Integration of Genome-wide Approaches Identifies IncRNAs of Adult Neural Stem Cells and Their Progeny In Vivo

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SUMMARY

Long noncoding RNAs (IncRNAs) have been described in cell lines and various whole tissues, but IncRNA analysis of development in vivo is limited. Here, we comprehensively analyze IncRNA expression for the adult mouse subventricular zone neural stem cell lineage. We utilize complementary genome-wide techniques including RNA-seq, RNA CaptureSeq, and ChIP-seq to associate specific IncRNAs with neural cell types, developmental processes, and human disease states. By integrating data from chromatin state maps, custom microarrays, and FACS purification of the subventricular zone lineage, we stringently identify IncRNAs with potential roles in adult neurogenesis. shRNA-mediated knockdown of two such IncRNAs, Six3os and Dlx1as, indicate roles for IncRNAs in the glialneuronal lineage specification of multipotent adult stem cells. Our data and workflow thus provide a uniquely coherent in vivo IncRNA analysis and form the foundation of a user-friendly online resource for the study of IncRNAs in development and disease.

INTRODUCTION

The mammalian genome encodes thousands of long noncoding RNAs (IncRNAs), and it is becoming increasingly clear that IncRNAs are key regulators of cellular function and development. Loss-of-function studies performed in cell culture indicate that IncRNAs can regulate gene transcription through the targeting and recruitment of chromatin-modifying complexes (Guttman et al., 2011; Huarte et al., 2010; Khalil et al., 2009; Tsai et al., 2010). While it is now evident that IncRNAs have important

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cellular and molecular functions, how they participate in development in vivo is poorly understood.

Emerging studies suggest that IncRNAs play critical roles in CNS development. For instance, in embryonic stem cells (ESCs), specific IncRNAs repress neuroectodermal differentiation (Guttman et al., 2011), and during in vitro differentiation of ESC-derived neural progenitor cells (ESC-NPCs), IncRNA expression is dynamic (Mercer et al., 2010). In the mouse brain, some IncRNAs are regionally expressed (Mercer et al., 2008), including among the six layers of the adult cortex (Belgard et al., 2011). In vivo functional data is limited, but mice null for the IncRNA *Evf2* have abnormal GABAergic interneuron development and function (Bond et al., 2009), and morpholino inhibition of two CNS-specific IncRNAs in zebrafish affects brain development (Ulitsky et al., 2011).

The subventricular zone (SVZ) of the adult mouse brain represents an ideal system for the study of IncRNAs in vivo. Throughout life, SVZ neural stem cells (SVZ-NSCs) generate large numbers of neuroblasts that migrate to the olfactory bulb (OB), where they differentiate into interneurons (Figure 1A). In addition, SVZ-NSCs are multipotent, capable of generating astrocytes and oligodendrocytes, the other major cell types of the CNS. In contrast to the embryonic brain, wherein multipotent precursor cells are inherently transient, continually changing their developmental potential and location over time and with organ morphogenesis, the adult SVZ retains its NSC population in a stable, spatially restricted niche, producing neurons and glia throughout life (Kriegstein and Alvarez-Buylla, 2009). This enduring population of multipotent stem cells and its well-characterized daughter cell lineages make the SVZ a particularly tractable in vivo model for molecular genetic studies of development. The SVZ has been used to elucidate key principles of neural development including the role of signaling molecules, transcription factors, microRNAs, and chromatin modifiers (Ihrie and Alvarez-Buylla, 2011). We have previously shown that the Mixed-lineage leukemia 1 (MII1) chromatin-modifying factor is required for the SVZ neurogenic lineage (Lim et al., 2009), and





recent studies indicate that MLL1 protein can be targeted to specific loci by lncRNAs (Bertani et al., 2011; Wang et al., 2011).

Here, we leveraged the SVZ-OB system to develop a greater understanding of IncRNA expression and function. First, we used Illumina-based complementary DNA (cDNA) deep sequencing (RNA-seq) and ab initio reconstruction of the transcriptome to generate a comprehensive IncRNA catalog inclusive of adult NSCs and their daughter cell lineages. This IncRNA catalog informed a subsequent RNA CaptureSeq approach, which increased the read coverage and read length for our SVZ cell analysis, validating the transcript structure and expression of many of these novel IncRNAs. Gene coexpression analysis identified sets of IncRNAs associated with different neural cell types, cellular processes, and neurologic disease states. In our analysis of genome-wide chromatin state maps, we identified IncRNAs

Figure 1. Outline of IncRNA Catalog Generation

(A) Schematic of sagittal section of adult mouse brain. SVZ neural stem cells give rise to migratory neuroblasts (red). These neuroblasts travel along the rostral migratory stream (curved arrow) before terminally differentiating and integrating into olfactory bulb (OB) neuronal circuits. Numbered schematics correspond to coronal brain sections highlighting dissected regions (yellow) used for RNA collection.

(B) Workflow for IncRNA catalog construction and characterization. See also Figure S1 and Table S1.

that-like key developmental genes-demonstrate chromatin-based changes in a neural lineage-specific manner. Using custom IncRNA microarrays, we found that IncRNAs are dynamically regulated in patterns reminiscent of known neurogenic transcription factors. To define IncRNA expression changes throughout the SVZ neurogenic lineage in vivo, we acutely isolated the major cell types of the SVZ with fluorescent-activated cell sorting (FACS) and probed IncRNA expression with our custom microarrays. We integrated these diverse experimental approaches to develop an online resource useful for the identification of IncRNAs with potential roles in SVZ neurogenesis (http://neurosurgery.ucsf.edu/ danlimlab/IncRNA). Furthermore, expression and shRNA-mediated knockdown experiments confirmed functional roles for IncRNAs identified by our integrative approach. Overall, our study demonstrates a generalizable workflow that assimilates genome-wide bioinformatic strategies with experimental manipulations for the identification of IncRNAs that regulate development.

RESULTS

Cataloging IncRNAs in the Adult Brain Neurogenic Zones

Because IncRNAs exhibit tissue-specific expression, previous mouse IncRNA databases

were not likely comprehensive for IncRNAs involved in adult neurogenesis. Thus, we identified IncRNAs expressed in the adult brain neurogenic niches by employing an RNA-seq and ab initio transcriptome reconstruction approach. First, we generated cDNA libraries of polyadenylated RNA extracted from microdissected adult SVZ tissue, which contains NSCs, transit-amplifying cells, and young migratory neuroblasts. To include the transcriptome of later stages of neurogenesis and neuronal function, we also generated cDNA libraries from the OB. Furthermore, we generated cDNA libraries from microdissected adult dentate gyrus (DG), the other major adult neurogenic niche, which locally contains all cell types of an entire neuronal lineage. Figure 1A shows a schematic of regions used for the cDNA libraries.

We used Illumina-based sequencing to obtain paired-end reads of these cDNA libraries from the SVZ (229 million reads),

OB (248 million reads), and DG (157 million reads). To broaden our IncRNA catalog, we also included RNA-seq data from embryonic stem cells (ESCs) and ESC-derived neural progenitors cells (ESC-NPCs) (Guttman et al., 2010). With this collection of over 800 million paired-end reads, we used Cufflinks (Trapnell et al., 2010) to perfom ab initio transcript assembly. This method reconstructed a total of 150,313 multiexonic transcripts, of which 140,118 (93%) overlapped with known protein-coding genes. Our IncRNA annotation pipeline (see Figure 1B and Experimental Procedures) identified 8,992 IncRNAs encoded from 5,731 loci (see Table S1 available online). There were 6,876 (76.5%) novel ones compared to RefSeq genes, 5,044 (56.1%) were novel compared to UCSC known genes, and 3,680 (40.9%) were novel compared to all Ensembl genes. Interestingly, 2,108 transcripts (23.4%) were uniquely recovered from our SVZ/OB/DG reads.

To substantiate the noncoding nature of our IncRNA candidates, we used the coding potential calculator (Kong et al., 2007) and found that over 80% of these transcripts have essentially no protein-coding potential (Figure S1A). Consistent with previous studies, IncRNAs were expressed at lower levels than protein-coding genes (2.49-fold difference; Mann-Whitney U, p < 0.0001) (Figure S1B), and their exons were less strongly conserved than protein-coding exons by PhastCons scores (Figure S1C).

The transcriptional start site (TSS) of some IncRNAs is proximal (<10 kb) to the promoters of protein-coding genes (Cabili et al., 2011; Hung et al., 2011), and we found that the TSS of 2,265 IncRNAs (25.2%) in our catalog were located within 5 kb of a protein-coding gene promoter (Figure S1D). Gene onotology (GO) analysis with the genomic regions enrichment of annotations tool (GREAT) (McLean et al., 2010) revealed that these protein-coding neighbors are enriched for homeodomaincontaining transcription factors, genes expressed in the brain, and genes that are typically repressed by Polycomb Repressive Complex 2 in ESCs (Figure S1E). While some IncRNAs had strongly correlated expression with their protein-coding neighbor, as a group they had no obvious correlation (Figure S1F), indicating that expression of this subset of IncRNAs is not likely related to local transcriptional activity of proteincoding genes.

To verify that the cDNA libraries of the SVZ and OB together represent a transcriptome enriched for adult neurogenesis, we first analyzed mRNA expression in the RNA-seq data. Differential gene expression identified 1,621 genes enriched >2-fold in the SVZ cDNA library as compared to the cDNAs from cells in the adjacent nonneurogenic striatum (76.4 million reads). As the primary site where NSCs and transit-amplifying cells proliferate, the SVZ was enriched for GO terms related to cell cycle and mitosis (Figures S2A and S2B). Neuroblasts migrate through the SVZ and into the OB, and, as expected, transcripts related to this migratory neuroblast stage of neurogenesis were enriched in these regions (Lim et al., 2006). The SVZ/OB expression profile included transcription factors known to play key roles in adult neurogenesis, such as DIx1, DIx2, AscI1, and Pax6 (Hsieh, 2012). Furthermore, in situ hybridization (ISH) data from the Allen Brain Atlas (Lein et al., 2007) confirmed the regional expression of many of these SVZ/OB-enriched genes (Figures S2C and S2D), and the SVZ/OB transcriptional profile (923 genes) was enriched for GO terms related to cell migration, development, and neurogenesis (Figure S2E).

IncRNAs Have Temporally and Spatially Unique Expression Patterns

To explore IncRNA expression patterns in multiple adult brain regions and embryonic forebrain development, we analyzed RNA-seq data of the six layers of the adult cortex (Belgard et al., 2011), adult whole prefrontal cortex (PFC), adult preoptic area (POA), whole embryonic day 15 (E15) brain (Gregg et al., 2010), and specific regions of the developing E14.5 cortex (ventricular zone, intermediate zone, and cortical plate) (Ayoub et al., 2011) (Figure 2A and Table S2). Unsupervised hierarchical clustering of expression profiles revealed region-specific and temporally related expression of both mRNAs and IncRNAs (Figures 2B and 2C). We calculated a specificity score for each transcript (Cabili et al., 2011) and found that the mean score was 0.57 (SD 0.21) for IncRNAs, while it was 0.45 (SD 0.17) for mRNAs $(p < 10^{-325})$, Wilcoxon rank-sum test); thus, IncRNAs exhibit greater brain region and temporal specificity than mRNAs, suggesting that they play important roles in the determination and/or function of specific neural cell types.

IncRNAs Are Associated with Specific Neural Cell Types and Neurological Disease States

To begin to infer functions for InCRNAs, we investigated the relationship between mRNA and InCRNA transcription by using gene coexpression analysis (GCA) to identify groups of transcripts, or "modules," whose variation in expression correlate across different brain regions and developmental time points. For mRNAs, module membership distinguishes sets of genes that correspond to specific cell types and biological processes (Oldham et al., 2008), and a similar "guilt-by-association" approach has been used to assign putative functions to InCRNAs based on their coexpression with protein-coding genes (Guttman et al., 2009).

Using RNA-seq data from 22 samples (Figure 2A and Table S2), we constructed transcript coexpression networks comprised of both mRNAs and lncRNAs. For the 56 modules of coexpressed transcripts, we performed enrichment analysis using gene sets from the Molecular Signatures Database (Subramanian et al., 2005) and other sources (Bult et al., 2008; Cahoy et al., 2008; Thomas et al., 2011; Zhang et al., 2010) to relate modules (described by "color," Figures 3A–3F) to specific adult neural cell types including cortical neurons (purple), striatal neurons (salmon), ependymal cells (green), and oligodendrocytes (grey60) (Table S3).

The dark red module (Figure 3E) was enriched for glial markers but also had a large number of known early neurogenic factors as prominent members (Table S3). This module was specifically associated with the ventricular zone of the embryonic brain, which contains radial glia, the stem cells of the developing brain and precursors of the adult SVZ-NSCs. We additionally identified a module (red, Figure 3F) specifically associated with the "stemness" transcriptional program and the cell cycle.

Interestingly, some modules were also associated with human disease, notably Huntington's disease (Thomas et al., 2011), Alzheimer's disease, convulsive seizures, major depressive disorder, and various cancers (Subramanian et al., 2005; Zhang





et al., 2010) (Figures S3A–S3F). For instance, the striatal neuron module (salmon) correlated with a gene expression set misregulated in Huntington's disease mouse models, suggesting a potential role for the 88 lncRNAs in this set in this neurodegenerative condition. Taken together, our coexpression analysis provides an important resource as a comprehensive annotation of lncRNAs to specific neural cell types in vivo and neurological disease states.

RNA CaptureSeq Verifies SVZ IncRNA Expression and Identifies Novel Splice Isoforms

Because many IncRNAs have not been previously annotated and are expressed at low levels, we employed a targeted RNA capture and sequencing strategy (CaptureSeq) to more robustly identify and characterize IncRNAs in the adult SVZ. With RNA CaptureSeq, cDNAs are hybridized to probe libraries tiled against the genomic regions of interest, eluted, and then sequenced (Figure 4A). Through this enrichment, the read Figure 2. mRNAs and IncRNAs Have Temporally and Spatially Unique Expression Patterns (A) Schematic summarizing regions, colored in yellow, used for this analysis.

(B) Hierarchical clustering results of all transcripts expressed across all samples.

(C) Hierarchical clustering results of IncRNAs expressed across all samples. The Pearson correlation coefficient was used as the distance metric. DG, dentate gyrus; STR, striatum; SVZ, subventricular zone; STR/SVZ, mixed dissection including both SVZ and striatal regions; OB, olfactory bulb; CTXA, cortical dissection laver 2/3: CTXB, cortical dissection laver 4: CTXC, cortical dissection laver 5: CTXD, cortical dissection layer 5; CTXE, cortical dissection layer 6; CTXF, cortical dissection layer 6b; POA, preoptic area; PFC, prefrontal cortex; E15, whole embryonic day 15 brain; VZ, ventricular zone of E14.5 cortex; SVZ/IZ, subventricular zone/intermediate zone of E14.5 cortex; CP, cortical plate of E14.5 cortex; ESC, cultured embryonic stem cells; NPCs, ESC-derived neural progenitor cells. See also Figure S2 and Table S2.

coverage of targeted transcripts is dramatically increased (Mercer et al., 2012). Furthermore, by using a 454 GS-FLEX Titanium instrument for sequencing, we obtained longer reads, which improve the delineation of rare splice isoforms.

For our RNA CaptureSeq probe library, we tiled across 100 MB of putative IncRNA loci and 30 MB of protein-coding regions as a control. We used this library to capture SVZ cDNA for sequencing (5,882,293 reads, median length of 356 bases per read). As expected, de novo assembly of sequences accurately reconstructed protein-coding transcripts and previously annotated IncRNAs (median identity of 90% for protein-coding RefSeq genes and median identity of 95% for annotated non-

coding RefSeq RNA). As an example, *Evf1* and *Evf2*, IncRNAs with roles in neural development, have overlapping genomic structures (Feng et al., 2006), and RNA CaptureSeq identified and distinguished both transcripts in the SVZ (Figure S4A). RNA CaptureSeq also eliminated sequencing bias related to transcript abundance (Figures S4B and S4C), and measured expression values were well correlated between CaptureSeq and conventional RNA-seq strategies (Figure S4D and Supplemental Experimental Procedures).

The enrichment and longer reads provided by RNA CaptureSeq enabled the identification of rare lncRNAs as well as uncommon splice isoforms in the SVZ transcriptome, yielding more than 3,500 lncRNAs that could not be detected by the short-read sequencing technology. For example, CaptureSeq identified an lncRNA transcript with an intron overlapping *Pou3f3*, a known neurogenic transcription factor (Figure 4B). In addition to this discovery of an lncRNA locus downstream of *Pou3f3*, RNA CaptureSeq also identified splice isoforms



Figure 3. IncRNAs Are Associated with Specific Neural Cell Types and Cellular Processes

(A–F) Top of each panel: heat maps depicting expression levels for six modules of coexpressed transcripts (rows) in 22 samples (columns) representing various brain regions and cell lines. Samples are labeled as in Figure 2. Red, increased expression; black, neutral expression; green, decreased expression. Middle of each panel: barplots of the values of the module eigengenes (Horvath and Dong, 2008), which correspond to the first principal component obtained by singular value decomposition of each module. Modules were characterized by performing enrichment analysis with known gene sets (see Table S3 and Supplemental Experimental Procedures). Bottom of each panel: pie charts indicating the abundance of IncRNAs within each module. Module members are defined as all transcripts that were positively correlated with the module eigengene at $p < 2.61 \times 10^{-8}$ (Supplemental Experimental Procedures). See also Figure S3 and Table S3.



Figure 4. RNA CaptureSeq Validates SVZ IncRNA Expression and Reveals Multiple Isoforms and Complex Locus Structures

(A) Schematic of RNA CaptureSeq procedure. We used Cufflinks's IncRNA assembly to define putative IncRNA loci and designed tiled probe libraries against these loci. The cDNA library was then hybridized to this biotin-labeled probe library, and after purification by streptavidin, the enriched population of IncRNAs was sequenced by 454 (Roche) long-read chemistry.

(B) Isotigs assembled at the *Pou3f3* locus revealed a distal transcriptional start site for a transcript that can be spliced into known noncoding RNA *2610017/09Rik*. (C) CaptureSeq-derived reads correctly assembled known protein-coding gene *Nr2f1* and identified four distinct TSSs for an IncRNA transcribed divergently from the *Nr2f1* promoter. The syntentic region in human reveals a similar organization of CpG islands and divergent transcriptional start sites for noncoding transcripts. Genes derived from RefSeq are colored purple, and genes from Ensembl are red. See also Figure S4.

that include exons of a previously annotated IncRNA (2620017I09Rik) that lies upstream of the Pou3f3 locus. Some IncRNAs are transcribed from multiple TSSs, which can be a challenge for transcript assembly. Adjacent to the locus of neurogenic transcription factor Nr2f1, CaptureSeq identified a

series of IncRNAs originating from four unique TSSs. This organization of protein-coding gene and multiple IncRNAs is conserved in humans, hinting at an evolutionarily conserved functional significance (Figure 4C). Thus, RNA CaptureSeq, in addition to providing a genome-wide validation of our SVZ IncRNA analysis, demonstrated previously underappreciated complexity to the structure of IncRNA loci. A complete annotation of CaptureSeq-derived transcripts is available at http:// neurosurgery.ucsf.edu/danlimlab/IncRNA.

Correlation between Histone Modifications and IncRNA Expression

Methylation of histone lysine residues is a critical determinant of transcriptional activity (Jenuwein and Allis, 2001). In previous work, IncRNA loci have been identified in part by the presence of H3K4me3 at the TSS (Guttman et al., 2009). For proteincoding genes, H3K4me3 enrichment at the TSS correlates with active transcription, whereas H3K27me3 is associated with a repressed state. Genes that are "bivalent" for both H3K4me3 and H3K27me3 are generally silenced but remain transcriptionally "poised" for activation or repression (Bernstein et al., 2006). To investigate whether IncRNA loci exhibit a similar correlation between promoter histone modifications and transcription, we performed chromatin immunoprecipitation sequencing (ChIPseg) for H3K4me3 and H3K27me3 in SVZ-NSC cultures and included sequencing data from ChIP-seg and RNA-seg studies of mouse ESCs, ESC-NPCs, and mouse embryonic fibroblasts (MEFs) (Mikkelsen et al., 2007).

In SVZ-NSCs, 3,671 (40.8%) IncRNAs were marked by either H3K4me3 or H3K27me3, and 928 (10.3%) were bivalent (Figure S5A). As has been described for protein-coding genes, these TSS chromatin modifications correlated strongly with IncRNA expression levels: IncRNAs monovalent for H3K4me3 exhibited higher expression levels than those by marked by only H3K27me3 or bivalent chromatin domains (p < 0.0001, Mann-Whitney U, Figure S5B). These data suggest that transcription of both IncRNAs and protein-coding genes utilizes similar chromatin-based regulatory mechanisms.

A Subset of Bivalent IncRNAs in ESCs Become Resolved to Monovalent H3K4me3 in SVZ-NSCs

In ESCs, bivalent domains identify key developmental genes. As ESCs differentiate into lineage-specific cell populations, many of these bivalent genes become activated (monovalent H3K4me3) or repressed (monovalent H3K27me3), reflecting the lineage specification and restriction of developmental potential (Bernstein et al., 2006). Thus, genes that are more likely to play roles in the neural identity of SVZ NSCs would be those that are bivalent in ESCs, activated in SVZ-NSCs, and also repressed (H3K27me3 monovalent or bivalent) in a nonneural cell type (MEFs). We found 302 protein-coding genes that meet these criteria, and analysis revealed that the most statistically significant GO terms for these activated genes pertain to early brain development (Figure S5C). For example, proneural Asc/1, Pou3f3, and Pou3f2 were bivalent in ESCs, H3K4me3 monovalent in SVZ-NSCs, and H3K27me3 repressed in MEFs (Figure S5D).

One hundred IncRNAs have a similar pattern of chromatinbased changes (Figure 5A). Furthermore, 76% of this set of IncRNAs was also monovalent for H3K4me3 in ESC-NPCs, suggesting that these IncRNAs are common to an early neural development transcriptional program. An example is *Incpou3f2*: this IncRNA locus is 2 kb upstream of the locus for known neurogenic transcription factor POU3F2. Both the *Inc-* *pou3f2* and the *Pou3f2* loci were bivalent in ESCs, monovalent for H3K4me3 in NSCs, and H3K27me repressed in MEFs (Figure 5B). Given the known relationship between chromatin modifications and the expression of key developmental regulators, we propose that this set of IncRNAs is enriched for those that play roles in early neural commitment in the adult SVZ.

IncRNAs Can Retain Bivalency in an Adult Stem Cell Population

Tissue-specific stem cells also retain bivalency at key loci, possibly reflecting retained gene expression plasticity (Cui et al., 2009; Lien et al., 2011). Protein-coding genes that are bivalent in ESCs and NSCs were highly enriched for GO terms related to neurogenesis (e.g., neuron differentiation, axonogenesis; Figure S5E). For instance, in adult SVZ NSCs, DIx1 and Dlx2 are transcription factors required for interneuron development, and these were bivalent in NSCs (Figure S5F). Thus, the identification of IncRNAs that are bivalent in both ESCs and NSCs might enrich for those involved in neuronal differentiation. These criteria were met by 583 IncRNAs (Figure 5C), such as three splice variants encoded from an IncRNA locus located ~50 kb upstream of protein-coding gene Odf3l1 (Figure 5D). We propose that IncRNAs bivalent in SVZ-NSCs are enriched for those that function in neuronal lineage specification.

IncRNAs Are Dynamically Regulated during Neuronal Differentiation

We next sought to define the dynamic changes in IncRNA expression during SVZ neurogenesis. SVZ-NSCs cultured as a monolayer can efficiently recapitulate key aspects of in vivo neurogenesis as assessed by immunocytochemistry (ICC, Figures 6A and S6A) (Scheffler et al., 2005). We generated cDNA libraries from SVZ-NSC cultures in self-renewal conditions and after 1, 2, and 4 days of differentiation and hybridized to both custom IncRNA (see Experimental Procedures) and standard gene expression arrays. Included in the set of upregulated transcripts were genes related to SVZ neurogenesis (e.g., Dlx1/2 and Dlx5/6) (Table S4). Also as expected, genes expressed at higher levels early in the SVZ lineage (e.g., Egfr and Nestin) were in the set of downregulated transcripts (Table S4). Like mRNA transcripts, IncRNAs also exhibited similar patterns of induction and repression (Figures 6B and S6B) over this 4 day differentiation time course.

In Vivo SVZ Lineage Analysis of IncRNA Expression

The adult SVZ contains three major cell types that represent a developmental continuum: (1) activated NSCs, which express glial fibrillary acidic protein (GFAP) and the epidermal growth factor receptor (EGFR), (2) transit-amplifying cells, which are EGFR positive but GFAP negative, and (3) migratory neuroblasts, which are CD24 positive (Figure 6C). In addition, the SVZ contains GFAP+ cells that do not express EGFR, and these have been termed "niche" astrocytes (Pastrana et al., 2009). We used these cell-specific characteristics to perform FACS to acutely isolate cell populations representing each stage of this neurogenic lineage and the niche astrocytes (Figure 6D). cDNA libraries for each SVZ cell type were

Cell Stem Cell IncRNA Regulation of Neural Stem Cells



Figure 5. IncRNA Loci Can Be Bivalent in Stem Cell Populations

(A) Venn diagram highlighting IncRNAs that were bivalent in ESCs, monovalent H3K4me3 in SVZ-NSCs, and H3K27me3 repressed (monovalent or bivalent) in MEFs. (B) The *Pou3f2* promoter and the promoter (yellow boxes) of a nearby IncRNA demonstrated a similar pattern of histone modifications (bivalent in ESCs, repressed in MEFs, and activated in SVZ-NSCs).

(C) Venn diagram demonstrating the number of IncRNAs that were bivalent in both ESCs and SVZ-NSCs.

(D) A novel IncRNA locus ~50 kb downstream of protein-coding gene Odf3/1. The promoter was bivalent in both SVZ-NSCs and ESCs. See also Figure S5.

generated and hybridized to our custom lncRNA and standard gene expression microarrays. Expression levels of both protein-coding genes and lncRNAs were visualized by heat maps organized by k-means clustering (transcripts) and unsupervised hierarcical clustering (cell types) (Figures 6E and S6C).

Cell Stem Cell IncRNA Regulation of Neural Stem Cells



To confirm the separation of SVZ cells, we examined differential mRNA expression. We found that 12,812 protein-coding probe sets were differentially expressed in a comparison between activated SVZ-NSCs and migratory neuroblasts (Table S4). As SVZ-NSCs become activated and differentiate into transit-amplifying cells, *Dlx1/2* and *Ascl1* become upregulated (Doetsch et al., 2002; Kim et al., 2007), and this was reflected in our transcriptional profiles (Table S4). As expected, the transcriptome of migratory neuroblasts was enriched for *Dlx1/2* downstream targets, including *Dlx5* and *Arx*, as well as markers of young neurons, including *Tubb3* (Table S4). Thus, these transcriptomes represent distinct stages of the SVZ neurogenesis and also distinguish niche astrocytes from NSCs.

Similar to the cell culture data (Figure 6B), we found sets of IncRNAs that showed transient increases in transit-amplifying cells, repression throughout differentiation, and significant induction in the terminally differentiated neuroblast population (Figure 6E). By integrating our chromatin state maps with this microarray expression data, we were able to begin to define an "IncRNA signature" for each stage of neurogenesis in vivo (Table S5).

Figure 6. Analysis of IncRNA Expression in the SVZ Lineage In Vitro and In Vivo

(A) Immunocytochemistry (ICC) of SVZ-NSC differentiation in vitro. In proliferation conditions, the culture is composed of neural precursor cells including GFAP+ (green) NSCs. After growth factor withdrawal, cells in these cultures differentiate into Tuj1+ neuroblasts (red, increasing numbers at 2 days and 4 days).

(B) Heat map representing expression of lncRNAs that were changed >4-fold from proliferation conditions to 4 days of differentiation. Color bars (orange, peach, light blue, and dark blue) at the right represent gene clusters resulting from k-means clustering, k = 4, Pearson distance metric.

(C) Schematic of the SVZ lineage. GFAP+, EGFR+ stem cells (blue) give rise to transit-amplifying (TA, green) cells. These TA cells give rise to Cd24+ low migratory neuroblasts (red).

(D) FACS plots for isolation of the SVZ lineage. Cells were dissociated from freshly dissected SVZ tissue from the hGFAP-GFP mouse and stained with EGF conjugated to the A667 fluorophore and a CD24 antibody conjugated to PE.

(E) Heatmap of IncRNAs differentially expressed throughout the SVZ lineage in vivo. Genes differentially expressed >2-fold between activated NSCs and neuroblasts were k-means clustered using the Pearson correlation metric, k = 5. Color bars at the right (dark blue, light blue, orange, peach, and green) represent gene clusters resulting from k-means clustering. See also Figure S6 and Table S4.

Identification of IncRNAs with Roles in Adult SVZ Neurogenesis

To facilitate the identification of IncRNAs with potential roles in SVZ neurogenesis, we constructed an online resource that allows the user to easily filter the IncRNA catalog for multiple variables including locus chromatin-state, expression in FACS-isolated SVZ cells, and

regulation during in vitro neurogenesis (http://neurosurgery. ucsf.edu/danlimlab/IncRNA). Using this resource, we filtered for those IncRNAs that were bivalent in ESCs and H3K27me3 repressed in MEFs. Of this set, which includes *Inc-pou3f2*, 100 IncRNAs were monovalent for H3K4me3 in SVZ-NSCs (Figures 5A and 5B), which would predict expression in the adult SVZ; indeed, ISH (Figures S7A and S7B) revealed Inc-pou3f2 expression in the SVZ, and, as predicted by the FACS microarray data, this transcript was not detected in the OB (Figure S7B).

Like *Inc-pou3f2*, IncRNA *Six3os* was also monovalent for H3K4me3 in SVZ-NSCs and downregulated in neuroblasts. Consistent with these observations, *Six3os* transcripts were detected in the SVZ but not the OB core (Figure 7A). To further investigate the role of *Six3os* in SVZ NSCs, we used lentiviruses to separately introduce two different short hairpin RNA (shRNA) sequences to knockdown *Six3os* (LV-sh-*Six3os*-GFP) in monolayers of SVZ-NSCs. After confirming *Six3os* knockdown in proliferating NSCs (Figure S7C), we assessed neuronal and glial lineages from LV-sh-*Six3os*-GFP-infected cells in comparison to controls infected with LV-sh-luc-GFP. After 7 days of differentiation, there were 2-fold fewer Tuj1-positive cells and 3-fold fewer





Figure 7. Functional Validation of IncRNA Candidates

(A) In situ hybridization (ISH) for *Six3os* using branched DNA probes. Positive signal is revealed by Fast Red alkaline phosphatase substrates, which appear as highly fluorescent, punctate deposits (left); DAPI nuclear counterstain is shown at the right. Blue boxes in SVZ and OB coronal schematics at left indicate regions shown at right. Scale bars represent 10 µm. V, ventricle; STR, striatum.

(B) Control (LV-sh-Luci-GFP) lentiviral infections in SVZ-NSC cultures after 7 days of differentiation. Top: immunocytochemistry (ICC) for Tuj1 (red) and GFP (green); middle: ICC for GFAP (red) and GFP (green); bottom: ICC for OLIG2 (red) and GFP (green).

(C) Analysis of *Six*3os knockdown in SVZ-NSCs after 7 days of differentiation. Two different constructs were used (sh-Six3os-1, sh-Six3os-2). Top: ICC for Tuj1 (red) and GFP (green); middle: ICC for GFAP (red) and GFP (green); bottom: ICC for OLIG2 (red) and GFP (green) after infection with a vector expressing shRNAs targeting *Six*3os (LV-sh-*Six*3os-GFP). Quantification of data is presented at right. Scale bars represent 10 μ m. Error bars represent SEM, five to six replicates for control group, two to three per experimental group. *p < 0.05, **p < 0.01, compared to sh-Luci, two-tailed t test.

(D) ISH with branched DNA probes for *Dlx1as* in the SVZ (top) and OB (bottom). Scale bars represent 10 μ m. V, ventricle; STR, striatum.

(E) Analysis of *Dlx1as* knockdown after 7 days of differentiation. Two unique targeting sequences were used (sh-Dlx1as-4, sh-Dlx1as-7). Top: ICC for Tuj1 (red) and GFP (green); middle: ICC for GFAP (red) and GFP (green); bottom, ICC for OLIG2 (red) and GFP (green). Quantification of data is presented at right. Scale bars represent 10 μ m. Error bars represent SEM, five to six replicates for control group, three per experimental group. *p < 0.05, **p < 0.01, compared to sh-Luci, two-tailed t test. See also Figure S7 and Table S5.

cells expressing OLIG2, a marker of the oligodendrocyte lineage (Zhou and Anderson, 2002). These decreases were accompanied by an increase in the number of GFAP+ cells (Figures 7B and 7C).

The DIx1/2 bigene cluster encodes IncRNA DIx1as (Liu et al., 1997; Dinger et al., 2008), and this locus was also bivalent in ESCs and H3K27me3 monovalent in MEFs. We used SVZ-NSC monolayer cultures to further investigate the chromatin state of the DIx1as TSS. In self-renewal conditions, DIx1as was bivalent, and after 30 hr of differentiation, H3K27me3 decreased, correlating with the start of *Dlx1as* upregulation (Figures S7D and S7E). Interestingly, we also found enrichement of the H3K27me3-specific demethylase JMJD3 (Agger et al., 2007) at the DIx1as TSS during differentiation (Figure S7F), suggesting that this chromatin-modifying factor plays a role in the activation of this IncRNA. Consistent with the transcriptional upregulation of DIx1as during SVZ neurogenesis in vitro, we observed robust Dlx1as expression in SVZ regions with migratory neuroblasts and the OB core (Figure 7D). We designed two knockdown constructs targeting the splice junction of Dlx1as and verified that these constructs target Dlx1as and not full-length Dlx1 transcript (Figure S7G). Knockdown of Dlx1as caused a decrease in expression of *Dlx1* and *Dlx2* after 2 days of differentiation compared to control (Figure S7H), suggesting that this IncRNA can regulate expression of its protein-coding gene neighbors. After 7 days of differentiation, we found a nearly 3-fold decrease in Tuj1+ neuroblasts, and an ~60% increase in the number of GFAP+ astrocytes. In contrast to knockdown of *Six3os*, the production of OLIG2+ cells was unaffected by *Dlx1as* knockdown (Figures 7B and 7E).

DISCUSSION

We performed an in-depth analysis of IncRNA expression of adult SVZ-OB neurogenesis, an excellent in vivo model system for the study of multipotent stem cells and neural development. Our use of two high-throughput sequencing-based approaches for the study of the IncRNA transcriptome (RNA-seq and RNA CaptureSeq) provided complementary data sets that together allowed the identification of thousands of novel IncRNAs, confirmation of rare transcripts, and resolution of previously unappreciated complexity of IncRNA loci.

Like the loci of genes encoding key developmental transcription factors, a subset of IncRNA loci showed changes of chromatin state during lineage specification. By integrating these chromatin state maps with data from custom microarrays and FACS purification of the SVZ lineage, our online resource (http://neurosurgery.ucsf.edu/danlimlab/lncRNA) and files (Table S5) facilitate the identification lncRNAs with potential roles in adult NSCs as well as neural development. Interestingly, we found that *Dlx1as* is required selectively for the SVZ neuronal lineage, whereas *Six3os* appears to play a role in both neuronal and oligodendrocyte differentiation. These data indicate that lncRNAs can play key roles in the glial-neuronal lineage specification of multipotent adult stem cells.

There were 2,265 IncRNAs that had proximal protein-coding gene neighbors (Figure S1D), and this gene set was enriched for homeodomain-containing genes. For instance, Six3os is proximal to the Six3 homeobox gene, and DIx1as is encoded from the Dlx1/2 bigene cluster (Liu et al., 1997; Dinger et al., 2008). Interestingly, Dlx1as knockdown caused a decrease in Dlx1/2 expression (Figure S7H), suggesting that this IncRNA plays a role in neuronal differentiation by regulating expression of its homeobox gene neighbors. In developing retina, knockdown of Six3os results in deficits in lineage specification through modulation of SIX3 activity (Rapicavoli et al., 2011); it will be interesting to investigate whether the defect in SVZ neurogenesis with Six3os knockdown (Figure 7C) similarly involves a change in SIX3 activity. Taken together, our genome-wide analysis and functional data further support the notion that IncRNAs and homeobox gene neighbors function cooperatively (Rapicavoli and Blackshaw, 2009).

A recent model of IncRNA action suggests that lineage-specific IncRNAs become activated during differentiation and guide histone modifications that create cell type-specific transcriptional programs (Guttman et al., 2011). MLL1 is a trithorax group (trxG) chromatin-modifying factor that is enriched at Dlx2 during SVZ NSC differentiation and is required for proper Dlx2 expression (Lim et al., 2009); however, how MLL1 is targeted to Dlx2 is not known. Interestingly, in mouse ESCs, IncRNA Mistral directly binds MLL1 and recruits it to Hoxa6 and Hoxa7 (Bertani et al., 2011) and IncRNA HOTTIP recruits MLL1 through an interaction with WDR5 to distal HOXA genes in human fibroblasts (Wang et al., 2011). Our work provides a useful resource for the identification of such IncRNAs. For instance, IncRNAs that immunoprecipitate with chromatin modifiers could be identified by hybridization to the IncRNA microarray and then filtered online by multiple other criteria (e.g., enrichment in neuroblasts, upregulation during neurogenesis, bivalency in ESCs, and repression in MEFs).

Our analysis of chromatin state maps and transcript expression suggest that histone modifications correlate with IncRNA expression in a manner similar to that of protein-coding genes. Some IncRNA loci were bivalent in both ESCs and SVZ-NSCs, and many of these IncRNA loci became transcriptionally active in SVZ neuroblasts, supporting their candidacy as key determinates of neurogenesis. In SVZ-NSC monolayer cultures, *Dlx1as* was bivalent and H3K27me3 repression decreased during neuronal differentiation (Figure S7D), correlating with the upregulation of *Dlx1as* transcription (Figure S7E). Interestingly, we also found enrichment of the H3K27me3-specific demethylase JMJD3 at the *Dlx1as* locus (Figure S7F), suggesting that active removal of repressive histone modifications plays a role in the expression of IncRNAs. Overall, our data raise the possibility that IncRNA loci, like protein-coding genes, are targeted by chromatin-modifying factors that have critical roles in development.

While this study attempted to be as comprehensive as possible, it is possible that some IncRNAs important for SVZ neurogenesis were not identified. The initial sequencing experiments were performed on microdissected tissues that contain several cell types. Even at our sequencing depth, transcripts that are expressed at low copy number in a small number of cells might not be detected. Despite this potential shortcoming, we were still able to identify thousands of previously unannotated IncRNA transcripts. Furthermore, our initial catalog proved sufficient for our primary objective, which was to integrate complementary data analysis strategies and experimental methods to identify IncRNA expression patterns coherent to an in vivo experimental model system.

The role of IncRNAs in development and disease is in the early states of investigation, and our analysis of the SVZ lineage provides a resource for the movement of this research into in vivo studies. More broadly, this work presents a generalizable workflow for the identification and categorization of novel transcripts, both coding and noncoding.

EXPERIMENTAL PROCEDURES

Brain Dissection for RNA Extraction

The brain from adult (older than postnatal day 60) male C57/B6 mice was removed from the skull and placed in ice-cold L15 media and a 0.5-mm-thick coronal slab was obtained. The lateral SVZ and striatum were then microdissected, avoiding contamination from the corpus callosum. FACS of SVZ cells was performed as described in Pastrana et al. (2009). DG was microdissected in ice-cold L15 media from 300- μ m-thick coronal sections obtained with a Vibratome.

High-Throughput Sequencing

Illumina libraries were prepared using standard protocols. Sequencing was performed with a Genome Analyzer IIx or Hi-Seq instrument (Illumina). CaptureSeq followed by 454 sequencing (GS-FLX Titanium, Roche) was performed essentially as described in Mercer et al. (2012).

Sequencing Data Processing

Sequencing analysis was performed on the Galaxy platform (Goecks et al., 2010), using TopHat (Trapnell et al., 2009) and Bowtie (Langmead et al., 2009) for read alignment. Table S2 describes all novel and previously published data sets used in this work. Cufflinks and Cuffdiff (Trapnell et al., 2010) were used for transcript reconstruction, quantification, and differential expression analysis. MACS was used to call ChIP-seq peaks (Zhang et al., 2008). For longer 454 reads, BLAT (Kent, 2002) was used to construct contigs.

IncRNA Catalog Assembly

We filtered .gtf files from Cufflinks using the UCSC genome browser to remove all single-exonic transcripts as well as any transcripts overlapping RefSeq genes, known protein-coding genes from other species using the "Other Ref seq" track and the "XenoRefGene" table. These filtered .gtf files were used as input for CuffCompare, and the complete Ensembl gene annotation .gtf file (downloaded from http://www.ensembl.org/info/data/ftp on May 23, 2011) was used as reference. We kept any transcript with the CuffCompare designation "I," which marks all novel intergenic sequences. To these sequences, we added annotated lincRNAs and processed transcripts from Ensembl. Finally, we excluded all genes known to RefSeq, giving the 8,992 transcripts in Table S1 a .bed file containing our putative lncRNAs. This file was merged with the Illumina iGenomes UCSC mm9 gtf annotation of all RefSeq genes (http://cufflinks.cbcb.umd.edu/igenomes.html on January 5, 2012), generating a final comprehensive annotation of noncoding and coding transcripts, which was then used for all subsequent analysis of RNA-seq data.

CaptureSeq Library and Expression Arrays

CaptureSeq and Expression Array designs are available upon request. For the CaptureSeq probe set, probes were designed to tile across \sim 100 MB of putative noncoding loci and \sim 30 MB protein-coding control loci, for a total of 6,287 genomic targets. For expression array probe selection, a FASTA file was generated representing all putative lncRNAs. Probe selection and design was performed by Roche Nimblegen according to their standard design rules.

In Situ Hybridization

In situs were performed using the QuantiGene ViewRNA ISH tissue Assay (Affymetrix). Probes were designed based on IncRNA sequences determined by Cufflinks. Images were taken on a DMI4000B fluorescent microscope (Leica).

ACCESSION NUMBERS

All data are deposited in NCBI GEO under accession number GSE45282.

SUPPLEMENTAL INFORMATION

Supplemental Information includes seven figures, Supplemental Experimental Procedures, and five tables and can be found with this article online at http://dx.doi.org/10.1016/j.stem.2013.03.003.

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