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Early Left-Planum Temporale Asymmetry in newborn monkeys (*Papio anubis*): A longitudinal structural MRI study at two stages of development



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ABSTRACT

The "language-ready" brain theory suggests that the infant brain is pre-wired for language acquisition prior to language exposure. As a potential brain marker of such a language readiness, a leftward structural brain asymmetry was found in human infants for the Planum Temporale (*PT*), which overlaps with Wernicke's area. In the present longitudinal *in vivo* MRI study conducted in 35 newborn monkeys (*Papio anubis*), we found a similar leftward *PT* surface asymmetry. Follow-up rescanning sessions on 29 juvenile baboons at 7-10 months showed that such an asymmetry increases across the two ages classes. These original findings in non-linguistic primate infants strongly question the idea that the early *PT* asymmetry constitutes a human infant-specific marker for language development. Such a shared early perisylvian organization provides additional support that *PT* asymmetry might be related to a lateralized system inherited from our last common ancestor with Old-World monkeys at least 25–35 million years ago.

1. Introduction

Language and its typical functional and structural asymmetric brain organization were initially considered as unique to Homo sapiens evolution (Crow, 2004), suggesting a specific "language-ready" brain dating back to 350 000 ago. Therefore, brain lateralization in several regions was hypothesized as one of the key features of the language-ready brain, as most humans show a greater cortical activation in the left hemisphere for most language functions (Vigneau et al., 2006). For instance, the left Planum Temporale (PT) - a region which overlaps with Wernicke's area - was found particularly activated in a variety of auditory language processing tasks like phonological auditory decoding (Shapleske et al., 1999) and including the main perception component of the audio-motor loop for phonological processing (Vigneau et al., 2006). In the pioneering work of Geschwind and Levitsky (1968), a leftward PT asymmetry was also found at the anatomical level, suggesting its relationship with functional brain asymmetry for language tasks (Josse et al., 2006; Tzourio-Mazoyer et al., 2018).

Additionally, the "language-ready" brain theory suggests that the infant brain is pre-wired for language acquisition (e.g. Dehaene-Lambertz et al., 2002). Indeed, studies have reported that all human infants seem to have an innate, inherited readiness for language acquisition, independently from culture. For instance, newborns are initially able to distinguish every phoneme before selectively discriminating only phonemes related to the language they are exposed to Kuhl et al. (2008).

During their first year, infants will be also sensitive to vocal sounds, their native prosody and vowels, infer the abstract structure of speech and connect words to their referents (Dehaene-Lambertz and Spelke, 2015).

The neural structure for such a language readiness remains unclear. Nevertheless, white and grey matter organization in infants reveals similar architecture in comparison with adults (Dubois et al., 2010). Interestingly, similar to adults, three-month-old infants' BOLD responses to speech showed a more pronounced activation of the *PT* in the left hemisphere (Dehaene-Lambertz et al., 2002), rasing the question whether or not the *PT* might be functionally lateralized from birth on. Addition-

Abbreviation: PT, Planum Temporale.

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ally, structural *PT leftward asymmetry* was also shown in *post-mortem* fetuses or infant brains in early development (Witelson and Pallie, 1973; Wada, 1975; Chi et al., 1977) and with *in-vivo* structural MRI images (Dubois et al., 2010; Hill et al., 2010; Glasel et al., 2011). Such a structural *PT* left asymmetry may be stablished during the last trimester of fetal life (Chi et al., 1977) and are later increasing during development, suggesting its links with language development (Chi et al., 1977).

However, several studies in nonhuman primates questioned the structural *PT* asymmetry as a human-specific marker for the brain specialization for language. Manual delineation of *post-mortem* brain (Gannon et al., 1998) and *in-vivo* MRI scans (Hopkins et al., 1998; Hopkins and Nir, 2010; Marie et al., 2018), showed larger left *PTs* in both apes and baboons, suggesting that this neuroanatomical feature is shared also with Old-World monkeys.

Whether such an early neuroanatomical feature extended to other nonhuman primate infants is unknown although this comparative question remains critical for determining its supposed human uniqueness trough evolution and its relation to a pre-wired brain for language acquisition.

Therefore, the aim of the present longitudinal *in-vivo* MRI study in nonhuman primates is to investigate, the structural neuroanatomical *PT* asymmetries in 35 baboon infants (*Papio anubis*) and its development across age through manual delineation of the region's surface. The earliest post-natal age class includes 33 newborns at the critical neurodevelopmental period below 3 months (as well as two 5-months old outliers) in which the synaptogenesis is maximal and the myelin, synapses and cell bodies are thus not fully mature (Scott et al., 2016). The follow-up MRI longitudinal scanning and *PT* delineation includes 29 of those 35 baboons at the older juvenile age class (i.e., from 7 to 10 months).

2. Methods

2.1. Subjects

Subjects ranged from 4 to 165 days of age (*Mean*: 32.63; SD: 6.13) and included 21 males and 14 females. Out of those 35 baboons, 29 were later rescanned a second time, ranging from 218 to 362 days of age (*Mean* = 278.62; SD = 30.11) (see table in supplementary methods with subjects' details).

All monkeys are housed in social groups at the Station de Primatologie CNRS (UPS 846, Rousset, France) and have free access to outdoor areas connected to indoor areas. All subjects are born in captivity from 1 (F1) or 2 generations (F2). Wooden and metallic structures enrich the enclosures. Feeding times are held four times a day with seeds, monkey pellets and fresh fruits and vegetables. Water is available ad libitum.

2.2. Animal handling

Minimally invasive medication was realized, and no premedication was needed. Mothers from the newborn subjects were captured with their infant the night before the scan at the Station de Primatologie for check-ups and were transported together the following day of the MRI session.

Upon arrival at the MRI center, the mother of the focal subject was sedated with an intramuscular injection of ketamine (3mg/kg) and medetomidine (30 μ g/Kg) as well as their focal infant if above 5 months old. Focal newborn below 5 months were not sedated and directly brought to the preparation room for the following procedures. Focal infants were then anesthetized under 6-8% sevoflurane induction with a mask. A catheter was then inserted into the caudal artery for blood-gas sampling, and tracheal intubation was performed for steady controlled ventilation using an anesthetic ventilator (Cato, Drager, Germany). Endtidal carbon dioxide was monitored and used to adjust ventilation rate (0.2 to 0.3 Hz) and end-tidal volume. The anesthesia inside the MRI machine was then maintained using 3% sevoflurane via a calibrated

vaporizer with a mixture of air 0.75 L/min and $\rm O_2$ 0.1 L/min). Peripheral oxygen saturation, heart rate and breathing rate, were monitored throughout experiments.

All animal procedures were approved by the "C2EA -71 Ethical Committee of neurosciences" (INT Marseille) under the number APAFIS#13553-201802151547729 v4, and has been conducted at the Station de Primatologie under the number agreement C130877 for conducting experiments on vertebrate animals (Rousset-Sur-Arc, France). All methods were performed in accordance with the relevant French law, CNRS guidelines and the European Union regulations (Directive 2010/63/EU).

2.2.1. Imaging protocol

From September 2017 to March 2020, *in-vivo* imaging was performed using a 3T clinical MRI scanner (MAGNETOM Prisma, Siemens, Erlangen, Germany) equipped with 80 mT/m gradients (XR 80/200 gradient system with slew rate 200 T/m/s) and a 2-channel B1 transmit array (TimTX TrueForm). For the sessions at t0 ("newborn" age class) and at t1 (i.e., from 7 to 10 months old), the animals were scanned in the supine position, with two 11 cm receive-only loop coils: one under the head and another one around the face of the animal. The holding of the two coils and the animal head was provided through the use of a pearltec bag (Vac Fix System) and some straps. Protection for noise reduction was attached around the ears. At the end of the MRI session, when fully awaked from anesthesia, baboons were carefully put back with their mother and then transported back at the Station de Primatologie for immediate (or delayed) reintroduction into their social groups under staff monitoring.

2.2.2. Structural acquisition protocol

T1w images were acquired using a 3D Magnetization Prepared Rapid Acquisition Gradient Echo (MPRAGE) (Mugler and Brookeman, 1990) sequence (0.4 mm isotropic, FOV = $103 \times 103 \times 102.4$ mm, matrix=256 × 256 slices per slab=256, sagittal orientation, readout direction of inferior (I) to superior (S), phase oversampling = 10%, averages=3, TR = 2500 ms, TE = 3.01 ms, TI = 900 ms, flip-angle = 8°, bandwidth=300 Hz/pixel, no fat suppression, pre-scan normalization). T2w images were acquired using a Sampling Perfection with Application optimized Contrast using different angle Evolutions (SPACE) sequence (Mugler et al., 2000) (0.4 mm isotropic, FOV = $154 \times 115.5 \times 102.4$ mm, matrix=384 × 288, slice per slab=256, sagittal orientation, readout direction I to S, phase oversampling=0%, averages=2, TR = 3200 ms, TE = 393 ms, bandwidth = 566 Hz/pixel, no fat suppression, echo train length = 790 ms and pre-scan normalization). The total acquisition time for structural scans was 65 min (35 min for T1w and 30 min for T2w).

2.3. Preprocessing of anatomical MRI

Anatomical T2w images of the first scanning session and anatomical T1w images of the second scanning session were noise corrected with the spatially adaptive nonlocal means denoising filter (Manjón et al., 2010) implemented in Cat12 toolbox (http://www.neuro.uni-jena.de/cat/) included in SPM12 (http://www.fil.ion.ucl.ac.uk/), which runs on MAT-LAB (R2014a).

Next, each image was manually oriented using ITK-Snap 3.6 according anterior and posterior commissures plane and the interhemispheric fissure plane.

2.4. Manual delineation of the PT

Manual delineation of the *PT* in the present study followed the same procedure than the previous MRI study on the *PT* asymmetry of adult baboons (Marie et al., 2018). However, because of the immature brains of the newborn subjects, T2w MRI signal was used instead of T1w MRI signal for manual delineation for the first longitudinal scans.

In fact, the T2w MRI signal is sensitive to the free water present in voxels. The proliferation of membranes due to synaptogenesis and the process of myelination will decrease the proportion of free water in the gray matter voxels and thus darken the images during maturation of the first human year post-natal. Therefore, a higher contrast is generated in comparison to T1w images in early immature brains, which helps for better delimitation (Dehaene-Lambertz and Spelke, 2015).

Following the procedures used in humans (Larsen et al., 1989), great apes (Hopkins et al., 1998; Hopkins and Nir, 2010; Cantalupo et al., 2003) and Old-World monkeys (Marie et al., 2018; Lyn et al., 2011), the surface of the *PT* homolog was measured in the coronal plane.

The T2w images of every subject were imported in ITK-Snap. In there, the region of interest was manually traced in the coronal plan on the individual native space with the ITK-Snap tool "Paint-Brush Mode" with feature round brush size 1, using a touchpad-driven pointer (Wacom Cintiq® 13HD).

Coronal planes were used because they display the full depth of the sylvian fissure, of which the Planum Temporale is its floor. As described by Marie et al. (2018), delineation of the *PT* was performed as followed: The posterior border of the *PT* was characterized by the last caudal slice displaying the Sylvian fissure. The anterior border was defined by the full closure of the Insula sulcus and grey matter. This technique was chosen due to the inconsistency of the presence of the Heschl's gyrus (see Marie et al., 2018; Lyn et al., 2011 for discussions).

For each slice, the delineation was traced on the most ventral boundary between the sulcus and the grey matter. In order to balance the rater's possible handedness bias, tracing for each subject was randomly undertaken either from the most medial to the most lateral pixel of the Sylvian fissure or from the most lateral to the most medial pixel. This step was repeated on the next slice, moving posteriorly until the Sylvian fissure fell out of view.

Next, a surface area was generated across all slices for each hemisphere independently in a given subject (see Fig. 1.A). For each subject, an Asymmetry Quotient (AQ) of the left (L) and the right (R) surface areas was computed AQ = (R-L) / $[(R+L)\times 0.5]$ with the sign indicating the direction of asymmetry (negative: left side, positive: right side) and the value, the strength of asymmetry. Further, as reported by Hopkins and Nir (2010) for humans and great apes, the AQ was also used to classify the subjects as left-hemispheric biased (AQ \leq -0.025), right biased (AQ \geq 0.025), or nonbiased (-0.025 < AQ < 0.025). A threshold of 0.025 represents a 2.5% difference in surface area between left and right PT.

A second rater, blind to the side, confirmed the measures of the PT in a subsample of 15 individuals for both hemispheres (interrater correlation coefficient for 30 PT tracing was r(30) = 0.94, p < 0.0001).

2.5. Statistics

Statistics were conducted with R 3.6.1 (R Core Team (2017). R: A language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria. URL https://www.R-project.org/.)

3. Results

3.1. PT structural asymmetry

We found a significant leftward asymmetry of the PT surface at a group-level in 35 newborn baboons (t0) according to a one sample t-test in the 35 subjects' AQ scores (see Fig. 1.B), Mean AQ = -0.058 \pm 0.067 SD; t(34) = -5.15, p < 0.0001. Categorization of individual AQ showed also a majority of leftward PT-biased individuals (see Fig. 1.C): 25 baboons exhibited a leftward hemispheric PT bias (71.4%) whereas 2 exhibited a rightward PT bias (5.7%) and 8 no PT bias (22.9%), a distribution quasi-identical than the one found in human infants (Wada, 1975). We found no difference of distribution between infant baboons and infant humans according to chi-square (p = 0.25 for three groups "Left

bias, Right bias, no bias" and p = 0.20 for two groups "Left bias, Right bias"). The number of leftward PT-biased baboons was significantly greater than the number of rightward PT-biased subjects according to chi-square test ($\chi 2 = 19.59$, p < 0.0001).

3.2. Age classes' comparison of PT lateralization's strength

Follow-up analysis among the 29 rescanned baboons when reaching 7 to 10 months of age (t1) showed a significant increased strength of the PT asymmetry (Mean Absolute AQ score, M. = 0.105 \pm 0.065 SD) in comparison to their earliest age class according to a paired sample t-test (Mean Absolute AQ score, M. = 0.073 \pm 0.049 SD), t(28) = -2.39, p = 0.024 (see Fig. 1.D) as well as a significant correlation between the two MRI sessions, r(29) = 0.55, p < 0.002.

3.3. Left, right PT surface areas

At t0, the mean PT surface areas were in the left hemisphere: $M.=50.34~\mathrm{mm^2}\pm9.27~\mathrm{SD}$ (in males $M.=51.07~\mathrm{mm^2}\pm9.80~\mathrm{SD}$; in Females $M.=49.24~\mathrm{mm^2}\pm9.17~\mathrm{SD}$); and in the right hemisphere: $M.=47.44~\mathrm{mm^2}\pm8.93~\mathrm{SD}$ (in males $M.=48.53~\mathrm{mm^2}\pm9.35~\mathrm{SD}$; in Females $M.=45.79~\mathrm{mm^2}\pm7.78~\mathrm{SD}$).

At t1, the mean *PT* surface areas were in the left hemisphere: $M. = 58.48 \text{ mm}^2 \pm 8.0 \text{ SD}$ (in males $M. = 59.23 \text{ mm}^2 \pm 8.07 \text{ SD}$; in Females $M. = 57.67 \text{ mm}^2 \pm 7.75 \text{ SD}$); and in the right hemisphere: $M. = 55.73 \text{ mm}^2 \pm 8.65 \text{ SD}$ (in males $M. = 57.07 \text{ mm}^2 \pm 9.99 \text{ SD}$; in Females $M. = 54.28 \text{ mm}^2 \pm 6.64 \text{ SD}$).

3.4. Age, sex, brain size effect

Multiple linear regression analyses showed that the right PT surface (p=0.001), the left PT surface (p=0.02) and age (p=0.033) predict PT asymmetry strength but not the subject's sex and brain volume. At t0 and t1, no significant differences of mean AQs, mean Left PT surface, and of mean Right PT surface were found between males and females.

4. Discussion

Our results showed that early post-natal nonhuman primate infants present a significant human-like neuroanatomical asymmetry of the Planum Temporale surface (PT) in favor of the left hemisphere. This finding is clearly consistent with early PT asymmetry found in human newborns and infants (Chi et al., 1977; Dubois et al., 2010; Glasel et al., 2011; Hill et al., 2010; Wada, 1975; Witelson and Pallie, 1973) although measurement methods, Left-Right-Ambi classification threshold and statistical power in terms of sample size differ as well as age class equivalence which overall make interspecies comparison challenging. Nevertheless, the distribution is quasi-identical to the ones reported in both human infants and human adults (Geschwind and Levitsky, 1968) but also in adult chimpanzees (Hopkins and Nir, 2010) and adult baboons (Marie et al., 2018). Our findings are also somewhat consistent with averaged-brain leftward asymmetries found in infant Rhesus macaques within large temporal clusters which seem to overlap with PT according to an automated source-based method (Xia et al., 2019). Such a similar age-related phenomenon was also described in human infants by Wada (1975). Interestingly, we found that the direction of individual PT asymmetry is consistent across age classes while its strength is increasing with age. In contrast, no sex or brain size effects were found on direction or strength of PT asymmetry. This finding is not consistent with the idea that increase in PT asymmetry in Hominidae evolution was due to increase in brain volume (Pilcher et al. 2001). Additionally, if strength of PT asymmetry is affected by sex in human adults (Hirnstein et al., 2019), it seems not the case in adult baboons (Marie et al., 2018), infant baboons and human infants (Dubois et al. 2010).

This finding in a non-linguistic species clearly questions the historical idea that such a maturational effect of the *PT* asymmetry's strength

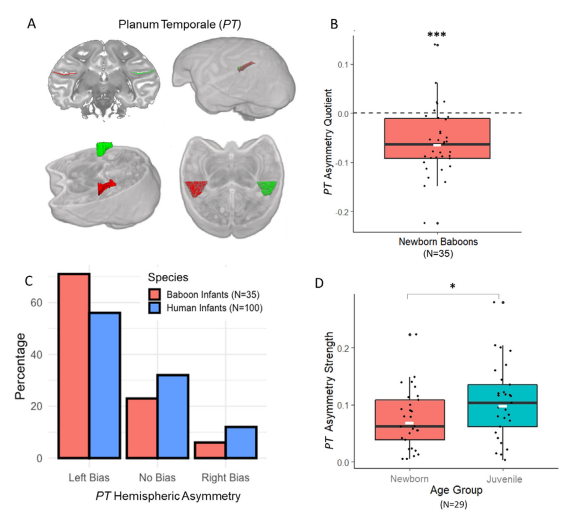


Fig. 1. (A) Representation of the asymmetric Planum Temporale (PT) in the baboon brain on a T2w image according to a coronal section, 3D brain render and oblique section oriented along the Sylvian Fissure (the left PT is in red and the right PT in green). (B) Mean Asymmetry quotient (AQ) for the Planum Temporale surface of the newborn infant baboons (N = 35). Negative Mean AQ score indicates leftward hemispheric asymmetry at a population-level. The long black line represents the median, the white short line the mean. ***P < 0.0001 (C) Subjects distribution (in percentage) as a function of the direction of their PT asymmetry in baboon infants (N = 35, in red) versus in human infants (N = 100, in blue, from Wada, 1975). (D) Variation of strength of the Planum Temporale surface's asymmetry (Mean absolute AQ score) among the 29 baboons scanned longitudinally at two early stages of development: Newborn (in red) *versus* Juveniles (i.e., from 7 to 10 months in blue). *P < 0.05.

is related to language development in human infants (Wada, 1975). One could ask whether the existence of the asymmetry shortly after birth has an innate, and thus, genetic component as hypothesized for human infants (Hill et al., 2010) or to what extent it is rather influenced by pre- and post-natal experience. In any case, the collective findings clearly provide additional support for the phylogenetic continuity between human and nonhuman primate species about such a brain asymmetric feature. Such a continuity extended at the earliest postnatal stage of development across both species may question the early *PT* asymmetry as a human newborn-specific marker of the language-ready brain.

Structural lateralization of such a language area may not solely account for a pre-wired brain for language acquisition as it was supposed for human babies (Dehaene-Lambertz et al., 2002).

However, it remains unclear which factor is driving such a common early asymmetric feature of the brain anatomy among human and nonhuman infants. One potential explanation is that early *PT* structural asymmetry might have nothing to do with development of language lateralization, given some studies in adults reported no match between structural and functional asymmetry of this region (Keller, 2011; Greve, 2013).

Nevertheless, the most recent study addressing this question in adults contradicts such a hypothesis (Tzourio-Mazoyer et al., 2018). Although the lack of match was confirmed between structural and functional asymmetry of the PT in a language task, structural PT asymmetry was found associated with functional lateralization of an adjacent auditory area at the end of the Sylvian fissure, suggesting its links with language lateralization.

Therefore, another potential explanation is that the early *PT* structural asymmetry in both human and nonhuman infants might predict the development of homolog communicative functions, which still form a foundation for core aspects of the human language system. Determining such common developing functions between species remains highly speculative, given the lack of longitudinal studies in infants on the emergence of brain-behavior relationships. Nevertheless, based on neuroimaging studies in nonhuman primate focusing on adults, it might be not excluded that shared properties of communicative systems in human and nonhuman primates could be related to *PT* structural asymmetry. For instance, previous studies in monkeys and apes have reported human-like functional lateralization for processing conspecific calls. However, its overlap with *PT* anatomical region remains unclear as well as the direction of the functional lateralization (i.e., toward left

versus right hemisphere) which are inconsistent across the literature (e.g. Poremba et al.; 2004; Gil-da-Costa and Hauser 2006; Petkov et al. 2008; Joly et al., 2012). Alternatively, some authors have proposed that properties of the communicative gestural system in nonhuman primates could constitute another potential functional candidate of PT specialization. In fact, whereas production of communicative manual gestures have been found highly lateralized in favor of the right-hand in both baboons and chimpanzees (Meguerditchian et al., 2013), a contralateral relation between PT structural asymmetry and hand preferences for communicative gesture was reported in adult chimpanzees (Hopkins and Nir, 2010; Meguerditchian et al., 2012). These latter findings have thus suggested that shared property between gesture signaling in apes and language system in humans might be both ultimately related to this asymmetry feature of the temporal lobe anatomy (Meguerditchian et al., 2012). Whether similar gestural functional specialization of the structural PT asymmetry exists in baboons has been not investigated yet although both chimpanzees and baboons have shown similar leftward structural asymmetry of the PT as well as similar rightward patterns of gestural communication's manual lateralization.

Further studies in our sample of infant baboons would help us determinate the potential relationship between these early *PT* structural asymmetry and development of manual lateralization of communicative gestures.

In conclusion, the present finding in nonhuman infants provides additional support to the hypothesis of a continuity between nonhuman and human primates concerning leftward structural *PT* asymmetry. Sharing such an anatomical feature of the brain at this earlier postnatal stage of development reinforced thus the idea of its common origins from our distant evolutionary ancestor, dated back 25–35 million years ago, although its potential link with the language-ready brain remains an open question.

Declaration of Competing Interest

Authors declare no competing interests.

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Author contributions

Y.B and A.M prepared the paper and the revision. Y.B. performed the tracing and analyses. C.B. performed the interrater tracing. J.S. parametrized the MRI sequences and optimized the MRI acquisition setup. B.N. designed the baboons' monitoring programs. L.V., L.R., R.L. and L.G. designed the specific procedures of welfare, anesthesia, monitoring and preparation of baboons in the MRI machine. J.L.A supervised and coordinated the MRI session. A.M. designed and supervised the study and MRI acquisitions.

Data and materials availability

All data is available in the supplementary materials.

Data availability statement

Our data will be available online upon publication and is attached to this submission in the supplementary material.

Supplementary materials

Supplementary material associated with this article can be found, in the online version, at doi:10.1016/j.neuroimage.2020.117575.

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