Transcriptional neoteny in the human brain

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In development, timing is of the utmost importance, and the timing of developmental processes often changes as organisms evolve. In human evolution, developmental retardation, or neoteny, has been proposed as a possible mechanism that contributed to the rise of many human-specific features, including an increase in brain size and the emergence of human-specific cognitive traits. We analyzed mRNA expression in the prefrontal cortex of humans, chimpanzees, and rhesus macaques to determine whether human-specific neotenic changes are present at the gene expression level. We show that the brain transcriptome is dramatically remodeled during postnatal development and that developmental changes in the human brain are indeed delayed relative to other primates. This delay is not uniform across the human transcriptome but affects a specific subset of genes that play a potential role in neural development.

Results and Discussion

General Pattern of Expression Changes During Brain Development. We analyzed gene expression levels in the dorsolateral prefrontal cortex (DLPFC) of 39 humans, 14 chimpanzees, and 9 rhesus macaques by using Affymetrix GeneChip Human Genome (GC HG-U133 Plus 2.0 microarrays (see Materials and Methods and SI Appendix, Table S1). For both humans and chimpanzees, the individuals’ age distributions cover the entire span of postnatal ontogenesis, with a particular focus on early life stages (Fig. 1A and SI Appendix, Fig. S1). In these individuals, we reliably detected and quantified the expression of 7,958 genes (Materials and Methods). Among these genes, we first analyzed the relative influence of factors—age, sex, and species—on total expression variation among individuals. Quantitatively, age explains the largest part of the total expression variation at 29%, followed by species at 17% and sex at <2% (Fig. 1B). Although the effects of age and species are highly significant (permutation test, P < 0.001), the effect of sex is not (P = 0.54) (SI Appendix, Table S2). Thus, in our dataset, age has by far the greatest influence on expression levels. Consistently, we find that a striking 71% of the 7,958 genes expressed in the human brain change significantly during postnatal development [at a false discovery rate (FDR) of 10%] (Fig. 1C). Functionally, these genes are significantly enriched in a range of biological processes that include cell adhesion, synaptic transmission, and axonogenesis (permutation test for overall enrichment, P = 0.002) (SI Appendix, Table S3).

Next, we estimated when during human and chimpanzee brain development these expression changes take place. We find that in both species gene expression changes occur most rapidly during the first few years of life. Approximately 50% of the total expression change observed between newborns and 40-year-olds occurred within the first year of life (Fig. 1D). Furthermore, the overall trajectory of age-related expression changes in the chimpanzee brain, although based on fewer samples, closely resembles the


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Data deposition: The data reported in this paper have been deposited in the Gene Expression Omnibus (GEO) database, www.ncbi.nlm.nih.gov/geo (GEO accession nos. GSE11528, GSE11512, and GSE15163).

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human one (Fig. 1D). Notably, we also find similar trajectories of genome-wide expression changes with age in the brains of 2 species of mouse (Mus musculus and Mus spretus) from birth until adulthood (Materials and Methods and SI Appendix, Fig. S2). Furthermore, the trajectories of age-related changes also show a high correlation on the individual gene level, both between humans and chimpanzees (median Pearson r = 0.90) and between humans and mice (median r = 0.83) (SI Appendix, Table S4). Thus, the pattern of expression changes with age observed here is not particular to humans and chimpanzees and most likely reflects fundamental changes shared among mammals in the brain's molecular and histological organization during postnatal development.

**Human–Chimpanzee Expression Differences.** We have found that age-related expression trajectories are generally conserved among humans, chimpanzees, and mice. Nevertheless, for individual genes the timing of expression changes may differ between species. To explore this issue, we tested each gene for expression differences between humans and chimpanzees by using multiple regression models. We find that 48% of age-related genes in humans are either expressed at significantly different levels or follow significantly different expression trajectories with age than in chimpanzees (at P < 0.0025, FDR = 10%) (Fig. 1C). Hence, despite the similarities between the expression profiles in human and chimpanzee brains described above, among all genes that change during prefrontal cortex development, approximately half change with age differently in the 2 species (Fig. 1C).

The proportion of genes classified as differently expressed between human and chimpanzee brains in this study is greater than previously reported (<10%) (18, 19) because of the larger sample size and different statistical criteria used here. Equalizing these factors, we find similar proportions of differently expressed genes (33% and 35%) as well as a significant overlap (50%, P < 10^{-10}) between this and a published dataset (19).

**A Test for Expression Heterochrony.** We next asked whether age-related expression differences between humans and chimpanzees reflect shifts in the timing of ontogenetic changes (i.e., heterochrony) between the 2 species. Heterochronic expression changes can be in the form of either delays (neoteny) or accelerations. Therefore, we designed a test to estimate both the direction and the amplitude of the shift in timing between expression profiles in the 2 species. In essence, this heterochrony test assumes that the species' ontogenetic trajectories are similar in shape (SI Appendix, Fig. S2) and then estimates whether introducing a difference in developmental timing significantly improves the fit between the 2 species' expression profiles (SI Appendix, Fig. S3).

Before applying this test to the human–chimpanzee comparison, we used it on a known developmental timing difference within the human brain: maturation of the human prefrontal cortex and of the caudate nucleus. The prefrontal cortex is among the last regions to mature during human brain ontogenesis (20, 21), but the caudate nucleus, a subcortical region of the brain, matures relatively early (20). To test whether we can detect a corresponding shift in gene expression timing, we compared gene expression patterns in the prefrontal cortex to those in the caudate nucleus in 13 human individuals aged 0–46 years old (Materials and Methods). We identified 2,979 genes as both age-related and differently expressed between the 2 brain regions. Applying the heterochrony test to these genes, we find that 2,261 of these genes show significant expression heterochrony between the 2 regions (at P < 0.05) and that for 58% of these genes the direction of the shift corresponds to slower maturation of the prefrontal cortex relative to the caudate nucleus (SI Appendix, Table S5). Thus, in line with anatomical observations, our test detects a substantial delay in postnatal development of the
prefrontal cortex compared with the caudate nucleus on the gene expression level.

**Testing the Neoteny Hypothesis of Human Evolution.** Next, to assess the differences in developmental timing between humans and other primates, we applied the heterochrony test to the human and chimpanzee expression data. To minimize possible sampling bias, we used a subset of 14 humans closely matching the 14 chimpanzees with respect to age and sex. Furthermore, we used the rhesus macaque as an outgroup to assign genes showing expression heterochrony to one of the following 4 phyloontogenetic categories: (i) human neoteny—expression changes occurred on the human lineage, and human expression corresponds to that in younger chimpanzees; (ii) human acceleration—expression changes occurred on the human lineage, and human expression corresponds to that in older chimpanzees; (iii) chimpanzee neoteny; and (iv) chimpanzee acceleration (Fig. 2).

Among the 3,075 genes expressed in all 3 species, we could confidently assign 299 to one of these 4 categories. Of these 299 genes, approximately the same proportions (15 to 25%) fall into categories ii, iii, and iv. In contrast, we find approximately twice as many genes (38%) in category i, human neotenic genes (Fig. 3A). This enrichment is robust against the choice of P value cutoffs, models used to describe age-related changes of individual genes, and criteria used to define neoteny (Table 1 and SI Appendix, Table S6). Furthermore, we observe the same results by using the complete human dataset, rather than a subset of 14 individuals. In addition, we obtain qualitatively the same results regardless of our assumptions concerning the gestation-time difference between humans and chimpanzees (Table 1). By comparing humans with humans, we also estimate the FDR of the test as ~10% (SI Appendix, Table S7).

To determine the generality and robustness of this result, we generated a second brain development dataset by using another prefrontal cortex region, the superior frontal gyrus (SFG), taking samples from 9 human, 9 chimpanzee, and 9 rhesus macaque individuals. All human and rhesus individuals differed from those used in the first dataset, and the experiment was conducted on a different microarray platform (Affymetrix Human Gene 1.0 ST). Applying the same analysis procedure to these data, we again find a significant excess of human neotenic genes compared with the other 3 phyloontogenetic categories (Fig. 3A and Table 1). The overlap between the human neotenic genes identified in the 2 brain regions is approximately twice as large as expected [Fisher’s exact test, P < 0.01] (SI Appendix, Table S8). Thus, there is a reproducible, human-specific, neotenic shift in gene expression during postnatal maturation of the human prefrontal cortex, causing adult humans to resemble juvenile chimpanzees in their expression profiles (SI Appendix, Fig. S4).

This shift, however, affects a limited portion of the total cortical transcriptome (~4%). By using simulations, we show that our test may fail to identify many real heterochronic changes (SI Appendix, Table S7), and therefore the actual percentage of genes affected by a human neotenic shift is likely to be higher. Still, based on the simulation results, we can confidently exclude the existence of a global neotenic shift affecting the entire transcriptome of the human frontal cortex (SI Appendix, Table S7). Thus, a parallel may be drawn between this result and previous morphological studies. Earlier formulations of the neoteny hypothesis of human evolution (e.g., that by L. Bolk) postulated that an ontogenetic shift affects human development in its entirety, including multiple organs and tissues (6). This notion was later modified and restricted the neotenic shift to brain growth (10, 11). Similarly, we find that the neotenic shift affects a limited group of genes expressed in the brain rather than the entire brain transcriptome. In a more general sense, this result suggests that mosaic evolution, as seen in brain structures among mammals (22), also applies to the evolution of gene expression patterns.
Properties of Human Neotenic Genes. We analyzed the genes affected by the neotenic shift in the human prefrontal cortex with respect to their histological location, function, regulation, and expression timing. First, with respect to their histological location, we used published gene expression data from human gray and white matter (23) and found that, in both brain regions, human neotenic genes are significantly overrepresented among genes expressed specifically in gray matter (P = 0.06 and P = 0.0001 in DLPFC and SFG, respectively) but not among genes expressed in white matter (P > 0.6) (Fig. 3B and SI Appendix, Table S9).

Second, with respect to function, we find that in both brain regions human neotenic genes show a tendency to cluster in biological processes related to growth and development (SI Appendix, Tables S10 and S11). When each dataset was considered separately, this result was not significant (P > 0.1).

Third, with respect to expression timing, we asked whether the observed neotenic shift affects human postnatal ontogenesis in its entirety or is limited to a particular age interval. If the latter is true, human–chimpanzee expression divergence should increase at the corresponding age. We thus compared the extent of expression divergence across age among human neotenic genes to expression divergence among genes in the other 3 phyloontogenetic categories as well as of all age-related and differentially expressed genes. For all gene categories, human–chimpanzee expression divergence is relatively small after birth and subsequently increases. Human neotenic genes, however, diverge more rapidly than genes in the other categories during early adolescence, showing the largest difference to other groups ~10 years of age (Fig. 3C). We reproducingly find the same result in the DLPFC dataset by using either all 39 or 14 selected human individuals. We find a similar increase in expression divergence around adolescence in the second dataset from the SFG (because of the limited sample size and wider age distribution in this dataset, we could not assess the timing and significance of this increase with confidence). Thus, at least in the DLPFC, the neotenic shift does not uniformly affect human life span but is particularly prominent in early adolescence.

Conclusion
By comparing the gene expression profiles in human, chimpanzee, and rhesus macaque prefrontal cortices throughout postnatal development, we have found that there is no uniform shift in the developmental timing between humans and other primates. We find instead a significant excess of genes showing neotenic expression in humans. This result is in line with the neoteny hypothesis of human evolution (6) and provides insight into the possible functional role of neoteny in human brain development. Specifically, we show that at least in one of the 2 cortical regions studied, the neotenic shift is most pronounced at the time when humans approach sexual maturity, a process known to be delayed in humans relative to chimpanzees or other primates (6, 24). Furthermore, the neotenic shift particularly affects a group of genes preferentially expressed in gray matter. Intriguingly, the timing of the shift also corresponds to a period of substantial cortical reorganization characterized by a decrease in gray-matter volume, which is thought to be related to synaptic elimination (21, 25, 26). The developmental pace of changes in gray-matter volume has been associated with the development of cognitive skills among humans (e.g., linguistic skills) (27) as well as with the development of disorders (e.g., attention-deficit/hyperactivity disorder) (28). Although the precise causes and consequences of the human neotenic shift remain unknown, together these observations suggest that ontogenetic timing differences between the human and the chimpanzee prefrontal cortex transcriptomes may reflect differences in sexual and cognitive maturation between the 2 species. According to this logic, delayed gray-matter maturation in the human prefrontal cortex may extend the period of neuronal plas-
The age distributions of the primates analyzed.

Of age, the relative ages of the mice used here (0, 14, and 56 days) approximate MG-430 2.0 GeneChip arrays. Because mice reach sexual maturation at 9.

A significant difference in RNA quality between species or within each species with samples, 5 of the 9 individuals were males, and there was no significant bias in the Gene 1.0 ST arrays. All humans and all but one rhesus macaque individual were rhesus macaques (0–28 years) and hybridized to Affymetrix GeneChip Human humans (ranging in age from 0–66 years), 9 chimpanzees (0–44 years), and 9

The binomial test P value for neoteny with the alternative hypothesis that >50% of genes that are assigned to the human lineage show delayed development in humans relative to chimpanzees.

The binomial test P value for human specificity with the alternative hypothesis that >50% of genes that show delayed development in humans vs. chimpanzees are assigned to the human lineage.

Stringent cutoffs: P < 0.01 in all 4 tests.

Relaxed cutoffs: P < 0.10 in all 4 tests.

ticity associated with active learning, thus providing humans with additional time to acquire knowledge and skills.

Materials and Methods

Sample Collection and Hybridization. For the first primate dataset, we dissected postmortem DLPCF samples containing 90–95% gray matter from 39 humans (ranging in age from 0–47 years; 67% males), 13 chimpanzees (0–44 years; 64% males), and 9 rhesus macaques (1–18 years; 44% males), and caudate nucleus samples from 13 humans (0–66 years; 78% males) (SI Appendix, Table S1). RNA extracts from the dissections were used to generate labeled cRNA and hybridized to Affymetrix GC HG-U133 Plus 2.0 arrays according to the standard protocol. Among the 3 species, human samples have more detailed sample information. Testing the effects of various sample characteristics—such as sex, RNA quality, postmortem delay, prolonged agonal state, and brain tissue pH—on gene expression levels above the background level in one third of the individuals in either species were considered expressed. Using different expression detection criteria did not affect our results (SI Appendix, “Supporting Materials and Methods”). In all experiments, probe sets with <8 probes remaining after the masking procedure were excluded from the analysis. Genes with detected expression levels above the background level in one third of the individuals in either species were considered expressed. Using different expression detection criteria did not affect our results (SI Appendix, “Supporting Materials and Methods”).

To analyze chimpanzee and rhesus macaque expression profiles in an unbiased way, in both primate datasets we masked the array probes that did not match the analyzed species’ genomes perfectly and uniquely by using the corresponding reference genomes (SI Appendix, Table S14). Similarly, for mice we identified MG-430 2.0 probes that did not match the M. spretus genome by using a statistical algorithm that detects discordant expression patterns among probes within a probe set (SI Appendix, “Supporting Materials and Methods”); these probes were masked.

For the GC HG-U133 Plus2.0 and MG-430 2.0 arrays, to group probes into probe sets, we used published chip definition files based on Ensembl genes (29). Expression levels were summarized, log-transformed, and normalized by using the “rma” function in the Bioconductor “affy” package (30). For the HuGene 1.0 arrays we used the “rma” algorithm adjusted to this novel array type to compute gene expression values and Affymetrix annotation files to map transcript clusters to Ensembl genes (SI Appendix, “Supporting Materials and Methods”).

In all experiments, probe sets with <8 probes remaining after the masking procedure were excluded from the analysis. Genes with detected expression levels above the background level in one third of the individuals in either species were considered expressed. Using different expression detection criteria did not affect our results (SI Appendix, Table S6). All original data files from the microarray experiments have been deposited in the National Center for Biotechnology Information Gene Expression Omnibus (GEO) with the accession numbers GSE11512 (GC HG-U133 Plus2.0 experiments) and GSE11528 (MG-430 2.0 experiments) (GSE15163 for the HuGene 1.0 ST experiments). The Chip Definition Files and R code used in the analysis can be found at www.picb.ac.cn/Comparative/data.html.

**Analyses of Age and Species Effects.** For analyzing age-related changes, we used base-2, log-transformed ages (log-age) starting from the inferred conception date. Log-transformed ages are frequently used to analyze developmental phenotypes with parametric models (e.g., 31–33). In our dataset, this transformation yields a more uniform distribution of errors across ages than the linear age scale, making it more suitable for statistical analysis (34). Nonetheless, an analysis based on the linear age scale results in the same principal findings (SI Appendix, Table

**Table 1. Expression heterogeneity test results**

<table>
<thead>
<tr>
<th>Dataset</th>
<th>DLPCF</th>
<th>SFG</th>
<th>Overlap</th>
</tr>
</thead>
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<tr>
<td>Numbers of humans vs. chimpanzees compared</td>
<td>14 vs. 14</td>
<td>39 vs. 14</td>
<td>14 vs. 14</td>
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<td>Assumed gestation-time difference in days</td>
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<td>0</td>
<td>20</td>
</tr>
<tr>
<td>Number of expressed genes</td>
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<td>3,075</td>
<td>3,075</td>
</tr>
<tr>
<td>Human-neotenic*</td>
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<td>154</td>
<td>121</td>
</tr>
<tr>
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<td>88</td>
<td>59</td>
</tr>
<tr>
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<td>46</td>
<td>86</td>
<td>59</td>
</tr>
<tr>
<td>Chimpanzee-neotenic*</td>
<td>74</td>
<td>118</td>
<td>78</td>
</tr>
<tr>
<td>P neoteny*</td>
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<td>1 × 10&lt;sup&gt;-5&lt;/sup&gt;</td>
<td>2 × 10&lt;sup&gt;-6&lt;/sup&gt;</td>
</tr>
<tr>
<td>P human specificity**</td>
<td>4 × 10&lt;sup&gt;-8&lt;/sup&gt;</td>
<td>7 × 10&lt;sup&gt;-6&lt;/sup&gt;</td>
<td>2 × 10&lt;sup&gt;-6&lt;/sup&gt;</td>
</tr>
<tr>
<td>Human-neotenic§</td>
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<td>97</td>
<td>61</td>
</tr>
<tr>
<td>Human-accelerated§</td>
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<td>61</td>
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<td>P neoteny§</td>
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<td>8 × 10&lt;sup&gt;-6&lt;/sup&gt;</td>
<td>6 × 10&lt;sup&gt;-6&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

*FDR = 10% for age and species effects; P < 0.05 for heterochrony and lineage (1-sided) tests.

*The binomial test P value for neoteny with the alternative hypothesis that >50% of genes that are assigned to the human lineage show delayed development in humans relative to chimpanzees.

The binomial test P value for human specificity with the alternative hypothesis that >50% of genes that show delayed development in humans vs. chimpanzees are assigned to the human lineage.

Stringent cutoffs: P < 0.01 in all 4 tests.

Relaxed cutoffs: P < 0.10 in all 4 tests.

Showed a slight but significant difference in RNA quality, accounting for this
Another factor related to age assignment is the difference in gestation time between species. Although humans are known to have a longer gestation time than chimpanzees, the exact extent of this difference is not certain and may vary among individuals. Therefore, to avoid any bias that can be caused by incorrect assumptions concerning the gestation-time difference, we assigned the same gestation time (280 days) to humans and chimpanzees in the main analysis. Such an assignment is conservative, because it ignores a shorter gestation period in chimpanzees and biases our results against finding human-specific neotony.

Using different gestation times for humans and chimpanzees did not affect our findings, including the lower percentage of detected neotenic shifts than would have been expected from a global shift (Table 1).

For computing the proportion of variance explained by age, species identity, or sex among all expressed genes, we used cubic regression models with log-age and linear regression models with species identity or with sex, respectively.

We used the multidimensional scaling algorithm “isoMDS” in the R “MASS” package (35) to calculate a 1D summary measure of global expression shown in Fig. 1D. We tested the age-related expression changes per gene by using polynomial regression models with log-age and species as factors and using the F test to assess significance (SI Appendix, Table S16). In both the age and sex tests, FDRs were computed by 1,000 random permutations of age or sex assignments and fixed at 0.05. Differently expressed genes between species were assigned to the human lineage if the distances among the 9 rhesus expression values and the human expression age curve were significantly greater than the distances to the chimpanzee curve (1-sided Wilcoxon test, P < 0.05) and vice versa for genes assigned to the chimpanzee lineage (SI Appendix, Table S17). Using multiple regression models for lineage assignment gives the same results (SI Appendix, Table S6).

Testing Expression Heterochrony. We identified and quantified differences in expression timing between humans and chimpanzees (or between the prefrontal cortex and the caudate nucleus) by testing whether a linear age transformation (i.e., an age shift) significantly minimizes the difference between the expression age trajectories of the 2 groups (see Fig. S3). To find the optimal age shift for each gene, we used a nonlinear least squares algorithm (NL2SOL) (“nl” function in the R “stats” package). The significance of the age transformation was assessed by using the F test (P < 0.05). We also tested whether the age transformation explains human–chimpanzee differences (or prefrontal cortex–caudate nucleus differences) at least as efficiently as a constant expression-level difference between two groups (SI Appendix, “Supporting Materials and Methods”).

We estimated the false-positive and false-negative rates by comparing humans with humans and by simulating transcriptome-wide age shifts on the orders of 10–30% and 70%, respectively (SI Appendix, Table S7). Note that current literature distinguishes among 3 types of heterochrony: a difference in the onset of development, a difference in the rate of development, and a difference in the duration of development (13, 37). In this context, our test evaluates either a shift in the onset of development or a difference in developmental rates without distinguishing between the two, a result of our limited sample size (SI Appendix, “Supporting Materials and Methods”). Because of this limitation, we use the term “neotony” to collectively describe all detectable situations in which adult human expression levels resemble expression levels of younger chimpanzees (SI Appendix, Fig. S4).

Characterization of Human-Neotenic Genes. We compared human-neotenic genes to genes identified in the other 3 ontogenetic categories in terms of (i) their enrichment in gene ontology groups under the biological process taxonomy (38) by using the FUNC tool (39), which applies a correction for multiple testing; (ii) their enrichment in genes expressed in specific brain structures; and (iii) changes in divergence between humans and chimpanzees across age, calculated as the absolute distance between human and chimpanzee expression age curves. In point ii, we identified 1,155 and 578 genes with gray- and white-matter-specific expression, defined as a 3-fold difference in expression between these structures by using published microarray data from human cortical gray and white matter (23) (SI Appendix, “Supporting Materials and Methods”). For all tests, we used phyloontogenetic gene sets defined based on FDRs as well as gene sets defined based on relaxed significance cutoffs (at P < 0.1) (see Table 1).

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