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Structural variability across the primate brain: A cross-species comparison

Running title: Structural variability across the primate brain

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ABSTRACT

A large amount of variability exists across human brains; revealed initially on a small scale by post mortem studies and, more recently, on a larger scale with the advent of neuroimaging. Here we compared structural variability between human and macaque monkey brains using grey and white matter magnetic resonance imaging measures. The monkey brain was overall structurally as variable as the human brain, but variability had a distinct distribution pattern, with some key areas showing high variability. We also report the first evidence of a relationship between anatomical variability and evolutionary expansion in the primate brain. This suggests a relationship between variability and stability, where areas of low variability may have evolved less recently and have more stability, while areas of high variability may have evolved more recently and be less similar across individuals. We showed specific differences between the species in key areas, including the amount of hemispheric asymmetry in variability, which was leftlateralised in the human brain across several phylogenetically recent regions. suggests that cerebral variability may be another useful measure for comparison between species and may add another dimension to our understanding of evolutionary mechanisms.

KEYWORDS

asymmetry, cross-species comparison, magnetic resonance imaging, grey matter, white matter, variability, evolution

INTRODUCTION

Comparative anatomy has revealed that the human brain is greatly enlarged as compared to other primates. This difference in size has been interpreted as an evolutionary expansion (Preuss 2011; Sherwood et al. 2012). However, the critical feature for the evolutionary changes underlying higher cognitive abilities may not be the size of the brain *per se*, but rather the relative expansion of some brain regions relative to others (Passingham and Wise 2012; Kaas and Stepniewska 2015). For example, some specific regions of prefrontal cortex and parietal cortex have shown relative expansion or specialisation along the evolutionary tree (Semendeferi et al. 2002; Schoenemann et al. 2005; Neubert et al. 2015), combined with a greater degree of lateralisation in the human brain (Nathan et al. 1990; Catani et al. 2007; Catani et al. 2010; Thiebaut de Schotten et al. 2011; Ocklenburg and Gunturkun 2012; Budisavljevic et al. 2015). Despite the importance of individual differences in evolution (Darwin 1859), no studies to date, to our knowledge, have compared inter-individual anatomical variability across species.

Traditionally, comparative anatomy studies have been based on *post mortem* measures: histological staining, blunt dissections and axonal tracing (Brodmann 1909; Vogt and Vogt 1919; Van Essen et al. 1986; Barbas and Pandya 1987). In recent years, *in vivo* magnetic resonance imaging (MRI) has become a useful tool not only for the investigation of human brain, but also for the direct comparison of human and non-human primate anatomy, both functionally and structurally (Croxson et al. 2005; Vincent et al. 2007; Mars et al. 2011; Mantini et al. 2012; Thiebaut de Schotten et al. 2012). Recent work demonstrates that anatomical inter-individual variability exists within humans with regard to surface anatomy (Whitaker and Selnes 1976; Toga and Thompson 2003; Uylings et al. 2005; Marie et al. 2015), cyto-architectural boundaries (Amunts et al. 1999), connectional anatomy (Catani *et al.* 2007; Thiebaut de Schotten *et al.* 2011), and vascular anatomy (van der Zwan et al. 1992). Similar results have been reported for the non-human primate brain (Van Essen and Dierker 2007; Zhang et al. 2013).

Here, we analysed the amount of structural variability across the human and macaque monkey brain. Our primary hypothesis was that the human brain would be more variable than the macaque monkey brain, particularly in regions that are relatively specialised in humans. Additionally, we hypothesised that a relationship exists between the anatomical variability and the evolutionary expansion in the human brain. These measures were acquired, where possible, with comparable equipment, parameters and resolution between species, in equally sized samples of humans and monkeys, and then analysed with identical methods.

MATERIALS AND METHODS

Subjects & MRI acquisition

Main analysis of T1-weighted images and diffusion-weighted images were obtained from two sets of 10 adult healthy controls (5 males and 5 females per set; age range 26 - 35 years at the time of data acquisition) from the Human Connectome Project (HCP) (http://www.humanconnectome.org, Q4 Release).

Two additional sets of 16 healthy controls (8 males and 8 females per set; age range 26 – 35 years at the time of data acquisition) from the Human Connectome Project (HCP) (http://www.humanconnectome.org, Q4 Release) were selected to determine the optimal number of participants to build a template.

T1-weighted images and diffusion-weighted images were obtained *in vivo* from 10 healthy rhesus monkeys ($Macaca\ mulatta$; 6 males and 4 females; age range 3.75-5.82 years at the time of data acquisition; mean 4.5 years). Studies of brain development would place these animals in the category of young adults ($Malkova\ et\ al.\ 2006$).

Human Connectome Project dataset

Data from the Human Connectome Project (HCP) dataset were acquired on a Siemens Skyra 3T scanner at Washington University in St. Louis. The scanner was equipped with a customized body transmitter coil with 56 cm bore size (Van Essen et al. 2013)

T1-weighted imaging

An axial three-dimensional (3D) T1-weighted imaging dataset covering the whole head was acquired for each participant (260 slices, voxel resolution = $0.7 \times 0.7 \times 0.7$ mm, TE = 2.14 ms, TR = 2400 ms, flip angle = 8°).

Diffusion-weighted imaging

A total of 111 near-axial slices were acquired with a multiband factor = 3 (Moeller et al. 2010; Ugurbil et al. 2013), isotropic $(1.25 \times 1.25 \times 1.25 \text{ mm})$ resolution and coverage of the whole head (TE = 89.5 ms, TR = 5520 ms). At each slice location, 18 images were acquired with no diffusion gradient applied. Additionally, 90 diffusion-weighted images were acquired, in which customized SC72 gradients were uniformly distributed in multiple Q-space shells (Caruyer et al. 2013). The acquisition of the diffusion weighting images was repeated three times with a b-value of 1000, 2000 and 3000 s/mm², respectively. Data were pre-processed using the default HCP pipeline (V.2), which includes correction for susceptibility, motion and eddy current distortions (Andersson et al. 2012; Sotiropoulos et al. 2013). Pairs of diffusion-weighted volumes were acquired with reversed right-to-left and left-to-right phase-encoding directions. This generates a pair of images where the applied diffusion gradient counterbalances distortions in opposite directions. From these pairs the susceptibility-induced off-resonance field was estimated using a method similar to that described in (Andersson et al. 2003) and corrected on the whole diffusion weighted dataset using the tool TOPUP as implemented in FSL (Smith et al. 2004). At each slice, diffusion-weighted data were simultaneously registered and corrected for subject motion and geometrical distortion using the tool EDDY as implemented in FSL. Finally FA maps were computed using DTIFIT (Behrens et al. 2003).

Rhesus monkey dataset

Data were collected from the monkeys under anaesthesia (protocols described in Mars *et al.* 2011; Sallet et al. 2011; O'Reilly et al. 2013). Protocols for animal care, magnetic resonance imaging, and anaesthesia were performed in accordance with the United

Kingdom Animals (Scientific Procedures) Act (1986). Anaesthesia was induced using ketamine (10 mg/kg intramuscularly; i.m.), xylazine (0.125–0.25 mg/kg i.m.), and midazolam (0.1 mg/kg i.m.) and maintained with isoflurane at a low concentration (0.9–1.7% expired; mean, 1.38%). Anaesthesia was supplemented with atropine (0.05 mg/kg, i.m.), meloxicam (0.2 mg/kg, i.v.), ranitidine (0.05 mg/kg, i.v.), and local anaesthetic (5% lidocaine/prilocaine cream and 2.5% bupivacaine subcutaneously as necessary). Physiological parameters including capnography, inspired and expired isoflurane concentration, SPO₂, core temperature, heart rate and blood pressure were monitored and kept constant to maintain normal physiological function.

All monkey MRI data were acquired in a 3 T MRI scanner with a full-size horizontal bore and a custom-built 4-channel phased-array coil with a single loop local transmit coil (Windmiller-Kolster Scientific, Fresno, CA, USA). These data have been used in previous publications (Mars *et al.* 2011; Sallet *et al.* 2011; Mars et al. 2013; O'Reilly *et al.* 2013; Sallet et al. 2013; Neubert et al. 2014; Noonan et al. 2014; Neubert *et al.* 2015).

T1-weighted imaging

T1-weighted images were acquired with a T1-weighted magnetization-prepared rapid-acquisition gradient echo sequence (128 slices, voxel resolution = $0.5 \times 0.5 \times 0.5$ mm, TE = 4.01ms, TR = 2500 ms). Three images were acquired and subsequently averaged for high signal-to-noise ratio.

Diffusion-weighted imaging

Diffusion-weighted images were acquired using echo-planar imaging (56 slices, voxel resolution = $1.0 \times 1.0 \times 1.0 \text{ mm}$, TE = 102 ms, TR = 8300 ms, acceleration factor = 2). Diffusion weighting was isotropically distributed along 60 directions using a b-value of 1000 s/mm^2 , with six volumes without diffusion weighting per run. Six averages were acquired per subject; three with left-right phase-encode direction and three with right-left, to facilitate correction for distortions along the phase encoding direction. As for the human datasets, the susceptibility-induced off-resonance field was estimated (Andersson et al. 2003) and corrected on the whole diffusion weighted dataset using the tool TOPUP

as implemented in FSL followed by registration, motion correction and eddy current correction EDDY (Smith *et al.* 2004). Finally FA maps were computed using DTIFIT (Behrens *et al.* 2003).

Template reconstruction

In order to quantify the variability our approach was first to produce an optimal template without spatial priors for each species and modality. We produced these templates using Advanced Normalization Tools (ANTs, http://stnava.github.io/ANTs/) (Avants and Gee 2004; Avants et al. 2010), which build a template iteratively combining affine and diffeomorphic deformations (Avants et al. 2008; Klein et al. 2009). Participants were first aligned together using affine transformation. Then the templates were built from all the subjects iteratively (n=4) using diffeomorphic deformations. Note that to avoid cross modality differences related to the voxel size, T1-weighted images were registered beforehand to FA maps.

Diffeomorphic deformations were performed using the SyN tool in ANTs (1). The effective resolution of the warp field used was the same as that of the fixed image (1.0 mm³ in the monkeys and 1.25 mm³ in the humans based on the resolution of the FA maps) and the smoothing of the warp field and neighbourhood correlation radius were the default values (3 voxels and 9x9x9). This results in a comparable warp field resolution for each set of data acquisition parameters. Obviously relative to the human brain the macaque monkey brain is roughly 1/12 of the volume (average monkey brain volume 100 cm³, adult human brain volume 1200cm³) and therefore our relative voxel resolution is not comparable; however the monkey data are at the current cutting edge for data acquisition *in vivo*. Importantly, our measures of variability are within-species numerically, with only the patterns of variability being compared between species, and thus this limitation is only minor in our hands.

Optimal number of participants to build a template and reproducibility.

The optimal number of participants was calculated producing T1 and FA templates from separate paired populations of equal gender distribution. This approach was repeated for

groups consisting of four, six, eight, ten, twelve, fourteen and sixteen subjects. Squared spatial Pearson's correlations between each pair (i.e. square of fslcc from FSL) was employed to calculate the percentage of shared variance (i.e. the similarity). **Supplementary Figure 1** indicates a steep increase of shared variance between templates produced from T1 and FA maps from 4 to 10 participants followed by a plateau from 10 to 16 participants. This result indicates that, using iterative diffeomorphic deformations, 10 subjects are sufficient to produce a good enough brain template to match the overall population.

Quantification of the variability

T1 and FA maps were registered to their corresponding templates and sets of 3 diffeomorphic deformation maps were extracted for each subject corresponding to the orthogonal projection of the deformations for each voxels (i.e. maps x, y, z). Variability was defined as the strength of the deformation (Euclidean distance) required to match each individual map to its corresponding template. The strength of the deformation was calculated for each voxel using the following formula (Gibbs 1881): $|(x, y, z)^T| = \sqrt{(dx^2 + dy^2 + dz^2)}$ where dx, dy and dz are the difference in each dimension.

Grey matter variability

FAST from FSL was employed to extract the grey matter ribbon from the T1 templates and consecutively employed as an inclusion mask to quantify the average grey matter deformation for each subject. Average grey matter deformation was projected on a 3D surface reconstruction using Anatomist (http://brainvisa.info). Surfaces are available on request from the authors.

White matter variability

We quantified for every subject the average white matter deformation in each voxel with an FA value > 0.2. Results were projected on the average tractography reconstruction of the human and the monkey dataset (Jones et al. 2002). Average tractography datasets are available on request from the authors.

Brain size effect

In order to control for the effect of brain size, we carried out the same estimation using the Jacobian determinant in order to report the proportional rather than the absolute value of the total deformation required to match the template (Ashburner 2007).

Cortical folding effect

In order to control from the brain curvature, areal expansion and morphological variability was estimated using Spearman rank-order correlation (Spearman 1904), and the stability of the results checked using robust regression.

Reproducibility

Reproducibility was assessed in humans using squared spatial Pearson's correlations to estimate the percentage of shared variance between the variability maps derived from the two sets of 10 adult healthy controls.

Comparison between variability in humans and areas of recent cortical expansion

To assess the phylogenetic signature of the morphological variability, we tested the association between the latter and an estimate of cortical expansion in humans compared to monkeys. We employed the macaque-to-human transformation map provided in Caret (http://brainvis.wustl.edu/) (Van Essen and Dierker 2007). This map (Fig. 6 top-left) was calculated by performing a surface-based registration between a macaque monkey and a human brain template. Crucially, the deformation was constrained to align 23 landmarks whose location can be accurately defined across species (e.g. the central sulcus or the location of the FEF). The geodesic distance between homologous regions within each species would differ not only according to the absolute size of the brain, but also according to the different cortical area separating the regions in either species. For instance, the distance between area MT and A1 is much higher in humans than in monkeys, while the reverse is true for the distance between FEF and the central sulcus (Van Essen 2004). These differences represent an estimate of the differential cortical expansion in humans and monkeys with respect to a common ancestor. For instance the

increased distance between MT and A1 reflects al relative expansion of high-order association cortex in the posterior temporal lobe and STS in humans. For simplicity, we refer here to this deformation map as a 'areal expansion map'. To assess the relationship between the areal expansion map and our map of anatomical variability in humans, we projected the former onto the each cortical voxel in the MNI152 template. We then correlated the values of either maps at each voxel on the cortex. This analysis was performed for each of the five lobes of the brain separately. The association between areal expansion and morphological variability was estimated using Spearman rank-order correlation (Spearman 1904), and the stability of the results checked using robust regression.

Asymmetries

Asymmetries in humans were quantified using regions of interest defined from the Harvard-Oxford cortical atlas (Zalla et al. 2004; Frazier et al. 2005; Makris et al. 2006; Goldstein et al. 2007). Asymmetries in monkeys were estimated employing regions of interest defined from the INIA19 Primate Brain Atlas (Rohlfing et al. 2012). Human and monkey T1 templates were registered to Harvard and INIA19 Brain atlases respectively and applied to each subject's variability map (i.e. Euclidian distance). An index of asymmetry was consecutively calculated for both species using the following formula: (Region in the right hemisphere – Region in the left hemisphere) / (Region in the right hemisphere).

Statistics

SPSS 22 software (SPSS, Inc., Chicago, IL, United States of America) was employed to carry on a repeated measure ANOVA with tissue studied (i.e. grey matter and white matter variability) as within-subjects and species (i.e. *Homo sapiens* or *Macaca mulatta*) as between-subjects factors. A one-sample t-test approach was applied to quantify the significance of the index of asymmetries for each region investigated. All statistics were Bonferroni corrected for multiple comparisons.

RESULTS

In both species, we analysed measures of grey (T1-weighted images) and white matter (fractional anisotropy; FA, as measured by diffusion-weighted imaging). By using non-linear deformation based morphometric analysis, we established the amount of intersubject variability for both measures, and then compared them between species.

Reproducibility of the findings. The amount of inter-subject variability for both measures was reproduced in two matched groups of ten human participants and squared spatial Pearson's correlations indicated a strong shared variance between the variability maps. T1-weighted variability maps shared 92.4% of variance and FA variability maps shared 91.2% of variance. This suggests that the variability maps we calculated with this method are highly reproducible across populations.

Monkey to human comparative variability in anatomy. The study had a 2 (tissue type: T1 vs. FA) x 2 (species: human vs. monkey) mixed factorial design, which was analysed with a repeated measure ANOVA revealing significant main effects for tissue type and species (**Figure 1**): Grey matter (T1) and white matter (FA) express different levels of variability with grey matter being much more inconstant than white matter in both species ($F_{(1,28)} = 22.272$; p < 0.001). The human brain was also statistically more variable than the monkey brain ($F_{(1,27)} = 385.494$; p < 0.001) with a significant interaction between tissue type and species ($F_{(1,27)} = 5.610$; p < 0.025).

When comparing the Jacobian determinant of the grey and white matter deformation, which represents proportional expansion and contraction independent of overall brain size, the human brain was not significantly more variable than the monkey brain ($F_{(1,27)} = 2.932$; p = 0.098) with no interaction between tissue and species ($F_{(1,20)} = 0.001$; p = 0.971) (**Figure 1**).

Grey matter variability. Both species showed distinct increased variability located in the upper visual field of primary visual area (V1), V4, lateral occipital sulcus (LOS), dorsal parieto-occipital sulcus (POS), middle and posterior parts of the superior temporal

sulcus (STS), posterior cingulate sulcus (CS), pars opercularis, frontal-eye-field (FEF) and the rostral prefrontal cortex (RPC) (**Figure 2**).

Both species showed lower variability in extrastriate visual areas (V2/V3) and ventral prefrontal cortex (VPC). However, while the temporo-parietal junction (TPJ) showed high variability in humans, the equivalent region in monkeys, the mid-STS, did not show similarly high variability. As compared to monkeys, humans showed specifically lower variability in the anterior portion of the lateral prefrontal cortex, while humans showed a "hot spot" of high variability in the dorsal medial frontal cortex (**Figure 2**).

White matter variability. Both species showed increased variability in the superficial white matter, which was particularly distinct for the white matter connecting the frontal eye field (FEF), primary motor areas (M1), primary sensory areas (S1) as well as extrastriate area 4 (V4) and striate area (V1) (Figure 3 and supplementary figure 2).

Both species showed lower variability in deep white matter structures including for instance, in the core of the corticospinal tract (CST), the cingulum (Cing), the fornix (FX) and the most medial portion of the inferior longitudinal fasciculus (ILF), and the optic radiations (**Figure 3 and supplementary figure 2**).

Grey variability in humans and cortical folding. Analysis of the correlation between grey matter variability findings in humans and surface curvature (Van Essen and Dierker 2007) (**Figure 4**) indicated a significant relationship with a small effect size between variability and cortical folding in the left (Spearman's rho = -0.135; p < 10^{-159}) the right hemispheres (rho = -0.065; p < 10^{-36}).

Grey variability in humans and areas of recent cortical expansion. Analysis of the correlation between grey matter variability findings in humans and macaque to human areal expansion (Van Essen and Dierker 2007) (**Figure 5**) revealed a significant positive correlation between variability and areal expansion in the temporal lobes and limbic

system bilaterally (left temporal lobe, Spearman's rho = 0.403; right temporal lobe, rho = 0.413; left limbic system, rho 0.320; right limbic system, rho = 0.517; p < 10^{-69} in all cases). There was a significant negative correlation in the occipital lobes bilaterally (left occipital lobe, rho = -0.208; right occipital lobe, rho = -0.366; both p < 10^{-55}). In the frontal lobes, there was a positive correlation between variability and areal expansion in the left hemisphere (rho = 0.137, p < 10^{-56}) but not in the right hemisphere (rho = 0.013, p = 1). In the parietal lobes, there was a positive correlation in the right hemisphere (rho = 0.060, p < 0.00001) but not in the left hemisphere (rho = 0.002, p = 0.629).

Because we found a significant relationship with a small effect size between variability and cortical folding, we repeated the correlation analysis between the grey matter variability findings and areal expansion, this time regressing out the mean cortical curvature measures. Our results remained highly significant with positive partial correlation between variability and areal expansion in the temporal lobes and limbic system bilaterally (left temporal lobe, Spearman's rho = 0.402; right temporal lobe, rho = 0.407; left limbic system, rho 0.311; right limbic system, rho = 0.517; p < 10^{-65} in all cases). There was a significant negative partial correlation in the occipital lobes bilaterally (left occipital lobe, rho = -0.237; right occipital lobe, rho = -0.417; both p < 10^{-55}). In the frontal lobes, there was a positive partial correlation between variability and areal expansion in the left hemisphere (rho = 0.130, p < 10^{-52}) but not in the right hemisphere (rho = 0.004, p = 1). In the parietal lobes, there was a positive partial correlation in the right hemisphere (rho = 0.071, p < 10^{-7}) but not in the left hemisphere (rho = 0.025, p = 0.369).

Cortical Asymmetries. Results are summarised in Figure 6 and supplementary tables 1 and 2. One sample t-tests revealed that, in humans, the Heschl's gyrus ($t_{(19)} = -3.879$; p = 0.045) as well as the superior (anterior portion $t_{(19)} = -5.190$; p = 0.002; posterior portion $t_{(19)} = -4.798$; p = 0.006) and middle (anterior portion $t_{(19)} = -4.978$; p = 0.004; middle portion $t_{(19)} = -5.650$; p = 0.001) temporal gyri were significantly more variable in the left hemisphere when compared to the right hemisphere. Conversely, the paracingulate ($t_{(19)} = 5.204$; p = 0.002) and lingual ($t_{(19)} = 4.861$; p = 0.005) gyri were significantly more variable in the right hemisphere.

In monkeys, solely the supramarginal ($t_{(9)} = 6.308$; p = 0.004) and angular gyrus ($t_{(9)} = 5.273$; p = 0.014) were significantly more variable in the right hemisphere.

DISCUSSION

In this study, we compared the amount of structural variability in the brain across two key species: humans and rhesus macaque monkeys. In addition, we explored the relationship between anatomical variability and areal expansion maps of the primate brain. Three main findings emerge from our work. Firstly, overall, the human brain is not proportionally more variable than that of the monkey, and both species have distinct similarities and differences in grey and white matter variability within specific regions. Secondly, there were some differences between species in specific regions that may underline functional differences between the two species. Thirdly, there was a significant correlation between our anatomical variability maps and the classical maps of evolutionary expansion for most of the lobes of the brain. Finally, the degree of grey matter variability shows significant differences between the left and the right hemispheres in the human brain, and very little difference between the hemispheres in monkeys. Based on these findings we concluded that variability is a useful measure for comparison between species, and cautiously suggest that this may reveal details about the relative evolutionary hierarchy of specific brain regions across species.

Our first finding revealed that the human brain only showed higher variability than the monkey brain when brain size was not controlled for (**Figure 1**). This is perhaps unsurprising given the strong role that brain size plays in primate evolution (Semendeferi *et al.* 2002; Sherwood et al. 2005) and its proposed relationship with higher-order cognitive functions (Navarrete et al. 2016). In light of the strong relationship between brain structure variability and performance in humans, we may expect brain variability to have a similar relationship with behaviour in non-human primates, particularly with regard to complex, recently evolved behaviour (Gilissen and Hopkins 2013; Hopkins and

Avants 2013; Phillips et al. 2013; Hopkins et al. 2014; Bianchi et al. 2016; Hecht et al. 2016).

Our second finding was that there was regionally specific localisation of variability in both species. This was particularly prominent in areas supporting higher-order cognitive function (Figure 2). For instance, the parieto-occipital sulcus is critically-placed to coordinate goal-directed actions in both humans and monkeys, and shows clear homologies between the two species. We also observed high variability in the pars opercularis in both species. Cortical stimulation of this area elicits orofacial movements in the primate brain (Huang et al. 1989) and phonemic paraphasia in the human brain (Duffau 2012). The posterior part of the STS in the monkey has been linked to the perception of species-specific vocalisations (Gil-da-Costa et al. 2004) and may be equivalent to Wernicke's area for speech perception and comprehension in humans (Wernicke and Eggert 1874). These areas were highly variable in monkeys and human respectively. The posterior cingulate sulcus, which also was highly variable, is a key hub of the default mode network and is therefore implicated in passive or internally-oriented tasks in both humans and monkeys (Mantini et al. 2012; Margulies et al. 2016). We also found high variability in the frontal eye fields (FEF) in both the human and the monkey brain. This complex region is believed to be highly expanded in the primate brain (Elston and Rosa 1998; Hill et al. 2010), notably contributing greatly to the voluntary orienting of attention, which is a shared function in both humans and monkeys (Buschman and Miller 2007; Mantini et al. 2012). In addition, a number of areas of the rostral prefrontal cortex that are implicated in both positive and negative outcome expectations, as well as outcome monitoring, have shown distinct functional similarities between humans and monkeys (Neubert et al. 2015). Correspondingly, in these areas also we found high variability in both species.

Areas of low variability within both species were in regions that were less recently evolved and exhibited preserved organisation, such as extrastriate visual areas (V2/V3), which are largely homologous between humans and monkeys (Van Essen et al. 2001; Orban et al. 2004).

It is important to mention that the pattern of variability follows gross anatomical landmarks to some extent, as demonstrated by the small effect size correlation we found between the grey matter variability and the surface curvature (Figure 4). This is interesting as a relationship between cortical folding patterns and histologically defined brain areas (Fischl et al. 2008) has been previously reported for some but not all areas in the brain. For instance primary visual areas are mostly localised in the calcarine sulcus but extend variably to neighbouring gyri in humans (Rademacher et al. 1993; Amunts et al. 2000) while there is considerable functional variation between primary visual areas in macaque monkeys (Van Essen et al. 1984). The consistency between cortical folding and cytoarchitectonic fields seems to decrease drastically with the brain hierarchy (Amunts et al. 2007); high order associative areas being less consistent with sulcal landmarks than primary areas (Fischl et al. 2008). Hence the variability of this effect across brain areas is consistent with the small effect size of the relationship between variability maps and the cortical folding. This may suggest a complex tripartite relationship between cytoarchitectonic fields, gross anatomical landmarks and interindividual anatomical variability.

We also identified very specific differences between species in particular grey matter regions (**Figure 2**). One difference between the species was a "hot spot" of high variability in dorsal medial frontal cortex, which was not evident in the monkey. Decision-making and abstract reasoning is arguably one of the higher functions of the human brain, allowing humans to imagine an outcome or create a representation of the unchosen option. For example, a region of human dorsal medial frontal cortex associated with the imagining of reward outcomes (Nicolle et al. 2012) did not correspond in a straightforward way to any region of the monkey cortex when comparisons between resting-state functional connectivity patterns were made across species (Neubert *et al.* 2015). While this region showed high variability in our data for the human brain, there was no clear equivalent in monkey, and is therefore consistent with its hypothesised uniqueness in humans. A second difference between the species was in the temporo-

parietal junction (TPJ), which showed high variability in humans, but the equivalent region in monkeys, the mid-STS (Perrett et al. 1992; Tsao and Livingstone 2008; Pinsk et al. 2009), did not show similarly high variability (**Figure 2**). The TPJ has thus shown a great deal of expansion from the monkey to the human brain (Van Essen and Dierker 2007). Despite its putative role in social cognition in monkeys (2011; 2014), the TPJ may process uniquely human attributes of social cognition, such as representing mental states (Saxe 2006; Besharati et al. 2016). The lack of high variability in monkeys may relate to the fact that in our study the monkeys came from a less varied social structure than the humans, but it may also reflect a possible link between the increased role of this region in social cognition and its inter-individual variability in the human brain.

Our third finding was a strong correlation between our grey matter variability maps with previously published evolutionary expansion maps (Van Essen and Dierker 2007) (**Figure 5**). This may indicate that the more variable the cortical areas are, the later they may have emerged on the evolutionary tree. Likewise, older, more ancient brain areas tend to be more invariant in their anatomy. The absence of white matter expansion maps in the literature has prevented us from verifying similar principles of white matter variability. Nonetheless, if we assume that there is a strong correlation between the variability and evolutionary expansion maps, our white matter variability maps may constitute an indirect estimate of white matter's phylogenetic dynamic (Figure 3). These results would suggest that white matter expanded from limbic specific and subcortical areas to later include more associative networks for higher cognitive functions. This interpretation corroborates previous phylogenetic models based on the measure of cortical cytoarchitecture (Yakovlev 1948; Maclean 1949, 1952; Mesulam 2000), neurosurgery (Duffau 2017) and functional connectivity (Margulies et al. 2016), which placed subcortical corticoid and allocortical areas (i.e. limbic zone) at the earliest level of the evolutionary tree followed by the mesocortex (i.e. paralimbic zone) and the isocortex (i.e. idiotopic and homotypical). This phylogenetic gradient is particularly evident on the medial surface of our variability maps (Figure 2, 3). Similarly, the distribution of anatomical variability on the lateral surface appears to follow evolutionary trends as depicted by the dual origin hypothesis (Dart 1934; Sanides 1964, 1970; Pandya et al. 2017). According to this hypothesis, brain evolution emerged from two systems, the hippocampocentric and olfactocentric divisions, that meet in the more recently developed regions located in the posterior part of the middle frontal gyrus (supporting, for example, working memory processes and decision making), posterior temporal lobe (i.e. language comprehension) and inferior parietal lobule (i.e. tool manipulation). We found these areas to be the most variable in terms of their cortical anatomy in the human brain as well as in their subjacent white matter anatomy, recapitulating the gradient suggested by the dual origin hypothesis. However, when interpreting our findings, we consider that we are comparing only two species, each one a product of millions of years of separation and separate evolutionary trajectories. Our findings bear a close resemblance to projected relative evolutionary expansion, and therefore may reflect a significant evolutionary finding, but additional species will need to be studied to make a strong conclusion about evolution.

Finally, we investigated the hemispheric differences in variability within the two species (**Figure 6**). Our findings are broadly consistent with the majority of studies comparing the degree of hemispheric asymmetry between the human and the monkey (Preuss 2011). We noted that the regions of greatest asymmetry were mainly those with the largest degree of hemispheric dominance in humans (Wang et al. 2014). For instance, language regions such as Heschl's gyrus and the superior and middle temporal gyri, were more variable in the left hemisphere. These results confirm, as previously demonstrated with fMRI (Tzourio-Mazoyer et al. 2015), that different brain phenotypes exist to support the phylogenetically recent hemispheric specialisation for language in humans. We also noted that the occipital lobe and the paracingulate gyrus were more variable in the right hemisphere than in the left. The paracingulate gyrus is highly variable in humans (Vogt et al. 1995) while, the variability in the occipital lobe might be related to inter-individual differences in the torque of the occipital lobe (LeMay 1976).

Several factors may have interacted with our findings on variability including genetic diversity, variety of environmental variety, variable personal experiences, and possible evolutionary processes. Differences in image resolution relative to brain size in humans

and monkeys, despite using cutting-edge methodology, may also have contributed to differences and may represent a limitation of our study. However, overwhelmingly, both species showed striking similarities in the pattern of grey and white matter variability, suggesting that the impact of these factors is minimal.

Three key anatomical features may also have confounded our results and represent possible limitations of our study. Firstly, although the size of cytoarchitectonic fields may have remained consistent, an anatomical shift of primary visual areas has caused them to move from the lateral surface to the medial wall along with evolution. This may have differed slightly across subjects and driven the negative correlation between recent macaque to human areal expansion and variability we found in the occipital lobes. Secondly, an obvious limitation for cortical variability is the degree of intersubject sulcal variability, for example in the paracingulate, central or precentral sulcus (Ono et al. 1990; Vogt et al. 1995; Van Essen 2004). However, we did not generally observe a higher variability in sulci, when compared to gyri or white matter. Thirdly, variation in the macroscopic gyral pattern could also underlie variability (e.g. Retzius 1896; Ono et al. 1990; Thompson et al. 1996; Baare et al. 2001; Yucel et al. 2001). For example, Heschl's gyrus exhibits a highly variable morphology (Brodmann 1909; von Economo and Horn 1930; Celesia 1976; Galaburda and Sanides 1980) that includes one to three gyri per hemisphere, with the number of gyri varying between hemispheres as well (Pfeifer 1920; von Economo and Horn 1930; Campain and Minckler 1976). Therefore an alternative explanation for our findings may be that the amount of variation in gyral shape patterns progressively increases across the macaque monkey, chimpanzee and human brain (Le May and Geschwind 1965; Chen et al. 2013) and may have led to the similarities and differences observed in both human and monkey brain variability. Alternatively, variability represents an important additional measure to the anatomical features described above, and may be an important tool for investigating evolutionary differences between species. Future studies will include a wider range of species in order to draw stronger conclusions about the evolutionary significance of structural variability.

Conclusions

Our three most striking findings were the high degree of similarity in variability between humans and monkeys, the strong correlation with macaque to human areal expansion maps, and the much greater degree of difference in variability between the hemispheres in humans than in monkeys, in specific functional areas. The latter supports the concept of increased hemispheric specialisation in the human brain, potentially one way in which evolution has led to a divergence between the human and monkey. Therefore, it may be that the same variability that makes us individually different from each other is also at the root of our differences from our ancestors and our closest evolutionary relatives. This suggests that cerebral variability may add another dimension to the study of evolutionary mechanisms.

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CAPTIONS

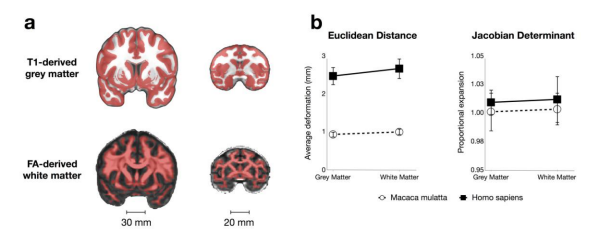


Figure 1 (colour): Quantification of the variability. a) T1 and FA maps templates computed for each species. Red indicated the mask used to calculate average deformation. b) Average deformation measured for grey and white matter with standard deviations. The left panel indicates absolute values quantified using Euclidean distance. The right panel indicates relative values estimated by the Jacobian determinant.

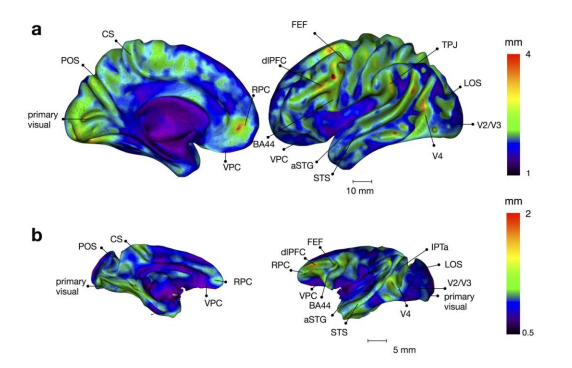


Figure 2 (**colour**): Grey matter variability. Three-dimensional projection of the average deformation measured for grey matter in humans (a) and monkeys (b). CS: central sulcus, POS: parieto-occipital sulcus, V1: striate cortex, V2: extra striate area 2, V3: extra striate area 3, V4: extra striate area 4, Cereb: cerebellum, VPC: ventral prefrontal cortex, RPC: rostral prefrontal cortex, LOS: lateral occipital sulcus, IPTa: inferior parietal tertiary association cortex, BA44: pars opercularis, FEF: frontal eye field, TPJ: temporo-parietal junction, aSTG: anterior superior temporal gyrus, STS: superior temporal sulcus. Please note that since neither a functional nor microscopic delineation of the microscopically or functionally defined cortical areas has been performed in this research, the localization of the functional areas reported in this illustration is an estimate.

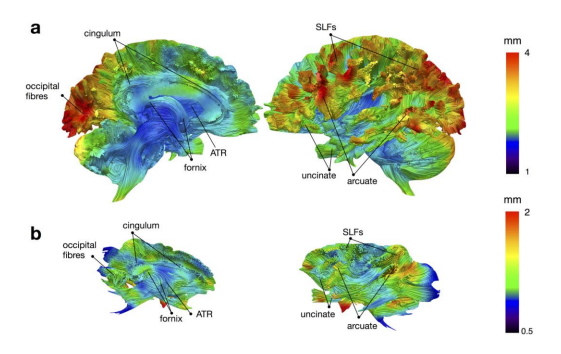


Figure 3 (colour): White matter variability. Three-dimensional projection of the average deformation measured for white matter in humans (a) and monkeys (b). SLFs: branches of the superior longitudinal fasciculus, ATR: anterior thalamic radiation.

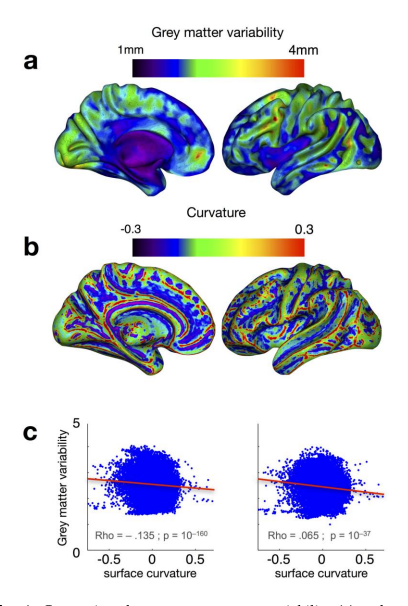


Figure 4 (colour): Comparison between grey matter variability (a) and cortical folding (b) in humans. c) Spearman rho correlations.

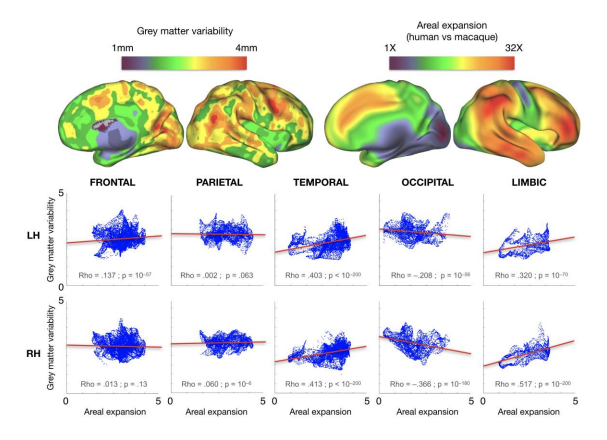


Figure 5 (colour): Comparison between grey matter variability and in humans and macaque to human areal expansion (2). Correlations are shown for each of the major subdivisions of the brain. Spearman rho and p values shown. LH: left hemisphere, RH: right hemisphere.

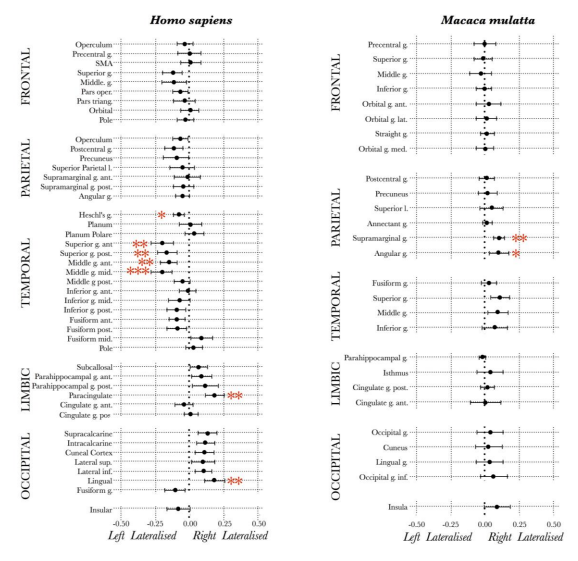
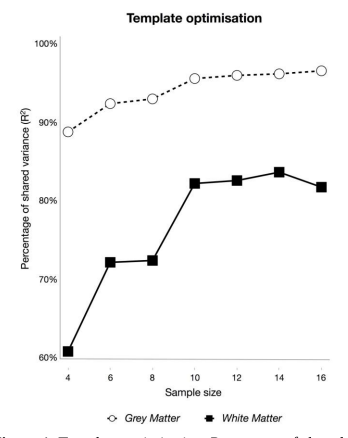
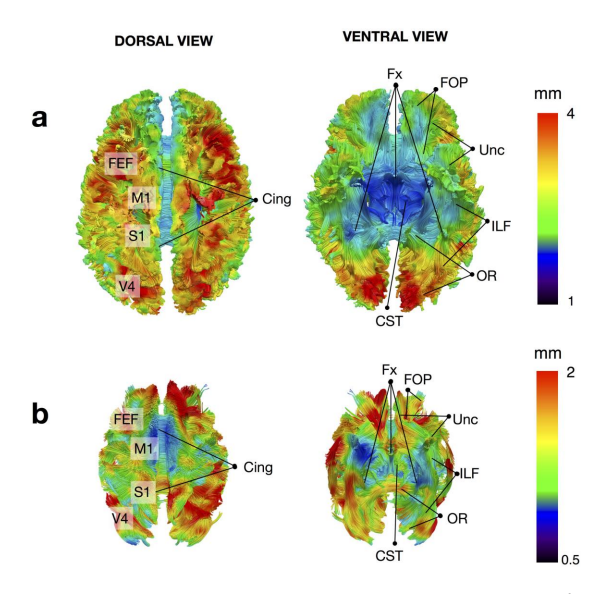


Figure 6 (BW): Lateralisation indices in grey matter variability in humans (left panel) and monkeys (right panel). Note that the scale for both species is different for visualisation purposes. ant.: anterior, mid.: middle, med.: medial, post.: posterior, sup.: superior, inf.: inferior, l.: lobule, g.: gyrus, SMA: supplementary motor area, oper.: opercularis, triang.: triangularis. * p < 0.05; ** p < 0.01; *** p < 0.001. Results are Bonferroni corrected for multiple comparisons.



Supplementary Figure 1: Template optimisation. Percentage of shared variance between the templates produced from the first and the second set of T1 and FA maps.



Supplementary Figure 2: White matter variability. Three-dimensional projection of the average deformation measured for white matter in humans (a) and monkeys (b). SLFs: branches of the superior longitudinal fasciculus, ATR: anterior thalamic radiation. Please note that since neither a functional nor microscopic delineation of the microscopically or functionally defined cortical areas has been performed in this research, the localisation of the functional areas reported in this illustration is an estimate.

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