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Development, Maturation, and Necessity of Transcription Factors in the Mouse Suprachiasmatic Nucleus

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The suprachiasmatic nucleus (SCN) of the hypothalamus is the master mammalian circadian clock. The SCN is highly specialized because it is responsible for generating a near 24 h rhythm, integrating external cues, and translating the rhythm throughout the body. Currently, our understanding of the developmental origin and genetic program involved in the proper specification and maturation of the SCN is limited. Herein, we provide a detailed analysis of transcription factor (TF) and developmental-gene expression in the SCN from neurogenesis to adulthood in mice (*Mus musculus*). TF expression within the postmitotic SCN was not static but rather showed specific temporal and spatial changes during prenatal and postnatal development. In addition, we found both global and regional patterns of TF expression extending into the adult. We found that the SCN is derived from a distinct region of the neuroepithelium expressing a combination of developmental genes: *Six3*, *Six6*, *Fzd5*, and transient *Rx*, allowing us to pinpoint the origin of this region within the broader developing telencephalon/diencephalon. We tested the necessity of two TFs in SCN development, *ROR α* and *Six3*, which were expressed during SCN development, persisted into adulthood, and showed diurnal rhythmicity. Loss of *ROR α* function had no effect on SCN peptide expression or localization. In marked contrast, the conditional deletion of *Six3* from early neural progenitors completely eliminated the formation of the SCN. Our results provide the first description of the involvement of TFs in the specification and maturation of a neural population necessary for circadian behavior.

Introduction

The suprachiasmatic nucleus (SCN) of the hypothalamus acts as the master mammalian circadian clock generating a near 24 h rhythm through the cyclic expression of key genes, including crucial transcription factors (TFs). The ability of the SCN to integrate external cues and translate rhythms throughout the body reflects a unique specialization of the nucleus and neurons within (Herzog and Schwartz, 2002). Properties underlying the production of 24 h rhythms have been well characterized; however, the genes involved in specification and development of the SCN are unknown.

The SCN is a heterogeneous group of neurons able to generate unified rhythmic output from individual neuronal oscillators (Webb et al., 2009). Expression of arginine vasopressin (AVP) and vasoactive intestinal peptide (VIP) divide the nucleus into

two subdivisions, termed dorsomedial/shell and ventrolateral/core, respectively. Although these gross divisions do not necessarily reflect differences in cellular properties, these areas have been shown to differ in their projection patterns (Leak and Moore, 2001), innervation (Moga and Moore, 1997; Abrahamson and Moore, 2001), and rhythmicity (Hamada et al., 2001; Yan and Okamura, 2002; Bendová et al., 2004; Karatsoreos et al., 2004).

There is an extensive literature describing the birth dating, onset of neurogenesis, and migration of neurons in the SCN of mammals. Initial experiments in rat estimated that the SCN is derived from neuroepithelium caudal to the optic recess during the third wave of hypothalamic development (Altman and Bayer, 1986). In addition, studies in mice have shown that SCN neurogenesis occurs in a ventrolateral to dorsomedial progression approximately 5 d before birth (Shimada and Nakamura, 1973; Okamura et al., 1983; Kabrita and Davis, 2008), consistent with previous findings in rat and hamster (Altman and Bayer, 1978, 1986; Davis et al., 1990; Antle et al., 2005). Additional information regarding the prenatal SCN remains limited, in part because of the difficulty in identifying SCN boundaries beyond cell density estimates before peptide expression. In addition, no single gene has yet been identified that unambiguously distinguishes all SCN neurons from adjacent neural populations.

Individual TFs can distinguish subsets of cells within heterogeneous populations and provide insight into both development and function (Dasen et al., 2005; Gray, 2008). To date, the study of TFs within the SCN has been either in their relation to clock function or as a consequence of focus on neighboring hypothalamic

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lamic regions. Investigation of TFs specifically within the SCN may provide insight into SCN development, the presence of genetically distinct areas, and relationships between the SCN and other brain regions.

We hypothesized that SCN function is influenced by TF expression and that these genes play a role in both the formation and maturation of SCN rhythmicity. Thus, we provide a detailed spatiotemporal description of TF and developmental-gene expression within the mouse SCN. We find that the SCN undergoes prenatal and postnatal maturation of gene expression. Furthermore, we find that patterns of TF expression reveal a specific gene cascade from which SCN neurons are derived and can be used to identify the developing SCN before published markers. In addition, we demonstrate a crucial role for an evolutionarily conserved TF in SCN formation.

Materials and Methods

Animals and housing. Male and female wild-type CD1 (Charles River), *Six3^{lox}* (Guillermo Oliver, St. Jude Children's Research Hospital, Memphis, TN), *RORα⁸* (002651; The Jackson Laboratory), and *Nestin-cre* (003771; The Jackson Laboratory) mice were used. Mice were maintained in a normal 12 h light/dark schedule under standard care conditions. Mutant animals were maintained on a C57BL/6 background (Charles River). All experiments were approved by the Animal Studies Committee at Washington University School of Medicine.

Genotyping. Mice were genotyped by PCR for disrupted *RORα* gene, *Six3* floxed, and/or wild-type alleles as described previously (Liu et al., 2006) or as outlined by The Jackson Laboratory. The presence of Cre was determined by the presence or absence of an amplified band using primers 5' GAGTGAACGAACCTGGTCCGAAATCAGTGC 3' and 5' GCATTACCGGTTCGATGCAACGAGTGATGAG 3'.

Tissue acquisition. Unless otherwise stated, all collections were done at approximately Zeitgeber time 8 (ZT8). Neonatal pups [postnatal day 0 (P0) to P2] or embryos [embryonic day 10.5 (E10.5) to E18.5] from timed pregnant females (morning of plug was E0.5) were anesthetized on ice and either transcardially perfused (\geq E16.5) or immersion fixed in 4% paraformaldehyde (PFA) in 0.1 M PBS, pH 7.4. Older mice were anesthetized with a ketamine/xylazine mixture before perfusion with 4% PFA. All tissues were postfixed in 4% PFA overnight at 4°C, cryoprotected in 20% sucrose in PBS, frozen in O.C.T. Compound Embedding Medium (Tissue-Tek), and stored at -75°C. Serial sections (14 or 20 μ m) were cut on a Hacker cryostat and thaw mounted on Superfrost Plus slides (Thermo Fisher Scientific). Five adjacent sets of sections were prepared from each embryonic and postnatal age and stored at -20°C.

ZT tissue collection. Tissue was collected as outlined above every 4 h for 24 h from animals (P21–P23) kept in a 12 h light/dark schedule starting at ZT2, 2 h after lights on (ZT0). For dark time points, animals were enucleated under red light before perfusion.

Probe synthesis. Plasmids for *in situ* hybridization (ISH) probes were acquired from published sources (Gray et al., 2004; Burns et al., 2008; Zhang et al., 2008) or purchased from Open Biosystems. Gene fragments from verified plasmids were linearized by direct amplification by sequence or vector-specific PCR (gene, GenBank accession number, region, forward primer, reverse primer); AVP, BC051997, GTAATACGACTCACTATAGG (T7), AATTAACCCCTCACTAAAGGG (T3); BMP7, BC010771, 281–993, GCCTCTGTCTTGTGCTGCCCTCC, CCTGGATGGGCAGACATCA; Dlx2, BQ443330, 75–797, CGGGCTCCGCTTCACACCTG, ACGGAGTGTCCTGGGAAAGTGA; Hmx2, BC023402, T7, ATTTAGGTGACACTATAG (SP6); Per1, BC039768, 532–1750, CCCAGAAAGAACTCATGACTG, GCATCTGGTAAAGCACCAGGG; Ptc1, CB723783, T7, T3; ROR α , BC003757, 273–1054, GCCACCTACTCCTGTCTCCTCG, GCAG-GCTCGCTAGAGTGGTGT; ROR β , BC024842, 331–987, CAG-GAACCGTTGCCAACACTGC, CCAGTACGTGGTGGAGTTCGC; Rx, BC024731, 539–1261, CCCGGGATTCTGCCCGGAGT, TTGGCCTTCAGGCGCAGAGC; sonic hedgehog (SHH), BC063087, 51–998, GAGACCAACTCCGATGTG, GAAAGCAGAGAACTCCGTGGC. Digoxigenin (DIG)-labeled antisense and sense RNA probes were made

using PCR products as template and T7, T3, or SP6 RNA polymerases (Roche). cRNA probes were purified using Quick Spin columns (Roche) and quantified by spectrophotometry. Probes were used at a concentration of 1–2 μ g/ml. Sense counterparts of all probes were tested to ensure probe specificity.

In situ hybridization. Slides were immersed in 4% PFA, permeabilized with proteinase K, and returned to 4% PFA before being washed in 0.1 M triethanolamine-HCl with 0.25% acetic anhydride. Once blocked in hybridization buffer at 65°C slides were incubated in hybridization buffer containing 1–2 μ g/ml DIG-labeled antisense cRNA overnight at 65°C. Slides were then washed in 2 \times SSC at 62°C, washed in 0.2 \times SSC at 65°C, blocked with 10% normal horse serum (NHS) in 0.1 M PBS, and incubated in alkaline phosphatase-labeled anti-DIG antibody (1:2000 in 10% NHS; Roche) overnight. Sections were washed and color was visualized using nitro blue tetrazolium and 5-bromo-4-chloro-3-indolyl phosphate (Roche). Staining was stopped after visual inspection. Sections were washed, fixed in 4% PFA, and coverslipped in 90% glycerol, Vectashield Mounting Medium (Vector Laboratories), or UltraCruz Mounting Media with 4',6'-diamidino-2-phenylindole (DAPI) (Santa Cruz Biotechnology).

Immunohistochemistry. Sections on slides were dried at room temperature and rinsed in 0.5% Triton X-100 in PBS (PBS-T) before incubating in 10% NHS in PBS-T. Slides were then placed into primary antibody overnight at 4°C. Primary antibodies used include the following: rabbit (Rb) AVP (1:800; Immunostar), Rb Ki67 (1:500; Thermo Fisher Scientific), goat (Gt) Lhx2 (1:500; Santa Cruz Biotechnology), Gt Lhx9 (1:750; Santa Cruz Biotechnology), mouse TuJ1 (1:1000; Covance), and Rb VIP (1:8000; Immunostar). Primary antibodies were removed, and sections were rinsed in PBS-T before incubation in secondary antibodies for 2 h at room temperature. Secondary antibodies used were either the appropriate Alexa Fluor 488 (1:1000; Invitrogen) or cyanine 3 (1:500; Jackson ImmunoResearch). Sections were washed and coverslipped with UltraCruz Mounting Media with DAPI.

Image acquisition. Images were acquired using a Nikon Eclipse 90i microscope, Photometrics Coolsnap HQ2 camera with a Prior Scientific ProScan II motorized translation stage, and acquired in Volocity (PerkinElmer Life and Analytical Sciences) or a Nikon Coolsnap V Ed slide scanner with SilverFast SE software (LaserSoft Imaging). Images were exported as eight-bit JPEG or TIFF files. All images were adjusted for clarity by filtering and/or modifying levels, as necessary, in Photoshop (Adobe Systems).

mRNA quantification and analysis. Percentage of SCN area containing either Lhx1 or ROR α expression was calculated by dividing area expressing mRNA by the total estimated SCN area marked with DAPI, in which area was calculated by outlining regions of interest within NIH ImageJ. Whole SCN quantifications of mRNA expression were performed by quantifying intensity of staining relative to background from 10 \times ISH images using NIH ImageJ. Margins of the SCN in E17.5 to adult animals were determined by cell density marked with DAPI nuclear stain. For statistical analysis of diurnal variation, one-way ANOVA was applied to relative intensity measurements obtained from ZT2, ZT6, ZT10, ZT14, and ZT18 for *Six3*, ROR α , and *Per1*. The differences of mRNA relative intensity between ZT6 and ZT18 were analyzed using Student's *t* test.

Results

Identification of SCN candidate transcription factors

The SCN is a specialized neural structure with an essential behavioral function. We hypothesized that aspects of SCN neurons and SCN function are a direct consequence of the TFs they express and that underlie specification. We set out to identify TFs that might be important for both the development and function of the adult SCN by reexamining a published analysis of nearly 1000 TF and TF-related gene expression patterns in mice at E13.5 and P0 (Gray et al., 2004). Two hundred twenty TFs were previously identified as restricted to populations within the hypothalamus of E13.5 and/or P0 mice. From these data as well as other published work (Rivkees et al., 1992; Schaeren-Wiemers et al., 1997; Jean et al., 1999; Wang and Lufkin, 2000), we identified 72 TFs as possibly expressed in discrete patterns of the postnatal (P0; sup-

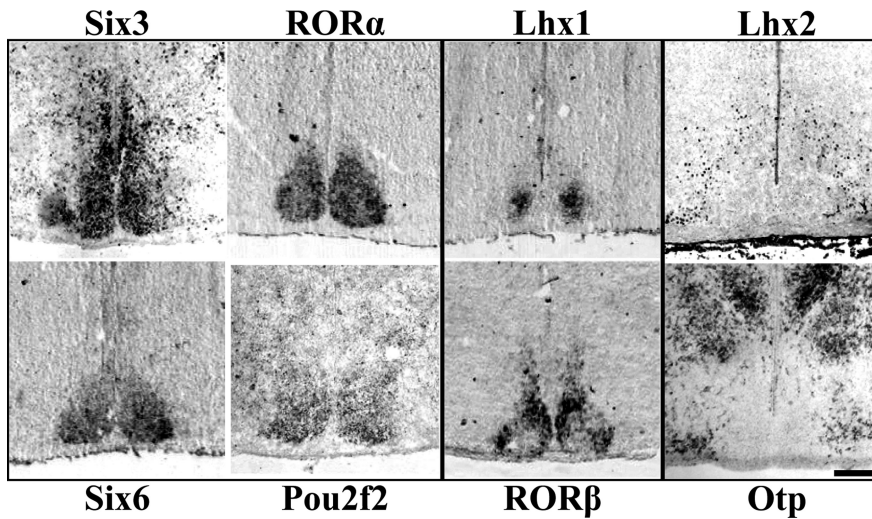


Figure 1. Diverse patterns of transcription factor expression in P0 mouse SCN. Representative ISH and IHC images of mid-SCN coronal sections showing specific expression throughout the entire SCN but not adjacent hypothalamus for Six3, ROR α , Six6, and Pou2f2, but restricted localization within the SCN for Lhx1 and ROR β . Lhx2 and Otp are expressed outside the SCN, indicating a transcription factor boundary. Scale bar, 250 μ m.

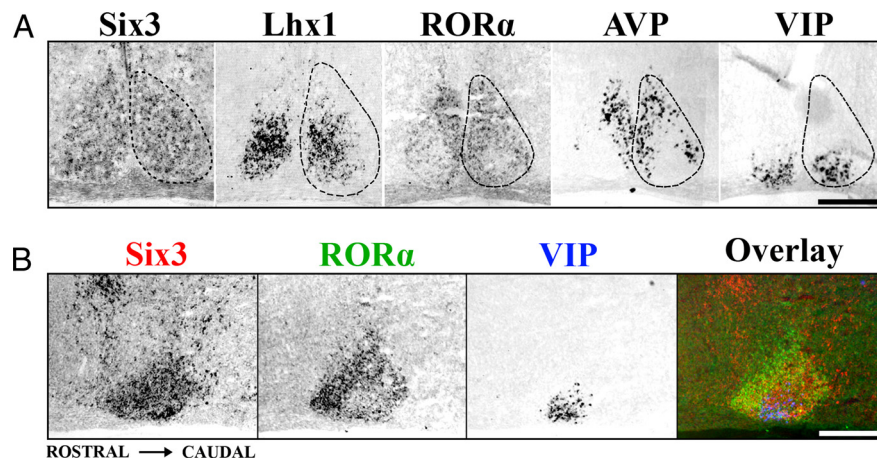


Figure 2. The adolescent mouse SCN shows discrete patterns of transcription factor expression. **A**, Representative ISH images showing localization of Six3, Lhx1, and ROR α expression within coronal sections of mid-SCN at P15. Note the discrete localization of ROR α to a region overlapping AVP but not VIP expression, whereas Lhx1 overlaps both. Dotted lines indicate the margins of the SCN. **B**, ISH images from adjacent sagittal sections of Six3 (red), ROR α (green), and VIP (blue) are overlaid in pseudocolor, showing the full rostral to caudal expression of Six3 throughout the SCN and the localized absence of ROR α from the VIP-expressing region. Scale bars, 250 μ m.

plemental Table 1, available at www.jneurosci.org as supplemental material) and/or prenatal (E13.5; supplemental Table 2, available at www.jneurosci.org as supplemental material) anterior hypothalamus that includes the SCN. Of these 72 TFs, 28 appeared potentially expressed within the postnatal SCN itself (supplemental Table 1, available at www.jneurosci.org as supplemental material). We confirmed the perinatal or adult hypothalamic expression of at least 68% of these 28 TFs using data from the Allen Brain Atlas (Lein et al., 2007).

SCN transcription factor expression is diverse and dynamic

Previous large screens of gene expression within the brain, although extremely informative, have generally lacked the spatial and/or temporal precision to determine the detailed boundaries of expression within the SCN (Gong et al., 2003; Gray et al., 2004; Lein et al., 2007; Shimogori et al., 2010). To directly address the

range of TF expression patterns within the early SCN, we selected a subset of nine postnatal TFs [Lhx1, Lhx2, Lhx9, Otp, Pou2f2 (Oct-2), ROR α (Rora), ROR β (Rorb), Six3, and Six6] for detailed ISH or immunohistochemical (IHC) analysis within the P0 mouse SCN. These genes were selected based on their relative levels of expression and specificity within the SCN region and/or previous work, suggesting a possible role in aspects of SCN function (Rivkees et al., 1992; Schaeren-Wiemers et al., 1997; Sato et al., 2004).

We found that TFs showed a range of expression patterns suggesting a diverse transcriptional environment (Fig. 1). Of the nine postnatal TFs analyzed, six were found specifically expressed in the SCN but not in adjacent hypothalamic nuclei. Compared with a cell density estimate of SCN margins, Six3, Six6, ROR α , and Pou2f2 were expressed throughout the entire SCN, whereas Lhx1 and ROR β both displayed regional specificity within the SCN. Conversely, we found two TFs (Lhx2 and Otp) expressed at lateral and dorsal boundaries but not within the SCN. Lhx9 was neither expressed within the SCN nor served as a boundary marker (data not shown).

To determine whether representative patterns of early postnatal TF expression are modified by postnatal maturation, we examined Six3, Lhx1, and ROR α further for expression at P15, just after eyes open. Six3 retained expression throughout the P15 SCN (Fig. 2A). Similar to expression at P0, Lhx1 displayed regionalized specificity within the SCN at P15. Expression of Lhx1 covered $52 \pm 2.6\%$ of the central SCN and partially overlapped both SCN ventrolateral/core and dorsomedial/shell (Fig. 2A). In contrast to expression at P0, ROR α was no longer present throughout the entire SCN; instead, expression was restricted to only the outer $71 \pm 2.2\%$ of the SCN, similar in distribution to AVP

and ROR β . ROR α expression did not overlap the VIP-expressing region, seen clearly in both transverse (Fig. 2A) and sagittal (Fig. 2B) sections through the SCN. We found that expression patterns of Six3, Lhx1, and ROR α at P15 were identical to those at P21 and adult (data not shown).

Six3 and ROR α show diurnal variation

The SCN can entrain to photoperiods (Sumova et al., 2006) and can be considered mature by P21. Also, by this time, the amplitude of circadian changes in expression level of clock genes has reached adult levels (Kováčiková et al., 2006). For clock genes, the differences in phase of expression are a consequence of the interactions in the transcription–translation negative feedback loop that underlie circadian rhythmicity. Much of our understanding of the phasic expression of genes within the SCN has relied on

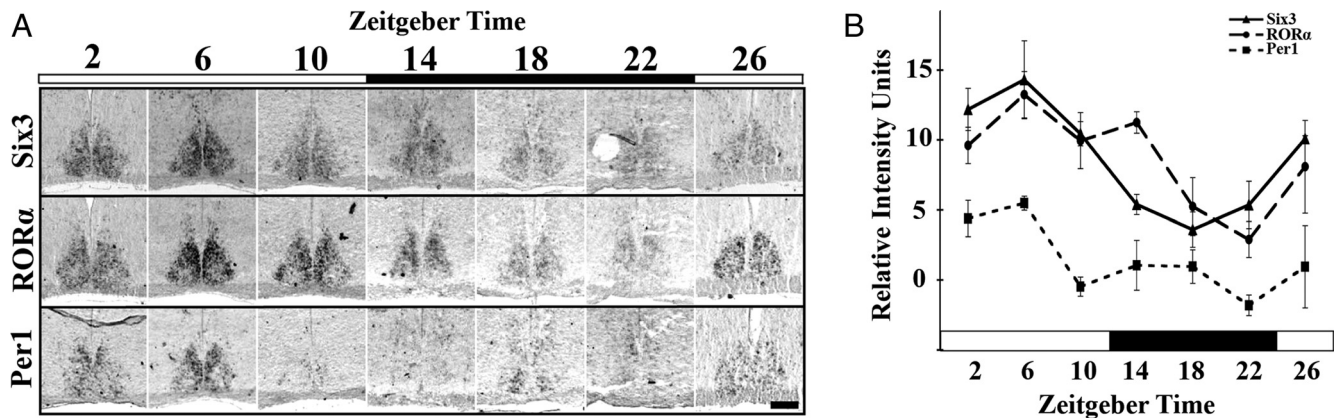


Figure 3. Six3 and ROR α show diurnal variation in expression. **A**, Representative ISH images showing expression of Six3, ROR α , and Per1 mRNA in the coronal mid-SCN in P21–P23 mice raised under a 12 h light/dark cycle and sampled every 4 h (ZT0 indicates lights on). In contrast to Per1 and Six3, ROR α is expressed only in the outer shell region. **B**, Quantification of averaged relative mRNA intensity in arbitrary units for Six3, ROR α , and Per1 over the same 24 h time period, showing diurnal variation for Six3 (one-way ANOVA, $F_{(5,47)} = 10.55, p < 0.0001$), ROR α (one-way ANOVA, $F_{(5,47)} = 6.64, p = 0.0001$), and Per1 (one-way ANOVA, $F_{(5,21)} = 4.56, p = 0.0089$), with peak at ZT6 and trough at ZT22 for ROR α and Per1 and ZT18 for Six3 ($n = 3$). Black bars indicate 12 h of dark. Error bars show SEM. Scale bar, 250 μm .

quantitative PCR from whole SCN lysates or radioactive ISH, both of which have limited spatial sensitivity (Sunkin, 2006).

Under standard 12 h light/dark conditions, the mRNA levels of central clock genes Per1, Bmal1, and Cry1 peak and trough at mid-day (ZT6) and mid-night (ZT18) (Oishi et al., 2000; Shearman et al., 2000), with Bmal1 and Cry1 in anti-phase of the mid-day peak of Per1. We tested with nonradioactive ISH whether Per1, Lhx1, ROR α , and Six3 mRNA displayed variations at these time points. We found that changes in Six3 and Per1 were significantly different between mid-day and mid-night ($p = 0.00064$ and 0.028 , respectively, Student's t test), whereas ROR α and Lhx1 levels were not ($p = 0.055$ and 0.1065 , respectively).

The spatial localization of ROR α at ZT8 in the mature SCN suggested that any change in phase of expression might correspond to changes in spatial localization as well as abundance. We tested the expression of Six3 and ROR α for changes in mRNA intensity and localization of mRNA over a diurnal period ($n = 3$ for each time point). We found that ROR α displayed temporal variation within the SCN (one-way ANOVA, $F_{(5,47)} = 6.64, p = 0.0001$), with peak at ZT6 and trough at ZT22, in phase with Per1 (one-way ANOVA, $F_{(5,21)} = 4.56, p = 0.0089$) (Fig. 3*A, B*), consistent with previous PCR-based analyses (Sato et al., 2004). ROR α was never expressed within the VIP-expressing region, which indicated a previously unrecognized spatial localization. Six3 also showed diurnal variations in expression, with peak at ZT6 and trough at ZT18 (one-way ANOVA, $F_{(5,47)} = 10.55, p < 0.0001$) (Fig. 3*A, B*).

Transcription factors identify the prenatal SCN

Although the SCN is not functionally mature until postnatal stages, early components of rhythmicity are present in late embryonic life. Some of the earliest rhythms were seen in day/night variations in metabolic activity as early as E19 in the rat (Reppert and Schwartz, 1984) as well as individual rhythmically firing neurons at E22 in the rat (Shibata and Moore, 1987), which corresponds to approximately E16.5 and E19.5 in mouse, respectively. The expression of Six3, Lhx1, and ROR α in both the early postnatal and mature SCN suggested that these same genes might provide insight into the development of the SCN during the prenatal stage in which cellular rhythmicity emerges.

To clearly identify the perinatal SCN, we determined the onset and localization of AVP and VIP by ISH and IHC. Consistent

with previous work in other rodents (Romero and Silver, 1990; Isobe et al., 1995; Ban et al., 1997), we found both AVP and VIP mRNA expression prenatally but protein-labeled cell bodies only at postnatal stages (data not shown). AVP mRNA expression was present in the SCN by E17.5 (two of three animals), while already strongly expressed in the supraoptic nuclei (SON) at this time (data not shown). VIP mRNA expression was present in the SCN by E18.5 (three of five animals) (data not shown). Within the SCN, both peptides were expressed in spatially restricted regions similar to that seen in the adult from their onset, suggesting that specification of peptidergic identity occurs before E17.5. The global expression of Six3 and ROR α in the SCN at P0 led us to theorize that their expression would identify the boundaries of the SCN during prenatal maturation. Thus, we analyzed the expression of Six3, Lhx1, and ROR α between the end of neurogenesis at E14.5 and birth (P0).

At E14.5, Six3 was expressed within a portion of the early hypothalamus in both the presumptive SCN region as well as dorsal areas. In contrast, Lhx1 was present in discrete bilateral regions distal to the third ventricle within the broader Six3 domain (Fig. 4). A small subset of the ventral most Lhx1/Six3 region was found to express ROR α in two of five animals examined, suggesting that the onset of ROR α (Fig. 4) occurs at this time point. At E15.5, before the onset of VIP or AVP mRNA expression, there was an increase in the area and approximate number of cells expressing Lhx1 and ROR α . Additionally, nuclear staining with DAPI revealed the first signs of aggregation into a cell-dense nucleus at this stage (Fig. 4, white arrow). However, both Lhx1 and ROR α expression extended beyond the newly forming, cell-dense area, which would be consistent with the presence of neurons that will become the SCN before the onset of clustering. We hypothesized that these genes could be used as prenatal genetic markers to define cells of the SCN more precisely than cell density estimates. By E17.5, ROR α expression encompassed the entire SCN and completely overlapped with Six3, whereas Lhx1 expression was reduced to a central portion of the SCN (Fig. 4). Six3 expression, however, also marked cells more dorsal and lateral to the SCN (data not shown). Postnatally, these dorsal populations retained their Six3 expression and comprised the anterior paraventricular nucleus (aPV) and subparaventricular regions of the hypothalamus (data not shown). By P0, Six3 and ROR α expression completely overlapped cell density esti-

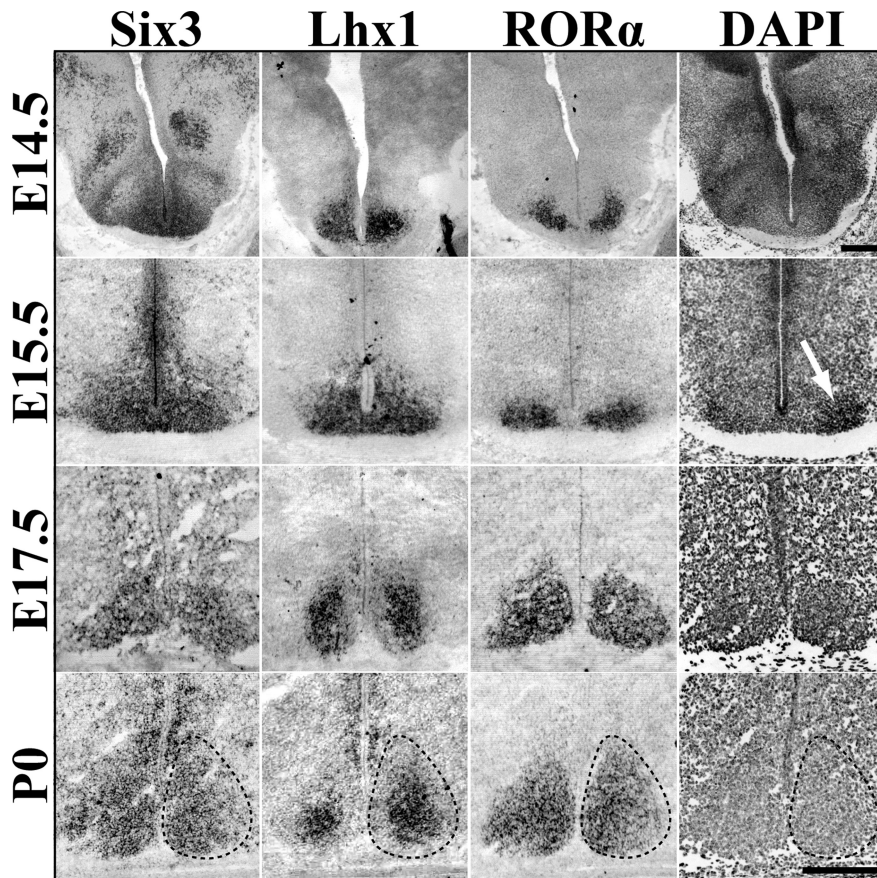


Figure 4. Dynamic transcription factor expression during SCN prenatal maturation. Representative ISH images of *Six3*, *Lhx1*, and *RORα* mRNA expression in the embryonic mouse SCN (E14.5 to P0), showing spatial specificity and prenatal maturation of transcription factor expression. Dotted lines indicate margins of SCN estimated from the cell density marker DAPI (example shown in rightmost panels). White arrow at E15.5 indicates onset of visible clustering of cells. Note change in *Lhx1* pattern from global to centrally restricted between E15.5 and P0. E17.5 *Lhx1* image was obtained from tissue cut at a different angle and thus the appearance of larger SCN compared with *Six3* and *RORα*. Scale bar, 250 μ m.

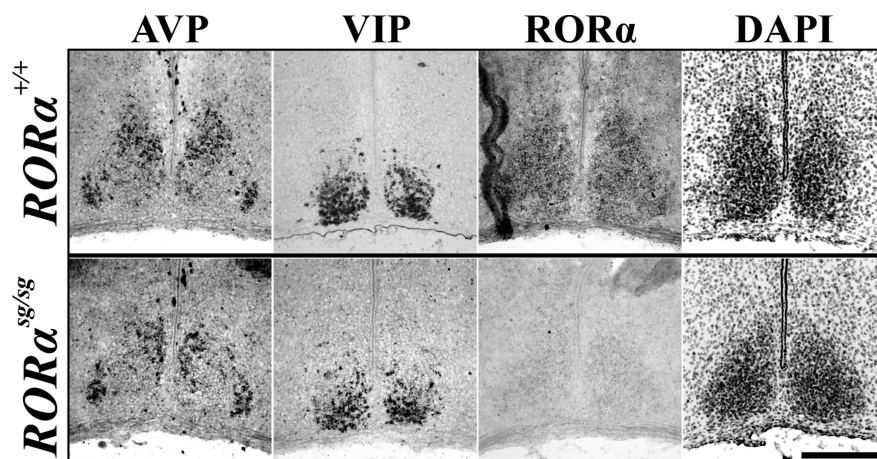


Figure 5. *RORα* is not necessary for normal SCN peptide localization. Representative ISH images of AVP and VIP mRNA expression, markers of SCN subdivisions, as well as transcription factor *RORα* and inverted DAPI nuclear stain in P15 *RORα* mutant (*sg/sg*, bottom) and wild-type littermate (+/+, top) mice. Note that segregation of peptidergic expression patterns was similar to littermate control. Scale bar, 250 μ m.

mates of SCN margins, whereas *Lhx1* continued to remain centralized (Fig. 4).

These data suggest that TFs can be used to identify the prenatal SCN before the onset of peptide markers, dense clustering of

cells, and onset of rhythmic activity. The data further suggest that the onset of *RORα* expression occurs after SCN neurogenesis is complete, consistent with literature describing the delayed onset of *RORα* expression within other brain regions (Nakagawa and O’Leary, 2003). In addition, these data indicate that the SCN is not fully specified by the end of neurogenesis and that SCN maturation occurs in stages both prenatally and postnatally.

***RORα* does not specify postnatal SCN structure**

In a molecular model of the circadian clock, *RORα* positively regulates *Bmal1*, thereby aiding in the stability of the core rhythm (Sato et al., 2004). Knock-outs of *RORα* show slight behavioral abnormalities in circadian rhythms, leading to a shortened circadian period (Sato et al., 2004). To date, it is unknown whether the change in circadian period is attributable solely to the interaction of *RORα* with *Bmal1* or whether there is an anatomical change within the SCN. Therefore, we investigated the role of *RORα* in the development of the SCN by comparing the localization of SCN markers in wild-type mice with *staggerer* mice (*RORα^{sg/sg}*) containing a deletion of the fifth exon in the *RORα* gene that causes formation of a premature stop codon and a functionally null protein (Hamilton et al., 1996).

We found that the expression of SCN markers in *RORα^{sg/sg}* mice were normal. The spatial mRNA expression patterns of AVP and VIP within the SCN as well as that of SCN boundary marker *Otp* in the hypothalamus (data not shown) remained unaltered in all P15 *sg/sg* mutants (four of four) compared with wild-type littermates (Fig. 5). Thus, although we found that the postnatal expression of *RORα* anatomically distinguished the core of the SCN from its shell, *RORα* expression was not necessary for the differentiation, presence, or segregation of peptidergic cells within the SCN.

Identification of the SCN neuroepithelium

The ability of *Six3* and *Lhx1* to delineate the prenatal, postmitotic SCN suggested that they might also precisely define the suprachiasmatic neuroepithelium within the early telencephalon/diencephalon boundary zone beyond previous anatomical estimates (Altman and Bayer, 1986). Therefore, we assessed the expression of these genes just before neurogenesis, at E10.5 and during neurogenesis (E11.5–E13.5) in mice. Specifically, we examined the region from the rostral eye fields extending caudally to the developing pituitary to identify the exact spatial origin of the SCN.

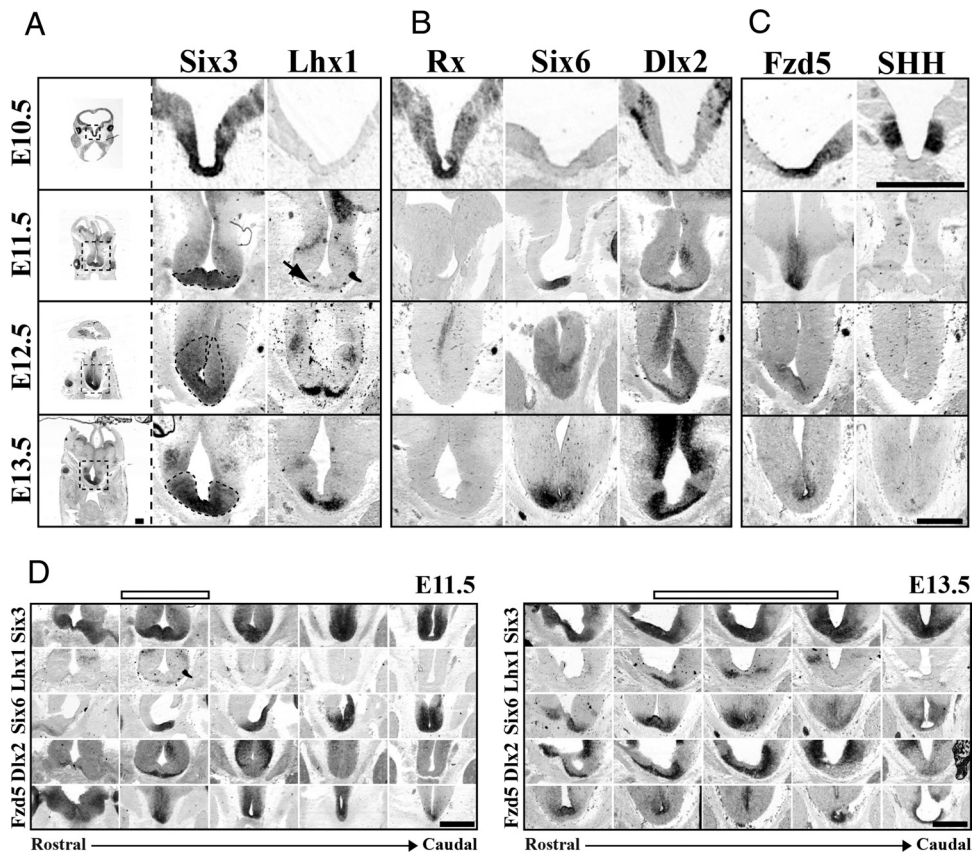


Figure 6. SCN is derived from a distinct progenitor domain. Representative images of Six3 and Lhx1 (**A**), Rx, Six6, and Dlx2 (**B**), and Fzd5 and SHH (**C**) mRNA expression (columns) in SCN progenitor domain from E10.5 to E13.5 (rows). Leftmost column shows entire coronal section, with dotted box indicating approximate region of enlarged images in corresponding row to the right. Dotted black lines in enlarged Six3 images indicate hypothesized early SCN margins outlined by Six3 expression from E11.5 to E13.5. Note at E10.5 that the region that will give rise to SCN neurons expresses a combination of Six3, Rx, and Fzd5. By E11.5, Rx expression is lost and Six6 expression begins. Also note the E11.5 onset of Lhx1 (black arrow) and Dlx2 on the most ventral aspects of tissue beneath the third ventricle. Scale bar, 500 μ m. **D**, Rostrocaudal location of SCN progenitor domain. ISH images depict patterns of Six3, Lhx1, Six6, Dlx2, and Fzd5 mRNA expression (rows) in transverse plane from rostral eye fields to caudal developing pituitary at E11.5 and E13.5. Open bars above indicate the developing SCN within the rostral-caudal axis. Note that, although expression of Six3 and Six6 is consistent along the ventral surface from rostral eye fields to the early pituitary, Lhx1 and Dlx2 mark discrete zones within this region at both E11.5 and E13.5. Please note in **D** that E11.5 depicts same transverse images as shown in **A–C**, and E13.5 shows the same Six6 and Fzd5 images shown in **A–C** now within the rostral-caudal context of the developing brain. Scale bar, 500 μ m.

At E10.5, Six3 marked a broad dorsoventral and rostrocaudal region of the developing floor plate (Fig. 6A and data not shown), consistent with previous reports (Oliver et al., 1995). At E11.5, Six3 was restricted to more ventral regions of the early telencephalon/diencephalon but remained expressed within the developing hypothalamus and pituitary (Fig. 6A, D). Lhx1 expression began at E11.5 in a very small band of cells at the ventral edge of the Six3-expressing zone (Fig. 6A, arrow), allowing for the identification of the early SCN within the telencephalon/diencephalon boundary region (Fig. 6D). Both a dorsal and rostrocaudal expansion in the number of cells expressing Lhx1 occurred from E12.5 to E13.5, whereas Six3 expression became more restricted but still surrounded the base of the third ventricle (Fig. 6A). The coincident onset of and increase in Lhx1 expression during the period of SCN neurogenesis suggests that Lhx1 is one of the first TFs to be expressed in fully postmitotic SCN neurons.

Relationship of early SCN to developmental transcription factors

Identification of the suprachiasmatic neuroepithelium suggested that the SCN was derived from a highly discrete region of the developing neural plate. Our initial reanalysis of brain TF expression identified 58 TFs with discrete expression in the area of the E13.5 anterior hypothalamus (supplemental Table 2, available at

www.jneurosci.org as supplemental material). We confirmed the embryonic hypothalamic expression of at least 66% of these 58 TFs using data from Genepaint and Eurexpress expression databases (Visel et al., 2004; Diez-Roux et al., 2011). These TFs included early forebrain genes Six6 and Dlx2. In addition, reduction in the amount of Rx (Rax) present in early neuroepithelium has been shown to disrupt SCN location and cell number by an unknown mechanism (Silver, 1977; Tucker et al., 2001). To clarify the origins of the SCN within the developing telencephalon/diencephalon, we compared the expression of Six3 and Lhx1 with that of early forebrain genes Six6, Dlx2, and Rx.

At E10.5, Rx expression overlapped that of Six3 in the early ventral floor plate (Fig. 6B) from the early eye fields through the early developing pituitary (data not shown), consistent with a previous analysis (Bailey et al., 2004). At E11.5, a rostrocaudal gap was evident in Rx expression. Rx persisted within the eye itself and in ventral floor plate just rostral to the developing pituitary (data not shown). Six6 expression was first seen at E11.5 in a ventrally restricted but overlapping pattern with Six3 within the entire rostrocaudal region examined, including within this Rx-negative gap (Fig. 6B, D). Dlx2 expression was first seen in a small band of ventral cells within the Six3/Six6-expressing and Rx-negative domain (Fig. 6B, D). Lhx1 was expressed in a subset of the Dlx2-expressing domain (Fig. 6D).

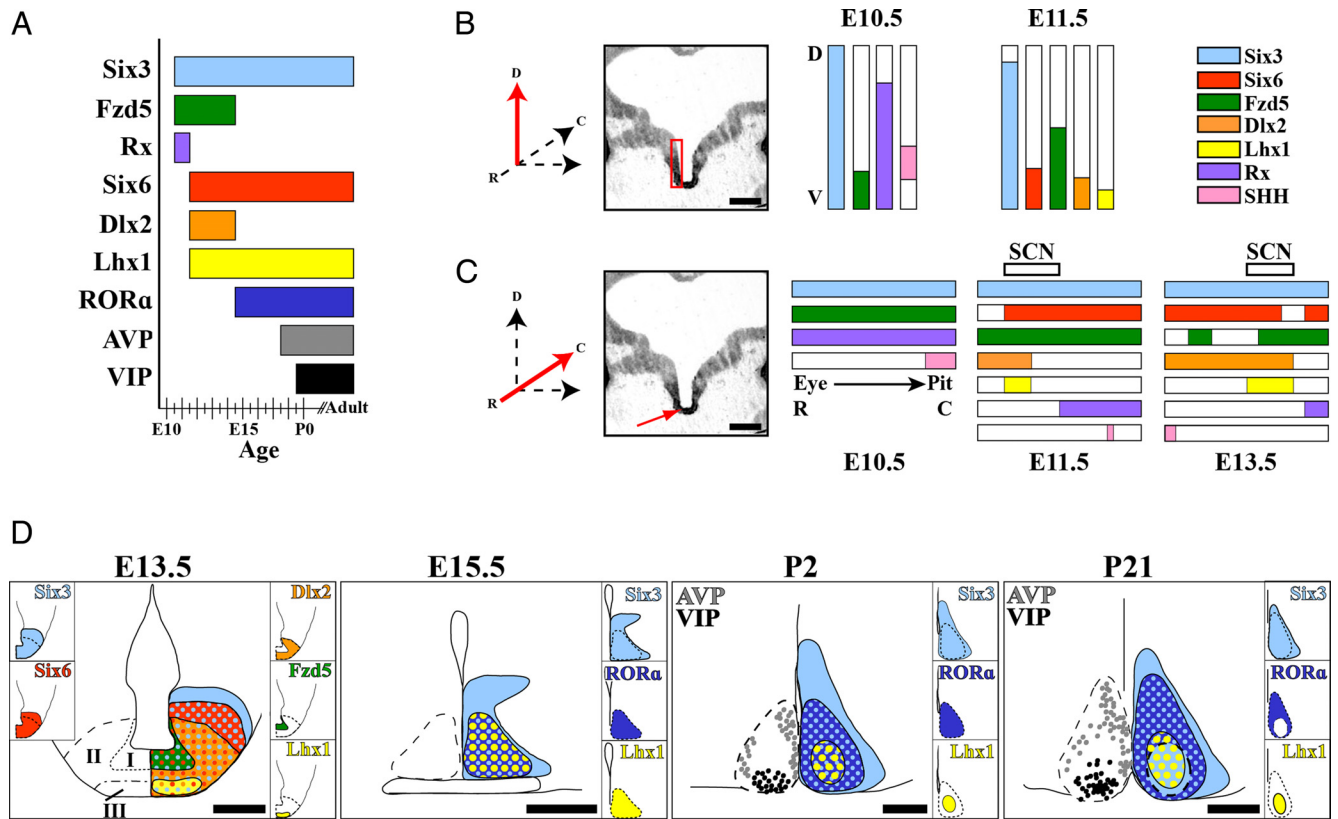


Figure 7. Summary of SCN development. **A**, Bar graph indicates onset and duration of transcription factor and peptide mRNA expression from E10.5 to adult in the mouse SCN. **B**, **C**, Diagrams summarize dorsoventral and rostrocaudal extent of developmental gene expression within the early telencephalon/diencephalon containing the hypothesized early SCN. Red arrows on orientation axes show directionality of diagram in relation to indicated red regions. **B**, Summary of dorsoventral (D–V) extent of mRNA expression (colored bars) within the same region of anterior hypothalamus, indicated by red box in image showing Six3 expression, at E10.5 (left) and E11.5 (middle). Scale bar, 250 μ m. Colors correspond to genes in legend (right). Note the absence of SHH but presence of Fzd5 within the ventral most domains, i.e., putative early SCN. **C**, Summary of rostrocaudal (R–C) extent of mRNA expression within the midline (red arrow) of the ventral developing telencephalon/diencephalon at E10.5 (left), E11.5 (middle), and E13.5 (right). Scale bar, 250 μ m. The most rostral point is marked by the retina (Eye) and caudal by the developing pituitary (Pit). Colors correspond to genes in legend in **B**. Open top bars at E11.5 and E13.5 indicate rostrocaudal extent of putative SCN progenitor domain within larger developing brain. **D**, Diagrams of SCN transcription factor localization indicate expression during neurogenesis (E13.5), within the prenatal postmitotic (E15.5), early postnatal (P2), and functionally mature SCN (P21). Dotted lines on left indicate the margins of the SCN. Insets indicate expression pattern of individual transcription factors. Colored dots indicate regions of coexpression. Note hypothesized zones of developing SCN at E13.5: I, premitotic/proliferative; II, late-mitotic/early-postmitotic; III, postmitotic. Also note the changes in localization of both Lhx1 and ROR α over time. Scale bars, 250 μ m.

By E12.5 and E13.5 as Six3/Six6 expression continued, the Dlx2-expressing region expanded rostrocaudally (Fig. 6D) and dorsally both beneath as well as along the lateral aspects of the third ventricle stratifying the region surrounding the base of the early third ventricle into three subregions (Fig. 6B). The ventral-most portions of Dlx2 expression overlapped with Lhx1 expression. The mid region retained only Six3/Six6/Dlx2, and the most dorsal aspects lacked Dlx2 (Fig. 6B). The overlap between Dlx2 and Lhx1 in the most ventral regions suggested that this combination marked newly postmitotic neurons of the SCN, consistent with findings in embryonic rat. The presence of an Lhx1-negative, Dlx2-positive region suggested that Dlx2 might be expressed during the transition period of cells during the last stages of mitosis to early stages of differentiation similar to its expression within the ventral thalamus (Andrews et al., 2003).

The absence of a single TF specific to the SCN suggested that SCN development might also rely on extrinsic morphogens and other signaling peptides to provide key regionalization signals (Burns et al., 2008; Liu et al., 2010). The expression of Six3, Six6, and Dlx2 in the presumptive suprachiasmatic neuroepithelium can be used to identify the immature SCN within the complex early signaling environment of the developing telencephalon/diencephalon. We found that, at E10.5, the Six3/Rx region also expressed Frizzled5 (Fzd5), a Wnt receptor, in a region surround-

ing the base of the third ventricle (Fig. 6C). This broad dorsal to ventral Six3/Rx/Fzd5 region was further demarcated by the expression of SHH. Six3/Rx/Fzd5 regions lateral to the ventricle expressed SHH, although it remained absent from the most ventral medial region at the base of the third ventricle (Fig. 6C). Unlike Rx, Fzd5 expression continued to overlap with Six3/Six6 at E11.5, further defining the SCN proliferative zone in both the dorsoventral (Fig. 6C) and rostrocaudal (Fig. 6D) dimensions. Fzd5 expression at E12.5 and E13.5 was still present beneath the third ventricle but was restricted to the dorsal region overlapping with the Six3/Six6 region only (Fig. 6C,D). Fzd5 expression did not overlap with the region containing postmitotic Lhx1 cells and Dlx2, which suggested the expression of Fzd5 likely marked regions of active proliferation. We found that, at E12.5, the most ventral Dlx2- and Lhx1-expressing regions overlapped with TuJ1, a marker of postmitotic neurons, whereas the more dorsal Dlx2-negative region expressed Ki67, a marker of proliferating cells (data not shown). Throughout this period of neurogenesis, the early SCN was completely devoid of bone morphogenetic protein 7 and lacked the common SHH receptor Patched1 (data not shown).

Our data suggest that the SCN is derived from a proliferative region expressing Six3, Six6, Fzd5, and transient Rx. Furthermore, early postmitotic cells of the SCN express Six3, Six6, Lhx1

and transiently express *Dlx2*. Figure 7 shows a summary of the relative patterns of developmental-gene expression during SCN development.

Six3 is required for SCN formation

In the present study, we found that *Six3* was continually expressed in the SCN and its progenitors from before neurogenesis to adult when expression was diurnal. This suggested that *Six3* has an important role in the SCN and its development. Therefore, we tested whether *Six3* is necessary for the formation and maturation of the SCN using a *Nestin-cre* transgenic to eliminate *Six3* (*Six3^{flox/flox}*) (Liu et al., 2006) specifically in progenitors within the neural plate before specification.

We analyzed the anatomy and expression of *Six3* mRNA in *Six3^{flox/flox}/Nestin-cre⁺* mutants ($n = 5$), *Six3^{flox/flox}/Nestin-cre⁻* ($n = 3$), and *Six3^{flox/+}/Nestin-cre⁺* ($n = 2$) littermate controls at E18.5–E19.5. We found the *Nestin-cre* transgenic exhibited a range of *Six3* loss in the developing mouse brain, from complete loss to no loss (Fig. 8), likely as a result of variation in the onset and duration of cre-recombinase expression. *Six3^{flox/flox}/Nestin-cre⁺* mice with complete *Six3* deletion ($n = 4$ of 5) showed holoprosencephaly and a noticeable enlargement and/or merger of the lateral and third ventricles not seen in littermate controls (data not shown). Furthermore, *Six3* ablations lead to loss of the pituitary (Fig. 8B). In contrast, retina and optic nerves remained intact (data not shown).

Six3^{flox/flox}/Nestin-cre⁺ E18.5 and E19.5 animals with complete bilateral loss of *Six3* expression showed no expression of *ROR α* or *AVP* in the ventral anterior hypothalamus (Fig. 8A, asterisks). Importantly, loss of *Six3* did not lead to the displacement of the SCN because no ectopic nuclei were present along the ventral surface either rostral or caudal to the optic chiasm, as assessed through a DAPI nuclear stain (data not shown). Loss of *ROR α* and *AVP* was SCN specific because the *aPV* and *SON* of *Six3^{flox/flox}/Nestin-cre⁺* mice retained *AVP* expression and the dorsal lateral geniculate (*dLG*) and ventroposterior medial (*VPM*) nuclei in both E18.5 and E19.5 mutants continued to express *ROR α* (Fig. 8B). In addition, in all mutants tested, the zona incerta (*ZI*), which normally expresses *Six3*, was unaffected (Fig. 8B).

To assess whether absence of SCN at E18.5 and E19.5 was attributable to dissolution of the nucleus after cells became postmitotic or the complete disruption of neurogenesis, we analyzed *Six3^{flox/flox}/Nestin-cre⁺* mice at E15.5 ($n = 5$) for early postmitotic SCN markers *Lhx1* and *ROR α* as well as for dense cell clusters. We found that loss of *Six3* leads to absence of both early postmitotic genes, consistent with a disruption in specification. E15.5 *Six3^{flox/flox}/Nestin-cre⁺* mice displaying partial *Six3* de-

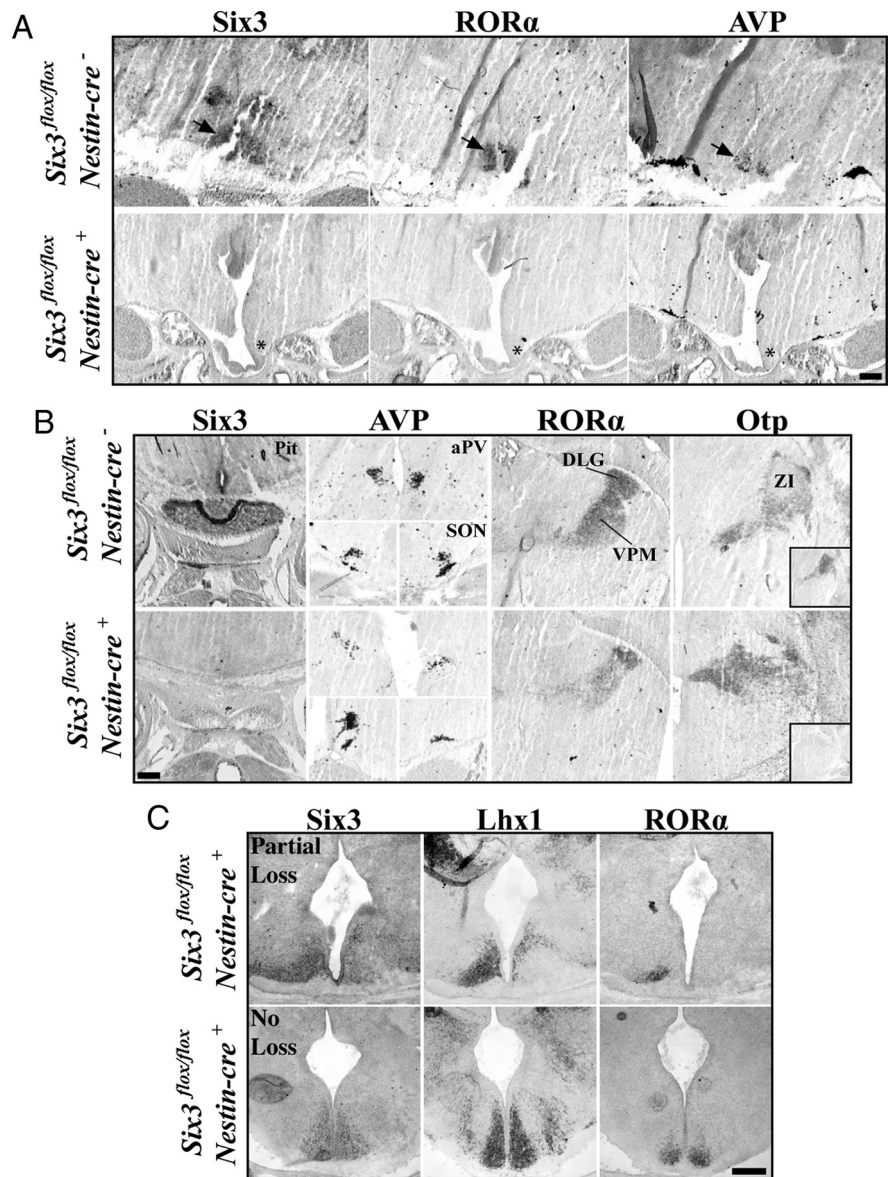


Figure 8. *Six3* is necessary for SCN formation. Expression of *Six3*, *ROR α* , *AVP*, and *Otp* in *Six3^{flox/flox}/Nestin-cre⁺* compared with *Six3^{flox/flox}/Nestin-cre⁻* littermate controls. **A**, Expression of *Six3*, *ROR α* , and *AVP* in SCN (arrows in top) are completely eliminated in conditional *Six3* knock-outs, shown at E18.5 (asterisks). **B**, Marker analysis of the pituitary, hypothalamic (*aPV* and *SON*), and thalamic (*dLG*, *VPM*, and *ZI*) nuclei show that complete disruption of *Six3* leads to loss of the pituitary, whereas nuclei surrounding the SCN remain intact in *Six3^{flox/flox}/Nestin-cre⁺* experimental animals. Bottom right inset in *Otp* column represents adjacent section for *Six3* expression. **C**, Variability in the targeted elimination of floxed *Six3* in E15.5 mice. Expression of *Six3*, *Lhx1*, and *ROR α* in *Six3^{flox/flox}/Nestin-cre⁺* mice that showed either partial loss (top) or no loss (bottom) of *Six3*. Top, Unilateral *Six3* deletions lead to unilateral loss of *ROR α* and *Lhx1*. Bottom, Mice that retain *Six3* expression still express both *Lhx1* and *ROR α* . Scale bars, 250 μ m.

letion showed a loss of *ROR α* and *Lhx1* only in regions lacking *Six3* (Fig. 8C). *Six3^{flox/flox}/Nestin-cre⁺* mice that retained expression of *Six3* showed prominent expression of both *ROR α* and *Lhx1* (Fig. 8C). Together, these results show the necessity of a transcription factor for proper SCN formation and development.

Discussion

Recently, several groups have directly analyzed the expression of large numbers of genes (>1000) within the developing and/or mature hypothalamus by *in situ* hybridization (Gray et al., 2004; Lein et al., 2007; Shimogori et al., 2010), looking for patterns of localization capable of defining small populations of neurons that may correspond to unique functional classes. All of these analyses

have shown broad genetic distinctions between hypothalamic nuclei. Other studies have also provided detailed analyses of the roles of several developmental genes in hypothalamic and pituitary ontogeny (Sládek et al., 2004; Davis et al., 2010). However, large anatomical screens, because of their breadth, often lack the spatial detail necessary for a more thorough analysis of specific brain regions. Here we provide a detailed analysis of developmental gene expression in the SCN, describe a specific gene cascade from which SCN neurons are derived, and identify a highly conserved transcription factor necessary for SCN formation.

Altman and Bayer (1986) hypothesized that the early suprachiasmatic neuroepithelium formed from a region just caudal to the optic recess. Our data narrow this description showing that the SCN is derived from a distinct region of neuroepithelium expressing a combination of the developmental genes *Six3*, *Six6*, *Fzd5*, and transient *Rx*, allowing us to pinpoint the suprachiasmatic neuroepithelium within the broader developing telencephalon/diencephalon (for summary, refer to Fig. 7A–C).

ZRDCT anophthalmic mice have decreased expression of *Rx* protein, and 30% of mutant mice show defects in SCN anatomy and function (Silver, 1977; Tucker et al., 2001). The defects vary widely with respect to disruptions in location and/or cell number, but dense clusters of cells are always seen, indicating formation of nuclei. These data along with our observation of transient *Rx* before SCN neurogenesis (Figs. 6D, 7A, B) suggests that *Rx* is not necessary for specification of SCN neurons but that loss may disrupt early progenitor domains (Liu et al., 2010).

Altman and Bayer further hypothesized that the suprachiasmatic neuroepithelium was divided into two regions, dorsal and ventral, from which the corresponding dorsomedial/shell and ventrolateral/core were derived. Our data are not consistent with the presence of two genetically distinct regions contained within the suprachiasmatic neuroepithelium. We suggest that the proposed regions outlined by Altman and Bayer are a consequence of temporal proliferation or differential response to non-cell-autonomous cues, such as morphogens or other early-signaling gradients, rather than distinct genetic differences in progenitor type.

We demonstrated that the onset of *Dlx2* and *Lhx1* expression at E11.5 is coincident with the documented onset of SCN neurogenesis (Kabrita and Davis, 2008). This suggests that *Dlx2* and *Lhx1* mark neurons as they exit the proliferative zone that will form the SCN. At E12.5/E13.5, the region that gives rise to the SCN contains both proliferating and postmitotic neurons. We demonstrate three zones of overlapping gene expression in the SCN at this age and propose that these distinct TF-expressing zones represent phases of SCN neuronal development: (I) dorsal *Six3/Six6/Fzd5*: premitotic/proliferative, (II) mid *Six3/Six6/Dlx2*: late mitotic/early postmitotic, and (III) ventral *Six3/Six6/Dlx2/Lhx1*: postmitotic (for summary, refer to Fig. 7D). Our proposed late-mitotic/early-postmitotic *Six3/Six6/Dlx2* zone is similar to the region of migratory “late-forming neurons” outlined by Altman and Bayer (1986). In addition, our data support the notion of a ventrodorsal gradient in SCN neurogenesis (Kabrita and Davis, 2008) because the expression of *Lhx1*, a postmitotic marker, was first seen at the most ventral aspects of the early SCN and progressively expanded dorsally as neurogenesis proceeded.

We found that TF expression within the postmitotic SCN was not static but rather showed specific temporal and spatial changes during both prenatal development and postnatal maturation. This was indicated by dynamic expression of *Lhx1* prenatally and *ROR α* postnatally, in which expression of both genes spanned the

entire SCN at early stages but became differentially regionalized through development. We propose that changes in localization of gene expression may indicate underlying structural or functional maturation of the SCN. This is supported by the central regionalization of *Lhx1* at approximately E17.5, coincident with the perinatal onset of peptide expression. Furthermore, dorsomedial/shell localization of *ROR α* , evident by P15, is concurrent with the proposed termination of retinal fibers into the SCN (Takatsuji et al., 1995) and onset of photoperiodic entrainment of clock genes (Kováčiková et al., 2005). It would be reasonable to presume that neurons or areas within the SCN may undergo additional specification or specialization as external innervation into the nucleus, such as postnatal retinal innervation (McNeill et al., 2011) and synaptogenesis progress. The identification of TFs with discrete localization within the SCN may reflect additional subdivisions and/or subsets of neurons with distinct functions.

Regional specificity of gene expression, similar to *ROR α* , has also been reported for *Period1* and *Period2*, with expression mainly limited to the dorsomedial/shell during subjective day in rat and hamster (Hamada et al., 2001). Interestingly, some evidence suggests functional division of the SCN into regions of endogenous rhythmicity (dorsomedial/shell) or induced rhythms (ventrolateral/core) (Sumova et al., 1998; Guido et al., 1999; Hamada et al., 2001; Yan and Okamura, 2002; Hamada et al., 2004; Ramanathan et al., 2006). Given the role of *ROR α* in stability aspects of the clock through regulation of *Bmal1*, it may be advantageous for the clock to lack stabilizing elements in an inducible region. Furthermore, the pan-SCN expression of *Six3* in phase with *Per1* may suggest a role for *Six3* in maintenance of clock gene expression or underlying circadian rhythmicity.

ROR α is one member of related orphan nuclear receptors that serve not only in the differentiation and development of multiple tissue types, including the cerebellum and thalamus (Becker-André et al., 1993; Carlberg et al., 1994; Hamilton et al., 1996; Matysiak-Scholze and Nehls, 1997; Nakagawa et al., 1997, 1998; Dussault et al., 1998; Vogel et al., 2000; Sato et al., 2004), but also in circadian function (Schaeren-Wiemers et al., 1997; Sumi et al., 2002). We found that loss of *ROR α* had no effect on either the presence or location of AVP and VIP neurons, indicating that it was not necessary for peptide expression or development of peptidergic segregation.

The role of *Six3* in eye development is highly conserved across vertebrates and in *Drosophila* (Simeone et al., 1994; Seo et al., 1999). *Six3* has been proposed to play a role in proliferation of retinal precursors (Del Bene et al., 2004), lens formation (Liu et al., 2006), neuroretina specification (Liu et al., 2010), regulation of rhodopsin expression (Manavathi et al., 2007), and early anterior patterning (Kobayashi et al., 2002; Lagutin et al., 2003; Gestri et al., 2005; Lavado et al., 2008). *Six3* has been shown widely expressed throughout the anterior brain at early stages, narrowing expression over time and remaining transient for many neuronal populations. We found that *Six3* was expressed within the suprachiasmatic neuroepithelium before the onset of neurogenesis into adulthood, suggesting an important role for *Six3* in the SCN. Moreover, expression of *Six3* in adult mice displayed diurnal variation. Although our data cannot discriminate between whether this variation is a response to rhythmic photic input or reveals an underlying circadian rhythm in *Six3* expression, they suggest a link between *Six3* and day/night changes.

We demonstrated that *Six3* is crucial to the development of the SCN. Using a Nestin-cre transgene to limit loss of *Six3* to neural progenitors, we tested the role of *Six3* in SCN specification. We showed that the loss of *Six3* during development leads to

elimination of ROR α and AVP expression as well as the absence of dense cell clustering perinatally, consistent with the elimination of the SCN. In contrast, extra-SCN structures, including hypothalamic SON and aPV, as well as thalamic VPM and Six3-expressing ZI, were still present. This suggests that Six3 may not be acting solely in global early patterning but may have an additional role in specification of the SCN. We propose that the absence of gene expression is a consequence of the failure to form the SCN initially and not a role for Six3 in the maintenance of SCN cell survival, because SCN-specific Lhx1 expression and cell density were also absent at E15.5. These data suggest a necessary role for Six3 in SCN development and that Lhx1 and ROR α represent lineage-specific SCN genes.

Overall, these data demonstrate that TF expression can provide insight into both the specification and development of the SCN. We find that TF expression can describe the early suprachiasmatic neuroepithelium and demonstrate that the SCN undergoes maturation both prenatally and postnatally. Maturation is evident by the changes in localization of gene expression within the SCN over time. It is currently unclear whether the specific loss of gene expression in distinct areas of the SCN is necessary for proper circadian function. However, we find that disruption of at least one TF, Six3, can have large-scale effects on SCN development. Profiling of genes with SCN-specific expression for spatial and temporal onset allows researchers, for the first time, to begin to understand the roles of these genes in the anatomical and functional development of this complex nucleus.

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