Regulation of chromatin accessibility and Zic binding at enhancers in the developing cerebellum

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To identify chromatin mechanisms of neuronal differentiation, we characterized chromatin accessibility and gene expression in cerebellar granule neurons (CGNs) of the developing mouse. We used DNase-seq to map accessibility of cis-regulatory elements and RNA-seq to profile transcript abundance across postnatal stages of neuronal differentiation in vivo and in culture. We observed thousands of chromatin accessibility changes as CGNs differentiated, and verified, using H3K27ac ChIP-seq, reporter gene assays and CRISPR-mediated activation, that many of these regions function as neuronal enhancers. Motif discovery in differentially accessible chromatin regions suggested a previously unknown role for the Zic family of transcription factors in CGN maturation. We confirmed the association of Zic with these elements by ChIP-seq and found, using knockdown, that Zic1 and Zic2 are required for coordinating mature neuronal gene expression patterns. Together, our data reveal chromatin dynamics at thousands of gene regulatory elements that facilitate the gene expression patterns necessary for neuronal differentiation and function.

Specialized cell function requires precise control of gene expression patterns. Chromatin regulation is involved in this process by establishing differential utilization of gene regulatory elements in cells of distinct fate lineages. Genome-wide studies have suggested that cell type–specific differences in gene expression are highly correlated with differences in both accessibility and activation state of distal gene enhancers1,2. These data have led to the hypothesis that developmental regulation of chromatin at enhancer elements mediates the process of cellular differentiation3.

Neuronal differentiation is comprised of multiple steps, beginning with the commitment of neural stem cells to become specified neural progenitors, which then leave the cell cycle to become postmitotic neurons. Prenatal patterning of the brain is critically dependent on temporally and spatially restricted expression of transcription factors that act at brain region–selective enhancer elements4,5. By contrast, after birth, sensory experience-driven synaptic activity has an instructive role in initiating programs of gene expression that underlie neuronal maturation. This allows processes such as synapse development and excitatory/inhibitory balance in neural circuits to be adapted to the environment6. However, whether chromatin-dependent regulation of enhancer function contributes to gene expression that mediates these later stages of neuronal maturation remains largely unknown.

To fill this gap in knowledge, we used the differentiation of CGNs in the postnatal mouse cerebellum to identify chromatin-based transcriptional mechanisms that drive the maturation of neuronal gene expression programs. CGNs, which comprise >99% of cerebellar neurons, are derived during early postnatal life from committed granule neuron precursors (GNPs) that proliferate in the outer portion of the external granular layer of the developing cerebellar cortex7. Following exit from the cell cycle, GNPs differentiate into immature CGNs that migrate to the inner granular layer, where they form synaptic connections and then mature. These changes in CGN differentiation and function are accompanied by known changes in neuronal gene expression8. Notably, primary GNPs isolated from the postnatal mouse brain recapitulate discrete and synchronized stages of CGN differentiation in culture, providing a means for experimental validation and genetic manipulation of gene regulatory mechanisms that mediate this process9,10.

DNase I hypersensitive (DHS) sites mark nucleosome-depleted regions that are universal hallmarks for gene regulatory elements, including promoters, enhancers, insulators and most transcription factor–binding sites11. We applied DNase-seq to globally map chromatin accessibility at key stages in the development of mouse cerebellum. We found that widespread changes in chromatin accessibility occurred as CGNs differentiated, marking dynamic enhancer elements that regulate the expression of genes necessary for proper neuronal function. In addition, we used our identification of these regions to determine a previously unknown role for the Zic transcription factors in coordinating the gene expression programs required for neuronal maturation.

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Cerebellar development involves extensive chromatin remodeling

We used DNase-seq to globally map chromatin accessibility at three key stages in the development of the mouse cerebellum: postnatal day 7 (P7), when the external granular layer (EGL) of the mouse cerebellar cortex has reached its maximal thickness as a result of the proliferation of GNPs (Fig. 1a); P14, when newborn postmitotic CGNs begin to populate the internal granule layer (IGL); and P60, when CGNs of the IGL express gene products that mediate mature synaptic functions (Fig. 1b, Supplementary Fig. 1a and Supplementary Table 1). We identified approximately 70,000 DHS sites at each of the three developmental stages and found that these elements mapped to promoters, gene bodies and intergenic regions (Supplementary Fig. 1b). The majority of DHS sites (~77%) were found outside of annotated gene promoter regions, highlighting the ability of DNase-seq to identify distal regulatory elements in the genome.

To determine how chromatin accessibility changes during cerebellar differentiation, we assessed quantitative differences in DHS signal intensity between the three stages. We identified 24,886 total DHS sites that significantly ‘opened’ or ‘closed’ (gained or lost signal) during postnatal development (FDR < 0.05; Fig. 1c, d, Supplementary Fig. 1c and Supplementary Table 2). Similar to all DHS sites, developmentally regulated DHS sites were found in both proximal gene promoters and at intronic, exonic and intergenic regions (Fig. 1e). Notably, we found evidence of extensive chromatin remodeling between P14 and P60, even though these samples were primarily comprised of CGNs at two postmitotic stages of neuronal differentiation. These data indicate that there is extensive chromatin remodeling over the course of neuronal differentiation in vivo both as proliferating neural progenitors leave the cell cycle and as postmitotic neurons mature.

Although an advantage of cerebellar cortex is that it is predominantly composed of granule neurons (~85%), this brain region also contains astrocytes (~15%) and other kinds of neurons such as Purkinje cells (~0.2%) (Supplementary Fig. 2a, b). To determine the effect of these other cell types on our chromatin analyses, we first...
quantified aggregate DNase-seq signal at the promoters of genes that are preferentially expressed in CGNs, Purkinje neurons or Bergmann glia of the adult mouse cerebellum. DNase signal was greatest at promoters of genes preferentially expressed in adult CGNs, suggesting that our DHS data represent chromatin state in this most abundant cell type (Supplementary Fig. 2c). Furthermore, when we compared the set of developmentally regulated DHS sites, these also had the strongest association with CGN-enriched genes (Supplementary Fig. 2d). ChiP-seq data sets obtained from mouse cerebellum were similarly enriched at CGN gene promoters over other cell types (Supplementary Fig. 2e.f).

As an additional way to control for cellular heterogeneity, we purified GNPs from the P7 cerebellum and differentiated these cells to CGNs in culture. These cultures are highly enriched for CGNs and depleted of glia and other kinds of neurons found in the cerebellar cortex in vivo. Once plated, freshly dissociated GNPs rapidly exit the cell cycle and synchronously differentiate, displaying characteristic neuronal morphologies by 3 d in vitro (+3 DIV) and forming synaptic connections by +7 DIV (Supplementary Fig. 3a). We performed DNase-seq in triplicate on freshly isolated GNPs or GNPs that were cultured for 3 or 7 d and identified 28,119 differential DHS sites at FDR < 0.05 (Supplementary Fig. 3b–d and Supplementary Table 3). Similar to our findings from in vivo cerebellar samples, we found that opening and closing DHS sites in cultured CGNs are primarily non-promoter distal elements (Supplementary Fig. 3e).

Although the majority (10,115 of 11,657 = 86.8%) of changes in DHS site accessibility occurred as GNPs left the cell cycle to become postmitotic +3 and +7 DIV samples. These data again indicate that postmitotic neurons can undergo substantial chromatin accessibility changes as they establish synapses and mature.

Comparing developmental regulation of DHS sites between the in vivo and culture systems revealed most sites exhibit matched direction of change (Supplementary Fig. 4a), but the magnitude of accessibility changes was generally lower in the cultured neurons. However, we also observed a cluster of DHS sites that closed gradually from P7 to P60 in vivo, but became transiently more accessible at +3 DIV before closing down at +7 DIV. Regulation of at least a subset of these sites accompanied transient increases in the expression of nearby genes. For example, Grin2b is a developmentally regulated NMDA-type glutamate receptor subunit that is most highly expressed in the neonatal brain, where it is important for the induction of long-term potentiation. Two transiently opening DHS sites in an intron of the Grin2b gene parallel the transient increase in Grin2b mRNA expression at +3 DIV in cultured CGNs (Supplementary Fig. 4b,c). These data suggest that, in addition to reducing cellular heterogeneity, this culture system complements our in vivo differentiation analysis by improving temporal resolution of chromatin regulation during early steps in CGN differentiation.

**Figure 2** Opening DHS sites mark late-acting neuronal enhancers. (a, b) Cumulative fraction of genes nearest to opened promoter-located, opened distal and all identified DHS sites with given fold-change in RNA-seq expression from P7 to P60 cerebellum (a) or from GNPs to +7 DIV (b). Rightward shift indicates genes increased in expression across developmental time. Significance assessed by Mann-Whitney U test (n = 3 biological replicates of pooled cerebellum). (c) Mean H3K27ac ChIP-seq signal (RPM, reads per million mapped) present at center of P7 to P60 opened DHS sites (4,053 sites) in either P7 cerebellum (orange line) or P60 cerebellum (green line). Gray = s.e.m. (n = 2 biological replicates of pooled cerebellum). (d) Percent of opened promoter-located, opened distal and all DHS sites overlapping H3K27ac peaks identified in P7 or P60 cerebellum. (e) UCSC browser image highlighting a DHS site found in the 3′ UTR of Cbln3 (black box), which opened during development and overlapped H3K27ac and H3K4me1 enrichment in adult cerebellum. (f) RNA-seq expression of Cbln3 across developmental time in vivo and in cultured CGNs. Error bars represent 95% confidence interval. FPKM, fragments per kilobase of exon per million mapped. (g) Luciferase reporter assay for enhancer activity conferred by a DHS site in Cbln3 at +3 DIV and +6 DIV. P = 0.0012 by two-sided Student’s t test (n = 3 transfections, t = 8.3). Error bars represent s.e.m.
Figure 3 CRISPR-VP64–based activation confirms enhancer activity of late-opening DHS sites nearby Grin2c. (a) DNase-seq and H3K27ac ChIP-seq signal in vicinity of Grin2c with two DHS sites that opened across development and increased in H3K27ac signal marked in gray. (b) RNA-seq expression of Grin2c across developmental time in vivo and in culture. Error bars represent s.e.m. with gRNAs, leftward shift indicates genes decreased in expression across developmental time. Significance assessed by Mann-Whitney test for dCas9-VP64 plus gRNAs versus dCas9-VP64 or gRNAs alone (n = 5 for dCas9-VP64 plus gRNAs versus dCas9-VP64 or gRNAs alone; n = 8 for site 1 controls, n = 10 for site 2 controls; t = 5.5 for site 1, t = 12.3 for site 2). Error bars represent s.e.m.

Supplementary Table 4). By clustering patterns of differentially expressed genes, we confirmed a number of previously described granule neuron gene expression dynamics and recovered gene ontologies descriptive of the differentiation process19–22 (Supplementary Fig. 5c–g and Supplementary Table 5). As with our DNase-seq analysis, we also performed RNA-seq on triplicate replicates of purified GNP s that were either freshly isolated or differentiated in culture for +3 or +7 DIV. We found 1,972 differentially expressed genes in the cultures across this time course (Supplementary Fig. 6a), and principal components analysis of RNA-seq expression values placed +3 and +7 DIV samples in between GNPs and P14 cerebellum along an axis consistent with developmental time (Supplementary Fig. 6b).

We postulated that the developmentally regulated chromatin accessibility changes that we observed might mark a combination of promoter and enhancer elements that become activated at specific developmental stages to drive target gene expression. We first mapped each DHS site to its nearest gene and analyzed the relationship between that DHS site and the change in its associated gene expression value between P7 and P60. Although all DHS sites together exhibited a normal distribution of nearby gene expression changes centered on a fold change of 1, both promoter and distal DHS sites that opened up between P7 and P60 were significantly associated with genes that had higher expression levels at P60 (P = 2.7 × 10−34 and P = 5.0 × 10−31, respectively, Mann-Whitney test; Fig. 2a). Similar to our in vivo data, we found a strong association between GNP to +7 DIV opening DHS sites and increased nearby gene expression in cultured CGNs (Fig. 2b).

The functional identity of DHS sites (for example, enhancers, promoters and silencers) can be predicted by overlap with specific histone modifications23,24. With the knowledge that most DHS sites are located outside of proximal promoter regions, and that our opening...
sites correlated with increased gene expression, we asked whether these sites colocalized with the enhancer mark H3K4me1 and the active enhancer/promoter modification H3K27ac. Comparing our developmentally regulated DHS sites to available mouse ENCODE ChIP-seq data from P56 cerebellum to P76 cerebellum revealed an enriched overlap between our opened DHS sites and the H3K4me1 (77%, P = 1.13 × 10^{-120}, hypergeometric test) and H3K27ac (50%, P = 8.82 × 10^{-88}) histone marks (Supplementary Fig. 7a). The combination of these two histone marks was particularly enriched in non-promoter opening DHS sites (45% overlap peaks of both marks), whereas promoter-located opening DHS sites were enriched for H3K27ac in combination with H3K4me3 (40% overlap peaks of both marks). This indicates that DHS sites that opened between P7 and P60 demarcate both active enhancers and promoters in adult cerebellum. To determine whether the opening of DHS sites over developmental time was associated with changing of histone marks, we performed ChIP-seq for H3K27ac in P7 and P60 cerebellum in duplicate. We found a strong correlation (Pearson’s r = 0.538) between global H3K27ac and DHS signal changes in the developing cerebellum, indicating that increased accessibility is associated with concomitant H3K27ac deposition (Supplementary Fig. 7b,c). Separating out H3K27ac peaks that map to proximal promoters further revealed that the global correlation is driven primarily by non-promoter sites (r = 0.278 promoters versus r = 0.621 non-promoters; Supplementary Fig. 7d,e). Matching these observations, P7–P60 opened DHS sites contained a mean H3K27ac signal that increased markedly over development (Fig. 2c). Furthermore, the set of opening DHS sites was markedly enriched for overlap with H3K27ac peaks compared with all DHS sites at P60 (Fig. 2d). Together, these data suggest that increased chromatin accessibility is associated with the activation of enhancer elements that promote developmentally regulated increases in CGN gene expression.

To determine whether the opening DHS sites indeed function as enhancers of mature CGN gene expression, we first used luciferase assays to test the function of a DHS site that becomes significantly more accessible both in vivo and in culture found in the 3’ UTR of the cerebellin 3 precursor (Cbln3) gene, which encodes a secreted C1q-domain protein that modulates the formation, refinement and maintenance of CGN to Purkinje cell synapses (Fig. 2e). This DHS site was associated with increased CGN gene expression. When cloned in front of a minimal promoter and transfected in CGNs, this element substantially increased luciferase expression (Fig. 2g). Furthermore, we found significantly more luciferase expression when we harvested neurons on +6 DIV compared with +3 DIV (P = 0.0012, Student’s t-test; Fig. 2g), indicating that the activity of this enhancer increases with the developmental maturation of the CGNs.

Next, to directly test the hypothesis that opening DHS sites represent enhancer elements in their native genomic context, we examined the locus surrounding the Grin2c gene on chromosome 11 in detail (Fig. 3a). In this interval, we identified two DHS sites that both opened and gained H3K27ac signal across postnatal development, one just upstream of the Grin2c promoter and the second in an intron of the nearby Tmem104 gene. Of the four annotated genes in this region (Tmem104, Grin2c, Fdxr and Fads6), all were expressed in cerebellum, but only Grin2c expression was highly upregulated in differentiating CGNs, where it is a NMDA-type glutamate receptor subunit that mediates mature synaptic functions (Fig. 3b, Supplementary Table 6). Grin2c expression can also be robustly induced by culturing CGNs for 7 d (+7 DIV) in the presence of brain-derived neurotrophic factor (BDNF), offering an opportunity to experimentally test transcriptional mechanisms of Grin2c regulation. Notably, BDNF-induced gene expression changes occurred largely independent of changes in chromatin accessibility, as we observed only 33 opened and 6 closed DHS sites in response to BDNF exposure (Supplementary Table 7).

To determine whether the two identified DHS sites function as enhancers of Grin2c in their endogenous chromatin context, we used

**Figure 5** Postnatal closing DHS sites are enriched for embryonic hindbrain enhancer activity. (a,b) Percent of hindbrain (a) or heart (b) localized VISTA Enhancer Browser staining (embryonic day 11.5) covered by all DHS sites identified in P7 cerebellum, heart, kidney or liver mouse tissues (matching number of top peaks). (c) Percent of VISTA Enhancer Browser tissue-localized enhancers overlapped by DHS sites that close from P7 to P60. Tissues ordered by overlap rank. (d) Closed DHS sites located closest to the Elavl2, Sall1 and Zfp423 genes, all of which decrease in expression from P7 to P60 (center). Hindbrain expression is driven by each DHS site at E11.5, as seen with LacZ (blue) staining of embryos on right. Black scale bar represents 1 mm. Images sourced from VISTA Enhancer Browser: hs643, hs152 and hs625. Error bars represent s.e.m.
a CRISPR RNA-guided method to recruit a synthetic dCas9-linked transcriptional activator to each of the two putative enhancers flanking *Grin2c* in cultured CGNs. Co-infection of dCas9-VP64 with CRISPR guide RNAs (gRNAs) targeting either site was sufficient to induce a selective upregulation of *Grin2c* expression compared with levels in control-infected neurons (Fig. 3c). By contrast, expression of three other nearby genes (*Rab37, Tnem104* and *Fdxr*) remained unchanged. Together, our CRISPR data indicate that two developmentally regulated DHS sites, including one located >50 kb away from the *Grin2c* promoter, act as gene-specific enhancers, highlighting the power of our comparative DHS analysis to locate physiologically relevant enhancers of important neuronal genes.

**Early-acting enhancers are deactivated before closing**

In contrast with the opening DHS sites, we observed only a modest association between DHS sites that close between P7 and P60 and the expression of their nearest gene (*P = 0.03* promoter located, *P = 0.09* for distal) or from GNPs to +7 DIV (*P = 0.003* promoter-located, *P = 0.10* for distal) (Fig. 4a,b). As expected, closing DHS sites largely did not colocalize with the enhancer markers H3K4me1 and H3K27ac in P56 brain (Supplementary Fig. 7a). However, these regions did exhibit somewhat greater H3K27ac ChiP-seq signal in P7 cerebellum than at P60, with decreases in this active enhancer mark being noted over the course of development at both promoter-located and distal DHS sites (Fig. 4c,d). These data suggest that at least a subset of the closing DHS sites are likely to be active enhancers in GNPs that are deactivated as CGNs differentiate. However, given that we observed noticeably less H3K27ac enrichment at P7 for closing DHS sites compared with the enrichment of H3K27ac at P60 for opening DHS sites, we considered the possibility that the loss of chromatin accessibility may slowly follow the functional deactivation of enhancer elements.

If this is the case, then the closing DHS sites may mark regions that function as enhancers in earlier stages in hindbrain development before the differentiation of GNPs. To test this hypothesis, we queried the VISTA Enhancer Browser, which contains enhancer elements that function as enhancers in earlier stages in hindbrain development. To test this hypothesis, we queried the VISTA Enhancer Browser, which contains enhancer elements that function as enhancers in earlier stages in hindbrain development33. These data provide important experimental support for our hypothesis that many of the DHS sites that close during postnatal cerebellar development serve as enhancers of genes that function early in hindbrain development and further suggests that functional deactivation of these enhancers precedes DHS site closure.

**Zics bind developmentally regulated DHS sites**

Developmental regulation of chromatin architecture is thought to determine gene regulatory element accessibility to transcription.
factor binding.\textsuperscript{1,34,35} We therefore sought to identify transcription factors whose regulatory functions may be dictated by chromatin accessibility changes by searching for enrichment of transcription factor motifs in DHS sites that opened or closed between P7 and P60 \textit{in vivo}, as well as those that opened or closed over 7 d of GNP differentiation in culture.

As expected, motifs for the MEF2 and NF1 families, which have been shown to be important for CGN differentiation \textsuperscript{9,36}, were enriched in both opening and closing DHS sites (Supplementary Fig. 8). However, we were surprised to find the zinc finger in cerebellum (Zic) transcription factor family motifs enriched in the set of opening DHS sites. Mutations in the human ZIC genes have been associated with cerebellar development disorders, suggesting their importance in CGN differentiation, and mouse knockout studies have indicated that the Zic5 function in GNPAs to prevent their premature exit from the cell cycle.\textsuperscript{37,38} However, the Zic transcription factors remain highly expressed in differentiated CGNs (Fig. 6a), raising the possibility that these factors also contribute to later stages in CGN maturation. Given our observation that the Zic motif was enriched in opening DHS sites, we hypothesized that the Zic transcription factors might change their gene targets over time by binding to developmentally regulated DHS sites.

To determine whether Zic transcription factors are differentially bound to DHS sites as CGNs differentiate, we performed ChIP-Seq from P7 or P60 cerebellum using an antibody specific for Zic1 and Zic2 (ref. 39) (Supplementary Fig. 9a, b). As a control, no significant peaks were observed with IgG pulldown from P7 or P60 cerebellum or when we performed Zic ChIP-Seq from P60 cortex, where Zic1 and Zic2 are not expressed (Supplementary Fig. 9c). Although 60% of ChIP-seq peaks overlapped between P7 and P60 (Supplementary Fig. 10a,b), we found over 15,000 peaks with significantly (FDR < 0.05) increased or decreased ChIP signal at P60 compared with P7 cerebellum (Fig. 6b and Supplementary Table 9). These data demonstrate that Zic transcription factors undergo dynamic changes in their DNA binding patterns in the developing cerebellum despite constant levels of expression.

Consistent with the enrichment of Zic-binding motifs in opening DHS sites, we found that ~65% (2,683 of 4,053) of the opening DHS sites overlapped with Zic peaks that displayed stronger ChIP signal at P60. By contrast, less than 1% (36 of 4,053) of the opening DHS sites overlapped with Zic peaks with decreased ChIP signal across this time period (Fig. 6c). We also found nearly 50% (2,735 of 5,503) of closing DHS sites showed decreases in Zic binding, whereas only ~3% (179 of 5,503) showed stronger Zic association over time. Increases and decreases in Zic binding over developmental time were correlated with H3K27ac signal, suggesting that Zic contributes to the enhancer activity of these elements (Fig. 6d). Taken together, these data suggest that the chromatin-regulated binding of Zic transcription factors to enhancers may contribute to differential regulation of gene transcription during CGN differentiation.

**Zics promote mature neuronal transcription**

To test whether Zic is required for regulation of gene expression during CGN development, we returned to examining the ~120-kb region surrounding the Grin2c gene, where we observed strong Zic ChIP-seq peaks in P60 cerebellum directly overlapping the two sites tested for enhancer function (Fig. 7a). To determine whether Zic is required for the developmental upregulation of Grin2c expression, we used lentiviral shRNAs to knockdown Zic1 and/or Zic2 expression in cultured CGNs (Supplementary Fig. 9a, b). Knockdown of either Zic1 or Zic2, or both together significantly reduced the expression of Grin2c compared with control-infected neurons (Fig. 7b). To test the role of Zic in the two DHS sites that we found to be \textit{Grin2c} enhancers specifically, we cloned each element into a minimal promoter luciferase reporter plasmid, transfected them into cultured CGNs, and measured luciferase activity 3 or 6 d later. Both elements demonstrated enhancer activity at +6 DIV (Fig. 7c) and knockdown of Zic1 significantly reduced the enhancer activity of both sites (P = 0.0026 and P = 0.0042, Student’s t test; Fig. 7c). Thus, the developmentally regulated recruitment of Zic to these regions is functionally important for these elements to induce transcription.

Finally, given the high degree of overlap between opening DHS sites and increased Zic1 and Zic2 binding as CGNs mature (Fig. 6c), we hypothesized that Zic might function to globally coordinate the maturation of gene expression programs in differentiating CGNs. Consistent with this hypothesis, we found a significant association between increased Zic binding and developmental upregulation of nearby gene expression (Fig. 8a). The relationship between Zic binding and gene expression was strongest when multiple differential Zic binding sites mapped to a single gene (Fig. 8b). To determine the requirement for Zic in CGN differentiation, we used RNA-seq to analyze the effects of Zic1 or Zic2 knockdown on global gene expression in cultured CGNs. Knockdown of either Zic1 or Zic2 drove significant changes in the expression of 81 and 147 genes, respectively (FDR < 0.10;
Mammalian genomes harbor over a million cis-regulatory elements in total. Previous studies comparing hundreds of diverse cell and tissue types demonstrate that the selective accessibility of these sites dictates cell-type identity and function. Developmentally regulated mammalian chromatin accessibility changes have previously been studied in the context of purified cell lineages like intestinal epithelium and CD4+ T cells. Here we focused our analysis on a single neuronal cell lineage to map the in vivo chromatin accessibility changes that occur during development. Our data reveal that chromatin accessibility is highly dynamic across both early and postmitotic stages of neuronal maturation. We found accessibility to be strongly linked to H3K4me1 and H3K27ac deposition outside of promoters, indicating that these chromatin accessibility changes identify regions that are enriched for poised and active enhancer elements. We demonstrated that this form of chromatin plasticity could contribute to transcription-dependent forms of learning and memory in the adult brain.

Figure 8 Zic1/2 promote the mature CGN transcriptional program. (a) Relationship between Zic binding changes and nearby gene RNA-seq expression changes from P7 to P60. Significance of shifts was assessed by two-sided Mann-Whitney U test. (b) Boxplots of gene expression change from P7 to P60 cerebellum were binned by number of Zic binding sites associated with each gene. Having multiple Zic binding changes nearby was more strongly associated with directional gene expression changes than single sites. Center line represents the median, box limits represent the first and third quartiles, and whiskers represent 1.5 times the interquartile range. (c,d) Relationship between cultured CGN development and Zic1 (c) or Zic2 (d) shRNA knockdown for genes marked significant in knockdown RNA-seq experiments (FDR < 0.10, n = 2 independent cultures). X axis shows fold change in gene expression between +3 DIV and +7 DIV in control cultures versus the expression change for that same gene at +7 DIV after knockdown of Zic1 (Fig. 8c) or Zic2 (Fig. 8d). These data reveal that genes that show reduced expression following Zic1 or Zic2 knockdown tended to be upregulated over the time course of CGN differentiation, whereas genes with higher expression following Zic1 or Zic2 knockdown tended to be downregulated over the course of differentiation. Thus, the net effect of Zic1 or Zic2 knockdown is a shift in the gene expression program toward a less mature pattern. Overall, these data indicate that Zic recruitment to regions of opened chromatin is essential for the maturation of gene expression programs in differentiating CGNs.

Supplementary Table 10). To determine how this set of Zic-regulated genes was related to the gene expression changes that accompany CGN differentiation, we plotted the expression change of each Zic-regulated gene between +3 DIV and +7 DIV in control cultures versus the expression change for that same gene at +7 DIV after knockdown of Zic1 (Fig. 8c) or Zic2 (Fig. 8d). These data reveal that genes that show reduced expression following Zic1 or Zic2 knockdown tended to be upregulated over the time course of CGN differentiation, whereas genes with higher expression following Zic1 or Zic2 knockdown tended to be downregulated over the course of differentiation. Thus, the net effect of Zic1 or Zic2 knockdown is a shift in the gene expression program toward a less mature pattern. Overall, these data indicate that Zic recruitment to regions of opened chromatin is essential for the maturation of gene expression programs in differentiating CGNs.

DISCUSSION Mammalian genomes harbor over a million cis-regulatory elements in total. Previous studies comparing hundreds of diverse cell and tissue types demonstrate that the selective accessibility of these sites dictates cell-type identity and function. Developmentally regulated mammalian chromatin accessibility changes have previously been studied in the context of purified cell lineages like intestinal epithelium and CD4+ T cells. Here we focused our analysis on a single neuronal cell lineage to map the in vivo chromatin accessibility changes that occur during development. Our data reveal that chromatin accessibility is highly dynamic across both early and postmitotic stages of neuronal maturation. We found accessibility to be strongly linked to H3K4me1 and H3K27ac deposition outside of promoters, indicating that these chromatin accessibility changes identify regions that are enriched for poised and active enhancer elements. We demonstrated that this form of chromatin plasticity could contribute to transcription-dependent forms of learning and memory in the adult brain.

We found that, although DHS site opening frequently coincided with an increase both in H3K27ac signal and transcription of nearby genes, DHS site closing correlated weakly with decreasing gene expression and predominantly occurred at elements that already lacked H3K27ac at P7. At least a subset of these closing elements functioned as enhancers in the cerebellar primordium during embryonic stages of brain development, suggesting that enhancer deactivation precedes the loss of accessibility. This evidence that an epigenetic signature is retained in the form of chromatin accessibility at some enhancers, even though they are no longer active, could permit the reactivation of gene expression programs otherwise thought to be cell type or developmental stage specific.

We observed a substantial difference in the magnitude to which DHS sites opened in CGNs that were differentiated in culture compared with in vivo. This suggests that full maturation of CGN chromatin accessibility requires cell non-autonomous factors in the developing cerebellum. These external influences could include interactions with the Purkinje neurons, which are important for the proliferation and migration of CGNs, as well as patterned neural activity arising from sensory input to the intact brain, which have a key role in refining cerebellar synapses. Further elucidating the role of extrinsic factors in the regulation of chromatin accessibility is likely to uncover mechanisms of gene-environment interactions relevant to both normal and abnormal brain development.

Differential DHS analysis allowed us to predict and confirm that members of the Zic transcription factor family preferentially bind DHS sites that open during CGN differentiation. Mice lacking Zic1 alone or in combination with Zic2 have a small cerebellum, indicating a requirement for the Zics in the normal proliferation of cerebellar GNPs.
Heterozygous loss-of-function mutations in ZIC1 and ZIC4 have been identified in patients with Dandy–Walker malformation, a congenital malformation of the cerebellum35, further demonstrating fundamental roles for these transcription factors in brain development. However, our evidence that Zics act in postmitotic CGNs to promote mature gene expression patterns was surprising in light of previous studies that have demonstrated a role for the Zics in preventing premature differentiation of dividing neuronal precursors43,44.

Zic family member expression remained constant throughout CGN development, yet these factors had distinct genome binding profiles at P7 and P60. Our evidence that sites of regulated Zic binding overlapped developmentally regulated DHS sites suggests that chromatin accessibility determines local Zic affinity. Developmental changes in chromatin accessibility are thought to be mediated by pioneer transcription factors that first recognize their sequence-specific DNA binding sites and then recruit chromatin-remodeling complexes45. Our data show that many gene regulatory elements that recruit Zic binding between P7 and P60 already overlapped the active enhancer mark H3K27ac at P7, although they further increase in H3K27ac signal over development. These data raise the possibility that other developmentally regulated factors bind these sites first, and then recruit Zic. The switch in Zic binding sites may explain how Zics get repurposed for two very different functions— inhibition of premature differentiation in GNPs and promotion of maturation in post-mitotic CGNs—at distinct steps in the differentiation of the same cell lineage. Although the Zics are widely expressed in the developing brain, CGNs are one of the few kinds of neurons that maintain Zic expression in adulthood, suggesting that other transcription factors may function at stage-specific enhancers to direct mature neuronal gene expression programs in other types of neurons.

Although previous studies have demonstrated that extra-cellular signal–regulated transcription factors can drive CGN maturation9,36,46,47, the contributions of chromatin accessibility dynamics to these processes have not been investigated. The idea that environmental stimuli can induce changes in chromatin accessibility has been called into question by studies that have shown global cell-type-specific gene expression. Nature 459, 108–112 (2009).

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METHODS

Methods and any associated references are available in the online version of the paper.

Accession codes. Sequencing data have been deposited at Gene Expression Omnibus ( GEO) under accession number GSE60731.

Note: Any Supplementary Information and Source Data files are available in the online version of the paper.

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AUTHOR CONTRIBUTIONS

A.E.W., G.E.C., C.L.F., R.W. and F.L. designed the study. F.L., C.L.F., R.W., L.S. and A.S. performed DNase-seq and ChIP-seq experiments. M.T.B., M.G.Y., C.M.V. and C.A.G. designed and performed targeted enhancer function assays. F.L. performed Zic knockdown experiments. C.L.F. performed all bioinformatic analyses. C.L.F., A.E.W. and G.E.C. wrote the manuscript with input from all authors.

COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

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ONLINE METHODS

Cerebellar immunostaining. C57BL/6Ncrl male or female mice (Charles River Labs) at P7, P14 or P60 were deeply anesthetized with isoflurane and their brains were dissected and flash frozen in OCT. Fresh frozen brains were cryostat sectioned in the sagittal plane at 18 µm, then fixed on slides in 4% paraformaldehyde (wt/vol) and rinsed with 1× phosphate-buffer saline (PBS). Slides were blocked in either 14% normal goat serum (vol/vol) or 10% normal donkey serum (vol/vol) (for primary antibodies raised in goat) and permeabilized in 0.3% Triton X-100 (vol/vol) before antibody incubation. Hoechst dye was used to label nuclei for identification of anatomical landmarks. Images were captured on a Leica DMi4000 inverted fluorescence microscope at 10× or 20× magnification with a Leica DFC-365 camera using Metamorph 7.0 software (Molecular Devices). We used the following primary antibodies: mouse anti-Ki67 (1:50; BD Pharmingen 550609), goat anti-Dcx (1:100, Santa Cruz sc-8066), rabbit anti-calbindin (1:1,000, Swant cat. #CB38). Secondary antibodies were used at 1:500: donkey anti-goat Cy3 (Jackson ImmunoResearch, 705-165-003), goat anti-rabbit Cy2 (1:125-246), goat anti-mouse Cy3 (115-165-146) and goat anti-mouse Cy5 (115-175-146). All animals were housed in groups of 4-5, given access to standard laboratory chow and water ad libitum and housed in a humidity and temperature-controlled room on a 14/10 h light/dark cycle. All experiments were conducted in accordance with an approved protocol from the Duke University Institutional Animal Care and Use Committee and guidelines from the National Institutes of Health for the Care and Use of Laboratory Animals.

Isolation of cerebellar nuclei. For in vivo samples, C57BL/6Ncrl male and female mice at P7, P14 or P60 were deeply anesthetized with isoflurane and decapitated for brain harvesting. The cerebellar cortex was dissected, flash frozen, and stored at −80 °C. For CGN cultures (described below), neurons were scraped into PBS and harvested by centrifugation at 1,000g for 5 min. Nuclei were extracted essentially as previously described59. Briefly, each cerebellum was dounced in 5 ml 2M sucrose, 1 mM MgCl2 and poured through a 100-μm filter. Filtered tissue was ultracentrifuged in 15 ml of 2M sucrose, 1 mM MgCl2 at 65,000g for 80 min. Supernatant was removed and the tight nuclear pellet was resuspended by pipetting in 1 ml buffer RSB from the DNase-seq protocol51. Nuclei were quantified by hemocytometer, yielding approximately 3–10 million per adult cerebellum (~30 mg tissue).

DNase-seq library generation. DNase-seq libraries were constructed from nuclear preparations as previously described51. Briefly, 15–30 million nuclei were digested with a range of recombinant DNase I enzyme (Roche) concentrations (between 1.2 and 12 U) for 15 min at 37 °C in 120 µl 1× DNase buffer. Digestions were checked by pulse field gel electrophoresis and material was pooled from three different DNase concentrations (extent of digestion matched between samples) in equimolar amounts following blunting-end reactions. Following ligation to adapters, MmeI digestion, streptavidin bead-based enrichment, and 14 cycles of PCR amplification, each library was sequenced for either 36 cycles on an Illumina GAIIx machine, or 50 cycles on a Hi-Seq 2000 platform to an average depth of 115 million aligned reads per sample. Nuclei from cerebellae of 4–6 mice or ~20 million cultured CGNs were pooled for each biological replicate, and three independent biological replicates were analyzed for each time point.

RNA isolation and sequencing. RNA was isolated from P7, P14 or P60 C57BL/6 mouse cerebellar cortex or cultured mouse CGNs using the Absolutely RNA kit (Harvard Medical School59), mouse anti-GFAP (1:100, Abcam cat. #ab10062) and rabbit anti-calbindin (1:1,000, Swant cat. #CB38). Secondary antibodies were used at 1:500: donkey anti-goat Cy3 (Jackson ImmunoResearch, 705-165-003), goat anti-rabbit Cy2 (1:125-246), goat anti-mouse Cy3 (115-165-146) and goat anti-mouse Cy5 (115-175-146). All animals were housed in groups of 4-5, given access to standard laboratory chow and water ad libitum and housed in a humidity and temperature-controlled room on a 14/10 h light/dark cycle. All experiments were conducted in accordance with an approved protocol from the Duke University Institutional Animal Care and Use Committee and guidelines from the National Institutes of Health for the Care and Use of Laboratory Animals.

Luciferase reporter assays. Putative enhancers were cloned in the vector pGL3 for ChIP-n. We cloned mm9 region chr14:56509910–56501851, with a Leica site upstream of lucifase. For the two regions in the Grin2c gene on Chr. 11, we cloned the following mm9 regions at the BamHI or SalI sites down-stream of the luc+ coding sequence: Site #1: chr11:115064061–115064510; Site #2: chr11:11528431–11528780. Luciferase reporters were transfected into cultured CGNs on +1 DIV by calcium phosphate precipitation as described9. Co-transfection of TK-renilla luciferase (Promega) was used to control for transfection efficiency and sample handling. As described in the text, in some cases plasmids encoding shRNAs targeting Zic1 in the vector pLKO.1 or empty pLKO.1 were co-transfected with the luciferase plasmids. Data presented are the average of at least three measurements from each of at least two independent experiments.

Chromatin immunoprecipitation. For ChIP we pooled cerebellum or cortex from three P7 mice or two P60 mice for each biological replicate. Brain samples were dounce in 1% formaldehyde (wt/vol) PBS buffer and kept at 25 °C for 15 min, washed twice with cold PBS, then lysed in 600 μl lysis buffer (1% SDS (wt/vol), 10 mM EDTA, and 50 mM Tris, pH 8.1). The crosslinked material was sonicated with a Bioruptor (Diagenode) with 30 s on/off cycles to an average size range of 150–350 bp as visualized by agarose gel electrophoresis. Sonicated supernatants were diluted tenfold in dilution buffer (0.01% SDS, 1.1% Triton X-100, 1.2 mM EDTA, 16.7 mM Tris-HCl, pH 8.1, 167 mM NaCl) before immunoprecipitation. 6 μl of antibody (anti-Zic 1/2 C terminus, courtesy of R. Segal, Harvard Medical School59, total rabbit IgG, Millipore 12-370, or anti histone H3K27Ac, Abcam Ab472) was first incubated with 100 μl of Dynabeads Protein G (Invitrogen 10004D) beads for 4 h at 4 °C, then the antibody conjugate was added to 6 ml of cell lysate for overnight IP. Standard TruSeq adapters were ligated for library preparation, which for Zic used the MicroPlex Library Preparation Kit (Diagenode), and for H3K27ac used the NEB Library Preparation Kit (NEB 6240S and NEB B7335s). 50 bp single-end sequencing was performed at the Duke Sequencing and Analysis Core Resource on a Hi-Seq 2000 machine. Two independent biological replicates were performed for each antibody, developmental time point, and brain region.

RNA interference. For the knockdown of Zic1 and Zic2 transcription factors, we purchased shRNAs targeting mouse Zic1 and Zic2 that were cloned in the lentiviral vector pLKO.1 (Thermo Scientific). Zic1 shRNA (TRCN0000096685), Zic2 shRNA (TRCN0000095251). Empty pLKO.1 was used as the control. Viral shRNAs were packaged as lentiviruses in HEK293T cells (ATCC) following standard procedures. Neurons were infected with lentivirus at a multiplicity of infection of 1 on +1 DIV, and samples were harvested for RNA or protein analysis on +7 DIV.
Quantitative RT-PCR. 800ng of total RNA was used for reverse transcription with oligo dT primers and Superscript II (Invitrogen). Quantitative SYBR green PCR was performed on an ABI 7300 Real-time PCR machine (Applied Biosystems) using the following intron-spansing primers (IDT): Grin2b, F: GAGCATAATTCCCGGCTACTC; R: AAGCCACCGTGTC- CCTATCTC; Zic1, F: CAGAGCCAGATCCGGACACTTATG; R: GAGACCTCGG- AACTGCGCTTGA; Zic2, F: GGCGGCGCAGTTCCACACGAGTA; R: TTTGCCACACCGGGAAAAACAG; Grin2c, F: TGGTGAGGCTTTCTTGCCTGT- CATRC; R: TGGCATTTAGACCTGTCGCAACAAA; Tubb, F: GGCGGGTGATAAG- GCATGAAA; R: CCCAGGCTTCTAGATCCACCA; Gapdh, F: CATGGCCCTTCCGCGT- TTCTC; R: TGTAGTCTACATATGCGAGGTT. Each sample was measured in triplicate, and all data were normalized to the expression of the housekeeping gene Gapdh.

Western blotting. 1 million CGNs or 10 mg cerebellar or non-cerebellar brain tissue was lysed in RIPA buffer to a final concentration of 10 mg ml⁻¹, then disrupted by sonication. 100 μg total lystate was run for SDS-PAGE and transferred to nitrocellulose for western blotting. Actin was used as a loading control. Primary antibodies were mouse anti-Zic1/2 C terminus; mouse anti-Actin (1:10,000; Millipore MAB1501). Secondary antibodies were goat anti-rabbit 1:770 (Biotium 200708) and goat anti-mouse 680 (Biotium 20065). Bands were visualized with fluorescent secondary antibodies using the Odyssey imaging system (LI-COR Bioscience).

Cas9-based RNA-guided gene activation. The CRISPR web interface from the Zhang lab at MIT (http://crispr.mit.edu) was used to design CRISPR guide RNA (gRNA) followed by a PAM site (NGG) that have minimal off-target matches. We used an enzymatically dead Cas9-VP64 (gRNA) followed by a PAM site (NGG) that have minimal off-target matches. We for increased resolution within peak calls. 250-bp windows with 50-bp overlaps were chosen of top 100,000 DHS peaks identified in the samples under consideration served by the first 5-prime base of each read) summed in the 250-bp windowed union of all other DHS sites were considered distal.

RNA-seq expression pattern clustering. RNA-seq data was analyzed in part with the cummeRbund R package (http://compbio.mit.edu/cummeRbund/) operating on Cuffdiff outputs. For cluster profiles, all significantly differential genes from the in vivo time course were clustered by IS distance to a priori defined expression vectors (for example, [0, 0, 0] for elevated expression in P14 relative to P7 and P60). The 100 best match genes from each vector were plotted and subjected to the Database for Annotation, Visualization and Integrated Discovery (DAVID) to find enriched Cellular Component and Biological Process gene ontologies. Principal components analyses (PCA) of gene expression were performed in cummeRbund.

Relationship between differential DHS and gene expression changes. To link differential DHS sites with gene expression changes, each DHS site was associated with its nearest UCSC mm9 reference gene. The fold change in each gene's FPkm values from P7 to P60 was computed. These percentages were compared to the percent overlap obtained from random sampling from all expressed genes (10,000 iterations).

Cell type–enriched gene expression comparisons. Purkinje cell, Bergmann glia, and granule neuron (CGN) enriched gene lists were obtained from. Sequenced RNA abundances captured with translating polysomes (TRAP-seq) in each cell type were compared with the DESeq package. This included 922 Purkinje cell, 2084 Bergmann glia, and 986 CGN enriched genes. For beadplots of DNsase-seq, H3K27ac, and H3K4me3 CHIP-seq data from P56 C57BL/6 cerebellum were retrieved from the Mouse ENCODE project August 2012 release from Ludwig Institute for Cancer Research (B. Ren laboratory). Available peak call files were used for determining overlap fractions with DHS sites in this study.

Motif-finding in DHS sites. Enriched motifs in differential DHS sites were identified using MEME-ChIP with a third-order Markov model background of all DHS sites. The top 10 Discriminative DNA Motif Discovery (DREM) hits by e-value were selected and aligned to known motifs contained in the JASPAR vertebrates and UniPROBE mouse databases.
and requiring unique alignments. bigWig coverage tracks were normalized by total number of mapped reads. Captured fragment length shifts were estimated for each library by cross-strand correlation maxima\(^{61,62}\) and used for peak calling with MACS v1.4.2 (ref. 63) with matched input controls and default \(P < 0.00001\) threshold for each replicate. Cross-strand correlation maxima were also used for quality control to assess signal to noise ratio of each ChIP-seq experiment and samples displaying low scores were omitted. For differential Zic binding detection, raw read counts mapped to the union of Zic ChIP-seq peak calls for all P7 and P60 replicates were used as input to DESeq\(^56\). Differential binding events between P7 and P60 were determined by DESeq negative binomial test at a FDR < 0.05 threshold. To correlate with expression changes observed, Zic binding sites were linked to the nearest expressed gene.

**Statistical analysis.** Individual statistical tests employed are noted in figure legends. Two-sided unpaired \(t\) tests were used for comparisons of normal distributions and Mann-Whitney or exact tests were used for non-normally distributed sequencing data. No statistical methods were used to predetermine sample sizes but our sample sizes are similar to those reported in previous studies\(^{44,49,64}\). Data collection and analysis were not performed blind to the conditions of the experiments. Order of sequencing sample collection was random but no other randomization of data was performed.

A Supplementary Methods Checklist is available.

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