DNA-mediated cooperativity facilitates the co-selection of cryptic enhancer sequences by SOX2 and PAX6 transcription factors

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ABSTRACT

Sox2 and Pax6 are transcription factors that direct cell fate decision during neurogenesis, yet the mechanism behind how they cooperate on enhancer DNA elements and regulate gene expression is unclear. By systematically interrogating Sox2 and Pax6 interaction on minimal enhancer elements, we found that cooperative DNA recognition relies on combinatorial nucleotide switches and precisely spaced, but cryptic composite DNA motifs. Surprisingly, all tested Sox and Pax paralogs have the capacity to cooperate on such enhancer elements. NMR and molecular modeling reveal very few direct protein–protein interactions between Sox2 and Pax6, suggesting that cooperative binding is mediated by allosteric interactions propagating through DNA structure. Furthermore, we detected and validated several novel sites in the human genome targeted cooperatively by Sox2 and Pax6. Collectively, we demonstrate that Sox–Pax partnerships have the potential to substantially alter DNA target specificities and likely enable the pleiotropic and context-specific action of these cell-lineage specifiers.

INTRODUCTION

A primary goal of genomics and associated high-throughput technologies is to decipher the cis-regulatory code that determines when, where and how sets of genes are expressed. This regulatory code directs biological processes such as embryonic development and differentiation and is the cause for phenotypic variance and evolutionary innovation. Likewise, aberrations in this code can lead to developmental abnormalities and/or disease progression. The location of cis-regulatory ‘enhancer’ DNA sequences that govern cellular identities is now known in many cell types and lists of transcription factors (TFs) that bind such enhancers and associated epigenetic signatures are expanding rapidly (1,2). Enhancers are often composed of clustered DNA elements with affinity for a set of TFs. The architecture of an enhancer, that is, the sequence of the DNA elements, their number, relative orientation and spatial arrangement is thought to determine enhancer activity (3,4). However, the process of how TFs in a combinatorial fashion select enhancers to regulate gene expression remains only superficially understood (5,6). TF interactions are dynamic, transient and often depend on cellular and genomic contexts (7). Both, complementary protein interaction surfaces and the sequence of composite DNA motifs mediate TF associations (8). So far, the structural architectures of very few TF–TF interactions have been described at atomic resolution. Some well-studied instances of heterodimeric TF partnerships include PPAR-γ–RXR-α (9), Pax5-Ets (10), AR–FoxA1 (11), Sox2-Oct4

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accommodation of diverse sets of binding sites by mechanisms (31). This modular domain structure in Pax6 enables the additional homeodomain C-terminal of the paired domain additionally latched into the minor groove via an extended main (DBD) consisting of N-terminal PAI and C-terminal structures in the legs of flies (27). The Pax gene family initiates eye development as remarkably testified when it eventually, δEF3 was found to be Pax6 (24). The chicken DC5 sequence can also effectively drive reporter gene expression in the Drosophila eye and cooperatively recruit homologous fly TFs, suggesting a deep phylogenetic conservation of the Sox2/Pax6 partnership and the regulatory circuit of lens specification (25). Pax6 had been known for some time to be a key regulator of many aspects of development, especially of neurogenesis (26). In addition, Pax6 initiates eye development as remarkably testified when it could be shown that its ectopic expression can induce eye structures in the legs of flies (27). The Pax gene family consists of nine members in mammals, all of which are key developmental TFs (28). Pax proteins are composed of a bipartite 128 amino acid paired (PRD) DNA-binding domain (DBD) consisting of N-terminal PAI and C-terminal RED subdomains binding the major groove of DNA and additionally latched into the minor groove via an extended linker (29,30). Some Pax TFs, including Pax6, contain an additional homeodomain C-terminal of the paired domain (31). This modular domain structure in Pax6 enables the accommodation of diverse sets of binding sites by mechanisms that can include an intricate interplay of the subdomains and alternate utilization of subdomains for the recognition of specific target sites (32–35).

In chicken embryos, Pax6 is initially broadly expressed in the head ectoderm and becomes subsequently restricted to the lens placode (24). Inductive signals derived from the optic vesicle induce Sox2/Sox3 proteins in the lens placode cells leading to the activation of δ-crystallin expression synergistically with Pax6 (24). A similar cooperation between Sox2 and Pax6 has been implicated in autoregulation of Sox2 by binding to an enhancer element termed N3, which is well conserved across vertebrates (36). Both, the N3 and DC5 enhancers are highly compact and possess canonical sox motifs, but highly degenerate pax motifs (36). Yet, this degeneracy is necessary for cooperative binding of Sox2 and Pax6 and for effective activation of target gene expression (24,36). Mutations of sox or pax half-sites as well as altered motif configurations abolished δ-crystallin expression (24). In mouse and humans, both, Sox2 and Pax6, have been demonstrated to be key regulators of neurogenesis (26,37–38). Sox2 and Pax6 are co-expressed in neural stem cells (NSCs), the developing eye and several other neuroectoderm-derived cells, providing ample opportunities for Sox2 and Pax6 to cooperatively bind to target DNA sites in the human genome, reminiscent of the chicken DC5 and N3 enhancers (Figure 1A). Additional Sox–Pax partnerships including Sox10 and Pax3 have also been described (39) and the co-expression of Sox and Pax family members is observed in a plentitude of cell types (Figure 1B). We therefore decided to interrogate the biochemical basis for the cooperation of Sox and Pax TFs to (i) uncover the sequence determinants for the cooperative interactions, (ii) understand the biochemical and structural basis for the cooperation of Sox and Pax TFs to (iii) ask if there is a Sox–Pax partner code that determines target gene selection, (iv) assess whether mutations within Sox2 and Pax6 causative of congenital human eye disorders alter Sox2–Pax6 cooperativity and (v) identify novel Sox/Pax target site in the human genome that supports cooperative interactions.

MATERIALS AND METHODS

Cloning, protein production and site-directed mutagenesis

Proteins encoding 79 amino acid HMG domains of mouse Sox2, Sox4, Sox5, Sox15, Sox17 and an extended 109 amino acid acid Sox2 were used with their amino acid sequences, cloning and purification procedures as published elsewhere (16). The extended 109 amino acid Sox2-HMG construct was chosen for TROSY studies to detect potential interactions in the extended C-terminal tail. This longer Sox2 construct was N153 isotope labeled, by growing Escherichia coli cells in M9 minimal media containing isotope-labeled ammonium chloride (Cambridge Isotope Laboratories) at 18°C and purified using chromatographic procedures as described previously (40). The following are the IMAGE IDs of the Pax cDNA clones used in the study: Pax2 (IMAGE clone ID: 40142579), Pax3 (IMAGE clone ID: 6518115), Pax6 (IMAGE clone ID: 4504106), Pax8 (IMAGE clone ID: 4239835) and Pax9 (IMAGE clone ID: 3707718). The paired domain sequences were GATEWAY BP cloned (Life Technologies) from their respective cDNA clones into the pDONR221 vector using primers with attB1 sites (Supplementary Table S1). The resulting pENTR constructs were verified by sequencing and recombined into the pETG40A and pETG20A destination vectors using the GATEWAY™ technology (Life Technologies). Recombinant Pax proteins were expressed and purified using a previously published protocol (41). Expression vectors with point mutations in Sox2-long (D123G) and Pax6 (R44Q and G36R) were generated using the QuikChange XL mutagenesis kit (Stratagene) according to manufacturer’s instructions (Supplementary Table S1). The incorporation of point mutations was verified by sequencing the expression plasmid.
Figure 1. Sox and Pax TFs are co-expressed in many cell types. (A) A scatterplot to compare SOX2 and PAX6 expression, highlighting co-expression in several human cell lines. (B) A clustered heatmap of 5′ RNA CAGE counts was plotted using gbase with row normalization (44). Red and blue color indicate high and low expression, respectively. Columns are 563 human FANTOM5 cell types. Rows are human SOX and PAX TFs.

DNA elements used

DNA sequences (Supplementary Table S2) labeled with 5′-carboxyfluorescein (FAM) or Cy5 were employed in the study. Single-stranded oligos were purchased from Sigma ProLogi and cognate double-stranded DNA elements were made by annealing the oligos in annealing buffer (20 mM Tris–HCl, pH 8.0; 50 mM MgCl2; 50 mM KCl), heated to 95°C for 5 min and subsequently ramping down at 1°C/min to 4°C in a polymerase chain reaction block. To study the effect of spacing between the sox and pax motifs, the following DC5 sequences with different spacing were used: -1 and -2 with reduced spacing and +1, +2, +3, +4 and +5 with increased spacing between the sox and pax half-sites. For affinity measurements, a previously crystallized DNA element (PDB ID: 6PAX) was used.

Electrophoretic mobility shift assays

Electrophoretic mobility shift assays (EMSAs) were performed with 5′-FAM or 5′-Cy5-labeled cognate double-stranded DNA elements (Supplementary Table S2). The binding buffer contains 10 mM Tris–HCl pH 8.0, 0.1 mg/ml bovine serum albumin, 50 μM ZnCl2, 100 mM KCl, 10% Ultrapure Glycerol, 0.10% IGEPAL CA-630 (octylphenoxy polyethoxyethanol) and 2 mM β-mercaptoethanol. For the differential assembly studies, typical binding reactions contain concentrations 100–300 nM of double-stranded DNA (>> Kd) with varying concentrations of Sox and Pax proteins. Protein concentrations were adjusted such that an equilibrium is reached and the four microstates are visible as distinctive bands on EMSA gels to allow reproducible quantification. All binding reactions were incubated for 1–2 h in the dark. Samples were loaded onto 1X Tris Glycine-PAGE gels (25 mM Tris pH 8.3; 192 mM glycine) and electrophoresed at 200 or 300V for 30–120 min at 4°C. The gel was imaged using a Typhoon 9140 phosphor imaging scanner or a Typhoon FLA-7000 PhosphorImager (FUJIFILM) and the fluorescent intensities of the free and bound DNA were quantified using the ImageQuant TL software. Cooperativity factors were calculated using the formula: $\omega = \frac{f_{\text{dimer}} f_{\text{freeDNA}}}{f_{\text{monomer1}} f_{\text{monomer2}}}$ where $f_{\text{dimer}}$ is the fractional contribution of dimeric protein complexes on DNA (Sox/Pax/DNA), $f_{\text{monomer1}}$ and $f_{\text{monomer2}}$ of monomeric (Sox/DNA or Pax/DNA) protein–DNA complexes and $f_{\text{freeDNA}}$ the fractional contribution of unbound DNA in EMSA gel lanes. The derivation of the formula has been described previously (16,42) and was further discussed in (43).

Homology modeling of Sox2–Pax6

The models of the Sox2–Pax6–DNA ternary complexes were build by superimposing B-DNA fragments corresponding to the DC5, N3 and DC5con sequences on the N-terminus and C-terminus of the structures of the Sox2/DNA and Pax6/DNA complexes, 1GT0 (13) and 6PAX (30), respectively. In addition, the DNA molecules at the C-terminus of the 1GT0 structure and at the N-terminus of the 6PAX structure were superimposed, so as to reflect the spacing between the Sox and Pax half-sites observed in the DC5, N3 and DC5con sequences. The sequence of the new DNA formed by concatenating the four DNA fragments and removing the overlapping bases was mutated to the correct DC5, N3, and DC5con sequences using chimera (https://www.cgl.ucsf.edu/chimera/). The energy of the resulting models of the ternary complexes was minimized in AMBER (www.ambermd.org) using a stepwise procedure in which restraints on the base pair geometries were gradually removed.
Bioinformatics analysis

Human FANTOM5 CAGE data (http://fantom.gsc.riken.jp/) were downloaded and condensed into a single table after averaging some duplicates and only the highest expressed promoter was kept for a given cell type. The clustered heatmap was produced after row normalization using gbase (44). ChIP-Seq peaks for PAX6 in human NSCs (45) or SOX2 neural progenitor cells (NPCs) (46) were intersected. Motifs were created manually taking EMSA data into account and co-bound SOX2/PAX6 sites were searched for motif ‘words’ using gbase (44). To produce genome plots using IGV, reads were re-mapped to the human genome (hg19).

Circular dichroism spectroscopy

The circular dichroism (CD) spectra were recorded on an Chirascan CD Spectrometer (Applied Photophysics Ltd) using a strain-free 10 mm x 1.0 mm rectangular cuvette using the following parameters: bandwidth, 1 nm; spectral range, 230–360 nm; step-size, 1 nm; time-pep-point, 0.5 s. In total, 1.6 μM dsDNA and 5 μM Pax6-PRD were mixed in 50 mM phosphate buffer, pH 8.0. Three repeat measurements were averaged.

Nuclear magnetic resonance experiments, data acquisition and processing

Nuclear magnetic resonance (NMR) experiments were performed using Bruker AVANCE II 600 and 700 MHz NMR spectrometers equipped with a standard TCI (Triple resonance cryoprobe) cryoprobe. The spectra were collected at 298 K. A series of 2D [15N,1H]-TROSY experiments was utilized to monitor the chemical shift perturbations (CSP) of the 15N, 1H spins. The chemical shifts were referenced directly (1H) relative to 4,4-dimethyl-4-silapentane-1-sulfonic acid (DSS). The spectral analysis was performed using CARA (47). For NMR experiments, a 31bp DC5 element was used with the following forward and complementary reverse strand sequence (DC5 fasta): 5′-TTCCATTGTGTTGCTCACCCTAGGATCC-3′; DC5 reverse: 5′-GGATCCATGTTGAGCGAAACACACATGAA-3′. The 31bp DC5 DNA element was dissolved in the following NMR buffer (50 mM K2HPO4/KH2PO4, 100 mM NaCl buffer, pH 7.0). The buffer components are kept consistent for all NMR experiments unless otherwise stated. The pH of the solutions was adjusted to 7.0 using NaOH. The final concentration of the DC5 stock solution was determined using NanoDrop at 260 nm. We have previously recorded 3D spectra of 1H, 13C and 15N labeled Sox2 (109 amino acid long) and assigned residues to the majority of resonances (40). The N15 labeled Sox2-HMG was added carefully to the DNA element (31-mer) in a stepwise manner to achieve final molar ratios of Sox2/DNA of 0.25, 0.5, 0.75 and 1.0, with the final DNA concentration of 0.2 mM. The 1D spectrum of the 31mer-DNA alone was used as a reference to observe perturbations inflicted by the binding of proteins to the DNA. A 2D [15N, 1H]-TROSY spectrum was collected at each titration point. The weighted CSP for backbone 15N and 1H N resonances were calculated by the equation (48): \[ \Delta \delta = |(\Delta \delta_{15N})^2 + (0.1 \Delta \delta_{1H})^2|^{0.5}. \]

RESULTS

Pax motif degeneracy is necessary for the cooperativity of Sox2 and PAX6

Sox2 and PAX6 are co-expressed in a number of human cell types, namely fetal eyes, NSCs, cerebellum and hippocampus (Figure 1A). A large number of other Sox and Pax family members are co-expressed in other cell types (Figure 1B). Given that Sox2 and PAX6 have been shown to act as ‘master regulators’ of pluripotency and neurogenesis, we surmised that Sox–PAX cooperativity is key for cell fate decisions. We previously set up quantitative EMSAs to estimate cooperativity factors (\(\omega\)), i.e. the ratios of the equilibrium binding constants of dimeric versus monomeric complexes of TFs binding to composite DNA enhancer elements (16,42). Three minimal native sequences extracted from the DC5 (24), N3 (36) and LE9 (49) enhancer elements were chosen to study the interaction between the Sox2-HMG and Pax6-PRD domains (henceforth termed Sox2 and Pax6).

In addition, we used a modified sequence of the DC5 element, DC5con (24) where the native low-affinity pax half-site is replaced with a high-affinity half-site as determined by SELEX (32,50) (Figure 2A). In agreement with previous studies (24), Sox2 and Pax6 were found to form ternary complexes on both the DC5 and N3 enhancer elements (Figure 2B). By contrast, using the LE9 enhancer (49), we observed competitive binding (Figure 2B). When the pax half-site of the wild-type DC5 element was replaced with DC5con, ternary complex formation was markedly weakened as evident from the observation that Sox2–DNA and Pax6–DNA complexes are still prominently visible in addition to the ternary complex suggesting ineffective complex formation (Figure 2B). To understand the structural basis for the cooperativity, we generated structural models of Sox2–Pax6 complexes on DC5con, DC5 and N3 enhancer sequences (Figure 2C). The models show that Sox2 and Pax6 do not interact directly through the globular core of their DBDs. However, they may form some protein–protein interactions through the C-terminal tail of Sox2 and communicate through DNA-mediated allostery. Indeed, when we calculated the cooperativity factor, we obtained values between 20 and 50 for Sox2–Pax6 cooperation on DC5 and N3 elements, respectively (Figure 2D). However, on the DC5con element, we measured additive binding (\(\omega = 1\)), demonstrating that degenerated pax half-sites as found in the native DC5 and N3 elements are essential for the cooperative interaction of Sox2 and Pax6. Next we incubated Sox2 and Pax6 simultaneously with Cy5-labeled DC5 and FAM-labeled DC5con to evaluate the distribution of Sox2 and Pax6 on alternative target sites. Binary Pax6–DNA complexes are detectable at low Pax6 concentrations on the high-affinity DC5con element. By contrast, ternary Sox2–Pax6–DNA complexes predominate on the high-cooperativity DC5 element and binary Pax6–DNA complexes are barely visible (Figure 2E). This suggests that in a cellular environment where large numbers of DNA motifs compete for TF binding, Pax6 is effectively recruited to its consensus binding site. However, activation-competent ternary Sox2–Pax6 complexes are more effectively recruited to composite motifs with cryptic DC5-like pax half-sites.
Figure 2. Pax motif degeneracy is necessary for cooperative ternary complex assembly. (A) A SELEX-derived Pax6 PWM (50) was downloaded from JASPAR and is displayed above DC5 (21) and N3 (36) enhancer sequences. The residues introduced into the DC5con element to generate a pax site resembling the SELEX consensus are shown in red. (B) EMSAs to monitor the formation of ternary Sox2–Pax6–DNA complexes on DC5con, LE9, DC5 and N3 elements. The cartoon on the left depicts different complexes obtained during EMSA. (C) A structural model of Sox2–Pax6–DNA complex is shown. The DNA is shown as gray surface. The Sox2-HMG derived from PDB-ID 1GT0(13) is shown in blue and the Pax6-PRD (PDB-ID 6PAX (30)) in orange. Pax6-PRD subdomains (DBD1 and DBD2) and the linker are marked. (D) Cooperativity factors are calculated after quantifying the four possible microstates seen on EMSA gels (free DNA, Sox2.DNA, Pax6.DNA and Sox2.Pax6.DNA) using established procedures (16) and are shown as boxplots using data from 9–13 replicate experiments. Values for the LE9 enhancer could not be reliably determined as the ternary complex bands were barely detectable. (E) EMSAs were performed by simultaneously incubating Sox2 and Pax6 with differently labeled DC5 and DC5con DNA. Gels are shown after scanning using Cy5 (670 nm band-pass emission filter) or FAM (520nm band-pass emission filter) settings on a Typhoon phosphorimag. In total, 300 nM of each Cy5-DC5 and FAM-DC5con were used along with 100 nM Pax6 (FAM and cy5 control, lanes 3,4,7,8) or 300 nM Pax6 (lanes 10–18). Moreover, 100 nM Sox2–109aa was used in the Cy5 and FAM control reactions (lanes 2,4,6,8) or with increasing concentrations from 25 to 600 nM (lanes 11–18).

The composite Sox2–Pax6-binding site is compact and rigidly spaced

Next we studied the effect of inter-motif distances on the efficiency for Sox2–Pax6 dimerization. The 2, 4 and 10 base pair spacers have previously been found to abrogate Sox2–Pax6 cooperation on the DC5 element (24). Here we extended these studies by interrogating the effect of motif spacing from -2 to 5 base pairs at 1-bp intervals (Figure 3A and B). We observed that even subtle changes to the spacing between the sox and pax half-sites had profound effects on complex formation (Figure 3A–C). Ternary complexes are barely visible and cooperativity factors turn negative upon reducing the spacing between the motifs by 1 and 2 bp. This indicates competitive binding possibly due to steric hindrance when the proteins are brought too close to each other. Upon increasing the spacing (+1,+2,+3 & +5), the cooperativity factor markedly decreases (~30-fold) reminiscent of the additive binding seen on the DC5con element (Figure 3B and C). The +4bp spacer is an exception as it retains a weak albeit still strongly reduced cooperativity as compared to the other non-native spacers (Figure 3C). Collectively, these experiments highlight that precise motif juxtaposition is necessary for the formation of a cooperative Sox2–Pax6–DNA complex. Even small perturbations to the
Figure 3. Effect of motif spacing and sequence features on Sox2–Pax6 cooperativity. (A,B) EMSAs using DC5 enhancer elements with systematically altered spacing between sox (blue) and pax (orange) binding elements. (C) Cooperativity factor measurements revealed a drop in the cooperativity by at least an order of magnitude for all artificial spacers as compared to the native element. (D) DC5 sequences were sequentially converted into DC5con-like sequences corresponding to the Pax6 consensus and cooperativity factors were determined (E). Whiskers in barplots indicate standard deviations. (F) Interactions between Asn14 and Gly15 and DC5con-like sequence as seen in the 6pax crystal structure and the DC5con model and perturbations of these interactions when Pax6 is bound to DC5 (G) or N3 (H) sequences.
enhancer architecture lead to drastic reductions in ternary complex assembly, suggesting rigid spatial constraints for the assembly of the complex.

**Nucleotide switches act in concert to enable cooperative Sox2–Pax6 interaction**

Next we aimed to identify the sequence features that convert the high-cooperativity DC5 element that facilitates dimer formation of Sox2 and Pax6, into a site detrimental to their co-recruitment. The pwm (position weight matrix) of the ideal Pax6 motif (JASPAR ID: MA0069.1) derived from SELEX assays (50) identifies four nucleotide positions with high information content that deviate from the degenerate pax half-site in the native DC5 and N3 elements (Figure 3D). These positions will be referred to as T3, G8, C9 and T11 (Figure 3D). In order to dissect the role of these four key nucleotide positions in the DC5 element, we sequentially converted the degenerate pax half-site in the native DC5 element to resemble the high-affinity DC5con pax half-site (Figure 3D). Interestingly, point mutations to any of the four key nucleotide positions in the DC5 element did not decrease cooperative binding (Figure 3E). In fact, some point mutations even led to an elevation of the cooperativity, including the G8 and C9 positions (Figure 3E). However, a G8C9 di-nucleotide mutation substantially lowered the cooperative binding (Figure 3E). The tri-nucleotide mutations T3G8C9 and G8C9T11 further enhanced this effect and successively approached a cooperative reminiscent of DC5con (Figures 2C and 3E). This suggests that the nucleotide positions do not linearly contribute to the complex formation but are interdependent. Our models of the three ternary complexes revealed that the interactions between the Pax6-PRD around nucleotides G8, C9 and T11 were found to be most optimal in DC5con (Figure 3F). The less optimal configuration of Pax6 bound to the DC5 (Figure 3G) and N3 (Figure 3H) sequences may be further improved by small reorientations of the Pax DBDs that may lead to configurations enhancing cooperativity with Sox2. Such configurations may also be triggered by changes in DNA structure induced by the presence of Sox2. However, it cannot be ruled out that, the DC5 and N3 sequences may induce an assembly of Pax6 in a conformation completely different from the one in our models, which resemble that in the structure of a previously solved Pax6-DNA complex (PDB: 6PAX) (30). Alternatively, the orientation of Pax6 could change to accommodate the DC5 sequence. Indeed, when we inspected the reverse-complement of the pax consensus sequence (con′) and of the consensus sequence co-crystallized with Pax6 (30) (6pax-rev), we observed a good agreement with DC5 (Figure 4A). In particular, nucleotides 3, 9 and 11 found to be critical for the ‘cooperativity switch’ (Figure 3) align perfectly between DC5 and 6pax-rev. We therefore decided to test if the DC5 element encodes a cryptic Pax consensus and mediates cooperative binding through a changed orientation of Pax6. To this end, we constructed DC5con-1 and DC5con-2 sequences containing a reverse complement of the sequence found in the 6pax crystal structure spaced appropriately so that the key nucleotides 3, 9 and 11 align with wild-type DC5 (Figure 4A). Structural modeling (refer Material and methods, Homology modelling section) suggests that Sox2 and Pax6 could co-bind such sequences (Figure 4B). However, EMSAs revealed that complex formation is inefficient on both tested DC5con′ sequences and monomeric complexes predominate (Figure 4C–F). Moreover, when the three nucleotides 3, 9 and 11 that match DC5 in DC5con′ but not in DC5con were mutated in DC5, the cooperativity remained high (Figure 4G and H). Collectively, these experiments suggest that Pax6 is unlikely to bind DC5 in a ‘flipped’ conformation (Figure 4B).

**Cooperative binding to specific enhancer sequences is a conserved trait in the Sox and Pax protein families**

We previously studied cooperative binding of Sox and Oct protein families to identify signature enhancer sequences. We observed a partner code with distinctive Sox-Oct combinations binding specific composite sox-oct DNA motifs (16). Selective binding relies on certain amino acids of the Sox HMG and interchanging these residues between Sox factors swaps their ability to dimerize with Oct4 on specific composite motifs and to direct cell fate decisions during pluripotency or endoderm induction (14–15,51). To test whether a similar partner code exists for Sox–Pax interactions, we purified representative members of the Sox and Pax families and conducted quantitative cooperative measurements. First we asked whether Pax6 selectively cooperates with Sox2 or if it also teams up with other Sox factors. To this end, we selected members from several subgroups of the 20-member Sox family. Specifically, we chose Sox4 of the SoxC group, Sox5 of the SoxD group, Sox17 of the SoxF group and Sox15 of the SoxG group and tested them alongside the SoxB1 group member Sox2. We found that Sox2, Sox4, Sox5, Sox15 and Sox17 cooperatively bound with Pax6 to DC5 and N3 enhancer elements in an indistinguishable fashion (Figure 5A and B). Next we conducted inverse experiments and compared the cooperative binding of Sox2 with Pax2, Pax3, Pax5, Pax6, Pax8 and Pax9 to the DC5 sequence. There are some sequence variations within the PRD paralogs, some of which were previously found to be critical for discriminative DNA binding (32) (Figure 5C). Nevertheless, we found that all Pax family members can cooperatively pair-off with Sox2 on the DC5 element, whereas the cooperativity is abolished on the DC5con element (Figure 5D and E). This suggests that all Pax PRDs possess biochemical features conducive for the cooperation with Sox2 on specific enhancers with cryptic Pax6-binding sequences. Therefore, cooperativity on DC5-like sequences is a biochemical trait conserved within Sox and Pax TF families. As Sox and Pax paralogs are co-expressed in diverse cell types (Figure 1B), it can be envisaged that there are ample opportunities for members of the two families to interact. However, the propensity to cooperatively recognize DC5-like sequence in vitro does not imperatively suggest that all co-expressed Sox/Pax pairs combinatorially regulate gene expression and further studies are necessary to establish new functional Sox/Pax partnerships. First, it has to be assessed whether co-expressed Sox/Pax pairs indeed co-bind enhancer sequences. Second, it has to be tested whether the binding event leads to the transactivation of a target gene.
Figure 4. Sox2 and Pax6 do not cooperate on DC5con' elements encoding a flipped pax consensus. (A) The sequence logo of forward and reverse pax consensus motifs is aligned with sequences of DC5, DC5con, DC5con' and forward and reverse sequences co-crystallized with Pax6 (PDB-ID 6pax). Switch nucleotides (Figure 3) are marked with asterisks. The nucleotides 3, 9 and 11 found to be part of the ‘cooperativity switch’ are identical between DC5 and DC5con'. (B) Structural model of a Sox2/Pax6 complex on the DC5con' sequence with Pax6 in a hypothetical flipped conformation. EMSAs to study the cooperativity of Sox2 and Pax6 on DC5 (C), DC5con (D), DC5con'-1 (E), DC5con'-2 (F) and mutations to make the DC5 sequence more closely match the DC5con sequence T3T11 (G) and T3C9T11 (H).
Figure 5. The potential to cooperate is conserved in the Sox and Pax TF families. Formation of ternary complexes between Pax6 and long (109aa) and short (79aa) versions of the Sox2-HMG and paralogous mouse Sox HMGs (79aa) of Sox4, Sox5, Sox17 and Sox15 on (A) DC5 and (B) N3 elements. Note that 79 amino acid Sox-HMG versions co-migrate with the Pax6-PRD bound to DNA, whereas the 109aa version migrates as distinctive band. (C) Multiple sequence alignment of mouse Pax paralogs marking secondary structure elements above the alignment. Numbering corresponds to human PAX6 uniprot-ID P26367 for easier reference to eye diseases mutations. Red bars mark the Asn17 and Gly18 di-peptide binding to the di-nucleotide switch and residues previously shown to provide discriminatory sequence recognition between Pax6 and Pax5 (32). (D and E) The Sox2-HMG (109aa) was assessed for its propensity to cooperate with Pax family members Pax2, Pax3, Pax6, Pax8 and Pax9.
DNA binding of clinically relevant Sox2 and Pax6 mutant proteins

Several eye diseases including blindness and aniridia are caused by heterozygous Pax6 mutations, several of which map to the paired domain (52–56). Likewise, Sox2 mutations were recently reported to be associated with visual disorders (57,58). As the biochemical consequences that lead to disease are not understood for most of those mutations, we tested whether DNA binding or protein-partnerships are perturbed. One selected mutation was found in a four-generation family proband in Australia that has bilateral anophthalmia and several milder visual system anomalies like typical optic fissure coloboma (57). A novel D123G mutation in the C-terminal tail following the Sox2 HMG domain was detected in this family that was attributed to anophthalmia (57). Since D123G maps to a region near the hypothetical binding interface between Sox2 and Pax6 (Figure 6A), we examined whether this mutation alters the propensity to form a ternary complex. However, we could not detect any changes in complex formation compared to the wild-type Sox2 (Figure 6B). Similarly, co-binding with Oct4 in the Fgf4 enhancer element was also not altered for the Sox2D123G mutation (Supplementary Figure S1). Next we selected the Pax6 missense mutations G36R and R44Q previously detected in aniridia patients (59,60). In our structural models, those mutations are in proximity to the putative Sox–Pax contact interface (Figure 6C). We therefore wanted to examine whether these two mutations could alter cooperativity with Sox2. We observed that G36R and R44Q markedly reduced affinity for DNA binding (Figure 6D). However, Pax6R44Q did not affect the cooperativity with Sox2 on the DC5 element (Figure 6E). The cooperativity factor for Pax6G36R could not be determined due to the low overall affinity for DNA. Therefore, diminished DNA binding of the Pax6 protein likely causes the development of aniridia in Pax6R44Q and Pax6G36R patients. Consistently, in co-crystal structures of the highly homologous Drosophila Prd protein bound to DNA it can be seen that the residue homologous to R44Q is in contact with the sugar phosphate backbone of DNA (29).

NMR and modeling imply a DNA-mediated allosteric mechanism for Sox2–Pax6 interaction

The mechanism of the cooperative recognition of DC5 DNA by Sox2 and Pax6 cannot be straightforwardly modeled based on available experimental structures. Our biochemical assays and structural models suggest that in particular the Pax6-PRD might adopt different conformations compared to the configuration seen when bound to its high-affinity element (29,30). Such conformational versatility has been described for the POU family that also contains a bivalent DBD. POU members were found to adopt strikingly different tertiary structure and altered stoichiometry depending on the sequence of the bound DNA element (61–63). To define the protein–protein interaction interface between Sox2 and Pax6, we performed NMR studies. First, we measured 1D 1H spectra of Sox2 alone, DC5, Sox2–DC5 and ternary Sox2–Pax6–DC5 complexes (Figure 7A). DC5 in complex with Sox2 showed significantly more dispersed imino proton resonances at the region between 12 and 14.5 ppm indicating bending of the DNA induced by Sox2 (Figure 7A). In addition, we observed significant chemical shifts and differential line broadening of the 1H doublet signals from the indole groups of the Sox2 tryptophan residues W13 and W41 in the 10–11 ppm region upon DNA addition, indicating a significant increase in molecular weight and effective complex formation. Addition of Pax6 to the Sox2–DC5 binary complex enhances differential line broadening in the 1H doublets of W13 and W41, indicating further molecular weight increase upon formation of the ternary complex (Figure 7A). However, binding of Pax6 to the Sox2–DC5 complex results only in relatively minor shifts of both Sox2 tryptophan 13N and 1H resonances (Figure 7B). Yet, the 1H resonance corresponding to W41 (eH) is observed to attenuate considerably while W13 (eH) remains stable (Figure 7A and B). These results suggest that incorporation of Pax6 into a ternary complex might affect the conformational dynamics of Sox2 and possibly affects how Sox2 bends DNA.

Comparative 13N,1H Sox2 spectra before and after DC5 addition revealed CSP and conformational-exchange induced line broadening for many peaks (Figure 8A and B). By mapping CSPs to our models of Sox2/DNA and Sox2/Pax6/DNA complexes (13), we found that residues undergoing CSPs are not restricted to the DNA contact interface, suggesting global structural rearrangements of Sox2 upon DNA binding (Figure 8C and D). The largest CSPs (> 0.1 ppm) observed are for K4, G16, K20, E24, N25, S34, K42, L43, A56 and K65 (Figure 8E). Residues that experience line broadening are predominantly located in the hydrophobic core of the protein. When Pax6 is added to form the Sox2–Pax6–DC5 ternary complex, only a few additional CSPs were observed in Sox2 (Figure 8B, D and E: K4, Q23, T80, L81). Yet, Pax6 addition leads to a considerable attenuation of peak intensity in the region of the major wing (helices 1 and 2), indicating that Pax6 may primarily affect the dynamics and stability of the Sox2–DC5 interactions rather than directly binding to Sox2 and altering its conformation. Notable exceptions are R15 and R19 that experience an increase in the peak intensity after Pax6 addition potentially indicating more favorable electrostatic interactions for these residues in the ternary complex. CD measurements showed that the DNA peak at 280 nm is slightly shifted in DC5 as compared to DC5con DNA (Figure 8F and G). Likewise in silico predictions (GBShape (64)) of the minor groove width suggest an altered pattern of groove compressions over the pax half-site (Figure 8H). Those structural differences could affect the propensity of the respective DNA sequences to mediate the co-recruitment of Sox2 and Pax6. Collectively, our data suggest that Sox2–Pax6 cooperation on DC5-like sequences is not mediated by complementary protein–protein interactions but rather by an allosteric, DNA-mediated mechanism that relies on a subtly adjusted DNA structure in agreement with our models of the ternary complexes. Collectively, our data suggest that, DC5-like sequences support a predominantly DNA-mediated mechanism of cooperativity, but DC5con-like sequences do not.
Figure 6. Eye disease mutations and Sox2–Pax6 interactions. (A) Structural model of a hypothetical Sox2–Pax6 complex. The D123G mutation (57) maps to a region in the C-terminal tail just off the portion visible in the crystal structure (indicated with arrow and dashed line). A domain plot indicating the Sox2 constructs used for cooperativity measurements are shown below. (B) A barplot comparing cooperativity factors for the interaction of Sox2 versus Sox2D123G with Pax6 on DC5 and N3 DNA elements. Whiskers indicate standard deviations from seven to nine measurements. (C) Location of two Pax6 mutations R44Q and G36R within the Pax6 PRD. (D) The missense mutations lower the affinity for DNA as compared to the wild-type PRD on a consensus Pax6-binding element (Pax6_xtal, Supplementary Table S1). (E) Cooperativity measurements for Pax6 and Pax6R44Q on DC5 and DC5con elements show that the R44Q mutation does not abolish cooperative complex formation. Whiskers indicate standard deviations from five to six measurements.

Identification of cryptic SOX2/PAX6 cooperativity loci in human NSC enhancers

The chicken DC5 and N3 sequences support cooperative and functionally important Sox2/Pax6 interactions. However, the cooperative interaction of SOX2 and PAX6 on related sequences has so far not been reported for human cells. We hypothesized that SOX2 and PAX6 also cooperate on DC5-like sequences during human neurogenesis. To test this hypothesis, we took advantage of recently published ChIPseq studies of Pax6 in human NSCs (referred to as neuroepithelial cells by the authors) (45) and SOX2 ChIPseq data from NPCs (46). While both cell types are not identical, they constitute the most closely related human cell types for which ChIPseq data are presently available. After intersection, we obtained 2353 regions co-bound by Pax6 and SOX2 (Figure 9A). De novo motif searches revealed that more than 30% of the shared loci contain consensus sox motifs (Figure 9B). However, known pax motifs are only present in ~3% of the putative NSC enhancers, suggesting that the vast majority of Pax6-targeted sites does not rely on high-affinity pax motifs in vivo. As only the N3 and DC5 sequences are known to support Sox2/Pax6 cooperation, there is no position-weight matrix or related binding models for capturing cryptic pax half-sites as observed in the DC5 or N3 enhancer. We therefore manually constructed degenerate DC5/N3-like consensus motif
The functional studies and the intricate behavior in qualitative binding assays encouraged us to quantitatively dissect Sox2 and Pax6 cooperativity and the molecular basis for this partnership. Collectively, we found that (i) cooperativity critically relies on degenerate *pax* sites and that consensus sites abrogate ternary complex formation; (ii) highly constrained spacing arrangements of *sox* and *pax* half-sites are critical for cooperative binding; (iii) key nucleotide switch positions in the degenerate *pax* half-sites act in concert to mediate cooperative assembly of Sox2 and Pax6; (iv) the mechanism of cooperative binding is evolutionary conserved within Sox and Pax families and, finally, (v) direct protein–protein interactions between Sox2 and Pax6 are sparse if present at all. Overall, these results suggest that stable Sox–Pax interactions rely on specific DNA sequences and not complementary protein–interaction interfaces. To rationalize the switch from a high-affinity DC5con-like to a high-cooperativity DC5-like binding configuration, we inspected the molecular environment of switch nucleotides in structural models. The ‘T3G8C9’ nucleotides are major features triggering the switch. T3 contacts Asn50 through van der Waals interactions (Figure 3F–H). G8 is directly contacted by Asn17 via hydrogen bond, G9’ (the base complementary to C9) is contacted by Gly18 through hydrogen bonds and finally T11 is contacted by the linker region residue Gly72 (30). Replacing G9’ by a DC5-like A9’ suggests a detrimental interaction with Gly18 as the N2 primary amine hydrogen donor is lost. Likewise, replacing G8 by C8 could de-stabilize the side chain Oδ of Asn-17 as it loses the H-bond donor amine at the position (Figure 3F, G and H). Replacing T11 to C (as in DC5) or G (as in N3) also leads to a less protein–DNA interactions in the region of the Pax6 linker, which was previously shown to be crucial for DNA binding (30). Hence, we surmise that on the high-affinity *pax* half-site, the ‘G8C9’ nucleotide in the DC5con sequence tightly tethers Pax6 to the minor groove of the DNA in a constrained configuration, in a manner that is independent of the influence of an adjacent Sox2-binding site. Hence, Sox2-Pax6 dimers are formed in an additive fashion. By contrast, a loss of this interaction by mutating G8C9 removes constraints and allows the Pax6 protein to adopt a conformation in favor of cooperative binding. The cooperative assembly on DC5-like sequences is likely facilitated by an allosteric, DNA-mediated mechanism. While structural models provide a rudimentary framework, the overall effect of these mutations in the degenerate *pax* half-sites on the binding topology and conformation of Pax6 is still ambiguous and will be made clear only when atomic resolution structures of Pax6 on DC5-like sequences become available (41).

A recent study on Sox2 and Pax6 cooperativity used plasmon resonance and gold nanoparticles tethered to the end of DC5 and DC5con sequences. While Sox2 alone induced indistinguishable bends, it was reported that the DC5 is more markedly deformed by Sox2–Pax6 dimers than the DC5con element (65). Apparently, Sox2–Pax6 binding induces structural changes to DNA in a sequence-dependent manner. Another study employed single molecule atomic force microscopy and reported cooperative binding as well as the requirement for Sox2-induced bending for effective complex formation (66). These studies are in accordance with the current findings.
with a model where DNA deformations supportive of cooperative complex formation occur on the DC5-like sequence but not on DC5con-like sequences. However, a R75E mutation mapping to the C-terminal extension of the Sox2-HMG was demonstrated to abrogate Sox2–Pax6 dimerization on DC5 (13). As this site interacts with Oct1 in the crystal structure of the Sox2/Oct1/Fgf4 ternary complex, the authors reasoned that this residue directly interacts with the Pax6-PRD as well (13). However, in our NMR experiments R75 could not be assigned due to pronounced line broadening. From our structural models, we observed that R75 is in proximity to the DNA backbone and is not far apart from the Pax6 PRD domain (Figure 5C). As our models on the DC5 and N3 enhancers suggest that the configura-
Figure 9. Identification of loci supporting Sox2/Pax6 dimerization during human neurogenesis. (A) Scheme illustrating the selection of candidate DC5-like sequences bound by Pax6 and SOX2 in human cells. (B) *De novo* motifs recovered using homer (http://homer.salk.edu/homer/chipseq/) in SOX2/PAX6 co-bound loci. (C) Genome plots spanning 100 kb (ACACA: acetyl-CoA carboxylase Alpha, EFNA5: ephrin-A5, UPP2: uridine phosphorylase 2) or 184 kb (FGFR2: fibroblast growth factor receptor 2) are shown for selected candidate loci predicted to recruit SOX2/PAX6 dimers in a cooperative fashion. Black bars indicate binding sites with DC5-like sequences used for EMSA validation. Binding site definitions were used as reported in the original publications (45,46). (D) Sequences of novel DC5-like composite motifs short-listed for EMSA validation. (E) EMSAs showing effectively formed SOX2/PAX6 dimers on newly identified NSC enhancer sequences.

ation of the ternary complexes needs to adapt to promote Sox–Pax cooperativity, it is possible that a glutamate at position 75 interferes with the adaptation. However, we cannot exclude that there might be some direct protein–protein interactions that do not lead to obvious chemical shifts in 2D-NMR. Yet, we surmise that a DNA-mediated mechanism is the main driver for the cooperative co-recruitment of Sox2–Pax6 to DC5-like sequences.

Genomic studies also point toward the versatile mechanism by which Pax6 engages chromatin. Analysis of the genome-wide binding of Pax6 by ChIPseq in human neuroectoderm revealed that less than 3% of the binding sites contain consensus *pax* motifs (Figure 9B). However, *sox* motifs were frequently found in proximity to Pax6-binding sites (Figure 9B). Moreover, missense mutations in either the PAI or RED subdomains differentially affect the genomic target selection of Pax6 (35). Apparently, some enhancers more strongly rely on PAI and others on RED for efficient genomic targeting. This suggests that Pax proteins employ different binding modes to recognize alternative sets of target sequences. Such differences could account for differential TF partner recruitment and profoundly affect the regulatory outcome of the binding event.

The striking observation that partner factors prefer sequences that are very different from their consensus motif creates another layer of complexity in understanding the principles underlying combinatorial regulation of transcription. A number of TF families are known to be able to accommodate diverse sets of binding sequences. Such binding modes pose specific challenges for motif scanning tools that rely on simple position weight matrices, which assume positional independence, or on consensus k-mer patterns from high-affinity sequence binders (67) or are too divergent to be captured by probabilistic models such as HMMs (68). Yet, given the critical roles of Sox and Pax TFs to direct cell fate choices and the frequent co-occurrence in the nuclei of diverse cell types, such unchartered non-consensus target sites are likely fundamentally important. Consistently, we detected eight novel target sites in human NSC enhancers where SOX2 and PAX6 cooperate. There-
fore, an unbiased assessment of how TF partnerships influence DNA sequence preferences as well as an understanding of the structural basis for DNA-induced allostery is required to decode how developmental programs are genetically hard-wired.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online.

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