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Genomic Resolution of DLX-Orchestrated Transcriptional Circuits Driving Development of Forebrain GABAergic Neurons

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SUMMARY

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AUTHOR CONTRIBUTIONS
Conceptualization, A.S.N., S.L., R.C.-P., and J.L.R.R.; Methodology and Investigation, S.L. (ChIP-seq, RNA-seq, ISH, AB characterization, luciferase assays), H.T. (enhancer CRISPR-i), J.D.P. (ISH, RNA-seq), S.N.S. (preliminary histone ChIP-seq), G.L.M. (preliminary DLX2 ChIP-seq), D.E.D., L.A.P., and A.V. (enhancers in mice), R.C.-P. (bioinformatics, modeling), and L.S.-F. (bioinformatics); Software, R.C.-P. and A.S.N.; Formal Analysis, S.L., R.C.-P., A.S.N., and J.L.R.R.; Writing – Original Draft, S.L., R.C.-P., A.S.N., and J.L.R.R.; Writing – Review & Editing, S.L., R.C.-P., H.T., J.D.P., V.G., D.E.D., A.S.N., and J.L.R.R.; Funding Acquisition, J.L.R.R. and A.S.N.; Resources, V.G. and M.T.M. (unpublished dCas9KRAB mice); Supervision, J.L.R.R. and A.S.N.

DECLARATION OF INTERESTS
J.L.R.R. is co-founder, stockholder, and currently on the scientific board of Neurona, a company studying the potential therapeutic use of interneuron transplantation. The other authors declare no competing interests.

SUPPLEMENTAL INFORMATION
Supplemental Information can be found online at https://doi.org/10.1016/j.celrep.2019.07.022.
DLX transcription factors (TFs) are master regulators of the developing vertebrate brain, driving forebrain GABAergic neuronal differentiation. Ablation of Dlx1&2 alters expression of genes that are critical for forebrain GABAergic development. We integrated epigenomic and transcriptomic analyses, complemented with *in situ* hybridization (ISH), and *in vivo* and *in vitro* studies of regulatory element (RE) function. This revealed the DLX-organized gene regulatory network at genomic, cellular, and spatial levels in mouse embryonic basal ganglia. DLX TFs perform dual activating and repressing functions; the consequences of their binding were determined by the sequence and genomic context of target loci. Our results reveal and, in part, explain the paradox of widespread DLX binding contrasted with a limited subset of target loci that are sensitive at the epigenomic and transcriptomic level to Dlx1&2 ablation. The regulatory properties identified here for DLX TFs suggest general mechanisms by which TFs orchestrate dynamic expression programs underlying neurodevelopment.

**In Brief**

Lindtner et al. reveal the regulatory wiring organized by DLX transcription factors in forebrain GABAergic neuronal specification, by integrating functional genomic, epigenomic, and genetic data on a transgenic mouse model. This network determines key sequence-encoded regulatory elements and implicates a combination of histone modifications and biophysical interactions.

**Graphical Abstract**
INTRODUCTION

The development and function of all forebrain GABAergic neurons depend on a transcriptional program that is distinct from other regions of the CNS (Long et al., 2007; Silberberg et al., 2016). Near the top of this transcriptional circuit lie the DLX homeodomain transcription factors (TFs) (Long et al., 2007). Four of the six Dlx genes encoded by the mammalian genome (Dlx1, Dlx2, Dlx5, and Dlx6) are expressed in the embryo by highly specific progenitor domains in the telencephalon, hypothalamus, and prethalamus (Eisenstat et al., 1999; Liu et al., 1997; Panganiban and Rubenstein, 2002). Within the embryonic telencephalon, these are the subpallial domains known at the ganglionic eminences (GEs), septum and preoptic area. The three GEs, the lateral (LGE), medial (MGE), and caudal (CGE), are the anlage for the projection neurons of the striatum, pallidum, and parts of the amygdala (Silberberg et al., 2016), as well as the interneurons that migrate to various structures including the neocortex and hippocampus (Lim et al., 2018).

Dlx RNA and DLX protein expression in the mouse GEs begin in progenitors at embryonic day 9.5 (E9.5) and follows a temporal program of Dlx2, Dlx1, Dlx5, followed by Dlx6 (Eisenstat et al., 1999; Liu et al., 1997). Their expression in the GEs overlaps in the secondary progenitor domain called the subventricular zone (SVZ), suggesting that the different DLX TFs compensate for each other. Indeed, while single Dlx mutants have relatively mild forebrain phenotypes that affect subsets of GABAergic neurons (Cobos et al., 2005; Qiu et al., 1995; Wang et al., 2010, 2011), Dlx1/2−/− and Dlx5/6−/− double mutants have strong forebrain phenotypes (Anderson et al., 1997a; Wang et al., 2010). Constitutive Dlx1/2−/− mutants have poorly differentiated basal ganglia and lack most cortical interneurons (CINs). Furthermore, Dlx1/2−/− mutants fail to express Dlx5 and Dlx6 (Anderson et al., 1997a), thus generating a Dlx-null state, allowing one to evaluate the transcriptome and epigenome in this “ground state” lacking all DLX proteins.

Previous studies of Dlx1/2−/− mutants identified GE RNA expression defects at E15.5 using array technology (Long et al., 2009b). These were coupled with RNA in situ hybridization (ISH) to define high-resolution analyses of specific developmental effects in distinct regions and layers of the GEs (Long et al., 2009a, 2009b). This analysis led to hypotheses about potential TF regulatory networks downstream of the DLX TFs but lacked critical information about which transcription start sites (TSSs) and distant regulatory elements (REs) (e.g., enhancers and silencers) are bound by DLX proteins during GE development.

Recent single-cell RNA-sequencing (scRNA-seq) experiments have illuminated the transcriptional changes in developing GE cells, highlighting the role of key TFs including the DLX proteins (Chen et al., 2017; Mayer et al., 2018; Mi et al., 2018). However, neither scRNA-seq nor classical genetic studies have captured mechanisms underlying transcriptional wiring. Thus, major questions remain regarding how GE cell fate is encoded at genomic, epigenomic, and transcriptomic levels by TFs such as the DLX proteins. To address this need and characterize the genomic transcriptional wiring orchestrated by DLX TFs, we integrated functional genomics and genetic approaches to dissect the role of DLXs in the developing GE. We identified DLX2, DLX1, and DLX5 binding sites and target genes across GE development and measured the responses of these loci to the absence of DLX.
proteins using transcriptomic and epigenomic methods and *in vivo* and *in vitro* studies of DLX-mediated RE function.

**RESULTS**

**DLX2, DLX1, and DLX5 Share Binding Properties in GEs and Regulate GABAergic Neuron Specification via Activating and Repressive Functions**

Toward elucidating how DLX TFs regulate transcriptional programs in the GEs, we used chromatin immunoprecipitation sequencing (ChIP-seq) and RNA-seq to compare DLX binding, chromatin state, and transcription in the GEs from wild-type and *Dlx1/2*−/− embryos (Figure 1A).

For ChIP, micro-dissections including MGEs, LGEs, and CGEs were prepared from wild-type (WT) E11.5, E13.5, and E16.5 embryos to compare DLX binding patterns during GE regional patterning and neurogenesis. We generated antibodies with specificity for DLX2, DLX1, and DLX5, as verified using immunofluorescence of E13.5 GE from WT, *Dlx2*−/−, *Dlx1*−/−, and *Dlx5*−/− mice and western blot (Figure S1). We were unable to obtain ChIP-grade DLX6-specific antibody. ChIP-seq was performed and libraries were prepared for each DLX at each time point along with input and negative controls including IgG as well as three distinct blocking peptides for DLX1, DLX2, or DLX5 ChIP-seq. These blocking peptides were the antigen used in the generation of the antibodies. Finally, we performed DLX2 ChIP-seq on WT and *Dlx2*−/− at E16.5; the results showed the disappearance of DLX2 specific peaks in the knockout (KO) (data not shown). Genome-wide binding patterns of all three DLXs showed high reproducibility across biological replicates and largely overlapping ChIP-seq enrichment patterns; *Gad2* and *Lhx6* loci are shown as examples (Figure 1B).

We generated a merged set of DLX2, DLX1, and DLX5 interaction sites that represented the union of all called peaks that passed strict significance thresholds (Table S3). DLX binding was enriched near loci associated with brain development in general and with more specific process, such as GABAergic specification. Target regions include promoter elements (18.1% peaks, 3,064 peaks, 7,671 total genes) and distal elements within intronic and intergenic regions (79.4% peaks, 13,440 peaks). All three DLXs showed similar patterns of an increasing number of genomic binding sites as development progressed from E11.5 through E16.5 (Figure 1C).

Many genomic regions showed DLX interaction at lower levels of enrichment. We compared the overlap of DLX binding targets for low-, medium- (the default set), and high-affinity peaks based on ChIP-seq enrichment (Figure 1F). We defined the level of affinity (stringency) by filtering out peaks above selected p values (10−5, 10−15, and 10−40 for low, medium, and high affinities, respectively) from the composition of the merged peak set across DLX ChIP-seq datasets. Using the low-affinity threshold, DLX binding was extensive across the genome (46,143 peaks), with reduced overlap among DLX2, DLX1, and DLX5 (27% common peaks). Overlap increased to 57% for medium-affinity (n = 16,919) and to 91% for high-affinity peaks (n = 4,570). Consistent with extensive binding overlap, there was high correlation of normalized ChIP-seq signal across DLXs (pairwise r^2 values > 0.7
for all comparisons; Figure 1D). Hierarchical clustering identified separation between E11.5 and the two later time points, followed by branching that separated DLX2 from DLX1 and DLX5.

Comparison of peak calls and genome-wide signal suggests that differences in ChIP-seq across DLXs are largely quantitative. Figure 1E is a heatmap representation separated for DLX2, DLX1, and DLX5 showing increasing binding across development across DLX TFs. Subthreshold enrichment across DLXs is present when comparing peaks that are only called in one or two of the three DLXs, further indicating that the majority of binding targets are shared (Figure S2). Notably, we identified a minority of DLX2, DLX1, and DLX5 peaks that showed specific DLX binding, although these peaks did not show enrichment for specific annotation classes. While our findings indicate that the great majority of DLX binding profiles overlap, corroborating the observed redundancy of these TFs, binding patterns may diverge in subsets of cells or later in development.

*De novo* motif analysis identified a core homeobox motif as the putative shared binding DNA target for all three DLXs (Figures 1H and S3). This primary TAATTA motif was strongly enriched at the center of DLX peaks and is similar to DLX binding motifs determined via SELEX analysis (Feledy et al., 1999). Highly similar homeobox motifs were also identified, suggesting that DLX TFs tolerate binding target sequence variation. Secondary motifs were also enriched within DLX peaks, suggesting combinatorial TF binding occurs at DLX target loci.

**Relationship between DLX Binding and DLX-Regulated Gene Expression**

We compared RNA-seq from WT and *Dlx1/2<sup>−/−</sup>* GE at E13.5 (n = 4 per genotype), identifying 328 differentially expressed (DE) genes at false discovery rate (FDR) < 0.05 (Figure 1G; Table S1). Downregulated genes (n = 181) were strongly enriched for terms associated with GABAergic neuron specification, while upregulated genes (n = 147) did not show enrichment for similarly specific terms (Table S2). Both upregulated and downregulated genes were enriched for annotation terms associated with general brain development. These DE results were largely consistent with previous microarray-based profiling comparing E15.5 WT and *Dlx1/2<sup>−/−</sup>* GEs (Long et al., 2009b).

There was a reduced distance between *Dlx1/2<sup>−/−</sup>* DE genes and DLX binding sites, although only for medium- and high-affinity peaks (Figure 1I), suggesting that the observed widespread lower-affinity DLX binding across the genome may be less functionally relevant than higher-affinity targets. Further supporting increased relevance of the higher-affinity DLX targets, there was a trend of increasing evolutionary conservation for DLX-bound promoter and distal elements as affinity increased for these peak sets (Figure 1J). The shared binding patterns and target motifs between DLX TFs suggest a model of competitive binding of these factors directing transcriptional activation and repression during basal ganglia and CIN development.
Loss of DLXs Leads to Changes in Chromatin State and Transcription at a Small Subset of Target Loci via DLX Binding to Distal and Proximal Regulatory Sequences

To test whether chromatin state is dependent of DLX binding, we used histone ChIP-seq to profile four histone post-translational modifications (PTMs) associated with REs and active or repressive state (H3K4me1, H3K4me3, H3K27ac, H3K27me3) in WT and Dlx1/2−/− E13.5 GEs (Figure 2A). Using WT ChIP-seq datasets, we segmented the genome into chromatin states via ChromHMM to analyze DLX binding context. Next, we identified regions that showed differential histone PTM signal in Dlx1/2−/− GE (Table S5). ChrX showed chromosome-wide differences, most likely from imbalanced sex representation in the pooled tissues used to generate these datasets. We thus restricted WT versus Dlx1/2−/− chromatin state genome-wide analysis to autosomes. Loci with significant changes in histone PTM signal in Dlx1/2−/− mutants were generally located nearby DLX peaks, suggesting that histone PTM and chromatin state changes in Dlx1/2−/− mutant GEs are directly associated with DLX binding.

In contrast to the widespread DLX binding across the genome, few DLX TF target loci (6.2%) exhibited significant changes in histone PTM signal in Dlx1/2−/− mutants (Figure S5). The finding that chromatin is unchanged in Dlx1/2−/− mutants for the majority of DLX TF binding parallels results from the intersection between DE genes and DLX TF binding, where most genes near binding sites do not show altered expression. These results suggest that, somewhat surprisingly, the majority of DLX TF binding events do not appear required for normal chromatin state or gene expression at E13.5.

We identified 510 DLX binding sites that were located within regions exhibiting loss of activating (H3K27ac, H3K4me3) and or gain of repressing (H3K27me3) signature and 574 sites with chromatin changes in the opposite direction (Table S3). We define the loci where DLX binding was required for establishment or maintenance of active transcription state as activating REs (a.REs) and the loci where DLX binding was associated with interfering with transcription as repressive REs (r.REs). As examples, the Slc32a1 (Vgat) and Otp loci feature putative REs (a.RE and r.RE, respectively) where DLX binding was present and show significant changes to histone PTM and gene expression in Dlx1/2−/− mutants (Figure 2B). Histone PTM signal averaged across a.RE and r.RE loci changed in all four histone PTMs (Figures 2C and S5). The nearest gene to a.RE and r.RE sites was more likely to show downregulation and upregulation, respectively, in the Dlx1/2−/− mutants compared to either all DLX binding sites or high-affinity sites (Figure 2D). Most genes with strong expression changes in Dlx1/2−/− mutants were located nearby one or multiple a.REs or r.REs, as represented by red or blue circles, respectively, overlaid on the volcano plot of all DE genes, with circle size indicating significance (likelihood ratio) of differential histone signal (Figure 2E).

There were differences in function of genes associated with a.REs and r.REs (Figure 2F). a.REs were enriched for genes associated with neuronal maturation and function. In contrast, r.REs were enriched near genes associated with brain patterning, neuron fate commitment, and TF and chromatin remodeler activity. Among r.RE targets, there was a strong enrichment for other TFs (e.g., Otp, Gsx1, Pax7). In comparison, a.RE targets were a combination of TFs (e.g., Dlx5, Sp8) and lineage-specific genes (e.g., Gad2, Slc32a1, and
Nrxn3). This suggests that DLX TFs activate and repress coherent gene sets representing the gene regulatory network required for transitioning from progenitor to post-mitotic neurons and for maturation of GABAergic cell types.

Genes associated with a.RE and r.RE loci were additionally enriched for disease ontology terms, with increased representation of autism among a.RE and glioma and intellectual disability for r.RE loci (Figure S6). a.REs and r.REs are additionally enriched for genes linked to autism via patient mutations (p values of 0.007 and 0.03, respectively, based on permutation test), and 25/99 (25%) of high-confidence autism spectrum disorder (ASD) risk gene TSSs are within 100 kb of an a.RE or r.RE (Figure S6; Table S10).

Validation of a.RE-Mediated Activation of Nrxn3 by DLX TFs

Nrxn3 mRNA expression was decreased in Dlx1/2−/− mutants, which was accompanied by change in local chromatin state at Nrxn3 putative regulatory regions bound by DLX TFs (Figure 3A). To verify our regulatory model of DLX-mediated activation or repression via binding to REs, we used in vivo and in vitro approaches to assess changes in Nrxn3 expression and DLX-dependent a.RE activity.

First, we performed Nrxn3 RNA ISH on E13.5 WT and Dlx1/2−/− mutants (Figure 3B). Nrxn3 expression in the SVZ of the GEs was strongly reduced in Dlx1/2−/− mutant. Next, we assessed the in vivo activity of an intragenic Nrxn3 element we identified (mm1203; 1,450 bp) that had robust DLX binding and properties of a DLX-dependent a.RE. For the mm1203 locus, there was H3K27ac and H3K27me3 enrichment in WT GE, and reduced H3K27ac and increased H3K27me3 in Dlx1/2−/− mutants. In a transgenic mouse assay, mm1203 (driving LacZ expression) had robust and reproducible (7/7) activity in the E12.5 telencephalon, with additional activity in the eye and a small domain in the hindbrain (Figure 3C). Tissue sections showed strong enhancer activity in the SVZ and mantle zone (MZ) of the LGE and in tangentially migrating cells (probably immature CINs), and weaker activity in the MGE.

Next, we performed luciferase transcription assays in postnatal day 19 (P19) tissue culture cells, a mouse teratocarcinoma cell line, to test whether DLX proteins could regulate the activity of mm1203. We generated a vector in which mm1203 is upstream of a minimal promoter and luciferase reporter gene (pGL4.23; Invitrogen). We transfected the mm1203 luciferase reporter vector alone or in the presence of plasmids expressing DLX2, DLX1, or DLX5 (Figure 3D). DLX2, DLX1, and DLX5 activated luciferase expression by 225-, 5-, and 6-fold, respectively, in the presence of mm1203.

Validation of Sequence-Encoded and Context-Dependent DLX Regulation of Arx

After verifying DLX-mediated activation of Nrxn3 expression, we next sought to verify sequence- and context-dependent activating and repressing function of DLX binding. To examine this, we tested the impact of DLX co-transfection with four Arx enhancers. The Arx locus is on chromosome X, and there are a number of shared DLX binding sites in the region (Figure 3E). Arx exhibits both cortical ventricular zone (VZ) and GE SVZ and neuronal expression in the developing telencephalon. There are four ultraconserved REs that regulate Arx expression (Colasante et al., 2008; Dickel et al., 2018). In transgenic mouse
assays, two of these Arx REs drove reporter expression in the GE (hs119 and hs121), whereas the other two were specific to the developing cortex (hs122 and hs123).

Only the GE enhancers exhibited decreased H3K27ac in Dlx1/2−/− mutants. This suggests that while the DLX TFs bind to both the GE- and cortex-specific REs, DLX binding only had an activating effect on the GE-specific REs. When tested in the luciferase co-transfection assays, hs119 was strongly activated by DLX2, DLX1, and DLX5 (290-, 94-, and 42-fold, respectively). hs121 showed specific activation by DLX2 and DLX1 (57- and 7-fold, respectively). On the contrary, neither of the cortical REs, hs122 and hs123, were activated by the three DLX TFs. These findings provide evidence that DLX-dependent activating and repressing activity in the GE is encoded by the primary DNA sequence of a.RE and r.RE elements.

### DLX-Dependent a.REs and r.REs Are Distinguished by Genomic Sequence and Context

After verifying the biological relevance of DLX-mediated a.RE and r.RE function regulating Nrxn3 and Arx, we compared Dlx1/2−/− sensitive elements (a.REs and r.REs) to each other and to DLX-bound loci that do not exhibit perturbation to chromatin state in Dlx1/2−/− mutants.

First, we tested differential representation of DNA binding motifs within a.RE and r.RE regions and between those RE regions to the ones that showed no histone PTM change to identify which motifs might distinguish activating from repressing function. For this, we considered motifs identified across all DLX peaks (Figure 1D) and added additional motifs identified specifically in a.RE (E2F3, ASCL1) or r.RE (FOXA1, GSC, OTX2) peaks (Figure S3). Using a cutoff of 1.2-fold enrichment between a.RE and r.RE regions, we identified motifs that showed directional enrichment (Figure 4A). In addition to the a.RE- and r.RE-specific motifs listed above, additional motifs showed stronger enrichment in a.REs, notably the family of basic-helix-loop-helix (bHLH) sites that includes ASCL1, AP4, NEUROG2, and OLIG2 motifs. While strongly enriched in both a.RE and r.RE peaks, there was additional increased enrichment of the primary DLX binding motif (TAATTA) in a.RE regions. Motifs that showed distinct a.RE versus r.RE enrichment were also enriched in a.RE or r.RE sequences compared to DLX-bound regions without histone signature changes.

We tested for features associated with genomic and epigenomic context of the DLX-bound loci that differentiate a.RE and r.RE elements. First, we tested for differences between promoter and distal binding patterns and across local chromatin states as identified by ChromHMM for all DLX peaks, high-affinity peaks, and a.RE and r.RE loci (Figure 4B). a.REs were enriched at distal regions (89%) and within regions with strong H3K27ac. In contrast, r.REs were skewed toward promoters (49%), and specifically toward bivalent (H3K4me3+, H3K27me3+) and active (H3K4me3+, H3K27me3−) promoters.

We assessed whether a.RE and r.RE loci and DE gene promoters showed increased local density of DLX peaks (Figure 4C). We found increased density of DLX peaks within 50 kb of DE gene promoters compared to randomly sampled matched gene sets (all gene mean, 1.7; DE mean, 2.8). Both a.RE and r.RE loci had increased density of local DLX peaks (Figure S5; Table S6). We tested whether a.RE and r.RE were more likely to be associated
with higher-affinity DLX binding. a.RE sites were enriched in the top quintile of DLX peaks, while r.RE loci did not show association with DLX peak strength (Figure S5). Finally, we examined correlation between a.RE and r.RE loci and histone PTM signal in WT GE, finding that a.RE and r.RE were enriched among regions with stronger histone PTM enrichment for H3K27ac, H3K27me3, and H3K4me3, but not for H3K4me1 (Figures 4D and S6).

We used logistic regression to evaluate the value of the sequence and context factors identified here to discriminate a.RE and r.RE loci (Figure 4E). For both a.RE and r.RE, chromatin context features were the strongest contributor to prediction accuracy, while DLX peak affinity and local DLX peak density were strong predictors for a.RE but not r.RE. DNA sequence motifs contributed to both models, although had less independent predictive power. Thus, sequence and context features of loci bound by DLXs were associated with the likelihood of sensitivity of these loci to DLX loss, enabling prediction of which DLX binding events have functional impact on chromatin state and gene expression in E13.5 GE without data from Dlx1/2−/− samples.

Interaction between Multiple DLX-Responsive a.REs and the Sp8 Promoter

We identified multiple loci featuring DLX binding in a >400-kb gene desert upstream of the Sp8 locus (Figure 5A). Sp8 is among the most significantly downregulated genes in Dlx1/2−/− mutants (logFC = −2.4; p = 2.24E-51), with ISH analysis showing reduction in the SVZ and MZ (Long et al., 2007, 2009b). Three of the DLX-bound loci in this region overlap with previously characterized GE enhancers (hs110, hs1007, and hs1226) (Figure 5B). hs110 and hs1007 are among the most statistically significant DLX-responsive a.REs, with near total loss of H3K27ac in Dlx1/2−/− mutants. H3K27ac in the region surrounding Sp8 also decreased in the Dlx1/2−/− mutant (Figure 5A). In contrast, for hs1226, only DLX2 had significant ChIP-seq signal and this region did not show H3K27ac changes in the Dlx1/2−/− mutant.

We tested the ability of these three REs to regulate Sp8 transcription in the LGE using CRISPR interference (CRISPR-i). We grew primary cultures from E13.5 dorsal LGE from mice expressing dCas9KRAB (V.G., M. Lebedinskaya, R. Wagner, R. Jaafar, M.T.M., unpublished data). When in proximity to a TSS, dCas9KRAB downregulates transcription through steric hindrance, histone methylation, and deacetylation (Gilbert et al., 2014; Rosenbluh et al., 2017; Thakore et al., 2015). Cultured cells were infected with a lentivirus encoding a guide RNA that targeted a specific RE or the Sp8 promoter. As a negative control we targeted a guide RNA (gRNA) to a limb RE in the same region (hs1148) that was weakly bound by DLX2. Transduction efficiency was >90%. 48 h later, we analyzed Sp8 RNA levels using qRT-PCR.

gRNA targeting Sp8 promoter reduced Sp8 RNA up to 6-fold (Figure 5C). All three gRNAs targeting GE enhancers also reduced Sp8 RNA. gRNA for hs110 (the highest-affinity DLX peak) downregulated Sp8 expression up to 4-fold, while gRNA for hs1007 and hs1226 reduced Sp8 expression by ~3-fold. gRNA for the limb enhancer (hs1148) did not change Sp8 RNA levels, providing evidence of gRNA/CRISPR-i-induced Sp8 downregulation in LGE cells that was specific to the DLX-bound GE enhancers (hs110, hs1007, and hs1226).
To directly test whether DLXs positively modulate gene expression via binding to hs110, hs1007, and hs1226, we used the luciferase transcription assay (Figure 5D). DLX2 activated hs110, hs1007, and hs1226-dependent luciferase expression 50-, 6-, and 4-fold, respectively. DLX1 and DLX5 also activated hs110 and hs1007, although to a lower extent than DLX2. Neither DLX1 nor DLX5 activated hs1226, correlating with the lack of DLX1 or DLX5 binding at this enhancer.

These results provide evidence that multiple DLX binding sites (hs110, hs1007, and hs1226) contribute to activation of Sp8 via DLX-mediated activation and physical interaction with the promoter. These findings are consistent with the model that DLX binding is critical for establishing robust transcriptional regulation for a.RE-associated loci.

**DLX TFs Mediate Spatial Activation and Repression of Core Regulatory Targets**

We used ISH to examine spatial changes in expression of genes perturbed in E13.5 GEs in *Dlx1/2−/−* mutants, testing 6 candidates for this study; results from 2 downregulated (*Arl4D and Grik3*) and 2 upregulated genes (*Tox and Lhx9*) are shown in Figure 6A. For all 6 targets (including those in Figure S7), ISH validated predicted change in expression in *Dlx1/2−/−* GE. Combining our results with published ISH analyses of *Dlx1/2−/−* mice at E15.5 (Long et al., 2009a, 2009b) generated a set of 50 predicted downregulated and 35 predicted upregulated genes. We scored each gene for increased or decreased expression in the VZ, SVZ, and MZ to test for consistent patterns of spatial change (Figure 6B). Genes with increased RNA in the *Dlx1/2−/−* mutants were consistently upregulated in ISH (29, 83%), with expression increased across GE layers, but most frequently in SVZ. Of the upregulated genes, 5 (*Otp, Gsh1, Ebf3, Gbx2, and Pax7*) were not expressed in WT GEs, and thus were ectopically expressed in absence of DLX TFs. No downregulated genes showed ectopic expression. Downregulated DE genes were validated with reduced expression via ISH (43 genes, 86%), but a different spatial pattern of strongest decrease in the MZ followed by SVZ. These results confirm RNA-seq changes and are consistent with DLX TFs’ role in controlling expression along the spatial axis of differentiation.

We next examined results from transgenic RE assays to characterize a.RE and r.RE activity as described above. We combined REs from the VISTA enhancer database (Visel et al., 2013) and two recent studies (Pattabiraman et al., 2014; Silberberg et al., 2016). Across all tested enhancers, both a.RE (24/30, 80%) and r.RE (31/39, 79%) had high rates of *in vivo* enhancer activity at E11.5. Spatial GE expression patterns in the brain were available from brain sections generated for 5 a.REs and 11 r.REs (Figures 6C, 6D, and S7). For these DLX-bound enhancers, a.REs and r.REs showed characteristic differences in activity in the telencephalon. a.RE activity in the GEs was restricted to SVZ and MZ, whereas r.RE activity in the GEs included the VZ (Figure 6D). Thus, changes in spatial expression of DLX TF-regulated genes were correlated with activity of DLX-mediated a.RE and r.RE loci.

**DISCUSSION**

The DLX TFs are master regulators of forebrain GABAergic neuronal development (Anderson et al., 1997a, 1997b; Cobos et al., 2007; Long et al., 2009a, 2009b; Pla et al., 2018; Yun et al., 2002). Our results show how DLX TFs organize a master regulatory
program by modulation of chromatin and transcriptional state, activating proneural transcriptional programs and repressing proliferative and alternative fate transcriptional programs.

Our integrated approach suggests a model for DLX function and enable construction of gene regulatory networks organized by DLX TFs in the GE (Figure 7A). As Dlx2/1/5 are co-expressed in individual cells, data indicate that they compete for binding to homeobox DNA motifs at shared genomic targets that expanded across development. DLX TF binding could drive either transcriptional activation (largely via interactions at distal elements) or repression (largely via binding near the TSS), while most DLX binding sites did not show a large effect with regard to changes in chromatin or transcriptional state in Dlx1/2−/− GEs. In contrast, a subset of loci showed chromatin and transcriptional states that were highly sensitive to the Dlx1/2−/− state.

Our analysis highlights emergent features of regulatory DNA sequence and chromatin context that in part explain variability among individual DLX binding targets in sensitivity to DLX loss. Using in vivo and in vitro approaches, we show that the sensitivity of REs to DLX-mediated activation or repression (i.e., a.RE and r.RE loci) is encoded by their DNA sequence. TF motifs associated with a.REs (e.g., ASCL1 and OLIG2) and r.REs (e.g., OTX2) are TFs with major functions in GE biology (Casarosa et al., 1999; Hoch et al., 2015; Long et al., 2009b; Petryniak et al., 2007). This is consistent with a model of combinatorial and/or competitive binding by TFs and co-regulators.

The transgenic RE and cell culture transcription assays performed here recapitulate our predicted regulatory activity and sensitivity to DLXs for Nrxn3, Arx, and Sp8 loci. In parallel, genomic context of DLX binding sites matters, as local chromatin state and inferred biophysical interactions were among the strongest predictors of epigenomic and transcriptional sensitivity to DLX loss. Specifically, a.RE and r.RE elements were associated with loci that featured clustered DLX binding and high rates of activating and/or repressing histone PTMs.

We propose that principles elucidated for the DLXs represent common characteristics of TFs that organize cellular differentiation and metazoan development. Predicting the impact of regulatory sequence variation, for example in evolution and disease, can be improved via statistical prioritization of TF binding events that are likely to have strong functional relevance. Previous studies found similar disagreement between the binding targets of a TF and transcriptional sensitivity of these targets to loss of the TF (Sandberg et al., 2016; Cusanovich et al., 2014), but they were not able to dissect the underlying nature of this disconnect. Our data are in line with the idea that enhancer-promoter and DNA-RNA-protein interactions underlie robust transcriptional regulation and phase separation for these critical genes (Boija et al., 2018; Hnisz et al., 2017). Our results suggest that the sequence-encoded RE function, in tandem with biophysical interactions, may explain how master TF binding establishes developmental transcription patterns.

The genes that are directly regulated by DLX-mediated a.RE and r.RE activity are strongly enriched for TFs, lineage-defining molecules, and disease-relevant loci. DLX regulatory
activity can be separated by DLX2/1/5 activity in the VZ, SVZ, and MZ. Figure 7B highlights the DLX TF gene regulatory network of TFs, non-TFs, and genes relevant to brain disorders. All of the genes depicted have DLX binding at the TSS or within the genomic region. For the majority of gene loci (45/71 TFs and 18/35 non-TFs), their RNA expression is altered (ISH and/or RNA-seq), and gene-associated RE epigenetic states are changed (a.RE or r.RE) in the Dlx1/2−/−. Below we discuss specific developmental programs that are orchestrated by DLX-dependent transcriptional circuitry.

DLX proteins are nearly ubiquitously expressed in the SVZ (SVZ1 and SVZ2; Eisenstat et al., 1999; Petryniak et al., 2007), where we propose they have a core function in promoting specification of forebrain GABAergic neurons. DLX2 and DLX1 promote activation of Dlx5 and Dlx6 (Anderson et al., 1997b), thereby generating cells with four degrees of DLX redundancy, creating an evolutionarily buffered foundation for ensuring the generation of forebrain GABAergic neurons.

DLX2, and subsequently DLX1, are co-expressed in subsets of VZ and most SVZ progenitor cells, where they promote proneural programs (Eisenstat et al., 1999; Yun et al., 2002). These include transcriptional repression of TF genes that regulate progenitor cell states, including Gsx2, Hes5, Pax6, Olig2, and Otx2. Some of these TFs also regulate forebrain regional fate (e.g. Gsx2, Otx2, and Pax6 [Hoch et al., 2015; Toresson and Campbell, 2001; Yun et al., 2001]) or promote oligodendrogenesis (Olig2 [Petryniak et al., 2007]). DLXs also promote proneural programs through repressing Notch1 and Delta1 (Dll1). In the absence of DLX proteins, the GEs have an ectopic expression of markers of the hypothalamus (Otp), midbrain (Pax7), pallium (Ebf3, Lhx9), as well as LGE ectopic expression of MGE genes (Gbx1, Gbx2) (Long et al., 2009b).

There is evidence for additional transcriptional pathways that drive GE specification and neurogenesis, which are driven by ASCL1 and GSX2. Ascl1, Dlx1/2 triple mutants lose characteristic GE molecular features (Long et al., 2009a; 2009b). Furthermore, removal of Gsx2 in the Dlx1/2 mutants rescues overexpression of Ascl1 (Wang et al., 2013). Neither the Ascl1 or Gsx2 locus show RE epigenetic changes (Figure 7B), suggesting that they function primarily upstream or parallel to Dlx genes.

DLX-driven transcription generates properties fundamental to GE cells, and subsets of their derived neurons, including expression of Gad1, Gad2, SLC32a1 (GABA regulatory genes), and other non-TFs including Ccr4, Ccr7 (Acrk3), Erbb4, Nrx3, and Robo2. DLX activates GE expression of multiple TFs including Arx, Lhx6, Lhx8, Maf, Matb, Sp8, Sp9, and Zeb2 (Zfhx1b), most with distinct functions in specific neuronal lineage. For instance, Maf and Matb regulate development of MGE-derived cortical interneurons; together, they repress the generation of somatostatin+ interneurons (Pai et al., 2019) Sp8&9 specify the identity and differentiation of striatopallidal projection neurons (Xu et al., 2018). DLX TFs additionally repress inhibitors of neuronal differentiation/maturation (e.g., Id2, Id4, and Ascl1).

We provide evidence for DLX-driven expression of neuronal features in the MZ. These include repression of hypothalamic genes (Hmx3/Nkx5−1, Otp), activation of striatal and
pallidal projection neuron genes (Gnal, Gucy1a3, Npas1, Rxr) and interneuron genes (Cxcr4, Cxcr7, ErbB4, Th). The Gene Ontology (GO) terms for genes dysregulated in Dlx1/2−/− (Figure 2F) include axon, synaptic transmission, and neuron apoptotic process; these provide evidence that DLX, through a.REs and r.REs, regulate multiple processes in maturing neurons. Indeed, Dlx mutants have defects in neuritic processes, synapse function, and neuronal survival (Cobos et al., 2007; Pla et al., 2018).

In addition to coordinating specific developmental programs, DLX-dependent transcriptional circuitry regulates many genes that predispose to neurodevelopmental disorders. For example, Arx, Maf, and Zfhx1b are targets of DLXs; mutations in these genes cause human intellectual disability that is linked to abnormal CINs in mouse mutants (Mowat et al., 2003; Niceta et al., 2015; Olivetti and Noebels, 2012). Defining DLX-dependent transcriptional circuitry provides a framework for understanding disease-causing pathways.

Our results demonstrate that while TF ChIP-seq provides critical interaction maps, chromatin interaction alone is not sufficient to elucidate the functional relevance of binding. By identifying the sequence and context constraints and showing that sensitivity to DLX TF ablation can be modeled without incorporating null mutant data, this study provides a path forward to move from TF binding catalogs, to a functional understanding of the gene regulatory wiring driving brain development.

STAR★METHODS

LEAD CONTACT AND MATERIALS AVAILABILITY

Further information and requests for resources should be directed to and will be addressed by the Lead Contact, John L. R. Rubenstein (John.Rubenstein@ucsf.edu).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Mice and genotyping—All procedures and animal care were approved and performed in accordance with National Institutes of Health and the University of California San Francisco Laboratory Animal Research Center (LARC) guidelines. Dlx1+/−, Dlx2+/− and Dlx1/2+/− mice were genotyped as in Qiu et al. (1995); Dlx5+/− mice were genotyped as in Depew et al. (1999). For RNA-seq experiments, an equal number of males and females were used. ChIP-Seq and native histone ChIP-Seq was performed on Mus musculus CD1 strain at developmental stage E11.5, E13.5 or E16.5. The embryos were not assessed genotypically for gender since we used a pool of embryos and therefore expect a roughly equal number of male and females.

Transgenic enhancer assays—Transgenic enhancer assays were performed at Lawrence Berkeley National Laboratory (LBNL) under the approval of the Animal Welfare and Research Committee (AWRC). Mice for transgenic assays were housed at the Animal Care Facility (ACF) of LBNL, monitored daily for food and water intake, and inspected weekly by the Chair of the AWRC and the head of the animal facility in consultation with the veterinary staff. The LBNL ACF is accredited by the American Association for the Accreditation of Laboratory Animal Care (AAALAC). Transgenic assays were performed in the Mus musculus FVB strain at developmental stage E11.5 or E12.5. The resulting embryos
were not assessed phenotypically for gender, which is not outwardly obvious at these ages. Therefore, we expect that a roughly equal number of male and female embryos were assessed.

**METHOD DETAILS**

**DLX antibody production**—Antibodies against DLX1, DLX2 and DLX5 were made in rabbits by GenScript. Antibodies against DLX2 were raised against amino acids (aa) 1–154 and N-terminally tagged with a Maltose binding protein (MBP) (Kuwajima et al., 2006). Antibodies against DLX1 were raised against aa 1–121 and N-terminally tagged with His-Tag. Antibodies against DLX5 were raised against aa 188–289, N-terminally tagged with His. Antibodies were purified using either high affinity M or lodoacetyl resin, based on whether the protein had the MBP- or His Tag, respectively. To assess the specificity of the antibodies, Western Blots were performed using E13.5 or E16.5 basal ganglia nuclear extract from either WT or the respective constitutive Dlx mutants (Figure S1A). Furthermore, the antibodies against DLX2/1 and 5 were also tested using immunohistochemistry assays of E13.5 telencephalon sections of WT and Dlx21−/−, Dlx1 or Dlx5−/− respectively (Figure S1B).

**Chromatin immunoprecipitation (ChIP)**—ChIP was performed using the DLX1, DLX2 and DLX5 antibodies described above. Basal ganglia were dissected in cold PBS from CD1 embryos (6 litters/Ab for all the DLX ChIP-seq at E11.5; 2 litters/Ab for the DLX2 ChIP E13.5; 3 litters/Ab for the DLX1 and DLX5 ChIPs E13.5; and 1 litter/Ab at E16.5). The basal ganglia consisted of the LGE, MGE and CGE progenitor and mantle zones; they were fixed in 1.5% formaldehyde at RT for 20 min, neutralized with glycine and washed gently in PBS. The fixed cells were lysed with a hypotonic buffer (50 mM Tris pH 7.5 / 0.5% NP40 / 0.25% Sodium Deoxycholate / 0.1% SDS / 150 mM NaCl) to obtain the nuclei; these were then lysed in 1% SDS buffer and the chromatin was sheared into 300–1000 bp fragments by sonicating for 40 cycles (30 s on and 45 s off) using a bioruptor (Diagenode). Immunoprecipitation (IP) reactions were performed with the sheared chromatin diluted 1/10 times with “dilution buffer” (0.01% SDS, 1.1% Triton X-100, 1.2mM EDTA, 16.7mM Tris-HCl, pH 8.1, 167mM NaCl, usually in 6 ml. Antibody was then added: 5 μg DLX specific Ab. Negative control CHIP reactions used either rabbit IgG (5μg) or blocking peptide (DLX antigen used for immunizing rabbits; 50x molar excess). Antibody/chromatin complexes were purified using Dynabeads (Invitrogen) and washed extensively in 4 different “wash buffers” (low salt (1st), 0.1% SDS, 1% Triton X-100, 2mM EDTA, 20mM Tris-HCl, pH 8.1, 150mM NaCl; high salt (2nd), 0.1% SDS, 1% Triton X-100, 2mM EDTA, 20mM Tris-HCl, pH 8.1,500mM NaCl LiCl (3rd), 0.25M LiCl, 1% IGEPAL CA630, 1% deoxycholic acid (sodium salt), 1mM EDTA, 10mM Tris, pH 8.1and TE, 4th).

Complexes were eluted in 1% SDS, 10mM Sodium bicarbonate buffer at 65°C for 10 min. Eluted chromatin was reverse cross-linked overnight at 65°C in the presence of 500mM NaCl, then subsequently treated with RNase (10 μg/ 200 μl reaction, 15 min at 37°C) and Proteinase K (10 μg/ 200 μl reaction, 60 min at 55°C) and cleaned using a ChIP DNA Clean & Concentrator kit (Zymo Research). The chromatin was quality controlled (QC) using
qPCR to check for enrichment of genomic DNA fragments that were expected, and not-
expected, to have DLX binding.

Libraries were prepared using an Ovation Ultralow DR Multiplex System (Nugen), size
selected in the range of 300 bp on a chip from BluePippin (Sage Science) and lastly QC
tested on a Bioanalyzer (Agilent).

The libraries were sequenced as single-ended 50 nucleotide reads on a HiSeq 4000
(Illumina) at Center for Advanced Technology (UCSF).

**Native histone ChIP**—Each native histone ChIP was performed starting with ~250,000
nuclei from WT and Dlx1/2−/−E13.5 basal ganglia. The native ChIP was performed as
described earlier described (Magklara et al., 2011). Briefly, nuclei were extracted and
digested with micrococcal nuclease (MNase, Sigma). A population of mono- and di-
nucleosomes were used in chromatin immunoprecipitation assays. Antibodies used were
specific to H3 monomethyl lysine-4 (H3K4me1, Abcam, ab8895), H3 trimethyl lysine-4
(H3K4me3, Abcam, ab8580), H3 trimethyl lysine-27 (H3K27me3, Active Motif, 39157) and
H3 acetylated lysine 27 (H3K27ac, Abcam, ab472). Immuno-precipitated DNA was washed,
isolated and cleaned as for the TF ChIP-Seq described above.

**gRNA design and lentivirus infection of primary dorsal LGE dCas9-KRAB
culture**—The dCas9-KRAB mice were generated in the FVB background with the
TARGATT™ site-specific knock-in technology (Tasic et al., 2011) by introducing into the
Hipp11 locus a construct expressing from the CAG promoter: puromycin resistance,
mCherry and the dead Cas9 (dCas9) protein fused to the KRAB (Krüppel Associated Box)
domain. The dCas9-KRAB protein lacks endonuclease activity but retains its ability to bind
to single guide RNAs (sgRNAs) for specific gene targeting and silencing through
heterochromatin formation induced by the KRAB domain (V.G., M. Lebedinskaya, R.
Wagner, R. Jaafar, M.T.M., unpublished data). Guide RNA oligonucleotides were annealed
and cloned into U6-stuffer-longTracer-GFP lentivirus vector. gRNA lentivirus were
produced in 293T cells through transfection of packaging plasmids psPax2, pmD2G
(Addgene) and gRNA lentivirus vector. Concentrated virus was used to infect primary dorsal
LGE culture deriving from dCas9-KRAB mice at the time of seeding. The RNA from the
culture was purified 48 hr later and assayed for target gene expression via RT-QPCR. List of
gRNAs used:

**SP8TSS:** TTGGAGCTACAATTGTGGCGGAAT
AAACATTCCGCCACAATTGTAGCT

**hs110:** TTGG CTTTGTACTGCGGCTCAATT
AAACAAATTGAGCCGCAGTACAAAG

**hs1007:** TTGG GTAACAGATGTTCGCTAATC
AAAC GATTAGCGAACATCTGTTAC
**hs1226**: TTGG GGCGAICTTATAGACAGCCCTC  
AAACGAGGCTGTCAAACTCGGCC

**hs1148**: TTGG CGGATCCAGCCGTCCTCCAC  
AAACGTGGAACGGCTCGATGTCCG

**qPCR and primers**—We used a SYBR green mix (PerfeCTa SYBR Green FastMix ROX, Quanta) including, buffer, polymerase, dNTPs and SYBR green for the qPCR reactions. We ran 10 μl reactions on a 7900HT Fast Real-Time PCR system (Applied Biosystems). We always used negative control primer-sets for the calculated 2^−ΔΔCT value and analyzed as in Vokes et al. (2007). For DLX ChIP-seq QC we used the following primers:

**Dlx112b**: F: CGGGCCCATCAAACACAC  
R: TGGCGGAAATTTGTCAATCTCAT

**Dlx15/6**: F: GCCATCAAATTGAAGCAG  
R: GACGGTTAAACGCTGCAATCAG

**Internal control upstream of Dlx2**: F: CAGGACTAAGCAGCCCTTTG  
R: TGACCCCAATGACTCTCCAC

For the dCas9-KRAB Sp8 RT-qPCR experiments we used two Sp8 gene specific primers for first-strand cDNA synthesis

**GSP1 for Sp8**: CAGAGGAGTGGATCCCAACC

**GSP2 for Sp8**: CAGGTAGCAGCAAGCATGGC

**Sp8-specific qPCR primers**: F: GCGCACAATTGCACCATACTC  
R: TTCTTCTCGGTTCCTCCCCTTCC

As internal control we used a gene specific primer against the Etv1 locus for first-strand cDNA synthesis:

**GSP for Etv1**: TGAACATGGGCTGTGGGGTT

**Etv1-specific qPCR primers**: F: CATCAGGACGGGATGCTTCA  
R: GAGGCCATGAAAAGCCAAACT

**RNA-seq**—Total RNA from four embryos (E13.5, 2 males and 2 females) of either WT or Dlx1/2−/− was purified using RNeasy Mini Kit (QIAGEN). The DNase-treated RNA was checked on Bioanalyzer and RNA-seq was performed at the UCSFSABRE Functional
Genomics Facility. Briefly, RNA sequencing libraries were generated using TruSeq Stranded Total RNA Library Prep Kit with Ribo-Zero Mouse, according to the manufacturer’s protocol (Illumina, San Diego, CA, USA). Fragment size distribution was assessed using the Bioanalyzer 2100 and the DNA high-sensitivity chip (Agilent). Concentration of the libraries was measured using the Kapa library quantification kit (Kapa Biosystems, Woburn, MA, USA). Libraries were multiplexed, sequenced on the HiSeq 4000 and generated, roughly 814 million single-end 50bp reads.

**In situ RNA Hybridization (ISH)**—18 μm frozen cryostat sections of E13.5 mouse head were dried, wash three times with PBS (5 min each) and steamed for 10 min in Sodium Citrate buffer (10 mM, pH6). Sections were cooled down to RT for 15 min then rinsed with PBS three times (5 min each). Acetylation was performed for 10min in Acetylation Buffer containing 1.3% triethanolamine, 0.17% HCl and 0.4% acetic anhydride in water. Sections were then rinsed with PBS three times (5 min each) and prehybridized by incubating with hybridization buffer (50% formamide, 5XSSC pH 4.5, 50ug/ml yeast tRNA, 1% SDS, 50ug/ml Heparin) for 1 hour in 67°C oven. After prehybridization in situ probes diluted in hybridization buffer at 500ng/ml were added for overnight incubation at 67°C. Next day slides were rinsed with pre-warmed 5X SSC pH 4.5, washed twice (30min each) with 0.2X SSC pH4.5 at 70°C and then wash once (5min) with 0.2X SSC pH4.5 at room temperature followed by a wash with NTT buffer (0.15 M NaCl, 0.1 M Tris pH 8.0, 0.1% Tween-20). Sections were blocked with NTT blocking buffer containing 5% heat inactivated sheep serum and 2% Blocking buffer (Sigma-Aldrich) for 1 hour at room temperature followed by an overnight incubation at 4C with Anti-Digoxigenin-AP antibody (1:5000 dilution in NTT blocking buffer). Next day sections were washed three times with NTT buffer (30 min each), followed by three 5-min washes with NTM buffer (0.15 M NaCl, 0.1 M Tris pH 9.5, 0.1% Tween-20, 50 mM MgCl₂, 2 mM Levamisole) and incubated with developing reagent BM Purple (Sigma-Aldrich) until desired intensity of the signal was reached. Development reaction was stopped with PBS. Sections were fixed in 4% PFA o/n and were coverslipped in Aqua-Mount (Fisher Scientific).

**Primers used to generate in situ hybridization (ISH) probes**

**Arl4d:**
F: CTTGACTGAAATGGCCCCTA
R: ACCGTCTCTTCTTGCTCGAC

**Grik3:**
F: CACCAACCCCAGTGTCTTCT
R: GTTGATAACCGCATCCGTCT

**Tox:**
F: AGATTGGAAACGCAGAAGGA
R: AGTGTAAGGCAGTCTGGACC

**Lhx9:** Rétaux et al., 1999

**Nrnx3:** Püschel and Betz, 1995
**Pax7**: Jostes et al., 1990

**Zbtb20**: F: CAGGTAATCCTGGCTCATGC  
R: GGCACCTACCCCTGTGTTCTC

**Immunofluorescent tissue staining**—Immunofluorescent labeling was performed on 18 μm (E13.5) cryosections with the following primary antibodies: rabbit anti-DLX1, rabbit anti-DLX2, rabbit anti-DLX5 (production of these described above). 546 Alexa-conjugated secondary antibodies (Life Technologies) were used. Sections were coverslipped with Aqua-Mount.

**Luciferase reporter assay in P19**—RE elements were cloned into a pGL4.23-*Luciferase reporter vector* (Promega) at the Xhol and BglII sites. P19 cells (ATCC) were plated in 96 well plates, transfected with a total of 200 ng DNA using X-tremeGENE HP DNA transfection reagent (Sigma-Aldrich). Luciferase assays were performed 48h after transfection using Dual Luciferase Reporter Assay System (Promega). A Renilla reporter was used for normalization. Statistical differences between experimental groups were determined with paired t test.

**List of primers used to clone RE elements into pGL4.23**

**mm1203**: F: GTGCCAACAGTGCTAGATTA  
R: TCTACAGGGCATTCCCAAAGC

**hs119**: F: GAATTTGAGCTGAAAACATTTCC  
R: TTGGATTCTGGGAAGAATCC

**hs121**: F: GCGAGGCGATGTAAAGAATG  
R: TTGAGTAGACAAACATGTTTTC

**hs122**: F: GATGCAAATACTTCTAGAATGG  
R: TAAAGTCAATCCCATGATAGTT

**hs123**: F: GCCCTATCTTATGGAAGGAAC  
R: TGCCACATGGAGATGAAGAG

**hs1007**: F: GCCTGTTTGTACATTAGCGAA  
R: TCAACAACAACAAAAAGACTAAA

**hs110**: F: ACCCATATTTTGTTGAGCT  
R: TGTAGCAGCATCTGAGGA
**hs1226**: F: CTGGAAAGGCCACC  
R: CAGGCTTTGCCTGA

**Western blot analysis**—Nuclear preparation from basal ganglia tissue from one litter of either E13.5 or E16.5 MGE was prepared using a NE-PER kit (ThermoFisher) with protease inhibitor (Sigma-Aldrich) and the total protein was separated in SDS-PAGE gels and transferred to nitrocellulose membranes. Western blotting was performed by standard procedures.

**Transgenic enhancer assays**—Enhancer names (mm/hs numbers) are the unique identifiers used in the VISTA Enhancer Browser ([https://enhancer.lbl.gov](https://enhancer.lbl.gov)). Candidate enhancer, mm1203, was PCR-amplified and cloned into an hsp68-promoter-lacZ reporter vector (Pennacchio et al., 2006). Transgenic assays were performed according to published methods (Kothary et al., 1989; Pennacchio et al., 2006).

Briefly, the enhancer-reporter vector was linearized and injected into the pronucleus of FVB strain single cell stage mouse embryos (E0.5). Embryos were implanted into surrogate CD-1 strain *Mus musculus* mothers and were then collected and stained for reporter gene expression at E11.5 or E12.5. Embryos were excluded from analysis only if they did not harbor the transgene or if they were not at the correct developmental stage. No comparisons were made between cohorts of transgenic embryos, so randomization and experimenter blinding were unnecessary and not performed. Sample sizes were determined empirically based on our experience performing > 2,000 transgenic enhancer assays. Only LacZ activity patterns that were observed in three or more different embryos resulting from independent transgene integration events of the same construct were considered reproducible.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

**Differential Gene Expression**—We determined the differential gene expression between the wild-type and *Dlx1/2−/−* samples by individually aligning the FASTQ reads from the RNA-seq experiment using STAR (version 2.4.2a; (Dobin et al., 2013), after QC’ing the files with FASTQC (version 0.11.6; Andrews, 2010) and counting reads with featureCounts (version 1.5.1; Liao et al., 2014). Following quality filtering and alignment, we analyzed the samples using a custom R script running edgeR (version 3.24.2) / limma (version 3.38.3; Phipson et al., 2016; Robinson et al., 2010), that used a statistical method that ascertained the differential gene expression levels.

**ChIP-seq Data Analysis**—FASTQ files containing the reads were QC’ed using FASTQC utility, and the remaining adapter sequences were trimmed using Trim Galore (version 0.4.5; Krueger, 2015). The resulting reads were aligned to the mouse (mm9) genome using Burrows-Wheeler Alignment (BWA; version 0.7.9a; Li and Durbin, 2009), duplicates removed with Samtools (version 1.8; Li et al., 2009). Peaks of enriched binding regions against both negative binding and input DNA control were called using MACS (version 2.1; Zhang et al., 2008) (Figure S4). Using DeepTools (version 2.5.3; Ramírez et al., 2016), we determined the Pearson correlation among the reads of all DLX dataset replicates (Figure S2). After removing the outlier replicates, the remaining ones were merged and analyzed
again, showing the correlation among the different DLXs at the different growth stages (Figure 1D). Narrow peaks of DLXs were overlapped and merged into one dataset using custom R scripts, where they were annotated for neighboring gene regulatory regions and filtered against repeat, blacklisted and gapped regions. Additional annotation was made for each DLX-bound peak overlapping evolutionary conserved regions, as well as VISTA transcriptional enhancers (Table S3).

Histone (H3K27ac, H3K27me3, H3K4me3, H3K4me1) broad peaks, also called using MACS2, were then overlapped with and annotated on the base peak set described above. Furthermore, MACS2 bdgdiff was used to assess the differential intensity of the histone PTMs as measured by ChIP-seq. We used the same approach to determine local H3K4me1 changes in WT versus KO, as a marker for transcriptional enhancer establishment or hinderance. We used a p value cutoff of 1E-4, with 350 as minimum length of peak for all narrow and broad peak calls; for differential histone peak calls we used a likelihood ratio cutoff of 1E+3. Command-line examples for the software used are provide in Table S7.

We used ngsplot (version 2.63; Shen et al., 2014) within custom scripts to display overall genome wide coverage, both as coverage plots and heatmaps, for a variety of stratifications (Figures 1E and 2C), with a fragment length of 150 bp. Moreover, we used DeepTools to make peak complexity plots (not shown). Peak complexity was further analyzed by assessing the number of peaks within each peak neighborhood and finding the statistical distribution of their means, both overall and within a.REs and r.REs (Figure S5); we ran a custom R script that performed permutation analysis at 95% confidence level to determine the statistical significance of those differences.

**Chromatin State Inference**—We inferred chromatin state change as a result of DLX binding, herein designated as a.REs and r.REs (activating and repressing regulatory elements, respectively), by the local occurrence of H3K4me3, H3K27ac, H3K27me3 signature mark changes in WT samples in relation to Dlx1/2−/− (KO) ones. Following determining significantly altered PTM regions in WT versus KO, we assigned a.REs to DLX-bound regions with increases in H3K4me3 or H3K27ac, or decreases in H3K27me3, in WT. Conversely, r.REs were assigned to regions with changes in the opposite direction; bound regions with dual assignment were disregarded. We inferred putative enhancer regions when the H3K4me1 was present in WT; the effect of DLX binding to putative enhancers was assessed by the differential likelihood of H3K4me1 in WT in comparison with Dlx1/2−/−.

We evaluated the correlation among histone marks at DLX TF-bound regions, intensity of histone ChIP-seq signal in WT, peak neighborhood complexity and differential gene expression, visually (Figure 5E) and by running ANOVA on the factors against gene expression fold change (Table S8).

Furthermore, we created an 8-state HMM model using our WT histone data and ChromHMM (Ernst and Kellis, 2017) and assigned chromatin states based on the emission probabilities. We further segmented the states into proximal and distal, as appropriate for the emission signal.
Motif Analysis—De novo motif discovery was performed using HOMER (version 4.9; Heinz et al., 2010) with standard parameters, 250 bp up- and downstream of DLX peaks. Upon selection of significant classes of overrepresented motifs on all DLX-bound regions based on FDR corrected (Benjamini-Hochberg method) p value < 0.05 and non-redundancy, we carried out enrichment analysis of those motifs on those regions, as well as regions stratified into distal and proximal sites, and a.REs and r.REs (Figures 1H and S3). All motif enrichment determination was also made with HOMER. Command-line examples for the software used are provided in Table S7.

We further analyzed the difference between motif enrichment in a.RE versus r.RE for HOMER known motif database using custom R scripts (Figure 4A) and found statistically significant differences between a.RE- and r.RE-relative enrichment by calculating means differences using both a bootstrapping approach and an unpaired two-sample Wilcoxon test (Table S9).

Gene Regulatory Network—We constructed a core gene regulatory networks (GRN) at the three growth stages with Cytoscape (version 3.6.1; Shannon et al., 2003) by assigning DLXs as source nodes and DE genes with annotated DLX-bound peaks at presumed REs, significant expression difference and chromatin state change (a.RE/r.RE) as target nodes (not shown). Node colors were assigned to as green for activation or downregulation in Dlx1/2−/− samples and red for inhibition or upregulation in KO, with intensity proportional to gene expression fold change. Edges were also colored as green and red as above, with widths proportional to the p values of the respective gene expression fold changes.

Gene Ontology Analysis—We conducted gene ontology analysis using the GREAT algorithm for the whole peak dataset and compared with the results for the peaks showing in a.REs and r.REs, by means of a custom R script running the rGREAT package (version 1.14.0; Gu, 2015). We found a number of differentially enriched GO terms in a.RE versus r.RE and selected a few relevant ones for our test system (Figure 2F); the complete results are on Table S4. The genes that differentially made the list for relevant disease GO terms are shown in Table S10 and are graphically depicted in Figure S6.

Modeling sequence and context features for predicting a.RE and r.RE loci—Logistic regression analysis was performed with a.RE, r.RE or no change class as the dependent variable and motif presence, histone PTM peak fold enrichment, DLX2 E13.5 peak fold enrichment, number of DLX peaks within 50KBP of peak, ChromHMM state and distal versus proximal location. a.RE or r.RE were compared against all other peaks. ROC curves were generated based on predicted versus observed results for the logistic regression model and area under the curve was calculated using discrete approximation.

DATA AND CODE AVAILABILITY

The genomic and epigenomic data generated in this study and presented in this publication have been deposited in NCBI database and are accessible through GEO Series accession number GSE124936 (https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE124936) and can be visualized in UCSC track hubs whose information is provided on Nord Lab.
Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Highlights

- DLX proteins modulate expression of GABAergic neuronal differentiation genes
- DLXs drive a complex regulatory network by transcriptional activation and inhibition
- Genomic and epigenomic context and sequence predict DLX effect on gene expression
- DLX regulatory wiring may reflect general mechanisms implicated in neurodevelopment
Figure 1. DLX2, DLX1, and DLX5 Genomic Binding in Embryonic GE
(A) Schematic of functional genomic dissection of DLX.
(B) DLX ChIP-seq coverage at Gad2 and Lhx6 loci; merged peak dataset represented inside a golden box on top of first DLX track.
(C) DLX peak counts by genomic feature at E11.5, E13.5, and E16.5.
(D) Heatmap showing pairwise Pearson correlation for genome-wide coverage values for DLX ChIP-seq.

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(E) Normalized coverage of ChIP-seq peaks. Each row represents a DLX binding region ± 10 kb. Numbers under the heatmap columns denote number of peaks called for each DLX/time point.

(F) Venn diagrams showing increasing percent of peaks shared across DLXs as peak stringency increases.

(G) Volcano plot showing E13.5 Dlx1/2−/− versus WT GE differential gene expression fold change and statistical significance.

(H) Shared primary binding DNA motif across DLXs (i) centered within ChIP-seq peaks (ii). Motifs that were strongly enriched within DLX peaks (iii); blue bars show motif frequency in DLX peaks and red bars in GC-matched background sequences (enrichment in parentheses).

(I) Distance from TSS to nearest DLX peak for DE genes by peak stringency.

(J) Median Phastcons scores for vertebrate-conserved elements at promoter and distal DLX peaks by stringency.
Figure 2. Chromatin State Is Dependent on DLX Binding for Key Regulatory Targets

(A) Schematic of epigenomic comparison of WT and DLX1/2−/− E13.5 GE.

(B) DLX and histone ChIP-seq coverage for Slc32a1 (Vgat, downregulated) and Otp (upregulated) loci; merged peak dataset represented inside a golden box on top of first DLX track, and differential histone PTMs shown as black bars under the peak where there was statistically significant difference.

(C) Average change in histone PTM signal in DLX1/2−/− GE at DLX-bound loci for regions featuring loss of activating or gain of repressive marks (a.RE) or gain of activating or loss of repressive marks (r.RE).
(D) Log₂ fold change of nearest gene expression for all DLX peaks, higher-affinity peaks, and a.RE and r.RE loci.

(E) Modified volcano plot showing intersection of all differential gene expression and a.RE and r.RE loci. Colored circles represent a.RE (red) or r.RE (blue) within 100 kb of gene TSS. Circle size shows magnitude of histone PTM change in Dlx1/2−/− GE.

(F) Differential enrichment of functional annotation terms comparing all DLX TF peaks and a.RE and r.RE loci.
Figure 3. *Nrnn*3 and *Arx* Are Regulated by DLX via RE Interactions

(A) ChIP-seq data for *Nrnn*3 locus; called DLX peaks and histone differential PTM as described in Figure 2B.

(B) ISH analysis of *Nrnn*3 expression in WT (upper panel) and *Dlx1/2*−/− (lower panel) forebrain at E13.5. (MGE, medial GE; LGE, lateral GE; CGE, caudal GE; SVZ, subventricular zone; VZ, ventricular zone; MZ, mantle zone). Images are representatives of 3 embryos. Scale bar represents 500 μm.
(C) *mm1203* sequence drives *LacZ* expression in E12.5 forebrain in a transgenic mouse enhancer assay. Dashed line indicates plane of section.

(D) *Dlx* transfection luciferase transcription assays in P19 cells showing activating effect of DLX on *mm1203*.

(E) ChIP-seq data for *Arx* locus; called DLX peaks and histone differential PTM as described in Figure 2B.

(F) Activity patterns in mouse transgenic assays for characterized *Arx* GE (*hs119, hs121*) and pallial (*hs122, hs123*) REs.

(G) *Dlx* transfection luciferase transcription assays in P19 cells showing specific activating effect of DLX on *Arx* GE REs. Luciferase data are represented as mean ± SEM (n = 5 for *Nrxn3* and n = 3 for *Arx*). Unpaired t test was used for the statistical analysis between the presence of enhancer in the reporter or no enhancer (*p < 0.05, **p <0.01, ***p <0.001, and ****p < 0.0001).
Figure 4. Genome Sequence and Context Define a.RE and r.RE Loci
(A) Differential motif enrichment in a.RE and r.RE loci. Motifs specific to a.RE or r.RE labeled in blue. Horizontal bar plot on left shows log2 fold enrichment difference between a.RE and r.RE with dotted blue lines shown for 1.2-fold enrichment difference. Heatmap on right shows a.RE and r.RE enrichment versus all DLX peaks.
(B) Distribution of all DLX peaks, higher-affinity peaks, and a.RE and r.RE loci by chromatin state. ChromHMM states defined by relative histone PTM enrichment as shown on left. Each dot in center plot represents one peak with density represented by shaded area.
Distal and proximal (promoter) region assignment shown as colored bars along the axis, with blue and yellow indicating distal and proximal regions, respectively; no chromatin state signal and repressed regions marked as both (black). Proportion of total peaks shown in bar plot at right.

(C) Distribution of the number of DLX peaks within 50 kb of TSS for all versus DE genes, with gene counts by DLX peak number histogram and representative example genes shown.

(D) Relationship between log₂ FC (fold change of gene expression Dlx1/2−/− versus WT) of the nearest TSS (x-axis) and histone H3K27ac signal in WT E13.5 GE (y-axis) and for all DLX peaks, a.RE, and r.RE. Symbol size shows density of DLX TF peaks within 50 kb. Distribution of H3K27ac enrichment for peak classes shown at right.

(E) Receiver operating characteristic (ROC) curve for logistic regression models predicting a.RE or r.RE status. Predictors include DNA motif presence, chromatin context (histone PTM, distal/proximal, ChromHMM state) and DLX (affinity and local peak density), and a full model with all features included.

***p < 2.2 × 10⁻¹⁶ in (C) and (D).
Figure 5. Sp8 Is Regulated by DLX Binding to Promoter-Interacting Distal Elements

(A) ChIP-seq coverage for Sp8 locus; called DLX peaks and histone differential PTM as described in Figure 2B.

(B) Transgenic RE assay results showing activity of three GE-active (hs110, hs1226, and hs1007) and limb-active (hs1148) REs.

(C) Downregulation of Sp8 mRNA caused by delivery of dCas9-KRAB and gRNA targeting putative Sp8 REs. y-axis shows the amount of Sp8 mRNA relative to control GFP gRNA (blue points) measured in RT-qPCR (n = 3 for gRNA-hs1148, n = 6 for the rest).

(D) DLx transfection luciferase transcription assay in P19 cells showing the effect of DLX on the Sp8 GE REs; data represented as mean ± SEM (n = 3). Unpaired t test was used for the statistical analysis between the presence of enhancer in the reporter or no enhancer (*p < 0.05; **p < 0.01, ***p < 0.001, ****p < 0.0001).
Figure 6. DLX TFs Regulate Spatial Patterns of Gene Expression

(A) ISH analyses using probes detecting *Arl4d* and *Grik3*, and *Tox* and *Lhx9* in WT and *Dlx1/2*<sup>−/−</sup> forebrain at E13.5 (MGE, medial GE; LGE, lateral GE; CGE, caudal GE; SVZ, subventricular zone; VZ, ventricular zone; MZ, mantle zone; open arrows, downregulation; black arrows, upregulations; red arrows, ectopic expression). Images are representatives of 2 embryos for each probe. Scale bar represents 500 μm.

(B) Summary of spatial changes in DE gene expression from ISH analysis comparing WT and *Dlx1/2*<sup>−/−</sup> forebrain.

(C) Transgenic RE assay activity patterns for representative a.RE and r.RE sequences.

(D) Summary of RE activity patterns within GE subregions (VZ, ventricular zone; SVZ, subventricular zone; MZ, mantle zone).
Figure 7. Regulatory Model and Gene Regulatory Networks Orchestrated by DLX

(A) Schematic model of DLX genomic function in developing GE.

(B) Curated gene regulatory network (GRN) for DLX-regulated transcription factors and lineage specification factors. The GRN is organized radially with regard to the laminar activity of genes (VZ/SVZ, SVZ, and MZ, shown in three different shades of gray). The effects of the Dlx1/2−/− mutation are indicated in three nested circles. The outer circle reports RNA changes measured by ISH (IS) (this study; Long et al., 2009a, 2009b). The middle circle reports histone PTM signal (H) changes. The inner circle reports RNA-seq (R)
changes. Red and green represent repressive and activating roles for DLX TFs on each assay, respectively. For histone changes, REs assigned to nearest TSS.
### KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| DLX2 (MBP-aa1–154)  | This paper | N/A; Available from the authors |
| DLX1 (His-aa1–121)  | This paper | N/A; Available from the authors |
| DLX5 (His-188–289aa) | This paper | N/A; Available from the authors |
| H3K4me1             | Abcam | Cat# ab8895; RRID:AB_306847 |
| H3K4me3             | Abcam | Cat# ab8580; RRID:AB_306649 |
| H3K27me3            | Active Motif | Cat# 39155; RRID:AB_2561020 |
| H3K27ac             | Abcam | Cat# ab4729:RRID:AB_2118291 |
| Anti-Digoxigenin-AP | Sigma-Aldrich | Cat# 11093274910; RRID:AB_2734716 |
| Goat anti-Rabbit IgG (H+L), Alexa Fluor 546 | Thermo-Fisher | Cat# A-11035; RRID:AB_2534093 |
| Normal Rabbit IgG   | Santa Cruz | Cat# sc-2027; RRID:AB_737197 |
| **Chemicals, Peptides and Recombinant Proteins** |        |            |
| DIG RNA labeling mix | Sigma-Aldrich | Cat# 11277073910 |
| T7 RNA polymerase    | Sigma-Aldrich | Cat# 10881767001 |
| DNease               | Sigma-Aldrich | Cat# 10104159001 |
| Sheep Serum          | Sigma-Aldrich | Cat# S2263 |
| Blocking reagent     | Sigma-Aldrich | Cat# 11096176001 |
| BM purple            | Sigma-Aldrich | Cat# 11442074001 |
| 37% Formaldehyde     | Ted Pella | Cat# 18508 |
| Dynabeads Protein G  | Thermo-Fisher | Cat# 10003D |
| Dynabeads Protein A  | Thermo-Fisher | Cat# 10001D |
| RNase, DNase free    | Sigma-Aldrich | Cat# 11119915001 |
| Protease K           | Sigma-Aldrich | Cat# 3115879001 |
| PerfeCTa SYBR Green FastMix ROX | Quanta | Cat# 95073-012 |
| MNase                | Sigma-Aldrich | Cat# N3755–50UN |
| Thermo Scientific Shandon Aqua-Mount Slide Mounting Media | Fisher Scientific | Cat# 14–390–5 |
| X-tremeGENE HP DNA transfection reagent | Sigma-Aldrich | Cat# 6366236001 |
| Complete EDTA-free Protease inhibitor | Sigma-Aldrich | Cat# 11873580001 |
| **Critical Commercial Assays** |        |            |
| RNA Clean and Concentrator | Zymo Research | Cat# R1015 |
| Ovation Ultralow DR Multiplex System | Nugen | Cat# 0344–32 |
| 2% Agarose Gel Cassette Blue Pippin | Sage Science | Cat# BDF2010 |
| High Sensitivity DNA Reagents | Agilent Technologies | Cat# 5607–4626 |
| TruSeq Stranded Total RNA Library Prep Kit with Ribo-Zero Mouse | Illumina | Cat# RS-122–2202 |
| KAPA Library Quantification Kit | Roche | Cat# 07960140001 |
| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| RNeasy Mini Kit     | QIAGEN | Cat# 74104 |
| NE-PER kit          | Thermo Fisher | Cat# 78833 |
| Dual Luciferase Reporter Assay System | Promega | Cat# E1980 |
| Deposited Data      |        |            |
| RNA-seq and ChIP-seq data | US National Center for Biotechnology Information | [http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE124936](http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE124936) |
| Recombinant DNA     |        |            |
| Hsp68-lacZ vector   | PMID 2557196 | n/a (available from DED, AV and LAP) |
| psPax2              | Addgene | Cat# 12260 |
| pmD2G               | Addgene | Cat# 12259 |
| Experimental Models: Organisms/ Strains |        |            |
| Transgenic enhancer assay: FVB | Charles River | [https://www.criver.com/](https://www.criver.com/) |
| ChIP-seq: CD1       | Charles River | [https://www.criver.com/](https://www.criver.com/) |
| dCas9-KRAB          | V.G., M. Lebedinskaya, R. Wagner, R. Jaafar, M.T.M., unpublished data | n/a |
| Software and Algorithms |        |            |
| STAR                | Dobin et al., 2013 | [https://github.com/alexdobin/STAR](https://github.com/alexdobin/STAR) |
| FASTQC              | Andrews, 2010 | [https://www.bioinformatics.babraham.ac.uk/projects/fastqc/](https://www.bioinformatics.babraham.ac.uk/projects/fastqc/) |
| featureCounts       | Liao et al., 2014 | [http://subread.sourceforge.net](http://subread.sourceforge.net) |
| edgeR               | Phipson et al., 2016 | [https://www.bioconductor.org/packages/release/bioc/html/edgeR.html](https://www.bioconductor.org/packages/release/bioc/html/edgeR.html) |
| limma               | Robinson et al., 2010 | [https://www.bioconductor.org/packages/release/bioc/html/limma.html](https://www.bioconductor.org/packages/release/bioc/html/limma.html) |
| Trim Galore         | Babraham Bioinformatics | [https://www.bioinformatics.babraham.ac.uk/projects/trim_galore/](https://www.bioinformatics.babraham.ac.uk/projects/trim_galore/) |
| BWA                 | Li and Durbin, 2009 | [http://bio-bwa.sourceforge.net/bwa.shtml](http://bio-bwa.sourceforge.net/bwa.shtml) |
| Samtools            | Li et al., 2009 | [http://samtools.sourceforge.net](http://samtools.sourceforge.net) |
| DeepTools           | Ramirez et al., 2016 | [https://deeptools.readthedocs.io/en/develop/](https://deeptools.readthedocs.io/en/develop/) |
| MACS2               | Zhang et al., 2008 | [https://github.com/taoliu/MACS/](https://github.com/taoliu/MACS/) |
| ngsplot             | Shen et al., 2014 | [https://github.com/shenlab-sinai/ngsplot](https://github.com/shenlab-sinai/ngsplot) |
| ChromHMM            | Ernst and Kellis, 2017 | [http://compbio.mit.edu/ChromHMM/](http://compbio.mit.edu/ChromHMM/) |
| HOMER               | Heinz et al., 2010 | [http://homer.ucsd.edu/homer/](http://homer.ucsd.edu/homer/) |
| Cytoscape           | Shannon et al., 2003 | [https://cytoscape.org](https://cytoscape.org) |
| rGREAT              | Gu, 2015 | [https://www.bioconductor.org/packages/release/bioc/html/rGREAT.html](https://www.bioconductor.org/packages/release/bioc/html/rGREAT.html) |