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Rewiring of human neurodevelopmental gene regulatory programs by human accelerated regions

Highlights

- Comprehensive *in vitro* and *in vivo* epigenomic profile of more than 3,100 HARs
- Nearly half of all HARs have features of neurodevelopmental enhancers
- Extensive human-specific rewiring of the *cis*-regulatory architecture of HARs
- Primate- and human-specific change in *PPP1R17* developmental expression patterns

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In brief

Human accelerated regions (HARs) are the fastest-evolving regions of the human genome. By interrogating the *in vitro* enhancer activity and *in vivo* epigenetic landscapes of more than 3,100 HARs during human neurodevelopment, we demonstrate that nearly half of all HARs act as neurodevelopmental enhancers. Further, we demonstrate *PPP1R17* to be a putative HAR-regulated gene likely involved in human-specific cortical neurodevelopment.



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Rewiring of human neurodevelopmental gene regulatory programs by human accelerated regions

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SUMMARY

Human accelerated regions (HARs) are the fastest-evolving regions of the human genome, and many are hypothesized to function as regulatory elements that drive human-specific gene regulatory programs. We interrogate the *in vitro* enhancer activity and *in vivo* epigenetic landscape of more than 3,100 HARs during human neurodevelopment, demonstrating that many HARs appear to act as neurodevelopmental enhancers and that sequence divergence at HARs has largely augmented their neuronal enhancer activity. Furthermore, we demonstrate *PPP1R17* to be a putative HAR-regulated gene that has undergone remarkable rewiring of its cell type and developmental expression patterns between non-primates and primates and between non-human primates and humans. Finally, we show that *PPP1R17* slows neural progenitor cell cycle progression, paralleling the cell cycle length increase seen predominantly in primate and especially human neurodevelopment. Our findings establish HARs as key components in rewiring human-specific neurodevelopmental gene regulatory programs and provide an integrated resource to study enhancer activity of specific HARs.

INTRODUCTION

Rapid evolutionary expansion of the human cerebral cortex is considered a vital step that enabled human-specific cognitive function. The increased size of the human brain is associated with a greatly prolonged period of corticogenesis (from about 7 days in mice to more than 130 days in humans) and increased capacity for neural progenitor cell (NPC) proliferation before neuronal differentiation (Lui et al., 2011; Pollen et al., 2015). Paradoxically, the NPC cell cycle is also lengthened in humans. Mouse

NPC cell cycle length ranges from 8–26.5 h, largely by variation of G1 phase (Arai et al., 2011; Takahashi et al., 1995), but NPC cycles in embryonic non-human primate can be up to 8-fold longer at 22–63 h (Betizeau et al., 2013; Kornack and Rakic, 1998), and NPCs in humans show a similarly long cell cycle (Pollen et al., 2015; Sub-ramanian et al., 2017). This lengthening of the cell cycle and neurogenic interval inevitably forces a substantially slowed overall pace of neurological development, generally referred to as a form of heterochrony, or neoteny (Geschwind and Rakic, 2013), that has dramatic implications for maternal and social behavior.

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Although the genetic basis underlying the evolutionary changes affecting the human brain remains largely unknown (Dehay et al., 2015; Lewitus et al., 2014; Lui et al., 2011; Namba and Huttner, 2017; Sousa et al., 2017), recent comparative genomic studies have identified over 3,100 DNA sequences with potential human-specific functions that have been hypothesized to play a role in this process. These regions, collectively termed human accelerated regions (HARs), are the fastest-evolving regions of the human genome (Bird et al., 2007; Bush and Lahn, 2008; Gittelman et al., 2015; Hubisz and Pollard, 2014; Lindblad-Toh et al., 2011; Pollard et al., 2006; Prabhakar et al., 2008). HARs are defined as genomic elements that are highly conserved across non-human mammals, suggesting that they perform essential functions, but also show rapid sequence changes selectively in the human lineage. HARs are mostly noncoding and are enriched for transcription factor binding motifs, suggesting that many HARs potentially function as human-specific regulatory elements that coordinate precise spatiotemporal patterns of gene expression (Capra et al., 2013; Hubisz and Pollard, 2014). Furthermore, prior studies have uncovered some HARs that have putative enhancer activity in human neuronal tissue, with the enhancer activity of some HARs differing from that of their chimpanzee orthologs (Boyd et al., 2015; Capra et al., 2013; Franchini and Pollard, 2017; Kamm et al., 2013; Uebbing et al., 2021), raising the theory that some HARs may contribute to human-specific brain development.

Because the particular pattern of nucleotide substitution that defines a HAR can arise by many mechanisms—including loss of function, biased gene conversion, or evolutionary drift (Pollard et al., 2006)—a critical challenge that remains is to define which HAR sequences represent true drivers of *bona fide* evolutionary changes. Prior studies using massively parallel reporter assays (MPRAs) have attempted to address this problem (Uebbing et al., 2021) but have been limited in terms of the number of HARs assessed as well as the inherent size limitations of synthetic oligo-based MPRAs. For example, synthetic oligo-based MPRAs are often limited to interrogating intact elements that are smaller than 150 bp in length (Uebbing et al., 2021), which represents only a small fraction of all HARs (Figure 1A).

To systematically identify the subset of HARs most likely to be contributing to recent evolution of the human cerebral cortex, we analyzed the in vitro and in vivo gene regulatory activity of over 3,100 HARs across multiple human and mouse cell types and tissues. To overcome the size limitations of standard MPRAs, we developed and applied an approach called CaptureMPRA, which leverages barcoded molecular inversion probes (MIPs) to capture target sequences of ~500 bp in length, enabling us to interrogate entire HAR elements and their surrounding DNA on individual probes. We interrogated humanchimpanzee differences in HAR enhancer function using this CaptureMPRA approach and integrated these data with in vivo histone marks and chromatin accessibility at HARs in human fetal neural cells to identify HARs with putative regulatory function in guiding human-specific brain development. Finally, we developed an easily searchable online resource (the HARHub) consisting of these data and published datasets of common and rare human HAR sequence variations (Doan et al., 2016) to serve as a resource for the scientific community. Our find-

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ings define many HARs that play key roles in neuronal gene regulatory programs, with nearly half of all HARs showing reproducible chromatin accessibility and enhancer activity in neural cells and tissue.

RESULTS

HAR elements demonstrate pervasive neuronal enhancer activity *in vitro*

To globally assess the regulatory potential of HARs in human neurodevelopment in a high-throughput manner, we first developed a new MPRA to which we refer as CaptureMPRA (caM-PRA). CaMPRA leverages barcoded MIPs to capture target sequences, in this case HARs, and then simultaneously evaluates enhancer activity at these sequences in a single *in vitro* experiment (Figure 1A). By using MIPs to capture target sequences of ~500 bp in length (Data S1), caMPRA overcomes many of the length restrictions associated with the standard MPRA design, enabling systematic evaluation of HARs, which are, on average, 269 bp in length. We then applied the caMPRA approach to all 3,171 reported HARs (Bird et al., 2007; Bush and Lahn, 2008; Gittelman et al., 2015; Hubisz and Pollard, 2014; Lindblad-Toh et al., 2011; Pollard et al., 2006; Prabhakar et al., 2008) as well as their paired chimpanzee orthologs.

Using 10,690 barcoded MIPs targeting flanking regions surrounding 3,171 HARs (Data S1) or tiling HARs more than 500 bp in length, we successfully captured 3,132 HARs and their paired chimpanzee orthologs from human and chimpanzee genomic DNA, respectively. To test the enhancer activity of these elements, we used two well-described neural cell lines, one human (SH-SY5Y) and one mouse (Neuro2A/N2A), to model human as well as non-human neural developmental activity in vitro (Figure S1). Overall, 49% of the elements demonstrated reproducible and significant enhancer activity in human neural cells (Figure 1B; Data S2), demonstrating the pervasive neural precursor-like enhancer activity of these HARs. Furthermore, HARs with enhancer activity were enriched for binding motifs of transcriptional regulators known to play vital roles in cortical neurogenesis (RBPJ, TBR2 [EOMES], PAX6, ATOH1, and TEAD) (Figure S2A), suggesting that many HARs act as neurodevelopmental enhancers.

Human-specific rewiring of the *cis*-regulatory architecture of HARs

To identify whether human-specific sequence divergence at HAR elements is associated with altered neural enhancer activity, we compared the enhancer activity of HAR elements with their chimpanzee orthologs (Figure 1B). Overall, 61% of the HARs with caMPRA enhancer activity showed significant alterations in enhancer activity between the human and chimpanzee orthologs. Of note, human-specific sequence divergence appears to be more commonly associated with increased enhancer activity exhibiting greater activity for the human than chimpanzee ortholog (Figure 1C). This indicates extensive evolutionary rewiring of the *cis*-regulatory activity of HAR elements in the human lineage.

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A Capture Massively Parallel Reporter Assay (caMPRA)



Figure 1. Rewiring of the cis-regulatory enhancer activity of HAR elements in the human lineage

(A) Schematic describing the caMPRA.

(B) caMPRA activity of all HARs and their chimpanzee orthologs in human neural cells. The p value was calculated using a two-sided paired t test. (C) Bar graph showing the number of HAR elements with increased or reduced activity of the human versus chimpanzee ortholog. *p < 0.001 (2-proportions z test).

(D) caMPRA activity of all HARs and their chimpanzee orthologs in human neural cells versus mouse neural cells. The p value was calculated using a two-sided paired t test.

(E) Bar graph showing the percentage of HAR elements with altered activity of the human versus chimpanzee ortholog versus the percentage of HAR elements with altered activity in human versus mouse neural cells. *p < 0.001 (2-proportions z test).

To disentangle the contribution of *cis*- versus *trans*-regulatory effects on rewiring of HAR element enhancer activity, we also compared the enhancer activity of each element between the human (SH-SY5Y) and mouse (Neuro2a/N2A) neural cell lines (Figure 1D). This revealed that over 94% of elements showed similar enhancer activity in human and mouse cells, indicating that differences in the neural *trans*-regulatory compartment between human and mouse do not significantly alter the overall enhancer activity of each HAR element. These findings demonstrate that the human-specific rewiring of HAR element enhancer activity is predominantly driven by *cis*-regulatory variation (Figure 1E).

HARs demonstrate *in vivo* enhancer features during human brain development

To determine whether HAR elements demonstrate features of *in vivo* enhancer activity in the brain, we profiled the chromatin state at HAR elements across diverse developmental cell types and tissues using DNasel hypersensitivity mapping (DNasel-seq) as well as chromatin immunoprecipitation sequencing (ChIP-seq) for enhancer-associated chromatin marks. Using fetal and adult DNasel-seq data collected as part of the Roadmap Epigenomics Consortium (Kundaje et al., 2015), we observed that 78% of HARs



Figure 2. HARs are enriched for human neurodevelopmental enhancer elements

(A) Genomic loci demonstrating tissue-specific DNasel-seq chromatin accessibility at three separate HAR elements.

(B) Top: the proportion of HAR elements that demonstrate DNasel hypersensitivity within 79 diverse human tissues based on whether the element is DNasel hypersensitive in brain tissues only, brain tissues plus other tissues, or non-brain tissues only. Bottom: for HAR elements that are DNasel hypersensitive in brain tissues only, shown is a Venn diagram of the number of elements that are DNasel hypersensitive in fetal versus adult brain tissue.

(C) Bar graphs displaying (top) the number of HAR elements overlapping DHSs in each tissue as well as (bottom) the total number of DHSs in each tissue. Fetal tissues are labeled in blue, whereas adult tissues are labeled in brown. *p < 0.001 (2-proportions z test).

(D) Schematic of fluorescence-activated nuclear sorting (FANS) of human fetal brain tissue.

(E and F) Genomic loci and density plots demonstrating FANS tissue-specific (E) DNasel-seq chromatin accessibility and (F) H3K4me1 and H3K27ac ChIP-seq signals at HAR elements.

(G) Venn diagram demonstrating the overlap of HARs with H3K4me1 or H3K27ac signal in NPCs or neurons.

(H) Bar plots showing the percentage of NPC and neuron DHSs that overlap with H3K4me1 or H3K27ac ChIP-seq peaks based on whether the DHS is also a HAR. *p < 0.001 (2-proportions z test).

demonstrated chromatin accessibility in at least one of the human cell or tissue types tested, with 56% demonstrating chromatin accessibility in human brain tissue (Figures 2A and 2B; Data S3). HAR elements were preferentially accessible in fetal as opposed to adult human brain tissue (Figures 2C and S2B), further implicating HARs as neurodevelopmental regulatory elements involved in human brain development. To test the potential for HARs to behave as cell-type-specific neurodevelopmental enhancers, we isolated NPCs and maturing neurons from human fetal brain tissue (gestational week [GW] 19–20) using fluorescence-activated nuclear sorting (FANS) and probed for chromatin features associated with enhancer activity (i.e., H3K4me1, H3K27ac, and chromatin accessibility) (Figures 2D and S3A; Creyghton et al., 2010). Specifically, nuclei from GW 19–20 human frontal cortex were isolated and labeled



Figure 3. PPP1R17 is a putative HAR-regulated neurodevelopmental gene

(A) Integration of caMPRA activity, chromatin accessibility, histone modification datasets, and long-range chromatin interaction data defines 63 HARs as likely contributors to human brain evolution.

(B) hg19 HiC and DNasel-seq data at the locus surrounding HAR2635, which is one of the HARs identified in (A). HAR2635-interacting loci are indicated by the green box and black arrows.

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using PAX6, SOX2, and NeuN fluorescently tagged antibodies. Labeled nuclei were subsequently sorted to selectively purify NPCs (SOX2⁺, PAX6⁺, NeuN⁻) and maturing neurons (SOX2⁻, PAX6⁻, NeuN⁺), and these purified populations were then subjected to ChIP-seq and DNaseI-seq. 504 HARs displayed chromatin features consistent with *in vivo* enhancer activity in GW19–GW20 NPCs or maturing neurons (as evidenced by overlap with H3K4me1 and/or H3K27ac ChIP-seq peaks) (Figure 2G). HAR elements overlapping DNasel hypersensitivity sites (DHSs) were significantly enriched for these enhancer marks compared with other DHSs in NPCs or neurons (Figure 2H). This further demonstrates the importance of HARs as active enhancer elements during human brain development.

The majority of H3K4me1 and H3K27ac ChIP-seq peaks were unique to NPCs or maturing neurons (Figures 2E and 2F; Data S4 and S5), indicating a substantial change in the regulatory DNA landscape during neuronal differentiation, with only 37% of H3K4me1- or H3K27ac-marked HARs overlapping these chromatin marks in NPCs and neurons (Figure 2G). Although NPCselective enhancer-marked HARs were enriched for transcription factor (TF) elements central for cerebral cortical patterning and NPC maintenance, such as LHX family proteins (Hébert and Fishell, 2008; Mangale et al., 2008; Monuki et al., 2001), neuron-selective enhancer-marked HARs were enriched for TF elements implicated in cell fate determination and cell differentiation, such as RFX proteins (Figures S3B and S3C). These findings implicate a substantial portion of HARs as regulatory elements involved in human-specific neuronal development.

PPP1R17 is a putative HAR-regulated neurodevelopmental gene

We next integrated our caMPRA, DNasel-seq, and ChIP-seq HAR data to identify a core set of HARs representing likely neurodevelopmental enhancers that might be contributing to the rapid evolutionary expansion of the human cerebral cortex. Overall, this approach identified 210 HARs with *in vitro* and *in vivo* features consistent with neurodevelopmental enhancer activity (Figure 3A; Data S6). Overlapping this set of 210 putative neurodevelopmental enhancer HARs with chromatin conformation data (HiC) from human fetal cortex (Won et al., 2016) revealed that these 210 HARs were significantly enriched for having longrange chromatin interactions with target genes in the fetal cortex (Figure S4A), providing further evidence of the *in vivo* enhancer activity of these HARs in human neurodevelopment.

In total, 63 HARs demonstrated neurodevelopmental enhancer activity in addition to long-range chromatin interactions with a target gene in the human fetal cortex (Figure 3A). Of these, the interaction between HAR2635 and *PPP1R17* stood out because the *PPP1R17* topologically associated domain (TAD) contains a *PPP1R17* promoter-proximal pri-

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mate-selective regulatory element as well as two HARs with strong fetal brain long-range chromatin interactions with the PPP1R17 gene promoter (Figures 3B-3E; Figure S4). This suggests extensive rewiring of PPP1R17 expression during nonhuman primate and human evolution. Targeted chromatin confirmation capture (3C) interaction analysis confirmed the selective interaction of the promoter of PPP1R17 with HAR2635 and HAR2636 in cultured neural cells (Figure 3C), suggesting that these two sequences likely play a role in regulation of PPP1R17 neural expression. In contrast, the nearby gene NEU-ROD6, which is highly expressed in postmitotic cortical neurons but not in progenitor cells (Bormuth et al., 2013; Han et al., 2018), does not show a strong long-range chromatin interaction with these HARs. Furthermore, although little is known about PPP1R17, it is highly expressed in NPCs in the developing human cortex (Noctor et al., 2004; Pollen et al., 2015; Zhong et al., 2018). It is also a reported repressor of protein phosphatase complexes 1 (PP1) and 2A (PP2A) (Endo, 2012), which are critical for the $G1 \rightarrow S$ transition during the cell cycle, a key step regulating NPC cycle length (Wlodarchak and Xing, 2016). This prompted us to further examine the pattern of expression of PPP1R17 in neurodevelopment as well as in other cell types across non-human primate and human evolution.

Primate-specific gain of *PPP1R17* expression in the developing cerebral cortex

To define the cell type specificity and regional distribution of PPP1R17 expression across mammalian evolution, we examined PPP1R17 expression in different regions of the developing brain in mouse, ferret, rhesus macaque, and human, comparing its expression patterns between lissencephalic mammals, gyrencephalic mammals, and primates. PPP1R17 is highly expressed in adult cerebellar Purkinje cells in all four of these mammals (Figures 4A and 4B) in a pattern that appears to be highly evolutionarily conserved. In contrast to this conserved pattern of cerebellar expression, PPP1R17 shows highly divergent expression in the developing cerebral cortex, where it is highly expressed in primates but not detected in ferret or mice at comparable ages (Figures 4C and S5A). PPP1R17 expression in the prenatal macague and human cortex is localized to the outer and inner subventricular zones (oSVZ and iSVZ, respectively) (Figure 4C) and is excluded from the ventricular zone and cortical plate. PPP1R17 expression co-localizes with TBR2, a marker of intermediate progenitor cells, and KI67, a marker of dividing cells (Figures 4C and S5B). In contrast, PPP1R17 is not expressed in the neocortex in mice or ferrets during development (Figure 4C). This gain of expression in the cortical SVZ in primates parallels several mouse-human differences in chromatin accessibility surrounding the PPP1R17 gene, including the gain

⁽C) Chromatin confirmation capture (3C) signal in SH-SY5Y human neural cells demonstrating long-range chromatin contacts between the *PPP1R17* promoter and HAR2635 and HAR3636. Data are maximum signal normalized.

⁽D) Genomic locus of HAR2635 showing evolutionary conservation and the DNasel-seq signal and single-cell ATAC-seq signal across multiple human fetal tissues and adult astrocytes.

⁽E) Genomic locus of the HAR2635-interacting gene *PPP1R17* showing the human fetal brain DNasel-seq signal and single-cell ATAC-seq signal in addition to evolutionary conserved elements and the DNasel-seq signal of contiguous sequence in the mouse fetal brain. A red box indicates the primate-selective regulatory element. Contiguous sequence elements between human and mouse are indicated by gray boxes.

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Figure 4. Primate-specific gain of PPP1R17 expression in the developing cortex

(A) Diagram of stained brain regions.

(B) Staining of human, macaque, ferret, and mouse cerebellum for PPP1R17 (green) and the Purkinje cell marker Calbindin (pink) demonstrates conserved cerebellar expression of *PPP1R17* across these four mammals.

(legend continued on next page)



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of a primate specific element as well as altered fetal brain chromatin accessibility at several surrounding conserved elements (Figure 3E).

Human-specific divergence in *PPP1R17* cell type distribution

In addition to primate-non-primate changes in early cortical expression, comparison of RNA sequencing (RNA-seq) data (Zhu et al., 2018) on different regions of the macaque and human brains at various stages of fetal and adult life revealed additional, striking human-macaque differences in PPP1R17 expression (Figures 5A and S6A), suggesting ongoing changes in expression within primates that we confirmed with immunofluorescence analysis (Figure 5B). Specifically, RNA-seq analysis showed highly concordant PPP1R17 expression in adult cerebellum and in fetal cortical NPCs between macaque and humans (Figures 5A and S6A). However, developing cortical expression of PPP1R17 was restricted to fetal development in humans and closely matched the time course of expression of TBR2. In contrast, in macaques, cortical expression of PPP1R17 was maintained well after neurogenesis ceased and even well into adulthood and, hence, diverged from the pattern of TBR2 expression (Figures 5A and S6A). The late cortical expression of PPP1R17 in adult macaques is driven not by expression in NPCs or neurons but, rather, by expression of PPP1R17 in cortical astrocytes (Figure 5B), indicating a further divergence in the cell type expression pattern of PPP1R17 in the primate lineage. These results demonstrate a primate-specific innovation in PPP1R17 expression in NPCs, especially intermediate progenitor cells, which coincides with an insertion of a new regulatory element into the promoter-proximal region of PPP1R17. Further changes in the cell type and developmental expression patterns of PPP1R17 between the macaque and human cortex also coincide with sequence changes at two HAR elements that connect to PPP1R17.

PPP1R17 regulates NPC cycle progression

To assess the putative role of *PPP1R17* in cell cycle regulation, we evaluated the effect of *PPP1R17* misexpression on mouse primary cortical neurospheres. Neurospheres electroporated with a *PPP1R17* construct were 36% smaller than those electroporated with a *GFP* construct (2-way ANOVA, p = 0.0003), with no increase in cell death (Figure 6A), implicating *PPP1R17* as a regulator of NPC proliferation. To test whether this effect was at least partially mediated through the G1 \rightarrow S transition, we next misexpressed *PPP1R17* in mouse neural cells that genetically encoded a G1 phase cell cycle indicator (Sakaue-Sawano et al., 2008). *PPP1R17*-misexpressing cells divided at half the speed of control cells (*PPP1R17* cells, 68.2 h; *GFP*, 32.7 h, p < 0.0001) and demonstrated a prolonged G1 phase in comparison with *GFP* controls (48.5% *PPP1R17* versus 33.8% GFP, p < 0.0001) (Figures 6B and S6B). These results implicate

PPP1R17 as a regulator of the G1 \rightarrow S transition and NPC cell cycle progression and indicate a potential role of PPP1R17 in regulating NPC cycle progression. These findings parallel the increase in cell-cycle length, neurogenesis duration, and heterochrony or slowed development seen predominately in the primate and human lineages.

An integrated resource for interrogating HAR function in humans

We collated the data generated in this paper into an easily searchable online resource through the UCSC Genome Browser (the HARHub: http://genome.ucsc.edu/cgi-bin/hgTracks?db= hg38&hubUrl=https://allendiscoverycenter-harhub.s3.us-east-2. amazonaws.com/HAR_hub/hub.txt; Figure S7). In addition to the caMPRA and epigenomic datasets (DNasel hypersensitivity, H3K27ac, and H3K4me1) generated here, this resource also contains previously published datasets of common and rare human HAR sequence variations obtained through targeted sequencing of HARs in the Homozygosity Mapping Collaborative for Autism (HMCA) (Doan et al., 2016). In addition, we also included traditional oligo-based MPRA results on a subset of HARs (Uebbing et al., 2021). This integrated HAR enhancer hub is a comprehensive resource for guiding future investigations of the function of HARs in the evolution of the human brain.

DISCUSSION

By interrogating the epigenetic landscape of more than 3,100 HARs during human neurodevelopment, we demonstrate that nearly half of all HARs act as neurodevelopmental enhancers and that the cis-regulatory architecture of these HARs has undergone extensive human-specific rewiring. Moreover, we demonstrate that human-specific sequence changes within HARs largely augment their enhancer activity within neuronal cells, further establishing HARs as human-specific neurodevelopmental enhancers. By leveraging caMPRA to evaluate the enhancer activity of entire intact elements for more than 90% of HARs, our study is the most thorough evaluation of HAR elements to date. This is in contrast with traditional oligo-based MPRAs, which are often limited to insert sizes of less than 137 bp in length (Uebbing et al., 2021), which represents less than 20% of intact HAR elements because the average HAR element is 269 bp in length.

The striking conservation within the *trans*-regulatory activity of HARs mirrors prior studies (Stergachis et al., 2014) and demonstrates that our assessment of enhancer activity using mouse and human neural cell lines is likely capturing the majority of HARs that act as neurodevelopmental enhancers. Furthermore, by combining our caMPRA data with chromatin data from the developing fetal brain, we establish that a substantial portion of HARs act as *in vitro* and *in vivo* enhancers and that, overall, HARs are enriched for being enhancers in the developing human

⁽C) Staining of human, macaque, ferret, and mouse developing cortex with PPP1R17 and SOX2 demonstrates cortical expression of PPP1R17 selectively within the two primate species, which appears to be localized to the outer and inner subventricular cortical germinal zones. VZ, ventricular zone; iSVZ, inner subventricular zone; OFL, outer fiber layer; SP, subplate; CP, cortical plate.

⁽D and E) Staining of the human fetal cortex with PPP1R17, SOX2, and TBR2, demonstrating (D) PPP1R17 expression localized to the subventricular zones and (E) PPP1R17, SOX2, and TBR2 colocalization.

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Figure 5. Human-specific divergence in PPP1R17 cell type distribution

(A) Top: Diagram of brain regions evaluated by RNA-seq at various developmental time stages in human and macaque brain development from fetal to adult. Bottom: *PPP1R17* expression in the cerebellum and cortex in macaques and humans as a function of developmental age demonstrates persistent expression of *PPP1R17* selectively in macaques well into adulthood.

(B) Staining of the human and macaque adult neocortex for PPP1R17 (green) and the astrocyte marker GFAP (purple) demonstrates predominant expression of PPP1R17 in cortical astrocytes in adult macaques.

brain. This firmly establishes that a subset of HARs represent genomic entities driving human-specific neurodevelopmental gene regulatory programs. In addition to their neural enhancer activity, many HARs also demonstrate evidence of enhancer activity in multiple other tissue types, indicating that HARs may also play a role in modulating human-specific gene regulatory programs for other organ systems. Of note, 11% of HARs lacked any evidence of *in vitro* or *in vivo* enhancer activity in any of the analyzed cell types (Data S3 and S6), indicating that these HARs may have gene regulatory element-independent functions, if they are functional at all.

Using our neurodevelopmental HAR enhancer data, we identified *PPP1R17* as a putative HAR-regulated gene that appears to play a central role within the primate cortical germinal zones. *PPP1R17* has undergone extensive rewiring of its cell type and developmental expression patterns in the primate and human cortex, which appears to be potentially mediated by HARs as well as a primate-selective regulatory element, although further studies evaluating the mechanistic role of these elements in *PPP1R17* gene regulation are likely needed. The primate-selective regulatory element directly upstream of *PPP1R17* encompasses several recently evolved retrotransposons that have been proposed as the most common mechanism for generating primate-specific regulatory elements (Jacques et al., 2013).

PPP1R17 cortical expression is restricted to progenitor cells within the oSVZ and iSVZ in the developing cerebral cortex of humans and macaques. These germinal zones are the primary neurogenic niches in human cortical development and have undergone a dramatic expansion in size during primate evolution (Lui et al., 2011; Reillo et al., 2011) because of augmentation of the transit-amplifying capacity of NPCs. Specifically, although rodent NPCs usually divide once (Haubensak et al., 2004; de la Torre-Ubieta et al., 2016; Noctor et al., 2004) or only a few times (Mihalas and Hevner, 2018) to produce two postmitotic neurons, human NPCs go through numerous additional rounds of symmetric division before terminal differentiation (Hansen et al., 2010; Pollen et al., 2015; Sessa et al., 2008). Our findings suggest that PPP1R17 may play a complex role not only in shortening the cell cycle but also in augmenting transit amplification and delaying neuronal differentiation, a process central to the increase in neuronal production in primate and human corticogenesis (Bystron et al., 2008). Although ferrets have a longer cell cycle than mice (Turrero García et al., 2016), we do not find PPP1R17 expression in their NPCs, suggesting that although PPP1R17 expression is sufficient to elongate the cell cycle, other mechanisms may also contribute.

Although humans and macaques demonstrate a gain of *PPP1R17* expression within cortical NPCs, humans and macaques differ in their expression of *PPP1R17* within the astrocyte lineage. Specifically, although macaques demonstrate robust *PPP1R17* expression in astrocytes throughout development, humans appear to lack significant astrocyte *PPP1R17* expression postnatally. In addition, humans also lack chromatin accessibility at HAR2635 within astrocytes (Figure 3D), suggesting that this HAR may play a role in guiding the cell specificity of *PPP1R17* expression in humans. It remains unclear whether this difference in astrocyte *PPP1R17* expression has a phenotypic effect because our understanding of

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Figure 6. PPP1R17 regulates cell proliferation and G1 phase in vitro

(A) Mouse primary neurosphere growth assay using cells transfected with a *GFP* or *PPP1R17* construct exposes a growth restriction for *PPP1R17*-expressing neurospheres, which were 36% smaller than GFP controls (2-way ANOVA, p = 0.0003). Error bars, SEM. Individual values represent wells containing neurospheres imaged at each time point; n = 48 GFP- and 37 PPP1R17-transfected wells.

(B) Cell cycle assay using Neuro2A cells with a G1 cell cycle marker in addition to a *GFP* or *PPP1R17* construct demonstrates (center) a longer doubling time (*PPP1R17* cells, 68.2 h; GFP, 32.7 h; p < 0.0001) and (right) a prolonged G1 phase for *PPP1R17*-expressing Neuro2A cells (48.5% for *PPP1R17* versus 33.8% for GFP, p < 0.0001, 2-way ANOVA). The total number of cells and percentage of cells expressing the G1 marker and GFP or *PPP1R17*-GFP were quantitated at 24 h and 48 h from 4 images per well for each well containing transfected Neuro2A cells (n = 13 GFP-transfected wells and 13 *PPP1R17*-transfected wells). Error bars, SEM.

differences in astrocyte neurobiology between primates is still in its infancy.

Although we show that PPP1R17 slows NPC cycle progression, and although the lengthening of NPC cycles is a well-documented feature of primates with larger brains, the evolutionary origin and presumed advantage of the remarkably extended neurogenetic period in primates remains enigmatic. Longer cell cycles are thought to reduce the rate of somatic mutation or maintain DNA quality control and may facilitate formation of the remarkable diversity of neuronal types that constitute the human brain (Boldog et al., 2018; Hodge et al., 2019), but they come at the potential cost of increasing heterochrony, a longer period of infantile helplessness, and the requirement for longer maternal care and changes in societal structure, suggesting complex and balancing forces. The deepening investigation of HARs and other evolutionarily dynamic genomic sequences may increasingly shed light on these complex evolutionary changes underlying distinct human brain cytoarchitecture and cognitive capacity.

STAR*METHODS

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SUPPLEMENTAL INFORMATION

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AUTHOR CONTRIBUTIONS

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DECLARATION OF INTERESTS

The authors declare no competing interests.

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Zhu, Y., Sousa, A.M.M., Gao, T., Skarica, M., Li, M., Santpere, G., Esteller-Cucala, P., Juan, D., Ferrández-Peral, L., Gulden, F.O., et al. (2018). Spatiotemporal transcriptomic divergence across human and macaque brain development. Science *362*, eaat8077.

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STAR*METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
PerCP-Cy5.5-conjugated anti-PAX6	BD Biosciences	Cat#561462, Clone O18-1330; RRID: AB_10715442
V450-conjugated anti-SOX2	BD Biosciences	Cat#561469, Clone O30-678; RRID: AB_10694256
Alexa Fluor 488-conjugated anti-NeuN	EMD Millipore	Product No. FCMAB317PE, Clone A60; RRID: AB_10807694
Anti-H3K27ac	Abcam	Prod#ab4729; RRID: AB_2118291
Anti-H3K4me1	Abcam	Prod#ab176877; RRID: AB_2637011
Goat anti-SOX2 (1:200)	Santa Cruz	Cat#sc-170320
Chicken anti-TBR2 (1:200)	Millipore	Cat#AB15894; RRID: AB_10615604
Rabbit anti-TBR2 (1:200)	Abcam	Prod#ab23345; RRID: AB_778267
Rabbit anti-PAX6 (1:200)	Covance	Cat#PRB-278P; RRID: AB_291612
Mouse anti-HOPX (1:200)	Santa Cruz	Cat#sc-30216; RRID: AB_2120833
Rabbit anti-PPP1R17 (1:200)	Novus	Cat#NBP2-13800
Mouse anti-PPP1R17 (1:200)	Novus	Cat#H00010842-B01P; RRID: AB_2067445
Mouse anti-SATB2 (1:200)	Abcam	Prod#ab51502; RRID: AB_882455
Rabbit anti-Ki67 (1:200)	Abcam	Prod#ab15580; RRID: AB_443209
Mouse anti-PCNA	Millipore	Cat#AB93501; RRID: AB_93501
Mouse anti-GFAP	Sigma	Cat#G3893; RRID: AB_477010
Rat anti-GFAP	Thermo Fisher	Cat#13-0300; RRID: AB_2532994
Alexa Fluor 488 Anti-Chicken Secondary Antibody	Thermo Fisher	Cat#A11039; RRID: AB_2534096
Alexa Fluor 594 Anti-Mouse Secondary Antibody	Thermo Fisher	Cat#A11032; RRID: AB_2534091
Alexa Fluor 594 Anti-Rabbit Secondary Antibody	Thermo Fisher	Cat#A11012; RRID: AB_2534079
Alexa Fluor 647 Anti-Rat Secondary Antibody	Thermo Fisher	Cat#A21247; RRID: AB_141778
DAPI (4',6-Diamidino-2-Phenylindole, Dihydrochloride)	Thermo Fisher	Cat#D1306; RRID: AB_2629482
Hoechst 33342	Thermo Fisher	Cat#62249
Normal Goat Serum	GIBCO	Cat#16210064-100ml
Bovine Serum Albumin	Sigma	Cat#A7906-100G
Fluoromount-G	Southern Biotech	Cat#0100-01
Bacterial and virus strains		
One Shot TOP10 Chemically Competent E. coli	Thermo Fisher	Cat#C404006
Biological samples		
Human brain tissue	This paper	N/A
Chemicals, peptides, and recombinant proteins		
Formaldehyde	Fisher Scientific	Cat#F79P-4
Glycine	Sigma	Cat#G7126-5KG
Triton X-100	Sigma	Cat#T9284-500ML
Paraformaldehyde	Electron Microscopy Sciences	Cat#15710
Fetal bovine serum	Atlanta Bio	Cat#S11150
Dulbecco's Modified Eagle Medium	Thermo Fisher	Cat#MT10013CV

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Continued		
REAGENT or RESOURCE	SOURCE	IDENTIFIER
Dulbecco's Modified Eagle Medium/F12	Thermo Fisher	Cat#11330032
Penicillin-Streptomycin	Thermo Fisher	Cat#15140122
AMPure XP beads	Beckman-Coulter	Cat#A63881
Retrievagen A	BD Biosciences	Cat#550524
Lysis Buffer	This paper	N/A
Wash buffer	This paper	N/A
Sonication buffer	This paper	N/A
AsiSI restriction enzyme	NEB	Cat#R0630
DNasel	Sigma-Aldrich	Cat#D4527
Mbol	NEB	Cat#R0147
T4 DNA Ligase	Thermo Fisher	Cat#EL0011
Critical commercial assays		
Lipofectamine LTX reagent	Thermo Fisher	Cat#A12621
TRIzol reagent	Invitrogen	Cat#15596018
NEBNext Single Cell/Low Input RNA	New England BioLabs	Cat#E6420S
Library Prep Kit for Illumina	U U	
Protein A Dynabeads	Thermo Fisher	Cat#10002D
Ovation Ultralow V2 kit	Nugen	Cat#0344-32
Neural Tissue Dissociation Kit (T)	Miltenyi Biotec	Cat#130-093-231
Deposited data		
Raw and analyzed data	This paper	GEO: GSE180714
Human reference genome build GRCh38/hg38	Genome Reference Consortium	https://www.ncbi.nlm.nih.gov/assembly/ GCF_000001405.26/
Human, chimpanzee, and macaque brain transcriptome data	Zhu et al., 2018	N/A
ENCODE Project and Roadmap Epigenomics Consortium	ENCODE Project Consortium, 2012; Kundaje et al., 2015	http://www.roadmapepigenomics.org/
UCSC-tools and genome browser	Kent et al., 2002	http://genome.ucsc.edu/
Pathway and gene set overdispersion analysis (PAGODA)	Fan et al., 2016	N/A
Single cell ATAC-seq data	Domcke et al., 2020	https://descartes.brotmanbaty.org/
Experimental models: Cell lines		-
Neuro2A	ATCC	Cat#CCL-131
SH-SY5Y cell	ATCC	Cat#CRL-2266
Experimental models: Organisms/strains		
CD1 Mice	Charles River	Crl:CD1(ICB)
Ferrets	Marshall Bioresources	N/A
Macaques	Yale University School	N/A
Oligonucleotides		
	CustomArray Rotholl M/A	https://www.customarravinc.com/
	Cusionianay, Botheli, WA	oligos_main.htm
Primers for 3C interaction assay, see Data S6	I his paper	N/A
Recombinant DNA		
Human genomic DNA	Promega	Cat#G1521
Chimpanzee DNA	Coriell	Cat#NS03489
pMPRA1 vector	Melnikov et al., 2014	Addgene 49349

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Continued		
REAGENT or RESOURCE	SOURCE	IDENTIFIER
pMPRA_donor2	Melnikov et al., 2014	Addgene 49353
pFucci-G1 Orange	Medical & Biological Laboratories Co., LTD.	Cat#AM-V9003M
PBCAG-eGFP	Gift from Joseph LoTurco (Chen and LoTurco, 2012)	Addgene 40973
pCyl50	Gift from Jae Yong Han (Lee et al., 2016)	N/A
pCAG-PB-GFP	This paper	N/A
pCAG-PPP1R17-PB-GFP	This paper	N/A
Software and algorithms		
MIPgen	Shendure Lab, University of Washington	http://shendurelab.github.io/MIPGEN/
R language	R Core Team, 2019 The R Project for Statistical Computing	https://www.r-project.org/
Python language, version 2.7	Python Software Foundation	https://www.python.org/
GraphPad Prism 8	GraphPad Software Inc., La Jolla, CA, USA	https://www.graphpad.com/scientific- software/prism/
Fiji	Schindelin et al., 2012	https://imagej.net/software/imagej
ImageJ	Schneider et al., 2012	https://imagej.net/software/imagej
IncuCyte Zoom	Essen Bioscience	IncuCyte S3 2018B
HOMER	Heinz et al., 2010	http://homer.ucsd.edu/homer/
Other		
HARHub	This paper	http://genome.ucsc.edu/cgi-bin/hgTracks? db=hg38&hubUrl=https://allendiscoverycenter- harhub.s3.us-east-2.amazonaws.com/ HAR_hub/hub.txt

RESOURCE AVAILABILITY

Lead contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Christopher A. Walsh (christopher.walsh@childrens.harvard.edu).

Materials availability

All unique/stable reagents generated in this study are available from the Lead Contact with a completed Materials Transfer Agreement.

Data and code availability

- HAR data have been deposited at GEO accession GSE180714. Processed ChIP-seq and DNasel-seq and HAR data are available at http://genome.ucsc.edu/cgi-bin/hgTracks?db=hg38&hubUrl=https://allendiscoverycenter-harhub.s3.us-east-2. amazonaws.com/HAR_hub/hub.txt. Accession numbers are listed in the key resources table. All the data is available from the lead contact upon request.
- All the code is available from the lead contact upon request.
- Any additional information required to reanalyze the data reported in this paper is available

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Cell lines

Neuro2A/N2A cells (ATCC, CCL-131) were grown in 10% fetal bovine serum, Dulbecco's Modified Eagle Medium, and 1% Penicillin-Streptomycin. SH-SY5Y cell (ATCC, CRL-2266) medium was 10% fetal bovine serum, Dulbecco's Modified Eagle Medium/F12, and 1% Penicillin-Streptomycin. Both cell lines were maintained in a 5% CO2 incubator at 37°C.

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Mice

All procedures were reviewed and approved by the Institutional Animal Care and Use Committee (IACUC) at Boston Children's Hospital (BCH). Timed pregnant CD1 mice (*Mus musculus*) were obtained from Charles River Laboratories in advance of in utero electroporation procedures after approximately 9-11 days of gestation. Mice were housed before and after procedures at BCH and maintained according to BCH and NIH animal research guidelines.

Ferrets

Wild-type ferrets (*Mustela putorius furo*) were obtained from Marshall Bioresources, with both males and females included in this study. No association between animal sex and findings was found. All animal work conducted as part of this study was in compliance with the NIH Principles for the Use of Animals, the NIH Guide for the Care and Use of Laboratory Animals, and the Animal Welfare Act, and all work was reviewed and approved by the Institutional Animal Care and Use Committee (IACUC) at Boston Children's Hospital (BCH).

Human and non-human primate tissue specimens and processing

Research performed on samples of human origin was conducted according to protocols approved under expedited category 5 with waiver of consent (45 CFR 46.110) by the institutional review boards of Beth Israel Deaconess Medical Center, Boston Children's Hospital, and Yale University School of Medicine. Fetal brain tissue was received after release from clinical pathology, with a maximum post-mortem interval of 4 h. Cases with known anomalies were excluded. Gestational ages (weeks 19, 20, and 23) were determined using fetal foot length. One subject was female, and the sex of other subjects was unknown. Tissue was transported in HBSS medium on ice to the laboratory for research processing.

Rhesus macaque (*Macaca mulatta*) brain samples were collected postmortem from both adult (11 years-old) and fetal (E70) specimens. All experiments using non-human primates were carried out in accordance with a protocol approved by Yale University's Committee on Animal Research and NIH guidelines.

METHOD DETAILS

CaptureMPRA library design, cloning, and transfection

Molecular Inversion Probes (MIPs) targeting flanking regions of all HAR sequences were designed using the MIPgen program (Boyle et al., 2014) and manufactured as a pool by CustomArray (Bothell, WA). Target genomic sequences from both human (Promega, G1521) and chimpanzee DNA (Coriell, NS03489) were captured as previously described (Cantsilieris et al., 2017) and purified using Agencourt AMPure XP beads at 0.9X concentration (Beckman-Coulter, A63881). The MPRA assay then was conducted as previously described (Melnikov et al., 2014). Briefly, the captures were cloned into modified pMPRA1 vector (Addgene, 49349). The only modification to the pMPRA1 vector was the addition of an AsiSI restriction site used for inserting our captured DNA fragments. The luciferase gene and minimal promoter (pMPRA_donor2, Addgene, 49353) was then cloned into the pMPRA1_Capture construct pool.

The transfection of the MPRA libraries was performed with five replicates per condition, and the experiment was conducted in triplicate. The cells (N2A and SH-SY5Y) were grown as adherent cultures for 24hr in 15 cm plates. Then, 20ug of MPRA constructs and 4ug of a fluorescent reporter construct per plate were transfected into cells with Lipofectamine LTX reagent (Thermo Fisher, A12621). The cells were checked for fluorescent reporter expression at 24hr and harvested 48hr post-transfection. mRNA was isolated using TRIzol reagent (Invitrogen, 15596018). Resulting Tag-seq amplified cDNA was deep sequenced using the Illumina Hi-Seq 2500 with 150-bp paired-end reads.

Fluorescence-Activated Nuclear Sorting (FANS)

Nuclear preparation and sorting were performed as previously described (Lodato et al., 2015; Thomsen et al., 2016). Nuclei were labeled in aliquots of 500 µL with antibodies ((PerCP-Cy5.5-conjugated anti-PAX6 (O18-1330; BD Biosciences), V450-conjugated anti-SOX2 (O30-678; BD Biosciences), Alexa Fluor 488-conjugated anti-NeuN (A60; EMD Millipore)) for 30 min at 4°C before FANS. Pools of 750,000 to 1,000,000 nuclei were collected per condition for downstream sequencing experiments unless otherwise noted.

RNA-Sequencing of FANS nuclei

1000 nuclei from each cell population were sorted in triplicate into a 96 well plate and cDNA libraries were immediately prepared using NEBNext Single Cell/Low Input RNA Library Prep Kit for Illumina according to manufacturer's instructions at 0.5X volume. Sorted nuclei were subjected to 18 cycles of PCR amplification following the reverse transcription step. Quality of amplified cDNA was assessed using Agilent 2100 Bioanalyzer (Agilent Technologies) Illumina-compatible cDNA libraries were subsequently prepared according to manufacturer's instructions at 0.5X volume with 20 ng of full-length amplified cDNA and 8 cycles of PCR enrichment. Prior to sequencing, library quality was assessed using the Agilent 2100 Bioanalyzer. 150 PE reads were generated using the Illumina HiSeqX Ten and analyzed as described below.

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Chromatin Immunoprecipitation (ChIP)

For ChIP of histone modifications, sorted nuclei were cross-linked via addition of 1% formaldehyde for 10 min at room temperature and quenched by the addition of 0.125 M glycine for 5 min at room temperature. Nuclei were lysed by 10 minute incubation in lysis buffer 1 (in mM: 100 HEPES-NaOH pH 7.5, 280 NaCl, 2 EDTA, 2 EGTA, 0.5% Triton X-100, 1% NP-40, 20% Glycerol) followed by washing in buffer containing 10 mM Tris-HCl pH 8.0, 200 mM NaCl. Chromatin was sheared using a Bioruptor (Diagenode) on high power mode for 50 cycles with 30 s pulses in sonication buffer (in mM: 10 Tris-HCl pH 8.0, 100 NaCl, 1 EDTA, 0.5 EGTA, 0.1% Na-Deoxycholate, 0.5% N-Lauroylsarcosine). For H3K27ac ChIP, 10 mM sodium butyrate (Fisher NC9851678) was added to all buffers.

Following sonication, chromatin was supplemented with 1% Triton and was incubated overnight with the following antibodies coupled to Protein A Dynabeads (Thermo Fisher, 10002D): H3K27ac (Abcam, 4729), H3K4me1 (Abcam, 176877). ChIPs were performed in three biological replicates for each cell population. Libraries were generated using the Ovation Ultralow V2 kit (Nugen, 0344-32) according to the manufacturer's instructions and PCR amplified for 13-16 cycles, depending on antibody. Library quality was assessed using the Agilent 2100 Bioanalyzer (Agilent Technologies). Seventy-five bp reads were generated on the Illumina Next-seq 500 and subsequently analyzed with our standardized ChIP-seq data analysis pipeline (below).

Chromatin Conformation Capture (3C)

3C PCR primers (Data S7) were designed according to a previous 3C method (Naumova et al., 2012). To obtain an interaction profile of the genomic region spanning HAR2635, HAR2636, and the PPP1R17 promoter, we designed several test primers at and around each HAR and a single anchor primer within the promoter region of PPP1R17.

3C targeting the PPP1R17 locus was performed in SH-SY5Y cells. Cells were cross-linked in 1% methanol-free formaldehyde for 10 minutes at room temperature (RT) followed by addition of glycine to a final concentration of 125 mM. Cells were incubated for 20 minutes in lysis buffer (10 mM Tris-HCl pH 8, 10 mM NaCl, 0.2% Triton X-100) on ice, followed by dounce homogenization. Nuclei were isolated by centrifugation and treated with 0.3% SDS followed by addition of 2% Triton X-100 for 1 hour at 37C each. Cross-linked chromatin was digested with 100U of Mbol restriction enzyme (NEB, R0147) at 37C overnight, followed by ligation under dilute conditions (2 ng/uL) using 75U of T4 DNA ligase (Thermo Fisher, EL0011) for 4 hours at 16C followed by 30 minutes at RT. Chromatin was reverse cross-linked via addition of Proteinase K. Purified 3C templates were amplified under standard PCR conditions. Quantification of interaction products were analyzed via agarose gel using ImageJ (Schneider et al., 2012), and relative interaction frequency was calculated from the relative intensity of each band.

Immunohistochemistry

Mouse brain tissue was fixed overnight at 4°C in 4% PFA. Fixed macaque and mouse tissue were sectioned at 50 um using a Vibratome. Tissue was permeabilized and blocked in 3% BSA and 0.3% Triton X-100 in PBS for mouse and ferret tissue, and 5% Normal Donkey Serum with 1% BSA for macaque and human tissue. Primary antibodies were diluted in blocking buffer and incubated overnight 1-2 nights at 4°C. Primary antibodies included goat anti-SOX2 (1:200, Santa Cruz sc-170320 and 1:150 R+D Systems AF2018), chicken (1:200; Millipore AB15894) and rabbit (1:200; abcam ab23345) anti-TBR2 (1:500 Abcam ab183991), rabbit anti-PAX6 (1:200; Covance PRB-278P, 1:200 Biolegend 901301, mouse anti-HOPX (1:200; Santa Cruz sc-30216), rabbit (1:200; Novus NBP2-13800) and mouse (1:200; Novus H00010842-B01P) anti-PPP1R17, mouse anti-SATB2 (1:200; abcam ab51502), rabbit anti-Ki67 (1:200; abcam ab15580, and 1:200 Invitrogen 14-5698-2), rabbit anti-TBR1 (1:500, abcam ab31940), and mouse anti-PCNA (1:200; Millipore AB93501). Sections were then stained with Alexa secondary antibodies and DAPI. Ferret and human brain sections were processed as previously described (Johnson et al., 2015, 2018). Imaging was performed using a Zeiss LSM 700 confocal microscope.

In Utero electroporation (IUE)

In utero electroporations were performed as described previously (Saito et al., 2018; Szczurkowska et al., 2016; Zhang et al., 2016) using pregnant mouse dams (embryonic age 12.5-16.5) under a Boston Children's Hospital Institutional Animal Care and Use Committee (IACUC)-approved protocol. Embryos were injected with approximately 2μ L of a 1μ g/ μ L mixture of plasmid DNA, delivered using a pulled glass micropipette inserted into the lateral ventricle of the brain. Subsequently, 50-V electric pulses (5 total, 1.1 s interval, 50ms duration) were passed through the brain over each injection site using paddle electrodes. Embryos were allowed to continue developing normally for 2-4 days after electroporation.

Plasmid cloning

Control (pCAG-PB-GFP) and human *PPP1R17* (pCAG-PPP1R17-PB-GFP) expression plasmids were cloned from PBCAG-eGFP (Addgene 49073), a generous gift from Joseph LoTurco (Chen and LoTurco, 2012). A P2A linker was inserted downstream of the eGFP coding region, and for the *PPP1R17*-expression plasmid, a 430bp region from the human exon 3 sequence was inserted, up-stream of the rabbit beta-globin polyadenylation signal and the *piggyBac 3'* terminal repeat sequence. Sequences were verified by Sanger sequencing.

Primary neurosphere growth assay

First, in utero electroporations were performed as described above with mice aged E12.5 and a mixture of control (pCAG-PB-GFP) or PPP1R17 (pCAG-PPP1R17-PB-GFP) plasmid and the PiggyBac transposase system to allow for genomic integration (Wu et al.,

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2006). Embryos were harvested after 48 hours in utero. The cortex of the embryos then were minced and dissociated with using the Miltenyi Biotec Neural Tissue Dissociation Kit, and the top 10% of GFP+ cells were sorted using Fluorescence Activated Cell Sorting (FACS). Cells were plated at 500 cells/well on a 96-well u-bottomed plate. Neurospheres were grown for 6-10 days *in vitro*, and images were collected by the IncuCyte imaging system (Essen Instruments, Ann Arbor, MI, USA) every 2 hours. Analysis was subsequently performed using IncuCyte Zoom software (Essen Bioscience) and GraphPad Prism (GraphPad Software Inc., La Jolla, CA, USA) from n = 48 GFP control and n = 37 *PPP1R17*-GFP wells containing neurospheres.

QUANTIFICATION AND STATISTICAL ANALYSIS

CaptureMPRA data analysis

To identify HAR barcodes, the sequence immediately preceding the adaptor sequence was isolated, and the occurrence of each barcode counted. To compare to the reference 10,690 barcodes, each barcode's Levenstein distance was calculated with respect to all reference barcodes using the stringdist package, and the minimum value was taken for each barcode. Multiple probes were summarized by taking the mean of each probe. Data was normalized to account for effective logFC differences across uneven library sizes by using the Trimmed Means of M normalization as implemented in the edgeR package. The RNA matrix was created by dividing each sample's RNA value by the respective plasmid DNA. The gplots package is used to create the heatmap on the log2 data. HARs with significant enhancer activity were identified using a one-sided one-sample t test of the three biological replicates to identify HARs with log2 mRNA/DNA > 0. This t test was performed independently using the human and chimpanzee orthologs as well as the data from N2A and SH-SY5Y cells, and those with a p value < 0.05 in at least one of these samples were considered to have significant enhancer activity. False discovery rates were also identified by shuffling the MPRA data between HARs and samples and recomputing the p values, to identify the proportion of enhancers identified at a given p value threshold that may be the result of a falsely identified enhancer. A two-sided paired t test was used to identify HARs with significantly different log2(mRNA/DNA) ratios between the human and chimpanzee orthologs.

Transcription factor binding motif analyses

Transcription factor binding motifs for all HAR sequences were performed using HOMER (Heinz et al., 2010) with the basic configurations: database version v4.4, significance p < 0.01. Enrichment of motifs was determined using options: randomly generated sequences and HAR input regions. ConTra v3 (Kreft et al., 2017) was used for visualization of TFBS changes across species with stringency: core = 0.95, similarity matrix = 0.85.

RNA Sequencing, alignment and DGE Analysis

FASTA and Homo_sapiens.GRCh38.93.gtf annotation files were retrieved from Ensembl and STAR version 2.5.2b was used to generate a genomic index from hg38 genome release 93 (-sjdbOverhang 149). Using default settings in STAR, bam files were generated for each sample independently and a gene count matrix was created using featureCounts (-p –s0) for paired-end, unstranded RNA-seq data. Raw gene counts for technical replicates were then summed together prior to normalization of biological replicates using the Median by Ratio method implemented by DESeq2 (version 1.20.0). Boxplots for genes of interest were generated with ggplot2 using gene counts after normalization of the Median by Ratio method. Normalized biological replicates were then analyzed to discover differentially expressed genes using DESeq2. Minimum requirements for a DGE was defined as a gene possessing ab-s(log2FoldChange) > 0.58 and a *P*-value < 0.05 after adjusting P values using the Benjamini-Hochberg method.

DHS resources and analysis

Human sorted nuclei were subjected to DNasel digestion and high-throughput sequencing, following previous methods (John et al., 2011). Additional DHS datasets were downloaded and analyzed from the ENCODE Project and Roadmap Epigenomics Consortiums (ENCODE Project Consortium, 2012; Kundaje et al., 2015). The number and proportion of all HARs that fell within DNase-I-sensitive regions (Hotspots) and DHSs (150 bp peaks) were identified as were calculated as previously described (John et al., 2011). In addition, the maximum DNasel density over each HAR was calculated using normalized DNasel density track sets and bedops (Neph et al., 2012).

ChIP Sequencing, alignment, and genome browser track generation

All experiments were sequenced on the Nextseq 500 (Illumina). Seventy-five base pair single-end reads were obtained for all ChIPseq datasets. All samples were aligned to the hg38 genome using default parameters for the Subread alignment software (subread-1.4.6-p3; Liao et al., 2013) after quality trimming with Trimmomatic v0.33 (Bolger et al., 2014) with the following command: java -jar trimmomatic-0.33.jar SE -threads 1 -phred33 [FASTQ_FILE] ILLUMINACLIP:[ADAPTER_FILE]:2:30:10 LEADING:5 TRAILING:5 SLIDINGWINDOW:4:20 MINLEN:45. Truseq adapters were trimmed out in ChIP-seq experiments.

To generate UCSC genome browser tracks for ChIP-seq data, all aligned bam files for each replicate of a given experiment were pooled and converted to BED format with bedtools bamtobed. The 75 base pair reads were extended in the 3' direction to 200 base-pairs (bp) (average fragment length for ChIP-seq experiments as measured by bioAnalyzer) with the bedtools slop command using the following parameters: -I 0 -r 125 -s. Published hg38 ChIP-seq blacklisted regions (ENCODE Project Consortium, 2012) were

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filtered out using the following command: bedops -not-element-of 1 [BLACKLIST_BED]. The filtered BED files were converted to coverageBED format using the bedtools genomecov command with the following options: -scale [NORM_FACTOR to scale each library to 20M reads] -bg. Finally, bedGraphToBigWig (UCSC-tools) was used to generate the bigWIG files displayed on browser tracks throughout the manuscript.

Hi-C data was visualized using the 3D Genome Browser (Wang et al., 2018). Specifically, both Fetal_Brain_GZ/Won2016-raw and IMR90/Lieberman-raw were visualized at 10kb resolution with a scale of 0 to 90.

ChIP-seq peak calling

To identify sites with enriched ChIP-seq signal (peaks), we applied the IDR pipeline (Li et al., 2011) using the MACS2 peak calling algorithm (Zhang et al., 2008) with the following parameters:-nomodel-extsize 200-keep-dup all. An IDR threshold of 0.01 was used for self-consistency, true replicate, and pooled-consistency analyses. The 'optThresh' cutoff was then used to obtain a final set of high-confidence, reproducible ChIP-seq peaks for each factor or chromatin modification at each stimulus condition tested.

ChIP-Seq peaks across all conditions and ChIP-ed factors were unioned to generate the comprehensive set of putative regulatory elements assessed in this study using the following command: bedops-everything [INPUT PEAKS] | bedtools sort -i - | bedtools merge -i - -c 4,5 -o collapse, collapse > [OUTPUT FILE]. This peak set was then intersected with HARs using bedtools intersect.

Gene ontology analysis

For HARs overlapping ChIP peaks, bedtools (bedtools closest) was used to determine the closest gene. Functional enrichment with gene ontology was implemented with the goseq package, setting an Benjamini-Hochberg FDR of 0.05. Fold change matrices were calculated using the gtools implementation of fold change.

Human single cell RNA-sequencing and ATAC-seq

Pseudotime course analysis of previously published RNA-seq data was executed using single-cell differential expression analysis (Kharchenko et al., 2014) and pathway and gene set overdispersion analysis (PAGODA) (Fan et al., 2016). Single cell analyses for Figure S6A were performed using previously published data from human neocortical development during mid-gestation (Polioudakis et al., 2019). Single cell ATAC-seq data for Figure 3 was obtained from https://descartes.brotmanbaty.org/ (Domcke et al., 2020). The track labeled "Glial cells" in Figure 3E corresponds to the "Astrocyte" cell population in Domcke et al. (2020). Similarly, the track labeled "Intermediate Progenitors" corresponds to the "Cerebrum Unknown.3" cell population in Domcke et al. (2020) based on the selective chromatin accessibility of several intermediate progenitor cell markers in this cluster, including *EOMES*.

Cell cycle assay

To obtain doubling time and quantification of time spent in G1, Neuro2A cells were seeded at low density (300-1000 cells/well) in a 96well plate format after chemical transfection by Lipofectamine with two plasmids: pFucci-G1 Orange (Sakaue-Sawano et al., 2008), and either the control (CAG-GFP) or *PPP1R17* misexpression vector (CAG-PPP1R17-GFP). Cells were imaged every 1-4 hours for 2-4 days at 20x magnification using the IncuCyte imaging system (Essen Instruments, Ann Arbor, MI, USA). Analysis of time-lapsed images was performed using 4 regions of interest per well (n = 13 wells each for GFP control and *PPP1R17*-GFP transfected cells) using IncuCyte Zoom software (Essen Bioscience) and GraphPad Prism (GraphPad Software Inc., La Jolla, CA, USA). Numbers of cells expressing both pFucci-G1 Orange (channel), and GFP control or *PPP1R17*-expression plasmids were quantified at each time point recorded. Doubling time was calculated using cells that were GFP-positive in both conditions.