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Epigenomic programming in early fetal brain development

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Aim: To provide a comprehensive understanding of gene regulatory networks in the developing human brain and a foundation for interpreting pathogenic deregulation. Materials & methods: We generated reference epigenomes and transcriptomes of dissected brain regions and primary neural progenitor cells (NPCs) derived from cortical and ganglionic eminence tissues of four normal human fetuses. Results: Integration of these data across developmental stages revealed a directional increase in active regulatory states, transcription factor activities and gene transcription with developmental stage. Consistent with differences in their biology, NPCs derived from cortical and ganglionic eminence regions contained common, region specific, and gestational week specific regulatory states. Conclusion: We provide a high-resolution regulatory network for NPCs from different brain regions as a comprehensive reference for future studies.

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Keywords: brain • cortex • DNA methylation • enhancer • epigenetics • fetal • ganglionic eminence • gestational week • neural progenitor cells • transcriptional network

Transcription is the driver of cell differentiation and specification involving multilateral interactions between epigenetic modifications and transcription factors. While significant progress has been made in defining transcriptional programs within distinct brain cell types [1-5], we still lack comprehensive knowledge of how these programs are shaped by mitotically inherited epigenetic states during brain development. Current models, based primarily on studies of the murine hematopoietic system, posit that specific histone epigenetic states are ‘primed’ in early development and subsequently engaged by the transcriptional machinery following cellular division and differentiation. In vitro models suggest that epigenetic priming and stage-specific transcription factor engagement may also occur during neuronal differentiation [6], and studies on CpG methylation during in vitro differentiation and in cortical tissue during fetal brain development revealed developmental-specific methylation states in both human and mouse [7-9]. An in vitro study on enhancer and transcriptome landscapes during early fetal cortex development using human-induced pluripotent stem cell-derived cortical organoids also revealed a major role of enhancer landscape in regulating radial glial cell growth and cortical neuron specification [10].

The early fetal period, gestational week 8 (GW8) to mid-gestation (GW18), is a critical period in neocortex development involving extensive neurogenesis and cellular migration [11]. This period is transcriptionally dynamic within and across different brain regions [5,12], and is associated with a directional loss of DNA methylation [13] and dynamic changes in enhancer gene regulation with gestational week [10]. However, our current understanding of
the dynamic DNA methylation states associated with fetal brain development are based on reduced representation techniques that have focused on CpG-rich regions within gene promoters leaving intergenic regulatory regions largely unexplored [14]. Furthermore, the spatiotemporal landscape of histone modifications during fetal brain development and their interplay with DNA methylation states remains to be defined.

Primary neural progenitor cells (NPCs) can be enriched from dissected fetal brain tissue and propagated as neurospheres in culture [15]. NPCs are heterogeneous populations of progenitor cells at different stages of pluripotency, including neuroepithelial cells, radial glial cells and other intermediate progenitors. During the neurogenesis period, NPCs originating from the germinative area including ganglionic eminence (GE) migrate to the cortical layers and NPCs originating from outer cortical layers are characterized by a more differentiated neuronal phenotype compared with GE-derived NPCs [16–18]. Indeed, NPCs derived from different brain regions have distinct cellular compositions based on their level of specification and differentiation [19–21]. However, the regulatory and transcriptional states that drive NPC development in different brain regions during early fetal development is largely unknown.

To address these gaps, we analyzed the transcriptional and epigenetic state of brain tissue and matched NPCs from four human fetuses. From these data we construct regulatory network models for primary human NPCs and the brain regions from which they were derived at three developmental stages. Comparisons of these models revealed that NPCs derived from different fetal brain regions and at different developmental stages are defined by distinct regulatory states. NPCs derived from different brain regions retain epigenetic signatures specific to these regions that are consistent with the differentiation stages of the cells. Consistent with in vitro models NPCs derived from later developmental stages are characterized by a directional increase in active regulatory transcriptional states. Lastly, transcriptional networks derived from fetal brain-derived NPC from monozygotic (MZ) twins display distinct regulatory states as early as midgestation. Our data provide the first comprehensive mapping of the epigenomes and transcriptomes of human fetal brain cells, and a foundation for further understanding of brain development and diseases.

**Material & methods**

**Human tissue samples & cell culture**

Human brain tissues were obtained from fetuses at GW13, GW15 and GW17 after informed consent. Gestational week was calculated from the first day of the last menstrual period. Mothers and the fetuses were healthy at the time of sample acquisition. Fetuses and tissue samples were assessed at the time of sample acquisition and were deemed to be phenotypically normal. As far as we are aware, the mothers did not suffer any complications during pregnancy. Samples were identified through unlinked codes in accordance with the federal Health Insurance Portability and Accountability Act guidelines. Specific brain regions were carefully dissected based on landmark and immediately frozen in liquid nitrogen or processed for cell culture. Cortical and GE regions were clearly identified and confirmed by HE section on adjacent tissue. Cells were extracted by gentle tissue dissociation with the Neural Tissue Dissociation kit with papain (Miltenyi Biotec Inc., CA, USA) and were plated at low density in ultra-low attachment cell culture flasks (Corning, NY, USA) in neural stem cell media (STEMCELL Technologies Inc., BC, Canada) supplemented with EGF (20 ng/ml), bFGF (10 ng/ml) and heparin (2 ug/ml). Cells growing as neurospheres were fed weekly by adding fresh media and passaged every 2 weeks by gentle manual dissociation. After 2–3 passages, neurospheres were harvest and frozen in liquid nitrogen.

**Whole-genome bisulphite sequencing**

Genomic DNA from NPCs were extracted, bisulphite treated and sequenced as previously described [22] (see also Supplemental Experimental Procedures). Sequence reads were directionally aligned to the human genome (GRCh37-lite) as described [14]. Differentially methylated regions (DMRs) were identified from differential methylation analysis using a C++ tool with p < 0.005 [23] and dynamic growth approach to concatenate adjacent differential methylated CpGs within 500 bp. Regions with less than three CpGs were filtered out. On average, DMRs are approximately 250 bp across all samples.

**Methylated DNA immunoprecipitation sequencing**

Methylated DNA immunoprecipitation sequencing (MeDIP-seq) was performed and fractional methylation calls extracted (see also Supplemental Experimental Procedures). Differential methylated CpGs were identified with fractional methylation call differences greater than 0.6 and hypermethylation greater than 0.75. DMRs were
concatenated from differential methylation CpGs with the same dynamic growth approach used for whole-genome bisulfite sequencing (WGBS). Regions with less than four CpGs were filtered. On average, MeDIP DMRs are approximately 100 bp across all samples.

**Methylation-sensitive restriction enzyme sequencing**

Methylation-sensitive restriction enzyme sequencing (MRE-seq) was performed as previously described [22] (see also Supplemental Experimental Procedures), and the signals were calculated for DMRs identified with WGBS for validation.

**Chromatin immunoprecipitation sequencing**

Chromatin immunoprecipitation sequencing (ChIP-seq) for H3K4me1, H3K4me3, H3K9me3, H3K27me3 and H3K36me3 were performed and raw reads aligned as described [22]. Enriched regions were identified with FindER v0.9.3b (www.epigenomes.ca/finder.html) (see also Supplemental Experimental Procedures). Differentially marked promoters were identified by calculating ChIP-seq signals for promoter regions (TSS +/− 1.5 kb), ranking promoters for all 52418 protein-coding and noncoding genes based on their signal level. Promoters with > 5000 differences in rank in pairwise comparisons were identified as differentially marked.

**RNA & miRNA sequencing**

RNA-seq and miRNA-seq were performed and raw reads aligned to a custom transcriptome (build from the National Center for Biotechnology Information GRCh37-lite reference and Ensembl v65 (GenCode v10) annotations), and reads per kilobase of transcript per million reads mapped (RPKM) calculated for genes and exons (see also Supplemental Experimental Procedures). Differentially expressed genes were identified by an in-house matlab tool as previously described [22] using entropy-based thresholds with false discovery rate (FDR) = 0.01. Isoforms were identified by employing the same differential expression analysis on exons and exon RPKM cutoff to identify exons expressed (>10% gene RPKM) in one sample but not the other (<1%).

**Single nucleotide variance identification**

Single nucleotide variants (SNVs) were called using SAMtools mpileup with −C50 to reduce the effect of reads with excessive mismatches, and BCFtools with −D100 to exclude sites with read depth greater than 100 for cortex and GE NPCs from all four subjects [24]. SNPs not included in dbSNP build 138 were filtered out. SNVs were compared between each of the six pairs among the four subjects to obtain discordant variants in NPCs. The overlap of the discordant variants from the two-cell types were taken as the discordant SNVs between the pair.

**Transcriptional regulation analysis**

GREAT (McLean et al.) and DAVID (Huang et al.) were employed to study biological processes and pathways associated with regulatory regions and genes using default parameters [57,61]. To study the transcription factor activities in regulatory regions, Homer was used to identify enriched transcription factors with q-value < 0.01 and percent of regions with motif > 20% [25]. Transcription factor target genes were identified with Homer annotated transcription factor binding site (TFBS) within a genomic window encompassing the transcriptional start site (TSS) +/− 1.5 kb. Transcriptional regulation network was visualized using Cytoscape [26].

**Results**

**Epigenome profiling of human fetal brain**

To explore the gene regulatory landscape of the developing human brain, dissected brain tissue and primary neurospheres derived from cortex and GE were obtained from four phenotypically normal human fetuses at GW13, GW15 and GW17 (Figure 1). Immunohistochemistry analysis of primary neurospheres cultures showed cell-type specificity with cortex-derived neurospheres displaying an increased neuronal phenotype compared with GE-derived neurospheres (Figure 2 & Supplementary Figure 1). To confirm the progenitor status of the neurosphere cultures, they were differentiated into neuroblasts (DCX+, CALRETININ+, TUJ1+) and astrocytes (GFAP+) (Figure 2 & Supplementary Figure 1). Massively parallel sequencing based assays were employed to generate ChIP-seq, WGBS, MeDIP-seq, MRE-seq and mRNA-sequencing (RNA-seq) datasets from the dissected primary brain tissue and the neurospheres derived from cortex and GE following the International Human Epigenome Consortium guidelines (Figure 1B & Supplementary Table 1) [27]. To validate our findings, we used an independent
cohort of 23 fetal brain samples ranging from GW8 to GW17 with Illumina 450K DNA methylation profiles [28], and 12 of these samples were also profiled through RNA-seq (European Genome-phenome Archive Accession Number: EGAS00001003368).

To establish genetic relationships between twin fetuses we called SNVs from RNA-seq reads [29] and performed pair-wise comparisons across subjects. On average, 11,227 discordant SNVs were identified across all pairs, except the GW17 twins for which only 1784 were found (Supplementary Figure 1 & Supplementary Table 2). Interestingly, discordant SNVs identified between the GW17 twins were homozygous in one twin and heterozygous in the other, hence can be either the result of genetic differences or the result of allele-specific expression. To distinguish these possibilities, we examined discordant SNVs using MeDIP-seq reads and found that all examined discordant sites were heterozygous across all three tissue and cell types, suggesting that the discordance observed in RNA-seq at
Figure 2. Characterization of fetal brain-derived neural progenitor cells. (A) Cells isolated from the cortex or GE are able to proliferate when cultured as neurospheres and maintain a stem/progenitor phenotype, confirmed by the expression of SOX2 and OLIG2. (B) When grown in presence of growth factors, NPCs express a mixture of stem cell markers (NESTIN+, SOX2+, OLIG2+), glial progenitors (OLIG2+, A2B5+), astrocytes (GFAP+) and neuroblasts (MAP5+) with cortex-derived NPCs presenting a more neuronal phenotype (MASH1 high, MAP5 high) while the GE-derived NPCs have a more glial phenotype (A2B5 high, GFAP more differentiated). (C) In 2% serum, NPCs are differentiate into a neuronal lineage, from an immature neuroblast (MAP2+, MAP5+) to a migrating neurons (Doublecortin DCX+, TUJ1+) to a postmitotic neuron (CALRETININ+), as well as into GFAP+ astrocytes.

FBS: Fetal bovine serum; GE: Ganglionic eminence; GW: Gestational week; NPC: Neural progenitor cell.

these positions was due to allele specific expression. We concluded that the GW17 twins (referred to as Subject1 and Subject2) were genetically identical and thus MZ.

NPCs derived from different fetal brain regions & at different developmental stages are epigenetically distinct

To identify common regulatory states of NPCs, we defined enhancer regions within the genomes of cortex and GE-derived NPCs as regions marked by H3K4me1 [30]. We selected H3K4me1 as it provides a broader view of both primed and active enhancer states and is used as standardized enhancer mark by the NIH Roadmap Epigenomics Mapping Consortium [14]. This analysis identified 39,033 core enhancers shared across all NPCs regardless of development stage or brain region from which they were derived (Supplementary Table 3). Intersecting NPC core enhancers with TFBSs predicted by Homer [25] revealed 13 transcription factors significantly enriched (q < 0.01 and > 20% enhancers with motif) (Supplementary Figure 2), including the master regulators of pluripotency NANOG and SOX3, as well as key regulators of brain development such as LHX3, a transcription factor involved in the specification of motor neurons and interneurons [31], and OLIG2, implicated in the early stages of oligodendrocyte
specification and maturation [32–37]. This core set of enhancers is predicted to play a role in defining the regulatory state of NPCs within the developing fetal brain (Supplementary Figure 2).

We next sought to establish whether cortical and GE tissue-derived NPCs retained distinct regulatory states following the in vitro expansion associated with their generation. We first identified DNA methylation signatures of cortex and GE NPCs derived at each developmental stage independently. We identified DMRs genome wide (see methods) between cortex and GE-derived NPCs by pairwise comparisons of WGBS datasets. DMRs were validated using MeDIP-seq and MRE-seq datasets [38,39] generated from matching NPCs (Supplementary Figure 3). On average, 1412 DMRs (average length 327 bp) covering 20,008 CpGs were identified between cortex and GE-derived NPCs, with 646 DMRs identified at GW13 and 2178 in GW17 suggesting increased divergence in DNA methylation with developmental age (Figure 3A & Supplementary Table 4). Directionality was also observed between NPCs with 3.5-fold more DMRs hypomethylated in cortex-derived NPCs compared with GE-derived NPCs. Consistent with differences in cell migration and fates between cortex and GE-derived NPCs [40], genes
associated with cortex-specific hypomethylated DMRs were found to be enriched in forebrain regionalization, while those associated with GE-specific hypomethylated DMRs were enriched in neuronal migration and differentiation (Figure 3B). These findings suggest that NPCs derived from distinct brain regions retain the epigenetic signatures specific to these regions.

To examine the transcriptional output of NPCs alone and in association with regulatory states, we generated RNA-seq libraries (Supplementary Table 1). Pairwise comparisons (see methods) identified 382 cortex NPC-specific and 456 GE NPC-specific genes differentially expressed in at least two subjects (hypergeometric p < 10^-26; Supplementary Table 5). Consistent with an increased divergence in DNA methylation with gestational week, NPCs derived from GW17 exhibited nearly twice the number of differential expressed genes as GW13 (Supplementary Figure 2). In addition, exon level differential expression analysis also identified twice as many putative gene isoforms in GW17 as in GW13, suggesting more divergent transcriptomes between cortex and GE NPCs in the later developmental age (Supplementary Figure 2). Associating differential expressed protein coding genes with H3K4me3 promoter density (TSS +/- 1.5kb; see methods; Supplementary Table 6) identified 159 differentially expressed genes with gains/losses in H3K4me3 promoter density, a majority of which (74.8%) showed expected orientation, in other words, gain of H3K4me3 with upregulation and vice versa. Gains and losses in H3K27me3 promoter density were associated with 508 differentially expressed genes with a majority (75.4%) in the expected orientation, in other words, gain of H3K27me3 with downregulation and vice versa.

Functional analysis of genes differentially expressed in both GW13 and GW17 between cortex and GE-derived NPCs revealed enrichment in genes implicated in neurogenesis and cell morphogenesis regulation (Supplementary Figure 2), including GFAP (Figure 3C), an astrocyte marker associated with brain disorders including gliomas [41–43]; NFIX (Figure 3C), an active transcription factor essential for the development of a number of organ systems including brain [44,45]; FEZF1, a transcription repressor involved in the axonal projection and proper termination of olfactory sensory neurons [46]; OTX1 and OTX2, essential in patterning the developing brain [47]; and VAX1, involved in forebrain and visual system development [48].

Taken together, our analysis between cortex and GE-derived NPCs supports a model of distinct epigenetic and transcriptional states in fetal brain regions as early as GW13 that become increasingly divergent with developmental age. We also provide evidence that NPCs derived from distinct brain regions retain region of origin signatures when cultured as neurospheres.

NPCs undergo epigenetic & transcriptional activation during early development

The classical view of epigenetic patterning during development posits a model whereby cellular differentiation is accompanied by epigenetic restriction [49]. To explore whether evidence of such epigenetic restriction could be found within the genomes of cells separated by 4 weeks of fetal brain development, hypomethylated DMRs between GW13 and GW17 were identified in cortex and GE-derived NPCs respectively (Figure 4A & Supplementary Table 4). In agreement with a previously reported developmentally associated genome hypomethylation in grossly dissected human fetal brain tissue [13], we observed 3.4-fold increase in the number of CpGs within hypomethylated DMRs in GW17 NPCs compared with GW13 (Figure 4A). Functional enrichment analysis of genes associated with GW17 hypomethylated DMRs revealed enrichment in cell fate commitment and brain cell differentiation (Supplementary Figure 4). Consistent with the enhancer activation in early cortical development [10], GW17-specific hypomethylated CpGs within DMRs were significantly enriched in core NPC enhancers (hypergeometric p < 10^-26), while GW13-specific DMRs showed no such enrichment (Figure 4B), suggesting that a loss of CpG methylation at active enhancers follows the establishment of histone mediated enhancer states during brain development.

To validate this observation, we examined the DNA methylation states of fetal brain regions across developmental stages in an independent cohort generated by 450K array (Supplementary Table 1). Consistent with our observation of a directional loss of methylation with development stages, we found an increase in the number of probes that were hypomethylated in GW17 (difference in beta value >0.15) compared with those that were hypomethylated in any of the GW13 individuals (Supplementary Figure 5). Furthermore, 241 out of the 667 probes within hypomethylated regions identified by WGBS were also hypomethylated in 450K array, a 12-fold enrichment of all hypomethylated probes identified in 450K array. Genes associated with these 241 probes were also highly enriched in neurogenesis and neuron development (Supplementary Figure 5), suggesting that the 450K array data were able to validate our findings and that WGBS provided a more comprehensive measurement of DNA methylation.
Figure 4 Temporal changes revealed a more active epigenetic state in the later developmental stages. (A) DMRs between GW13 and GW17. We identified more hypomethylated DMRs in GW17 (blue) than GW13 (red) in both GE-derived NPCs (inner circles) and cortex-derived NPCs (outer circles). (B) Fold enrichment of DMRs between GW13 and GW17 in core NPC enhancers. GW13 hypomethylated DMRs, shown in red, were not enriched in the enhancers, while GW17 hypomethylated DMRs, shown in blue, were enriched. (C) Transcription factors exclusively enriched (Benjamini corrected $p < 0.01$, left panel and percent of enhancers with motif $>20\%$) in GW17 unique enhancers, and their transcription levels in cortex-derived NPCs (middle panel) and GE-derived NPCs (right panel). GW13 expression values are shown in red, and GW17 are shown in blue. Out of the five transcription factors, OLIG2, FOXO1, and AR were expressed. (D) Methylation differences between GW17 and GW13 in enhancers with the particular transcription factor binding sites showed hypomethylation in GW17 (left panel). The vertical line, indicating hypomethylation greater than 0.2, was used as a cut-off for hypomethylated enhancers. Fold change of transcriptional levels between GW17 and GW13 for the nearest genes of the hypomethylated enhancers identified in the left panel showed upregulation in GW17 (right panel). The vertical line indicates twofold increase in expression.

DMR: Differentially methylated region; FC: Fold change; GE: Ganglionic eminence; NPC: Neural progenitor cell; RPKM: Reads per kilobase of transcript, per million mapped reads; TF: Transcription factor.
However, only 47% of DMRs identified from WGBS are probed in 450K array and of them 66% have less than three probes within the region, leaving the majority of the enhancer landscape unmeasured.

We next explored gene promoters (TSS+/−1.5 kb) differentially marked by H3K4me3 and H3K27me3 between GW13 and GW17. H3K4me3 differentially marked the promoters of 526 protein-coding genes with 2.6-fold more genes gaining H3K4me3 in GW17 compared with GW13. We observed the opposite trend in H3K27me3 with a sixfold decrease in H3K27me3-marked gene promoters in GW17 compared with GW13 (Supplementary Table 6), consistent with a more active promoter histone modification landscape in GW17. A subset of the genes associated with the differentially marked promoters were also differentially expressed with a majority (73%) in the expected orientation and significantly enriched in genes related to neurogenesis and axonogenesis (Supplementary Figure 4).

To explore the transcription factor regulatory network of GW13 and GW17 NPCs, we overlapped gestational week specific enhancers, differentially marked by H3K4me1 between GW13 and GW17, with TFBSs and identified five transcription factors exclusively enriched in GW17 specific enhancers, three of which were expressed in both cortex and GE-derived NPCs (Figure 4C). The three expressed transcription factors play important roles in brain development: OLIG2, FOXO1, involved in cell specification [50]; and androgen receptors (AR), involved in sex differences in brain morphology and behavior [51]. Enhancers containing these TFBSs were also enriched in hypomethylated CpGs [52] at GW17 (Figure 4D), and genes associated with these hypomethylated enhancers [25] were transcriptional upregulated in GW17 compared with GW13 (Figure 4D). The OLIG2 promoter itself was also hypomethylated and its transcript upregulated in both cortex and GE NPCs at GW17 (Figure 5A & Supplementary Figure 5). Moreover, 60% of OLIG2 target genes were both hypomethylated in their promoters and upregulated (Figure 5B), and were enriched (FDR <0.05) in genes implicated in neurogenesis in both cortex and GE-derived NPCs (Figure 5C). Among the predicted 132 target genes, 25 were differentially expressed between GW13 and GW17, of which 22 genes were upregulated at GW17 (Figure 5D). These upregulated genes were highly enriched (FDR <0.05) in genes critical for axon guidance and cell migration, including EPHA3 [53], NTNI [54], NTNG1 [55] and SLIT2 [56], reinforcing the critical role of OLIG2 in regulating the transcriptional and epigenetic programming of neurogenesis in early human development.

Taken together, our analysis suggests that epigenetic and transcriptional activation, rather than restriction, is the dominant signature associated with the specification of transcriptional programs associated with early brain development.

**Temporal transcriptional programs are specific for NPCs derived from different brain regions**

Unsupervised clustering of protein-coding gene expression showed that GW13 NPCs from different brain regions clustered together and with GW15 GE-derived NPCs, while GW15 cortex-derived NPCs clustered with GW17 cortex NPCs (Figure 6A). These relationships were also observed by independent clustering of the expression of exons, noncoding genes and miRNAs (Supplementary Figure 6). It suggests that NPCs derived from different regions were transcriptionally similar at GW13 but that by GW15 region-specific signatures emerge. Differentially expressed genes between GW13, GW15 and GW17 were identified (Supplementary Table 5), and categorized into eight profile groups based on their temporal expression pattern (upregulated, downregulated and not differentially expressed) between GW13-GW15 and GW15-GW17. While overall, we observed a trend for increasingly gene expression with GW in NPCs (Figure 6B), differentially expressed genes shared by both cortex and GE-derived NPCs showed no discernable pattern across gestational weeks (Figure 6C). In contrast an increase in expression was observed between GW13-GW15 in cortex-derived NPCs (Figure 6C). In GE-derived NPCs, a similar upregulation in gene expression was observed, however, it occurred between GW15 and GW17 (Figure 6C). Several key factors in brain development followed this pattern (Figures 6D), including OLIG2, as well as LHX4, a transcription factor critical for nervous system development [58]; WNT5A, involved in regulating axon growth and guidance [59]; and FOXH1, playing key roles in patterning of the forebrain [60]. To validate our results, we investigated the expression levels of the differential expressed genes between different GW in the independent validation cohort of 12 fetal brain samples, and found that these genes were stably expressed through GW8 to GW13 but were significantly upregulated from GW13 to GW17 (t-test p < 0.01) (Supplementary Figure 5).

Taken together, these analysis provide a high-resolution view of the key transcriptional and epigenetic network changes occurring during early fetal brain development and supports a model of increased transcriptional activity associated with different developmental stages in NPCs derived from different brain regions.
Figure 5. Regulatory network of OLIG2. (A) UCSC genome browser tracks showed OLIG2 hypomethylated in promoter region and upregulated in GW17. (B) Heatmap of enhancer signal level (H3K4me1) of GW17 specific enhancers with OLIG2 binding sites (left), their corresponding DNA fractional methylation (middle) and transcription levels of their nearest genes (right) showed hypomethylation of the enhancers and upregulation of the predicted target genes in GW17. (C) DAVID [57] GO biological processes enrichment analysis of OLIG2 target genes indicating significant enrichment for brain development related biological processes. (D) Cytoscape network of OLIG2 target genes that are differential expressed. A total of 22 genes were upregulated in GW17 (red) while only 3 were downregulated (green). The size of the circles corresponds to the fold change of expression levels between GW17 and GW13. Among the upregulated genes, EPHA3, NTN1, NTNG1 and SLIT2 were associated with axon guidance and cell migration.

FDR: False discovery rate; GE: Ganglionic eminence; GO: Gene ontology; GW: Gestational week; mC: Methyl-cytosine; RPKM: Reads per kilobase of transcript; UCSC: University of California, Santa Cruz.

Epigenetic & transcriptional differences between MZ twins arise before midgestation
The genomes of MZ twins are genetically identical providing a unique opportunity to study developmental processes that give rise to epigenetic variations during normal development within the womb. To investigate epigenetic differences arising between MZ twins during fetal brain development, we analyzed DNA methylation...
Figure 6. Stage-specific differential expression. (A) Unsupervised clustering of expression levels of protein coding genes. NPCs derived from different regions clustered together at GW13. However, by GW15, GE-derived NPCs still clustered with GW13 NPCs, while cortex-derived NPCs branched off and clustered with GW17 cortex NPCs. (B) Number of differentially expressed genes between different gestational weeks (left panel: GW13 vs GW17, middle panel: GW15 vs GW17, right panel: GW13 vs GW15) in cortex-derived NPCs (red), GE-derived NPCs (blue) and shared by two cell populations (purple). Bars pointing up showed upregulation in the later stages, and bars pointing down showed downregulation, suggesting a general trend of upregulation in later stages. (C) Patterns of expression for genes differentially expressed between gestational weeks in cortex-derived NPCs (top panel), in GE-derived NPCs (middle panel) and in genes shared by both cortex and GE-derived NPCs (bottom panel). Genes are divided into eight expression profile groups according to differential expression analysis of GW13 vs GW15 and GW15 vs GW17, represented by the arrows of eight different colors. The thickness of the line and the number in each panel corresponded to the number of genes within each category, and the dashed lines means no genes in that category. Cortex-derived NPCs showed dominant transcription activation in GW13-GW15, while GE-derived NPCs showed similar activation but in GW15-GW17. (D) Heatmap for expression values of stage-specific differential expressed genes in cortex NPCs and GE NPCs showed stage-specific expression patterns for key regulators of brain development.

DN: Downregulated; GE: Ganglionic eminence; GW: Gestational week; NPC: Neural progenitor cell; ST: Stable, not differentially expressed; UP: Upregulated.
and histone modification data from brain tissue, and cortex and GE-derived NPCs of the GW17 MZ twins. MZ twin DMRs were identified through pairwise differential methylation in each tissue and cell type respectively (see methods). In dissected brain tissue and cortex-derived NPCs, there was significant asymmetry in the orientation of these DMRs with an average of twofold more hypomethylated DMRs present in subject two compared with subject one (Figure 7A & Supplementary Table 4). As all samples were processed in the same batch using standardized protocols, we considered it unlikely that the asymmetry is a result of sampling bias or technical differences (see methods). To explore the possible functional significance of these asymmetric MZ twin DMRs, we examined their relationship to protein coding genes. On average, we identified 405 DMRs (12% of total) within the promoters of protein-coding genes, a threefold enrichment compared with that expected by chance (hypergeometric p < 0; Supplementary Figure 7). We associated DMRs to genes, and gene enrichment analysis of the resulting gene one (Figure 7A & Supplementary Table 4). As all samples were processed in the same batch using standardized methods). In dissected brain tissue and cortex-derived NPCs, there was significant asymmetry in the orientation of twin DMRs were identified through pairwise differential methylation in each tissue and cell type respectively (see

...tissue, consistent with the more neuronal phenotype that we observed in cortex-derived NPCs.

...increase in active epigenetic and transcription states with developmental stage, the more active epigenetic state of tissue specific comparisons suggest that cortex-derived NPCs may be at a more epigenetically and transcriptionally active state compared with GE-derived NPCs. Taking into account temporal analysis of NPCs which suggests an increase in active epigenetic and transcription states with developmental stage, the more active epigenetic state of the cortex-derived NPCs suggests that cortex-derived NPCs are developmentally more advanced than GE-derived NPCs, consistent with the more neuronal phenotype that we observed in cortex-derived NPCs.

Targeted CpG methylation array studies have demonstrated that the genomes of human fetal brain tissue undergo DNA hypomethylation with age [13], and that the bulk of temporal transcriptional changes occur at
Figure 7. Epigenetic and transcriptional differences between monozygotic twins arise early in fetal brain development. (A) Total DMR length (top panel), number of enriched bases in enhancers (middle panel) and number of differential expressed genes (bottom panel) between MZ twins. Subject1-specific hypomethylated DMRs/enhancers and upregulated genes were shown in red, and Subject2-specific ones were shown in blue. (B) DMRs between MZ twins in three tissue and cell populations from GREAT [61] functional enrichment analysis all showed enrichment in Homeobox protein domains. (C) UCSC genome browser MeDIP-seq tracks showed hypermethylation in Subject2 compared with Subject1 at the promoter region of CCDC169 in dissected brain tissue as well as cortex and GE-derived NPCs. (D) DAVID [57] functional enrichment analysis for genes differentially methylated in their promoters and differentially expressed between the MZ twins showed enrichment in brain developmental processes.

DE: Differential expression; DMR: Differentially methylated region; FDR: False discovery rate; GE: Ganglionic eminence; GOBP: Gene ontology – biological process; GOMF: Gene ontology – molecular function; MeDIP-seq: Methylated DNA immunoprecipitation sequencing; MZ: Monozygotic; NPC: Neural progenitor cell; UCSC: University of California, Santa Cruz.
prenatal stages with the activation of critical regulators during early-mid fetal development [10,12]. Our analysis confirmed a developmentally associated hypomethylation between GW13–17 and extended this to show that this hypomethylation, in particular in enhancer regions, was associated with increased transcription factor binding activities of critical transcription factors, such as OLIG2. The activation of these transcription factors may lead to activation of genes critical for axon guidance and cell migration, suggesting an essential role of epigenetic regulation in increased cell migration activities during this critical developmental period. Furthermore, by comparing the transcription profiles in two distinct NPC populations, we found that the transcriptional activation occurred in waves, and at different developmental stages in NPCs derived from different brain regions. This pattern supports a model where NPCs from both brain regions share a common epigenetic state, possibly inherited from a common progenitor cell, and that cortex-derived NPCs diverged earlier and became more differentiated than GE-derived NPCs, providing additional insight into the spatiotemporal developmental trajectory within the fetal brain.

Postnatal morphological differences in the gross anatomy of the brain tissue of human MZ twins are well documented [68–70]. Previous studies on postnatal MZ twins suggested that the epigenetic differences between the twins arise primarily through exposure to divergent environments [71], and may contribute to phenotypical discordance in development and disease onset [72–75]. However, these studies were unable to find distinct epigenetic signatures at the age of three due to technical limitations [76]. Whether or when these differences arise in prenatal MZ twins and their consequences are still poorly understood. Our analysis of the epigenomes of prenatal MZ twins provides evidence of epigenetic differences as early as GW17 that are associated with the activities of master regulators of brain development such as the WNT family of proteins.

Access to normal phenotyped and cryopreserved fetal brain specimens is challenging and there is a paucity of epigenomic profiles available for comparative research. In the present study we have comprehensively profiled the epigenic states of biospecimens derived from four normal human fetuses. To validate our observations of a gestational week associated DNA hypomethylation signature and associated induction in gene expression we analyzed an independent cohort of 23 fetal brain samples spanning from GW8 to GW17. However, we were unable to obtain additional relevant epigenomic profiles to validate predicted gestational week enhancer states and in particular our findings of significant epigenomic variation between MZ twins and between NPCs derived from different brain regions. Future studies using targeted approach in a larger cohort are needed to validate these additional findings.

Our analysis provides an unprecedented annotation of epigenetic and transcriptional states of early fetal brain development. Analysis across developmental stages supports a model of epigenetic and transcriptional activation with developmental age, and suggested that cortex-derived NPCs may be at a more advanced epigenetic state than GE-derived NPCs. Our results reveal epigenetic differences between MZ twins during early fetal periods. We provide a high-resolution regulatory network for NPCs from different brain regions, providing a comprehensive reference for future studies in brain development and disease.

**Conclusion & future perspective**

Our analysis of epigenetic states of normal human fetal brain regions and derived NPCs support a model of temporal epigenetic activation and the establishment of distinct epigenetic states within NPCs derived from different brain regions and across monozygotic twins. These data provide a comprehensive set of epigenomic features for further exploration of the epigenetic state changes associated with normal fetal brain development and its dysfunction in disease.

**Supplementary data**

To view the supplementary data that accompany this paper please visit the journal website at: www.futuremedicine.com/doi/suppl/10.2217/epi-2019-0319

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Summary points

• We provide a high-resolution regulatory network for neural progenitor cells (NPCs) from different brain regions.
• NPCs derived from different fetal brain regions are epigenetically distinct and become increasingly divergent with developmental age.
• NPCs derived from distinct brain regions retain region of origin signatures when cultured as neurospheres.
• NPCs undergo epigenetic and transcriptional activation during early development.
• OLIG2 plays a critical role in regulating transcriptional and epigenetic programming of neurogenesis in early human brain development.
• NPCs derived from distinct brain regions acquire transcriptional activation at different developmental time points.
• Epigenetic differences between monozygotic twins arise as early as GW17.
• Epigenetic differences between monozygotic twins are associated with differential expression of genes critical to brain development.

Epigenomic programming in early fetal brain development

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Ethical conduct of research

The authors state that they have obtained appropriate institutional review board approval or have followed the principles outlined in the Declaration of Helsinki for all human experimental investigations. In addition, for investigations involving human subjects, informed consent has been obtained from the participants involved.

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