rodents (Rakic 1995; Zecevic et al. 2005; Kriegstein et al. 2006; Martinez-Cerdeño et al. 2006). After controlling for differences in developmental schedules, isocortical neuronal production extends for an unusually long time in humans and macaques compared with mice (Clancy et al. 2001; Workman et al. 2013; Cahalane et al. 2014; Charvet et al. 2017a). The extension in the duration of isocortical neurogenesis leads to an amplification of upper layer neurons (i.e., layer II–IV) and a concomitant expansion of long-range cortico-cortical association pathways in adult primates (Charvet et al. 2015, 2017a, 2019; Krienen et al. 2016). Given these data, it may be expected that there are differences in radial glial organization between humans and mice during neurogenesis because an extension in the duration of cortical neurogenesis should entail an extended period in which fibers are radially organized to allow newly born neurons to migrate from the progenitor pool to the CP. Extending the duration of neurogenesis might also lead to major changes in cortical circuitry by potentially altering the timing of initial axon extension of pioneer neurons and/or extending the duration over which cortical association fibers grow. Many of the callosally projecting neurons are situated in the upper layers (i.e., layer II–IV) (Isseroff et al. 1984; Meissner et al. 1991; Greig et al. 2013), which have undergone a dramatic expansion in primates (Charvet et al. 2017a, 2019), and a concomitant increased expression of select genes encoding ion channels, neurofilament, and synaptic proteins in upper layers (Zeng et al. 2012; Krienen et al. 2016). In addition, callosally projecting neurons express a suite of genes, some of which are differentially expressed in macaques compared with mice (Fame et al. 2011, 2016a). These studies highlight major modifications in callosal organization between primates and rodents. Thus, we can use diffusion MR tractography to investigate deviations in developmental processes giving rise to the evolution of cortical association pathways in the human lineage.
Materials and Methods

We first discuss sample acquisition, scanning parameters, and tractography to compare the developmental time course of fibers from diffusion MR tractography in humans (n = 5) and in mice (n = 5; Figs 1-3). We then discuss how we measured and compared corpus callosum growth trajectories in humans, macaques, and mice (Fig. 4). Corpus callosum scans for humans (Shi et al. 2011; Cohen et al. 2016; Khan et al. 2018a, 2018b), macaques (Young et al. 2017), and mice (Chuang et al. 2011) were acquired from multiple sources. Scanning parameters and sample size, as well sources are listed in Supplementary Table 1.

Sample Acquisition for Diffusion MR Tractography

Mice

Five intact mouse brains at E 10.5, 14.5, 16.5, 17.5, and 4 postnatal weeks (PW) were obtained from ongoing developmental or genetic research projects at Boston Children’s Hospital (BCH) and Massachusetts General Hospital (MGH) under approved protocols at the hospitals. Mice were deeply anesthetized (xylazine/ketamine 100 mg/mL, 60 mg/kg) and either sacrificed by transcardial perfusion with 0.9% saline followed by 4% paraformaldehyde (PFA) (>postnatal day 5) or decapitated and immersion-fixed (<postnatal day 3). Fetal brains were immersion-fixed after the removal of the skull. Postnatal animals were perfused with 4% PFA followed by immersion fixation for at least 2 weeks in 4% PFA. Brains were placed in Formblin for imaging.

Humans

A total of five brains at GW 17, 20, 21, 22, and 30 were imaged for diffusion MR tractography. Samples from GW 17-22 were scanned ex vivo, whereas the GW 30 individual was scanned in vivo. Postmortem brains were obtained from the Department of Pathology, Brigham and Women’s Hospital (BWH, Boston, MA, USA) and the Allen Institute Brain Bank (AIBB, Seattle, WA, USA) with full parental consent. The known primary causes of death were complications from prematurity. The brains were grossly normal, and standard autopsy examinations of all brains undergoing postmortem HARDI revealed no or minimal pathologic abnormalities at the macroscopic level. All brains utilized for this study were analyzed under institutional review board-approved protocols.

Tissue Preparation for HARDI in Humans

At the time of autopsy, all brains were immersion-fixed, and the brains from the BWH were stored in 4% PFA, and the brains from the AIBB were stored in 4% periodate-lysine-paraformaldehyde (PLP). During MR image acquisition, the brains from the Brigham and Women’s Hospital were placed in Formblin solution (Aussmont; e.g., Takahashi et al. 2012) and the brains from the AIBB were placed in 4% PLP. These different kinds of solutions in which the brains from different institutes were placed tend to change the background contrast (i.e., a dark background outside of the brain using Formblin and bright background using PLP), but they do not specifically change diffusion properties [e.g., fractional anisotropy (FA) and apparent diffusion coherence (ADC)] within the brain.

Scanners

Different scanner systems were used to accommodate brains of various sizes (same as in e.g., Takahashi et al. 2012; Xu et al. 2014; Miyazaki et al. 2016; Wilkinson et al. 2017). Magnetic resonance (MR) coils that best fit each brain sample were used to ensure optimal imaging. Mouse specimens as well as human specimens from BWH [2 fetal specimens (GW 17, 20)] were imaged with a 9.4T Bruker Biospec MR system. Ex-vivo specimens from the AIBB were imaged with a 3 T Siemens MR system [2 fetal specimens (GW 21, 22) at the A. A. Martinos Center, MGH, Boston, MA, USA, because they did not fit in the 4.7T bore. In order to obtain the best signal-to-noise ratio and highest spatial resolution, we used custom-made MR coils with one channel on the 4.7T and 3T systems (Takahashi et al. 2012; Kolasinski et al. 2013; Xu et al. 2014). The in vivo subject (GW 30) from BCH was imaged on a 3T Siemens MR system at BCH, Boston, MA, USA, when the individual was asleep and not anesthetized. At a 3T scanner, the use of high b values such as 8000 s/mm² is problematic because it causes artifacts in that the tractography appears inconsistent with previous findings from tract-tracers. We reason that the issue of different acquisition systems is not significant for tractography because both systems yielded similar patterns.

Scanning Protocols

Scanning protocols are listed in Supplementary Table 1. For the human brains at GW 21 and GW 22 from BWH and for mouse specimens, a 3D diffusion-weighted spin-echo echo-planar imaging sequence was used. Sixty diffusion-weighted measurements (with the strength of the diffusion weighting,
the DTI tractography pathways. This feature is a recognized limitation with the traditional DTI technique because one simply connects each local maxima on an ODF, we applied the streamline algorithm with the principal eigenvector on a tensor to produce the DTI tractography pathways. This feature is a recognized limitation of DTI (Mori et al. 1999). Hence, in the current study, we used HARDI, which can theoretically detect multiple local maxima on an orientation distribution function (ODF). Using each local maxima on an ODF, we applied the streamline algorithm with Q ball to initiate and continue tractography (Tuch et al. 2003), thus enabling us to identify crossing pathways within a voxel.

Diffusion MR trajectories were propagated by consistently pursuing the orientation vector of least curvature. We terminated tracking when the angle between two consecutive orientation vectors was greater than the given threshold (40°), or when the fibers extended outside of the brain surface. For visualization purposes, the color-coding of tractography connections was based on a standard red–green–blue (RGB) code using directions of the middle segment of each fiber. In some cases, this RGB code relies on the direction of every fiber segment. In other cases, the RGB code applies to the average direction of fibers.

Reconstruction and Identification of Diffusion MR Tractography
We reconstructed white matter tracts in each brain with the Diffusion Toolkit and TrackVis (http://trackvis.org). We used a FACT algorithm for diffusion MR tractography (Mori et al. 1999), as in previous publications (for humans, Takahashi et al. 2010, 2011, 2012; Schmahmann et al. 2007; D’Arceuil et al. 2008; Das and Takahashi 2018; for mice, Rosen et al. 2013; Lodato et al. 2014; Fame et al. 2016a; Kanamaru et al. 2017), which connects fibers using a local maximum or maxima. The FACT algorithm is limited in its ability to resolve crossing pathways when used with the traditional DTI technique because one simply connects the direction of the principal eigenvector on a tensor to produce the DTI tractography pathways. This feature is a recognized limitation of DTI (Mori et al. 1999). Hence, in the current study, we used HARDI, which can theoretically detect multiple local maxima on an orientation distribution function (ODF). Using each local maxima on an ODF, we applied the streamline algorithm with Q ball to initiate and continue tractography (Tuch et al. 2003), thus enabling us to identify crossing pathways within a voxel.

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Regions of Interest to Select Fibers
We first extracted fibers from whole brain tractography, and we used slices or hand-drawn regions of interest (ROIs) to highlight pathways of interest in humans and mice. Whole brain tractography captures fibers coursing through the whole brain and permits to detect how overall organization of fibers might vary between species. For whole brain tractography and slice filters, we had a minimum length threshold set to about 1 mm for a visualization purpose. A single slice filter (either set along the horizontal or sagittal direction) permits viewing fibers coursing through the slice of interest. To study fibers in contact with the medial walls of the developing cortex, multiple sagittal slice filters of varying thickness with “either end” restrictions were used. Thickness of slice filters and placement were dependent on which pathway we aimed to capture during development (e.g., corpus callosum and anterior commissure). No threshold as to minimum or maximum FA was used to include or exclude fibers. We did not constrain the ROIs to the developing white or gray matter to highlight select commissural fibers (e.g., corpus callosum and anterior commissure) and track their developmental time course in both species. ROIs were variably placed to highlight commissural fibers of interest. By changing the size of the ROIs, we ensured not to miss pathways of interest.

Variation in Corpus Callosum Area Between Primates and Rodents
It has been previously reported that the corpus callosum is expanded in primates compared with other mammals (Manger et al. 2010). We aimed to confirm that the corpus callosum is indeed expanded in primates compared with rodents. To that end, we use structural MRI and atlases to measure the corpus callosum area from midsagittal sections in small-brained monkeys and large-brained rodents in order to control for variation in overall brain size. We supplement these data with previously published data (see Supplementary Table 1; Manger et al. 2010; Yuasa et al. 2010; Charvet et al. 2015, 2019; Schilling et al. 2017). We use phylogeny-generalized-least-squares statistics (PGLS) to obtain phylogenetically controlled slopes of the logged values of the area of corpus callosum versus the logged values of brain size (Pagel 1999; Freckleton et al. 2000). We test whether taxonomy (i.e., primate versus rodent) accounts for a significant percentage of the variation in corpus callosum area. These phylogenetically controlled generalized linear models were performed via restricted estimated maximum likelihood with the software program R and the library package ape (Pagel 1999; Freckleton et al. 2000). The phylogeny, which includes branch lengths for these species, was taken from Bininda-Emonds et al. 2007.

After controlling for variation in brain size, primates possess a larger corpus callosum compared with rodents (see Supplementary Table 1). A PGLS model with brain size and taxonomy are both significant independent variables accounting for 98.2% of the variance in corpus callosum area ($F = 348; R^2 = 0.981; P = 1.137 \times 10^{-10}; \lambda = 0$) and an Akaike information criterion (AIC) score of $-25.51$. A PGLS model that uses brain size alone to predict corpus callosum area only accounts for 96.8% of the variance ($F = 393.3; P = 1.539 \times 10^{-10}; \lambda = 0$), with an AIC score model of $-18.50$. The addition of taxonomy as an independent variable in the PGLS model results in a lower AIC score than the model without taxonomy as an independent variable. These findings confirm that primates possess a larger corpus callosum than rodents when controlled for brain size.

Growth Trajectories of the Human, Macaque, and Mouse Corpus Callosum Area
Comparative analyses of growth trajectories can pinpoint what developmental programs might account for variation in cortical association pathways in humans versus mice. To quantitatively investigate whether the corpus callosum growth follows a similar trajectory across primates and rodents, we measured the area of the corpus callosum from...
midsagittal sections of FA images from mice, macaques, and humans at several stages of development (see Supplementary Tables 1 and 2). We here select FA images from diffusion MR scans, rather than structural MR scans, as we had done when comparing the corpus callosum of adult primates with those of rodents. We select FA images to track the development of the corpus callosum because the corpus callosum is easily identifiable from its bright white appearance from midsagittal sections at successive stages of development and across species (Fig. 3). The FA images range from E 17 to postnatal day 60 in mice (n = 42), from GW 24 to 18 years of age in humans (n = 37), and from two PW to 36 months after birth in macaques (n = 102).

Multiple datasets were used to capture corpus callosum areas in humans (see Supplementary Table 1; Shi et al. 2011; Cohen et al. 2016; Khan et al. 2018a, 2018b; Fig. 4). To corroborate our findings of corpus callosum growth trajectories in humans, we also consider corpus callosum growth trajectories from a previously published dataset that relied on structural MRI scans to capture the growth of the corpus callosum in humans (n = 80). Ages range from shortly after birth to 25 years of age (Sakai et al. 2017). The FA images of mice are made available by John Hopkins University and the human brain scans were previously published (Chuang et al. 2011). The FA images of macaques are from the UNC-Wisconsin Neurodevelopment Rhesus Database (Young et al. 2017). We manually measured the area of the corpus callosum in mice, rhesus macaques, and humans from midsagittal sections at different ages (Fig. 5A). We excluded one outlier for macaques because the corpus callosum measurement from FA images was inconsistent with that obtained from the T1 structural MR.

Statistical analyses used to compare the growth of the corpus callosum between species were performed with the software programming language R. We tested whether a linear plateau would fit the data in each species with the library package easynls (model = 3; linear plateau). We subsampled the number of individuals from which the corpus callosum was measured and repeated the same analyses to ensure that the identified ages in which the corpus callosum ceases to grow in both species are not driven by outliers and that the sample size is sufficient to detect when the corpus callosum ceases to grow. We tested for a nonlinear regression (model 3) with the log-10-transformed values for corpus callosum area and the log-based 10 values for age expressed in days after conception (Fig. 5B–E). Tested samples range from 3 to total sample size obtained for each species. For each sample size, we tested for nonlinear regressions 10 times and we only consider ages in which the corpus callosum ceases to grow from models that detect a significant cessation of corpus callosum growth (P < 0.05). We extract the 95% confidence interval (CI) from these data to identify the range of possible ages as well as the percentage of variance explained by the models.

We assess whether the corpus callosum growth is extended in primates compared with rodents after controlling for variation in developmental schedules. The timing of developmental transformations (i.e., events) can be used to identify corresponding ages across species (Clancy et al. 2001). Controlling for variation in developmental schedules permits identifying whether certain development processes occur for an unusually long or short period of time in humans (Finlay and Darlington 1995; Clancy et al. 2001, 2007; Nagarajan and Clancy 2008; Workman et al. 2013; Charvet et al. 2017a; Charvet and Finlay 2018). Because the translating time resource extends up to 2 years of age in humans and its equivalent in other species, we extrapolated ages at later time points as we had done previously (Charvet et al. 2019). After controlling for variation in developmental schedules between species, we assess whether the corpus callosum ceases to grow later than expected in humans and macaques relative to mice.

Results

We describe the overall organization of pathways in the forebrain of humans and mice during development with a particular focus on radial and emerging commissural fibers. We also observe that diffusion MR tractography identifies pioneer axons originating from the medial wall of the developing cerebral cortex in both humans and in mice.

Radial Glia Organization of the Cortex in Humans and Mice

We explore the development of fiber organization in developing mouse brains from E 10.5 to FW 4 (Fig. 1, see Supplementary Fig. 2) and in humans from GW 17 to 30 (Figs 2 and 3). In mice, many of the fibers are observed coursing across the medial to lateral direction in the forebrain throughout development (Figs 1 and 2). According to the translating time model, a mouse at E 16.5 is roughly equivalent to humans at GW 12–16 and a mouse at E 17.5 is equivalent to humans at GW 14–18 (Fig. 2). We therefore compare mice at E 16.5–17.5 with humans at GW 17–21. We use whole brain tractography and coronal slice filters meant to capture fibers coursing through coronal slices shown in Figure 2 to identify species differences in the overall pattern of fiber organization in the two species. At GW 17–21, fibers are principally oriented radially with fibers extending from the proliferative pool towards the CP (Fig. 2A–C). These fibers likely represent radial glial cells known to guide the migration of neurons from the proliferative pool towards the CP. At E 16.5 (Fig. 2D,E) as in E 17.5 (Fig. 2F), we observe fibers coursing principally across the medial to lateral direction through the cortex. Some of these fibers likely correspond to thalamo-cortical, cortico-thalamic, or callosally projecting pathways. Thus, diffusion MRI highlights that mice and humans matched for age differ in their cortical organization. Radially organized fibers are evident in the human cortex but this is not the case in mice, presumably because radial fibers dissipate earlier in mice than in humans. Indeed, cortical neurogenesis wanes around E 18 in mice (Caviness et al. 2003, 2009), but cortical neurogenesis is thought to persist beyond GW 20 in humans (Zecevic et al. 2005; Malik et al. 2013; Charvet et al. 2017a).

Commissural Development in Mice and Humans

In mice, at the earliest gestational age studied (E 10.5), pathways crossing the midline were observed in anterior and ventral regions of the brain (Fig. 1A–D). We did not observe callosal fibers at this age. At E 14.5 (Fig. 1E–H), we observed short pathways crossing the midline in the presumptive corpus callosum (Fig. 1G,H). These fibers emerge or terminate within the cingulate cortex, and project contralaterally through the dorsal midline of the telencephalon (i.e., the developing corpus callosum). We consider these fibers emerging or terminating within the cingulate cortex to represent pioneer axons coursing through the corpus callosum because they are reminiscent of previously described pioneer axons that act as scaffolds to guide development of other axons crossing the dorsal midline (Koester and O’Leary 1994; Rash and Richards 2001). At E 16.5,
Figure 1. Maturation of commissural fibers as assessed from diffusion MR tractography in mice at various ages of development (E 10.5, E 14.5, E 16.5, E 17.5, and postnatal week 4: PW 4). (A) At E 10.5 (A–D), the dorsal view of brain fibers from whole brain tractography (A) and horizontal slice filters (B) shows that many of the pathways course across the anterior to posterior direction in the dorsal telencephalon as well as in the diencephalon. We identify pathways coursing through the midline in the ventral telencephalon by placing ROIs to identify commissural fibers (C, D). (B) At E 14.5 (E–H), we observe callosal pathways as well as putative pioneer axons. (C) At E 16.5 (I, L), we observe pathways coursing across the medial to lateral axes through the developing cortex. (I) Whole brain tractography through the mouse brain shows that the telencephalon primarily contains fibers coursing medial to lateral. The dorsal view of a horizontal slice through the telencephalon also highlights fibers coursing through the dorsal to ventral and medial to lateral direction within the developing cortex. (K, L) Placing ROIs to identify commissural fibers identifies callosal and putative pioneer pathways. Yellow arrows in 16.5E denote putative pioneer axons, while blue arrows denote callosal pathways. At E 17.5 (M–P), horizontal (M, O) and sagittal views (N, P) highlight callosal and putative pioneer pathways coursing through the dorsal midline. ROIs were used to identify callosal and pioneer pathways. At postnatal week 4 (Q–T), ROIs highlighting commissural fibers show callosal fibers as well as the hippocampal commissure fibers coursing across
fibers course across the medial to lateral direction throughout the developing cortex (Fig. 1I). We also observe a population of anteriorly located callosal pathways extending across the midline (Fig. 1K,L, yellow arrows). Some of these fibers originate or terminate within the cingulate cortex (Fig. 1K,L, blue arrows), whereas other fibers originate or terminate from more lateral regions of the developing cerebral cortex. Thus, fibers reminiscent of pioneer axons are first observed emerging or terminating from the cingulate cortex, followed by fibers spanning more lateral regions of the cortex. Given this developmental sequence, neurons emerging from the cingulate cortex may act as a scaffold for later-developing neurons that originate from the lateral regions of the cortex. As development progresses, the corpus callosum becomes increasingly evident at E 17.5 (Fig. 1M–P) and PW 4 (Fig. 1Q–T).

We next compare corpus callosum development in mice (Fig. 3A–C) and humans (Fig. 3E–J) matched for age. In mice, the corpus callosum is evident at E 16.5 and E 17.5 with pathways extending across the medial to lateral axis (Fig. 3A–D). At E 17.5, a larger callosal structure is evident in the rostral cortex compared with the caudal cortex (Fig. 3C,D). In humans, at GW 17 (Fig. 3E, pink arrow), at GW 20 (Fig. 3G), and through GW 22 (Fig. 3H) and GW 30 (Fig. 3I), we observe some pathways extending from the superior medial wall of the developing cortex projecting towards the contralateral hemisphere through the corpus callosum. Interestingly, other fibers arising from more lateral regions of the cortex also project towards the corpus callosum at GW 22 (Fig. 3H, pink arrow), at GW 21 (Fig. 3F, green arrow), and at GW 30 (Fig. 3I, white arrow). Some of these fibers course within the subventricular and ventricular zones (Fig. 3E, blue arrow). A few fibers extend into the gray matter of the medial wall of the developing cortex at GW 17 (Fig. 3E, red arrows), GW 21, and GW 22 (Fig. 3H, orange arrows). We also observe variation in the corpus callosal structure across the anterior to posterior axis with seemingly larger callosal structures located in the frontal cortex than in more posterior regions of the cortex at GW 17 (Fig. 3E). This variation in the corpus callosal structure across the anterior to posterior axis of the cortex observed in the GW 17 human is reminiscent of those observed in mice at E 17.5. Thus, in humans as in mice, there is a tendency for more anterior callosal pathways to be expanded than in posterior regions. As development progresses, the corpus callosum increasingly extends across the medial to lateral axis in both species.

Corpus Callosal Growth Trajectories Across Species

To test whether there are deviations in the timing of corpus callosum development between humans and mice, we measured the area of the corpus callosum at successive ages across primates (i.e., humans and macaques) and mice (Figs 4 and 5). Ages in humans and macaques are mapped onto mouse age to control for variation in developmental schedules across species. We use the translating time model to find corresponding ages, which relies on the timing of developmental transformations to find corresponding ages across species (Clancy et al. 2001; Workman et al. 2013; Charvet et al. 2017a; Charvet and Finlay 2018).

In both humans and in mice, the corpus callosum grows and ceases to grow (i.e., plateaus) after birth (Fig. 4). In humans, the corpus callosum grows and reaches a plateau in growth at about 1.75 years of age in humans, which is equivalent to 35 days after conception in mice (easynls fit model = 3; adj \( R^2 = 0.74; P < 0.05; n = 37\); Fig. 4A). In macaques, the corpus callosum continues to grow past the predicted ages for those of mice (postnatal day 10) but our nonlinear model does not capture a linear plateau in growth with these data (easynls fit model = 3; n = 102; Fig. 4B). In mice, the corpus callosum reaches a plateau in growth at around 28.5 days after conception, which is 10 days after birth (easynls fit model = 3 adj \( R^2 = 0.89; P < 0.05; n = 42\); Fig. 4C).

To ensure that the identified ages in which the corpus callosum ceases are robust to variation in sample size, we subsampled the number of individuals from which the corpus callosum area was measured in humans and mice. We then extracted identified ages in which the corpus callosum significantly ceases to grow with the easynls model 3 package (Fig. 5). We only extracted results from the model if the model accounts for a significant percentage of the variance (\( P < 0.05\)), and we identify ages in which the corpus callosum ceases to grow in both humans and mice (Fig. 5B,C). The corpus callosum ceases to grow between 1.66 years of age (i.e., 878 days after conception; 5% CI) to 1.85 years of age (i.e., 948 days after conception, 95% CI; \( n = 16–37\); Fig. 5B). In mice, the sample size in which the models account for a significant percentage of the variance ranged between 17 and 42, and the corpus callosum ceases to grow between PD 9.3 (i.e., 27.8 days after conception; 5% CI) and PD 10.2 (i.e., 28.7, 95% CI; Fig. 5C). Regardless of sample size, the nonlinear regression accounts for more than 80% of the variance in humans (Fig. 5D) and in mice (Fig. 5E). Taken together, these observations demonstrate that the identified age in which the corpus callosum ceases to grow in each species is robust to variation in sample size and is not driven by outliers.

To further ensure that our results align with those of others and that the corpus callosum ceases to grow at around 1–2 years of age in humans, we supplement our analysis with a previously published dataset that captures the growth of the corpus callosum in humans (Sakai et al. 2017; see Supplementary Fig. 3). Such an analysis shows that the corpus callosum growth plateaus at 1.87 years of age (easynls fit model = 3; adj \( R^2 = 0.78; P < 0.05; n = 86\); see Supplementary Fig. 3). Although it is possible that there is small amount of corpus callosum growth past those ages, these two different datasets both demonstrate that most of the corpus callosum growth ceases around 1–2 years after birth in humans such that the corpus callosum continues to grow for an extended period of time in humans compared with mice.

We next confirm the protracted growth of the corpus callosum statistically, and we select the most recent and largest dataset capturing developmental transformations across rodents and primates, including humans (Charvet et al. 2017b). Neural transformations that occur on 27.5–28.5 days after conception in mice (\( n = 4\)), which is when the corpus callosum ceases to grow in mice, occur between GW 27 and about 2.6 months after birth in humans (lower 95% CI: GW 26.5; upper
Figure 2. Diffusion MR tractography of cortices at GW 17–21 (A–C) and mice at E 16.5–E 17.6 (D, F) shows major differences in the organization of fibers spanning the cortex. In humans, many fibers course radially within the developing cortex. At GW 17 whole brain tractography (A) and 1 mm thick coronal slices through the anterior and posterior regions of the cortex (B) show fibers coursing from the proliferative zones to the CPs (white arrowheads). A similar situation is observed at GW 21 (C) where coronal slices set to capture fibers coursing through these slices identify radial fibers spanning the proliferative pool and the CP in the anterior and posterior cortex (white arrowheads). In contrast, diffusion MR scans of mice at E 16.5 (D, E) and at E 17.5 (F) show fibers principally organized across the medial to lateral axis (white arrowheads). (D) Whole brain tractography of mice show fibers coursing across the medial to lateral and dorsal to ventral direction through the developing cortex. (E) Coronal slices through the rostral and caudal cortex at E 16.5 capture fibers coursing tangential to the cortical surface rather than radially (white arrowheads). (F) At E 17.5, a similar situation is observed in the mouse where fibers are observed coursing across the medial to lateral direction (white arrowheads). Humans at GW 17–21 and mice at E 16.5–17.5 were selected for comparison because they are at roughly equivalent ages of development (Workman et al. 2013). The following abbreviations are used: A: anterior; P: posterior; R: rostral; C: caudal; M: medial, L: lateral; D: dorsal; and V: ventral.
Figure 3. Diffusion MR tractography captures the maturation of the corpus callosum in mice (E 16.5: A, B; E 17.5: C, D) as well as human brain from GW 17 to 30 (E–J). (A) ROIs were used to highlight brain commissural and pioneer fibers in humans and in mice. Sagittal slices set through the E 16.5 mouse brain show that fibers are principally organized across the medial to lateral axis. (B) ROIs set through the dorsal midline highlight callosal fibers at this age at E 16.5 (B) and at E 17.5 (C). The corpus callosum is more prominent at E 17.5 than at E 16.5. At E 17.5, more fibers are evident in the rostral cortex (C) than in more caudal regions (D). At GW 17 in humans (E), putative pioneer and callosal pathways are also observed across the anterior to posterior axis with more prominent callosal fibers evident in anterior regions than in posterior regions. (E) The yellow arrow highlights callosal pathways from the lateral brain regions, and the pink arrow shows callosal pathways from the upper medial wall. (F) The blue arrow shows callosal pathways from the ventricular region, and red arrows show possible residual pioneer axons at the cingulate cortex. The corpus callosum gains increasing prominence with age (F–J). (H) At GW 21, callosal pathways are evident and from the upper medial wall and lateral brain regions (green arrow). (I) Callosal pathways from the upper medial wall, cingulate cortex, and lateral brain regions (green arrow) at GW 21 and GW 22. (F) At GW 30, we observe callosal pathways from the cingulate cortex, upper medial walls, and lateral walls of the brain (white arrows). (J) Dorsal views of horizontal slices through the corpus callosum from GW 17 to GW 30 show that the corpus callosum becomes increasingly prominent with age.
Figure 4. The growth of the corpus callosum is protracted in humans (A) and macaques (B) compared with mice (C). In order to control for variation in the length of developmental schedules, age in humans and macaques is mapped onto mouse age in days after conception according to the translating time website (Clancy et al. 2001; Workman et al. 2013). (A) The area of the corpus callosum reaches a plateau in growth between 1 and 2 years of age in humans (i.e., 1.75 years after birth). (B) The corpus callosum shows sustained growth in macaques across the examined ages and does not plateau. (C) The corpus callosum reaches a plateau in growth at 28.5 days after conception in mice, which is 10 days after birth in mice. The corpus callosum grows for much longer than expected in humans and macaques when human and macaque age is mapped onto that of mice. Vertical bars identify when the corpus callosum ceases to grow, as well as birth.

95% CI: 2.03 months after birth). Yet, in humans, the corpus callosum growth ends at 1.75 years of age, which is past the 95% CI generated from these data. Taken together, these findings demonstrate that the corpus callosum continues to grow for significantly longer than expected in humans compared with mice.

Discussion

In this study, we tracked the development of fibers in the human and mouse cortex. Our findings demonstrate the ability of HARDI to track fiber development in different species. During cortical neurogenesis, fibers of the developing human cortex are primarily radially aligned, but this is not the case in age-matched mice. We focus on the developmental timeline of corpus callosum development, and we find that the corpus callosum grows for an extended period of time in humans compared with mice. We discuss the strengths and limitations of diffusion MR tractography, as well as the potential implications for the observed conserved and deviant developmental programs giving rise to variation in the cortical structure and connectivity patterns in the human lineage.

Diffusion MR Scanning for Basic Development and Health

Although ultrasound remains the predominant modality for evaluating disorders related to pregnancy, fetal MRI has been increasingly used to track normal and abnormal human development (Mitter et al. 2015; Jakab et al. 2017). Diffusion MRI is still in its infancy, and more work is needed to identify what developmental processes can be observed from diffusion MR tractography. HARDI is a new innovative approach to identify developmental programs in the human fetal brain, which can also be used to identify conserved and deviant developmental programs between humans and model organisms such as mice. It is important, however, to acknowledge the reliability of HARDI tractography when reporting diffusion imaging studies. As this method is in its infancy, HARDI does suffer from certain pitfalls (Maier-Hein et al. 2017). Additionally, the low number of specimens available for study and the differences in imaging procedures used to capture tractographies across development and between species make it challenging to draw quantifiable conclusions from HARDI tractography. It is currently difficult to obtain sufficient postmortem fetal brains to resolve this problem. Due to these limitations, we integrate our data with comparative analyses of callosal growth trajectories. However, studies such as these are essential as they identify key developmental processes (e.g., pioneer axons) in the human fetal brain, which should increase the utility of this work for disease detection and prevention.

Development of Radial Organization in Humans and Mice

Radial glia have been identified as an important player in developmental mechanisms accounting for the expanded isocortex of primates in adulthood (Rakic 1978, 1995), and we observe major differences in radial organization across the isocortex of humans and mice at corresponding ages. Fibers coursing across the mouse cortex from E 16.5 to 17.5 are principally organized across the medial to lateral axis. These fibers, which extend across the medial to lateral axes of the developing cortex, likely represent cortico-thalamic or thalamo-cortical pathways. This is in contrast with humans at corresponding ages where fibers in humans (GW 17–20) are primarily organized across the medial to lateral axis. These fibers, which extend across the medial to lateral axes of the developing cortex, likely represent cortico-thalamic or thalamo-cortical pathways. This is in contrast with humans at corresponding ages where fibers in humans (GW 17–20) are primarily organized radially across the isocortex. The radial organization of the developing cortex has previously been characterized with diffusion MR tractography in humans. Radially aligned fibers are evident at GW 17 and begin to diminish around GW 21 with residual fiber pathways still observed at GW 40 (Miyazaki et al. 2016). As development
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Figure 5. (A) Midsagittal planes through FA images of human brains at GW 40, 20 months, and 9 years of age show that the corpus callosum has a bright white appearance in contrast to surrounding brain regions. We have contoured the area of the corpus callosum in white as examples of corpus callosum measurements. To assess whether corpus callosum measurements are robust to variation in sample size, we subsampled the number of individuals from which the corpus callosum area was measured in humans and in mice. We select nonlinear regression models that account for a significant percentage of the variance. Identified age in which the corpus callosum ceases to grow ranges between 1.6 and about 1.8 years of age in humans (B) and between postconception day 27 and 29 in mice (C). Regardless of sample size, our model accounts for more than 80% of the variance in humans (D) and in mice (E). Horizontal dashed lines represent 95% CI from the age in which the corpus callosum ceases to grow in humans (B) and in mice (C), and the percentage of variance explained in humans (D) and in mice (E).

The finding that the human fetal cortex shows a predominance of fibers oriented radially compared with that observed in mice can be integrated with known deviations in developmental programs between primates and rodents. Previous work use thymidine-based methods to identify when cortical neurons progress, radial glia as well as other proliferative cells wane as cells exit the proliferative zone, and newly born neurons assume their final position within the cortex (Rakic 1974; Miyata et al. 2001; Zecevic et al. 2005; Takahashi et al. 2012; Malik et al. 2013; Arshad et al. 2015; Charvet et al. 2017a).
are generated (i.e., born) in macaques and rodents (Rakic 1974; Bayer and Altman 1990). A comparative analysis of thymidine-based studies showed that cortical neurogenesis is extended in macaques compared with examined rodents (Clancy et al. 2001). It is not the onset of cortical neurogenesis, but rather, it is the end of cortical neurogenesis that is protracted in primates relative to rodents (Clancy et al. 2001; Cahalane et al. 2014). That is, superficially located cortical neurons (i.e., layers II–IV neurons) undergo neurogenesis later in macaques considering the timing of other developmental transformations (Clancy et al. 2001; Workman et al. 2013). The extended duration of cortical neurogenesis is concomitant with an expansion in proliferative cells during cortical neurogenesis (Smart et al. 2002; Fish et al. 2008; Martinez-Cerdeño et al. 2012; Romero and Borell 2015). The extension in the duration of neurogenesis is concomitant with an expansion of upper layer neuron numbers in the primate cortex relative to those of rodents in adulthood (Charvet et al. 2017a). The notion that primates extend the duration of cortical neurogenesis for an unusually long time is further supported by a comparative analysis of cortical proliferative pool trajectories across humans, macaques, and mice (Charvet et al. 2017a). These data show that the growth of the cortical proliferative pool is protracted in humans and macaques compared with mice after controlling for variation in developmental schedules. Therefore, the duration of cortical neurogenesis extends for longer than expected relative to rodents (Charvet et al. 2011).

Extending the duration of neuron production should lead to an extension in the duration in which neurons are produced and migrate to the CP. Accordingly, mice at E 16.5 and E 17.5 should not possess many radial aligned fibers because neurogenesis is largely complete at these ages (Caviness et al. 2003, 2009; Yuzwa et al. 2017; Preissl et al. 2018). Indeed, single-cell-based sequencing methods have shown that the relative number of radial glia drops from approximately 14–0% between E 14 and E 18.5 (Yuzwa et al. 2017; Preissl et al. 2018; Loo et al. 2019). This is in contrast with humans at GW 17–21, which still possess a large proliferative pool and self-renewing radial glia, as well neurons exiting the proliferative pool to migrate towards the CP (Rakic 1974, 2003b; Zecевич et al. 2005; Malik et al. 2013; Arshad et al. 2015; Charvet et al. 2017b). We here assume that these radial fibers observed in humans are radial glia because these tractographies have been confirmed immunohistochemically in humans (Xu et al. 2014). However, the observed species differences in radial organization between humans and mice reported in the present manuscript have not been confirmed immunohistochemically, and several possibilities exist as to why a radial organization predominates the human fetal cortex but not mice at equivalent ages. For instance, the human fetal cortex may be composed of more or larger radial glia compared with age-matched mice. More work is needed to identify the developmental time course of radial glia development in humans and in mice. Nevertheless, the observation that putative radial glia are evident in humans but not in age-matched mice aligns with known deviations in the duration of cortical neuron production between primates and rodents.

Development of Commissural Fibers in Mice and Humans

We track the developmental sequence of corpus callosum development in mice. We observed fibers reminiscent of pioneer neurons emerging or terminating from the cingulate cortex as early as E 14.5. Similar fibers were also observed at E 16.5 (McConnell et al. 1989; O’Leary 1994; Ren et al. 2006). At GW 17, short fibers originating from the upper medial wall of the developing cortex and, more sparsely, from the cingulate cortex course towards the contralateral hemisphere in humans (Charvet et al. 2017a). The sparse fibers from the cingulate cortex likely correspond to residual pioneer pathways, which are known to originate in the cingulate cortex and cross the midline (Rakic and Yakovlev 1968; Koester and O’Leary 1994; Ren et al. 2006). As development progresses, the corpus callosum becomes increasingly evident in both humans and mice.

Conservation and Variation in the Timing of Corpus Callosum Development

We compare developmental timing of the corpus callosum in mice, macaques, and humans (Schwartz and Goldman-Rakic 1991; Clancy et al. 2007; Workman et al. 2013). According to the translating time model, early callosal developmental milestones such as the emergence of the corpus callosum and myelination onset are conserved in their timing across humans and mice (Workman et al. 2013). In the present study, we do not observe major differences in callosal development between the two species at relatively early stages of its development, although these findings do not necessarily entail that there are no early differences across species. We do, however, find that there are clear differences in the growth trajectories of the corpus callosum between primates and mice such that deviations in the duration of callosal growth account for the expansion of the corpus callosum in primates.

After controlling for variation in overall developmental schedules across species, the corpus callosum continues to grow for an extended period of time in humans. According to our nonlinear regressions, we find that the corpus callosum ceases to grow between 1 and 2 years of age, which is consistent with the work of others who have shown that the corpus callosum ceases to grow several months to a few years after birth in humans (Clarke et al. 1989; Giedd et al. 1999; Tanaka-Arakawa et al. 2015; Sakai et al. 2017; Vannucci et al. 2017). In mice, the corpus callosum area only grows for a few days, which
is also consistent with previous work (Wahlsten 1984; but see Chuang et al. 2011). Although these nonlinear regressions capture when most or all of the corpus callosum ceases to grow, it is possible that there is still some corpus callosum growth passed 1–2 years of age in humans and PD 10 in mice. We found that the macaque corpus callosum grows throughout the examined postnatal ages, and these findings align with previous work tracking the growth of the corpus callosum in macaques (Scott et al. 2016). We have not here identified the underlying developmental processes responsible for variation in callosal growth trajectories between species, and several possibilities exist. The human corpus callosum might prolong the duration with which corpus callosum axons increase in number or axons increase in thickness. Alternatively or in addition, the duration with which axons myelinate may be protracted in humans relative to mice. Although we do not resolve the underlying cellular process responsible for variation in corpus callosal growth trajectories across species, our findings demonstrate that the protracted development of cortico-cortical pathways is linked to their expansion in the primate lineage (Charvet et al. 2017b, 2019).

The corpus callosum is responsible for a variety of complex behaviors, and protracted growth of long-range cortico-cortical pathways such as the corpus callosum may be responsible for protracted behavioral development observed in humans relative to mice. Some behaviors such as weaning are deviant in their timing in humans relative to other species (Hawkes and Finlay 2018). However, very few studies have compared the timing of behavioral development between humans and other species. Integrating data on the timing of long-range cortico-cortical pathways with behavioral milestones may serve to identify the behavioral implications of extending the maturation of long-range cortico-cortical pathways in the primate lineage.

Conclusion
The timing of developmental transformations may be a powerful mechanism through which variation in cortical association pathways emerge. Several hypotheses have focused on the timing of axonal development as a source of evolutionary changes in connectivity patterns (Innocenti 1995; Striedter 2005). It has been proposed that neurons that reach their targets early can outcompete other targets, thereby expanding their representation within the brain (Catania 2001). A previous study showed that the star-nose mole’s fovea matures earlier than adjacent appendages. Thus, early axon extension might favor competing for targets, and their overrepresentation in the mature isocortex (Catania 2001). On the other hand, prolonging the duration of axon extension and refinement might afford more time to outcompete targets, which would accordingly increase the territory of select pathways in the adult brain (Deacon 1990; Striedter 2005). These two seemingly contradictory hypotheses emphasize the importance of timing in generating variation in connections between species. Our study supports the view that an extended duration with which corpus callosum pathways grow accounts for evolutionary changes in their structure in the adult brain. Thus, evolutionary changes in a cortical association pathway emerge by modifying the time with which they develop.

Supplementary Material
Supplementary Material is available at Cerebral Cortex online.

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