Six1 and Six2 of the Sine Oculis Homeobox Subfamily are Not Functionally Interchangeable in Mouse Nephron Formation

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The vertebrate Six1 and Six2 arose by gene duplication from the Drosophila sine oculis and have since diverged in their developmental expression patterns. Both genes are expressed in nephron progenitors of human fetal kidneys, and mutations in SIX1 or SIX2 cause branchio-oto-renal syndrome or renal hypodysplasia respectively. Since ~80% of SIX1 target sites are shared by SIX2, it is speculated that SIX1 and SIX2 may be functionally interchangeable by targeting common downstream genes. In contrast, in mouse kidneys, Six1 expression in the metanephric mesenchyme lineage overlaps with Six2 only transiently, while Six2 expression is maintained in the nephron progenitors throughout development. This non-overlapping expression between Six1 and Six2 in mouse nephron progenitors promoted us to examine if Six1 can replace Six2. Surprisingly, forced expression of Six1 failed to rescue Six2-deficient kidney phenotype. We found that Six1 mediated Eya1 nuclear translocation and inhibited premature epithelialization of the progenitors but failed to rescue the proliferation defects and cell death caused by Six2-knockout. Genome-wide binding analyses showed that Six1 selectively occupied a small subset of Six2 target sites, but many Six2-bound loci crucial to the renewal and differentiation of nephron progenitors lacked Six1 occupancy. Altogether, these data indicate that Six1 cannot substitute Six2 to drive nephrogenesis in mouse kidneys, thus demonstrating that the difference in physiological roles of Six1 and Six2 in kidney development stems from both transcriptional regulations of the genes and divergent biochemical properties of the proteins.

Keywords: gene duplication, vertebrate evolution, Six1 and Six2, nephron progenitor, maintenance, differentiation

INTRODUCTION

The duplication of developmental regulatory genes is one of the evolutionary driving forces leading to the diversity and complexity of higher eukaryotes. One of such multigene families encodes for the sine oculis (SIX) homeodomain transcription factors, which can be divided into three subfamilies—Six1/2, Six3/6, and Six4/5 based on their sequence and structure conservation (Kawakami et al., 2000; Kumar 2009). Each of these subfamilies is respectively duplicated from each of the three SIX genes in Drosophila—so (sine oculis), ptx (also known as DSix3), and DSix4 (Seo et al., 1999). These factors are found in diverse organisms ranging from flatworms to humans.
and are crucial for cell lineages that give rise to cranial sensory organs, brain, kidney, muscle and gonads. The three Drosophila SIX genes are thought to have arisen by duplication of a single ancestral gene, an event that occurred prior to the evolution of the Bilateria (Kawakami et al., 2000). Since the genome of most invertebrates contains only a single gene for each subfamily, the complexity of the SIX subfamilies has arisen by a further duplication of each SIX gene at the onset of vertebrate evolution.

The SIX family proteins are characterized by the presence of two evolutionarily conserved domains, the SIX-domain specific (SD) that mediates interactions with partner proteins and DNA binding homeodomain (HD). Both the N-terminus adjacent to the 5′ end of the SD and the C-terminus flanking the 3′ end of the HD are considerably variable in length with a very low degree of sequence conservation across all SIX family members (Kawakami et al., 2000; Kumar 2009). Evidence from multiple studies has revealed a remarkable similarity in the binding sites for Six1/2/4/5 (TCAGGTTC) (Suzuki-Yagawa et al., 1992; Kawakami et al., 1996; Spitz et al., 1998; Harris et al., 2000; Sato et al., 2002; Brodbeck et al., 2004). Analysis of the divergent C-terminal domains of the SIX proteins found that its presence increases the affinity of the HD for DNA (Hu et al., 2008). However, how the structural diversity of the C-terminal regions confers specificity in regulating target genes remains poorly understood.

Six1 and Six2 of the so subclass show spatiotemporally overlapping expression patterns in many tissues during development (Oliver et al., 1995; Abitua et al., 2015; Horie et al., 2018). For instance, SIX1 and SIX2 are coexpressed in human fetal nephron progenitors (O’Brien et al., 2016). Heterozygous SIX2 missense mutations were identified in patients with renal hypoplasia characterized by reduced kidney size and/or maldevelopment of the renal tissue following abnormal organogenesis (Weber et al., 2008), while SIX1 mutations result in Branchio-Oto-Renal (BOR) or Branchio-Oto (BO) syndrome (Ruf et al., 2004; Okada et al., 2006; Kochhar et al., 2008), an autosomal dominant disorder characterized by abnormal development of the second branchial arch, otic with or without renal anomalies (Melink et al., 1976; Fraser et al., 1980; Abdelhak et al., 1997a; Abdelhak et al., 1997b; Vincent et al., 1997; Kumar et al., 2000; Vervoort et al., 2002; Kochhar et al., 2007). Because of the overlapping expression of SIX1 and SIX2 in the fetal nephron progenitors, previous studies used chromatin immunoprecipitation followed by deep sequencing (ChIP-seq) to address how much diversity exists among their transcriptional targets and found that ~80% of SIX1-bound sites are shared by SIX2 (O’Brien et al., 2016). This led to the speculation that SIX1 and SIX2 may be functionally interchangeable in targeting common downstream genes in the nephron progenitors, despite SIX2 binding to more sites. However, it is difficult to test this possibility and decipher the difference in their biochemical activities in the human kidney.

In contrast, the expression of Six1 and Six2 in the mouse kidney only transiently overlaps in the metanephric mesenchyme (MM) before the onset of ureteric bud (UB) branching (Li et al., 2003; Xu et al., 2003; Self et al., 2006; Nie et al., 2011; Park et al., 2012; O’Brien et al., 2016). Kidney organogenesis in mice commences when the MM induces the UB to outgrow from the nephric duct and invade into the MM at ~ E10.5-E11.0. The UB tip cells then induce the MM to form the cap mesenchyme (CM, also called nephron progenitors) surrounding the UB tip and subsequent reciprocal interactions between the CM and the UB tip cells lead to repeated UB branching to form the nephron tubules and collecting duct system. Throughout development, high levels of the Six2 expression are maintained in the nephron progenitors (Self et al., 2006; Park et al., 2012; O’Brien et al., 2016). Consistent with these differential expression patterns, germline Six2 deletion in mice leads to renal hypoplasia due to depletion of the nephron progenitors (Self et al., 2006), while Six1 knockout causes renal agenesis due to malformation of the MM (Li et al., 2003; Xu et al., 2003; Kobayashi et al., 2007; Nie et al., 2011; Xu and Xu 2015).

The non-overlapping expression between Six1 and Six2 in mouse nephron progenitors prompted us to examine if the forced expression of Six1 in the nephron progenitors can rescue Six2-deficient kidney phenotype by crossing a Six1 knockin into Eya1 locus (Eya1Six1+) mouse model (Nie et al., 2010) with Six2−/− mice (Self et al., 2006). Eya1 is coexpressed with Six2 in the MM progenitors throughout kidney development and it acts upstream of Six2, but their gene products interact and Six2 mediates Eya1 nuclear translocation (Xu et al., 1999; Xu et al., 2014). To our surprise, analysis of Eya1-Six1+/Six2−/− mice revealed that Six1 failed to rescue Six2-deficient kidney phenotype. We found that Six1 was able to mediate nuclear translocation of Eya1 and inhibit rapid premature differentiation of the progenitors that occurred in Six2−/− mice but failed to substitute for Six2 to renew and maintain the nephron progenitors. Hence, the Six1 and Six2 proteins are not functionally interchangeable in the nephron progenitors. We further performed ChIP-seq in ~E13.5 Eya1-Six1+/− kidneys to investigate if Six1 targets Six2-bound regulatory regions that are essential for the maintenance of nephron progenitors, as suggested in humans. Our analyses revealed that Six1 only selectively occupied a small subset of Six2 target sites, but many Six2-bound loci that are crucial to the renewal and differentiation of nephron progenitors lacked Six1 occupancy. Thus, these data demonstrate that Six1 and Six2 have not maintained equivalent biochemical properties since their divergence early in vertebrate evolution.

**MATERIALS AND METHODS**

**Animals and Genotyping**

All animal protocols were approved by Animal Care and Use Committee of the Icahn School of Medicine at Mount Sinai (protocol #06-0822).

The Eya1-Six1+/− (Nie et al., 2010) and Six2−/− (Self et al., 2006) were maintained at the Icahn School of Medicine at Mount Sinai Animal Facility. Mice were bred using timed mating, and noon on the day of vaginal plug detection was considered as E0.5.

**Histology, Immunohistochemistry, and in situ Hybridization**

Histology, immunohistochemistry, and in situ hybridization were carried out according to standard procedures. Briefly, kidneys...
were fixed in 4% paraformaldehyde (PFA) for overnight at 4°C, dehydrated, and embedded in wax or OCT. Paraffin or frozen sections were generated at 6µm of thickness. We used six embryos for each genotype at each stage for each probe and the result was consistent in each embryo. Probes for ISH were reported previously (Xu et al., 2014). Cy3-, Cy2-, Cy5- and FITC-conjugated secondary antibodies were used and Hoechst was used for nuclear counter-staining.

Primary antibodies: Anti-Six1 (12,891, Cell Signaling), -Six2 (MBS610128, Mybiosource), anti-Wt1 (sc-192, Santa Cruz Biotechnology), -Eya1 (25-067, Prosci Inc. and MABE1047, Sigma), and -PH3 (ab10543, Abcam).

**TUNEL Assays**

The TUNEL assay was performed using the Apop Tag kit for *in situ* apoptosis fluorescein detection (S7110, Millipore-Sigma) following the manufacturer’s instructions.

**Cell Counts and Statistical Analysis**

TUNEL- or PH3-positive MM progenitor cells on the peripheral side of branching UB were counted from 28 to 35 sections of 5 different kidneys and values represent average number of TUNEL⁺ or PH3⁺ cells (±standard deviations) per section (6µm). Two-tailed Student’s *t* test was used for statistical analysis.

**Chromatin Immunoprecipitation Followed by Deep Sequencing**

For Six1 ChIP, we used 40 kidneys from ~E13.5 Eya1Six1 embryos. ChIP-seq was performed according to previous protocols with some modifications. Briefly, the kidneys were cross-linked with 1% formaldehyde at room temperature for 30 min and then homogenized and lysed in cold lysis buffer (50 mM HEPES–KOH, pH 7.5, 140 mM NaCl, 1 mM EDTA, 10% glycerol, 0.5% NP–40, 0.25% Triton X-100, 1 × protease inhibitors), followed by spinning at 2000 g at 4°C, resuspending in cold wash buffer (10 mM Tris–HCl, pH 8.0, 200 mM NaCl, 1 mM EDTA, 0.5 mM EGTA, 1 × protease inhibitors) and spinning at 2000 g at 4°C in a benchtop centrifuge. Samples were then resuspended in 1 ml cold sonication buffer (10 mM Tris–HCl, pH 8.0, 2 mM EDTA, 0.1% SDS) and sonicated to 200–500 bp fragments using a Covaris S220 Focused-ultrasonicator. Sonicated chromatin was cleared by pelleting insoluble material at 13,000 RPM at 4°C, followed by preclearing with protein A/G beads and incubation with 1–2 µg antibody overnight (anti-Six1, Cell Signaling #12891) or 1–2 µg rabbit IgG as a negative control. Chromatin–antibody complexes were precipitated with protein A/G beads at 4°C for another 5 h. Immunoprecipitated complexes were subjected to series of wash steps with low salt buffer (20 mM Tris–HCl 8.0, 150 mM NaCl, 2 mM EDTA, 1% Triton X-100, 0.1% SDS), high salt buffer (20 mM Tris–HCl pH 8.0, 500 mM NaCl, 2 mM EDTA, 1% Triton X-100, 0.1% SDS), LiCl wash buffer (10 mM Tris–HCl pH 8.0, 250mM LiCl, 1mM EDTA, 1%NP–40, 1% sodium deoxycholate) and TE plus NaCl, followed by elution and reverse crosslinking overnight at 65°C. The quality control of ChIPed DNA was performed with Qubit 2.0 Fluorometer using dsDNA HS assay Kit (Q32854, ThermoFisher Scientific) and Agilent 2200 TapeStation System using High Sensitivity D1000 Reagents (5067–5585, Agilent). The pulldown and input control sequencing libraries were generated using the ThruPLEX DNA-seq Kit (R400429, Rubicon Genomics) and sequenced on Illumina NextSeq 500.

**Peak Calling, Annotation, and Motif Enrichment Analysis**

Quality controls using FastQC (v0.11.2) (www.bioinformatics. babraham.ac.uk/projects/fastqc) were generated and raw sequencing reads were then aligned to the mouse mm10 genome using default settings of Bowtie (v2.2.0). Peak-calling was performed using MACS (v2.1.1) with various *p*-value cutoffs. The peak bed files were generated from peak calling against genomic input control or IgG control with the default setting (10⁻⁵ cutoff). The common peaks from these two bed files were used for subsequent analyses. The overlapping peaks of bed files were identified using the bedtools from Galaxy (https://usegalaxy.org/).

Motif enrichment analysis was performed using the Homer package (v4.8.3) (Heinz et al., 2010). The peak annotation and gene ontology analysis was performed using GREAT program (great.stanford.edu). The bamCoverage and coverage heatmap were generated by centering and scaling corrected ChIP-seq coverage using mean and standard deviation, and plotting normalized coverage across 3-kb centered on the Six1-enriched region.

**Data Access**

ChIP-seq data sets have been submitted to the NCBI Gene Expression Omnibus (GSE189131).

**RESULTS**

**Forced Expression of Six1 in Nephron Progenitors Fails to Rescue Six2⁻/⁻ Kidney Phenotype**

The nephron progenitors on the peripheral side of the branching UB tip self-renew to replenish to generate a sufficient number of nephrons. In contrast, the progenitors on the ventral side of the branching UB tip undergo a mesenchyme-to-epithelial transition to form pretubular aggregates (PTAs), which then epithelialize to form renal vesicle (RVs)—the primordia of nephron tubules. Six2 is highly expressed in the multipotent nephron progenitors, and Six2 knockout leads to rapid RV formation and deletion of the nephron progenitors, resulting in hypoplastic kidneys (Self et al., 2006). To examine the functional equivalence between Six1 and Six2 during nephrogenesis, we tested whether forced expression of Six1 in the nephron progenitors can rescue Six2-deficient kidney phenotype by crossing Six2⁻/⁻ mice with a Six1 knockin mouse model expressing Six1 under Eya1 transcriptional regulatory control (Eya1<KM>) (Nie et al., 2010).
Eya1 is coexpressed with Six1 in the uninduced MM (Xu et al., 2003; Xu and Xu, 2015) and with Six2 in the CM, and it is upstream of Six2 (Xu et al., 2014). We previously reported that renal agenesis associated with Six1-deficiency can be rescued in Eya1Six1/+;Six1−/− mice (Nie et al., 2010). However, to our surprise, forced expression of Six1 failed to rescue Six2-deficient hypoplastic kidneys at E14.5–17.5 (n = 8, Figures 1A,B). Quantitative analysis of kidney size showed that the length of Eya1Six1/+ kidney was ~104 ± 5% (n = 6, p = 0.0375) at E14.5 and ~101 ± 3% (n = 8, p = 0.0418) at E17.5 of the wild-type, while the length of Eya1Six1/+;Six2−/− kidney was ~106 ± 5% (n = 8, p = 0.0357) at E14.5 and ~114 ± 7% (n = 8, p = 0.0208) of Six2−/− littermate. Histological analysis and immunostaining for Wt1, which is expressed in the CM, PTAs, RVs and differentiating podocytes, revealed that both Eya1Six1/+;Six2−/− and Six2−/− mice lacked the nephron progenitors in the outmost nephrogenic zone where the UB branching morphogenesis normally takes place (Figures 1C,D). Similar observation was obtained when we used Eya1CreER to induce Six1 expression in the MM by tamoxifen administration (Eya1CreER;R26-Six1mCherry) and tested its ability to restore kidney development in Six2−/− (data not shown). Thus, these findings demonstrate that the

![FIGURE 1](https://example.com/fig1.png)

**FIGURE 1** | Forced expression of Six1 in the MM cells under Eya1 transcriptional control cannot rescue the Six2-deficient kidney phenotype. (A,B) Kidneys of wild-type, Eya1Six1/+; Six2−/− and Six2−/− at E14.5 and E17.5. (C) Hematoxylin and Eosin-stained sections of wild-type, Eya1Six1/+, Eya1Six1/+;Six2−/− and Six2−/− kidneys at E14.5. Arrows point to glomerulus-like structures. (D) Anti-Wt1 immunostaining on sections showing depletion of Wt1-labeled nephron progenitors (white arrows) in the outmost region of both Eya1Six1/+;Six2−/− and Six2−/− kidneys. Panels on the right are higher magnification of the panels on the left. Abb. a, adrenal gland; k, kidney; ub, ureteric bud; ur, ureter. Scale bars: 100 μm.
expression of Six1 in the MM cells under the Eya1 transcriptional regulatory control cannot functionally substitute for Six2 to maintain the nephron progenitor cells during mouse kidney development.

**Six1 Inhibits Rapid Differentiation of the Nephron Progenitors Caused by Six2 Knockout**

We next examined whether there is a partial rescue at earlier stages, from the first “T” bud stage at E11.5. As reported previously (Self et al., 2006), the progenitors within the entire MM surrounding the branching UB underwent premature differentiation to form PTA- or RV-like structures in Six2−/− embryos at E11.5 (Figure 2D) and RV-like structures surrounded the entire UB at E12.5 (Figures 2H,I). In contrast, Eya1Six1/+;Six2−/− MM progenitors surrounding the branching UB appeared less dense than those in Eya1Six1/+ or wild-type littermate controls at E11.5 (Figures 2A–C), but they did not commit to an overall RV fate, as shown by histological analysis and anti-Wt1 immunostaining (Figures 2C,G,K). This observation was consistent with the expression pattern of Wnt4, which is one of the earliest markers labeling differentiating nephron structure PTAs at E11.5 and RVs at E12.5 in the MM (metanephric mesenchyme) surrounding the branching UB (ureteric bud) (arrows in D,H). Arrows in panels C and G indicate the absence of PTAs or RVs within the MM cells of Eya1Six1/+;Six2−/− (Figures 2C,G,K). Anti-Wt1 immunostaining on sections of wild-type, Eya1Six1/+;Six2−/−, and Six2−/− kidneys at E12.5. Note that the kidney development in panel K was slightly delayed. (M–T) Whole-mount in situ hybridization showing Wnt4 expression in differentiating progenitors in wild-type, Eya1Six1/+ and Eya1Six1/+;Six2−/− and its global activation in Six2−/− MM at E11.5 and E12.5. Arrows in C and G indicate the absence of Wnt4 activation in the MM progenitors at UB tips. Arrows in P and T indicate ectopic Wnt4 activation in the MM of Six2−/− embryos. Scale bars: 100 μm. FIGURE 2 | Forced expression of Six1 in the MM progenitors appears to inhibit their rapid epithelization induced by Six2 knockout. (A–H) Hematoxylin and Eosin-stained kidney sections of wild-type (A,E), Eya1Six1/+ (B,F), Eya1Six1/+;Six2−/− (C,G) and Six2−/− (D,H) at E11.5 and E12.5. Six2−/− displayed aggregates at E11.5 and RVs at E12.5 in the MM (metanephric mesenchyme) surrounding the branching UB (ureteric bud) (arrows in D,H). Arrows in panels C and G indicate the absence of PTAs or RVs within the MM cells of Eya1Six1/+;Six2−/−. (I–L) Anti-Wt1 immunostaining on sections of wild-type, Eya1Six1/+;Six2−/−, and Six2−/− kidneys at E12.5. Note that the kidney development in panel K was slightly delayed. (M–T) Whole-mount in situ hybridization showing Wnt4 expression in differentiating progenitors in wild-type, Eya1Six1/+ and Eya1Six1/+;Six2−/− and its global activation in Six2−/− MM at E11.5 and E12.5. Arrows in O and S indicate the absence of Wnt4 activation in the MM progenitors at UB tips. Arrows in P and T indicate ectopic Wnt4 activation in the MM of Six2−/− embryos. Scale bars: 100 μm.
but the second branching was either incomplete or abnormal. Thus, while the UB development in Eya1Six1/+;Six2−/− is arrested at the second branching, Six1 expression in Six2−/− MM appears to prevent premature mesenchyme-to-Epithelial Transition.

Six1 Cannot Substitute for Six2 to Maintain the Nephron Progenitors

As we noticed that the MM in Eya1Six1/+;Six2−/− is not as dense as in Eya1Six1+/+, we sought to examine whether Six1 can rescue cell survival or proliferation defects in Six2−/− embryos. TUNEL (terminal deoxynucleotidyl transferase dUTP nick end labeling) assay revealed a noticeable degree of apoptosis throughout the MM areas, as a marked increase in the number of TUNEL+ cells in the MM was detected in both Eya1Six1/+;Six2−/− and Six2−/− (Figures 3A,C). It should be noted that increased apoptosis was also observed in the branching UB of Six2−/− embryos at E11.5 (Figure 3A). In contrast, anti-PH3 (phosphohistone H3)-labeled mitotic cells in the MM were decreased in Eya1Six1/+;Six2−/− and more decreased in Six2−/− littermate embryos (Figures 3B,D). These data suggest that while Six1 appears to prevent premature epithelialization of the progenitors, it is not able to substitute the essential role of Six2 in nephron progenitor renewal and survival, thus leading to depletion of the nephron progenitors and arrest of nephrogenesis during second UB branching in Eya1Six1/+;Six2−/− embryos.

Six1 Mediates Nuclear Localization of Eya1 in the Nephron Progenitors

Loss of either Eya1 or Six2 in the nephron progenitors leads to increased cell death and reduced proliferation (Figure 3) (Self et al., 2006; Xu et al., 2014). We therefore performed ISH to confirm Six1 expression in the MM progenitors on the peripheral side of the branching UB in Eya1Six1/+ at E11.5, compared to no expression in wild-type or Six2−/− littermates (Figure 4A). In Eya1Six1/+;Six2−/−, the MM appeared smaller in size as outlined by the Six1 expression (Figure 4A). We previously reported that although Eya1 mRNA is expressed in Six2−/− MM progenitors, Eya1 protein is localized in the cytoplasm of Six2−/− MM cells and its nuclear translocation depends on Six2 activity (Xu et al., 2014). In Eya1Six1/+;Six2−/− embryos, the Eya1+ domain was comparable to that seen in Six2−/− littermates, both of which were smaller in size than in control littermates (Figure 4B). Notably, however, anti-Eya1 immunostaining revealed nuclear localization of Eya1 in Eya1Six1/+;Six2−/− MM progenitors, in contrast to its cytoplasmic localization in Six2−/− MM cells (Figure 4C). Thus, Six1 is also able to mediate nuclear translocation of

**FIGURE 3** | Six1 fails to rescue defective progenitor cell proliferation and cell death associated with Six2-deficiency. (A,C) TUNEL labeling on kidney sections at E11.5 showing increased apoptosis in the MM of Eya1Six1/+;Six2−/− and Six2−/− mutants (arrows in A), and quantification of TUNEL+ cells in the MM (C). Note that increased apoptosis was also observed in the UB of Six2−/− embryos. (B,D) Anti-PH3 (Phosphohistone H3) staining showing reduced PH3+ mitotic cells in the MM of Eya1Six1/+;Six2−/− and more reduced in Six2−/− mutant than in wild-type or Eya1Six1/+ controls (arrows in B) and quantification of PH3+ mitotic MM cells (D). Scale bars: 100 μm. For quantification, TUNEL+ or PH3+ MM progenitor cells were counted from 28 to 35 sections of 5 different kidneys and values represent average number of TUNEL+ or PH3+ cells (±standard deviations) per section (6 μm). Data from Eya1Six1+/+, Eya1Six1/+;Six2−/− or Six2−/− were compared with wild-type. ***p-value < 0.001 and **p-value < 0.01.
Eya1 in the progenitors in the absence of Six2. This observation is consistent with the previous finding that Six1 or Six2 but not Six3 can mediate nuclear translocation of Eya1 when coexpressed in HEK293 cells (Buller et al., 2001).

We further examined whether Eya1 expression can be maintained in the MM progenitors of Eya1Six1/+;Six2−/− embryos at later stages. As shown by whole-mount ISH (Figure 5A), Eya1 expression was strongly detected in the MM progenitors of wild-type or Eya1Six1/+ littermate kidneys at E12.5, but only residual Eya1 transcripts were observed in Eya1Six1/+;Six2−/− kidneys. Consistent with our previous observation (Xu et al., 2014), anti-Eya1 immunostaining revealed higher levels of Eya1 protein expression in the CM and lower levels in the differentiating RVs of wild-type or Eya1Six1/+ kidneys (Figure 5B). Co-immunostaining with a Six1-specific antibody confirmed co-expression of Six1 with Eya1 in Eya1Six1/+ but not in wild-type kidneys (Figure 5B). However, in Eya1Six1/+;Six2−/− kidneys, neither Eya1 nor Six1 protein expression was observed. Thus, Six1 cannot replace Six2 to maintain Eya1 expression in the MM progenitors in Eya1Six1/+;Six2−/− at E12.5.

Genome-wide ChIP-seq analysis identifies selective binding of Six1 to a subset of Six2 targets essential for nephron progenitor cell maintenance but no co-occupancy of Six1 with Six2 to genes involved in nephron differentiation such as Wnt4, Fgf8, and Pax8.

Our data suggest that Six1 cannot drive developmental programs regulated by Six2 to expand and maintain the nephron progenitors, thus implying that Six1 cannot target Six2-occupied sites in the nephron progenitors. In the human kidney, while ChIP-seq identified more SIX2 binding sites than SIX1 (~81% (1307 of 1610 peaks) of SIX1 peaks overlapped with SIX2 peaks (~20.8% of 6276 SIX2 peaks) (O’Brien et al., 2016). However, the functional cooperation and relative contributions of SIX1 versus SIX2 to progenitor cell maintenance and differentiation remain unknown. To directly address the potential functional differences between the mouse Six1 and Six2, we performed ChIP-seq to map Six1-bound regions genome-wide in Eya1Six1/+ kidneys. Immunohistochemistry with a Six1-specific antibody confirmed Six1 protein expression in the CM of Eya1 Six1/+ at E16.5 (Figure 6A), indicating the stability of the overexpressed Six1 protein in the nephron progenitors. We then used kidneys ~ E13.5 for Six1 ChIP-seq to avoid differentiating structures, and the peak bed files were generated from MACS peak calling against both genomic input DNA and IgG ChIP-seq controls with the default setting (10−5 cutoff). Peak calling identified 148 and 2527 Six1 peaks respectively from the two different datasets, and we focused on the 2527 peaks (2206 genes) for subsequent analyses and comparison with our recent Six2 ChIP-seq data sets on wild-type kidneys at the same developmental stage (Li et al., 2021)
Only 190 Six1 regions (~7.5%) overlapped with Six2 peaks (Supplementary Table S3), but ~41% of putative Six1 target genes (906 out of 2206 genes) were shared with Six2 (Supplementary Table S4). While a significant fraction of Six1 peaks were located near the promoter, ~29% were found between 5–500 kb of TSSs (transcriptional start sites) and the majority of these peaks were distributed within intronic and intergenic regions (Figure 6D). Homer de novo motif search revealed that Six1 or Six2 motifs were not among the top mostly enriched motifs in the Six1 peaks (Figure 6E). Motifs for transcription regulators RELB, Stat3, Tcf21, Osr2, and Smad4 are among the top five enriched motifs. Gene Ontology (GO) analyses using GREAT did not identify nephron-specific terms, but revealed overrepresentation of genes associated with chromatin modification, histone modification, and transition of mitotic cell cycle (Figure 6F).

Eya1 is a critical factor for the maintenance of the nephron progenitors and interacts with Myc and Six2 (Xu et al., 2014), but Six2 is likely involved in maintaining high levels of Eya1 expression as Six2 binding to multiple regions at the Eya1 locus were identified in E13.5 and E16.5 kidneys (Park et al., 2012; O’Brien et al., 2016; Li et al., 2021). Our data suggest that Six1 cannot replace Six2 to maintain Eya1 expression in the progenitors (Figure 5), we therefore asked if Six1 can target the Six2-bound regions at the Eya1 locus to replace the Six2 function (Supplementary Table S2). Among the 190 regions shared between Six1 and Six2, Six1 enrichment was identified at ~112-kb upstream of the Six2 promoter (Figure 6G), representing one of the several Six2 targeted sites with the active histone mark H3K27ac deposition. Similarly, Six1 enrichment was identified at a distal region ~38-kb downstream of the Mycn promotor (Figure 7A), which is one of the several Six2-bound sites at the Mycn locus and is a highly conserved region associated with H3K27ac-deposition. Thus, Six1 appears to selectively target a small subset of Six2-bound sites at genes essential for nephron progenitor maintenance, which may explain why the number of mitotic MM progenitor cells was more reduced in Six2−/− than in Eya1Six1/+;Six2−/− (Figure 3D). The lack of Six1 co-occupancy at Six2-target sites of the Eya1 locus may at least partially lead to the disappearance of Eya1 expression in Eya1Six1/+;Six2−/−.

Previous studies have suggested a role for Six2 in inducing nephron differentiation as Six2 targets regions associated with H3K27ac-deposition at genes essential for nephron differentiation, such as Fgf8, Pax8, and Wnt4 (Park et al., 2012; O’Brien et al., 2016). In contrast, Six1 binding to these putative Six2 target genes was not observed (Supplementary Table S2). Interestingly, however, co-occupancy of Six1 with Six2 was identified at the promotor regions of the enhancer of zeste homolog Ezh1 and Ezh2, which had H3K27ac-deposition (Figure 7A). Ezh1 and Ezh2 are polycomb histone methyltransferases, and both are required for the renewal potential of nephron progenitors and co-regulate chromatin accessibility (Liu et al., 2020). Since inactivation of both Ezh1 and Ezh2 triggers unscheduled activation of...
Wnt4-driven differentiation and results in early termination of nephrogenesis (Liu et al., 2020), Six1 binding to these two genes could maintain the expression of both Ezh1 and Ezh2 in the absence of Six2, which in turn prevents ectopic activation of Wnt4 in the nephron progenitors in Eya1Six1+/−;Six2−/− kidneys.

We further compared with human SIX1 ChIP-seq data in the fetal kidneys and found only 58 (~2.3%) of mouse Six1 peaks overlapped with human SIX1-binding sites and ~27% of putative SIX1 target genes shared with human SIX1 (Figure 7B, Supplementary Table S5, S6), despite the high conservation between Six1 and SIX1. Previous studies
identified only ~8% of mouse Six2-binding sites but ~50% of putative Six2 target genes overlapped with human SIX2 targets (O’Brien et al., 2016). Sequence comparison between these proteins revealed that the C-terminal domain is quite divergent in length between Six1 and Six2 with a low degree of sequence similarity, and the C-terminus of human SIX2 is shorter than the mouse Six2 (Figure 7C). This structural difference may affect the stability of forming complexes with specific partner proteins, thus resulting in Six1 selectively binding to only a small subset of Six2 target sites.
DISCUSSION

Six1 and Six2 genes are expressed in the MM progenitors during mouse kidney development, but Six1 is expressed before Six2 transcription and overlaps with Six2 only transiently before the onset of UB branching. Six1−/− embryos fail to form a functional MM competent for inducing UB branching, whereas Six2−/− mice exhibit later defects associated with expansion and maintenance of the nephron progenitors. Previous studies found that Six1 also cooperates with Six4 to regulate the formation of the MM as this structure is not formed in Six1−/−;Six4−/− mice (Kobayashi et al., 2007; Xu and Xu 2015), while Six4−/− mice are viable and normal (Ozaki et al., 2001). In this study, we overexpressed Six1 in the MM progenitors under Eya1 regulatory transcriptional control and examined the functional interchangeability between Six1 and Six2. Our data show that Six1 cannot fully rescue the Six2-deficient kidney phenotype, thus indicating that the different physiological roles of Six1 and Six2 relate to differences in both transcriptional regulations of the genes and divergent biochemical properties of the proteins.

From the sequence alignment of Six1 and Six2, it is evident that these two proteins share the highly conserved SD and HD regions (Figure 7C), except for the C-terminal domains that differ in length with a low degree of similarity. Since the HDs have very low intrinsic sequence specificity (Treisman et al., 1992), such structural differences adjacent to the HDs are likely to modify the specificity of the proteins by influencing the stability of complexes formed with particular partner proteins, thus affecting the ability of Six1 to compensate for Six2 loss. Based on our data, Six2 appears to have distinct and non-interchangeable roles in the multipotent nephron progenitors. Although Six1 can mediate nuclear translocation of Eya1 in the absence of Six2 (Figure 4), it apparently cannot act as a substitute for Six2 in maintaining Eya1 expression and driving the expansion and survival of the progenitors. This could be due to distinct stabilities of transcriptional complexes formed with partner proteins rather than differences in DNA-binding site preference, which explains why Six1 only selectively targets a small subset of Six2-bound regions and why Six1 cannot bind to Six2-target sites at the loci of Eya1 and Mycn. Consistent with this view, our previous studies testing the efficiency of co-expression of different members of the Eya family with various members of the SIX protein family in inducing hair cell development found that the combination of Eya1/Six1 acts most efficiently to activate a specific downstream regulatory program controlling inner ear hair cell or spiral neuron induction in a cochlear explant system (Ahmed et al., 2012a; Ahmed et al., 2012b). We found that ~90% of Eya1/Six1-cotransfected cochlear nonsensory epithelial cells became Myo7a+ hair cells and that this function of Eya1 or Six1 is non-interchangeable with Eya2 or Six2, as only ~3% of Eya1/Six2- or ~6% of Eya2/Six1-cotransfected cochlear nonsensory epithelial cells became Myo7a+ hair cells (Ahmed et al., 2012a). Thus, we speculate that the divergent C-terminus of Six1 and Six2 may confer differential specificity in transcriptional complex formation and subsequent DNA recognition. Six1 may fail to collaborate with Six2-interacting transcription factors or chromatin regulators to generate a permissive chromatin context in the multipotent progenitors necessary for their renewal and survival. Although we do not understand why the mouse Six1 does not bind to many Six2 target sites that are co-occupied by human SIX1/ SIX2, the simplest explanation is that there may be species-specific factors that affect the ability of Six1/SIX1 to target Six2/SIX2 sites. This may also explain why only ~8% of mouse Six2-binding sites overlapped with human SIX2 peaks (O’Brien et al., 2016). Interestingly, while conserved, the C-terminus of Six2 also differs in length between the two species (Figure 7C). Human SIX2 is five amino acid residues shorter than mouse Six2, shortening the length difference between human SIX1 and SIX2. Thus, a comparative study with different SIX family proteins in transactivating nephron progenitor-specific target genes may give useful information about the importance of the C-terminal region.

Previous studies indicated that Six2 is likely to induce nephron commitment and differentiation by regulating the expression of target genes involved in nephron differentiation, such as Fgf8, Pax8, and Wnt4 (Park et al., 2012). Although Six2−/− nephron progenitors undergo premature differentiation, it is still unclear how Six2 acts to counter nephron differentiation. In contrast, Six1 does not appear to have an equivalent function as Six2 in inducing differentiation, because Six1 binding to the Six2 target sites at the Fgf8, Pax8 or Wnt4 was not detected. Our observation that the rapid commitment to an RV fate occurring in Six2−/− did not happen in Eya1Six1+/−;Six2−/− suggests that Six1 inhibits premature nephron differentiation in the absence of Six2. We thus speculate that the nuclear Eya1 in Eya1Six1+/−; Six2−/− may play a role in countering nephron progenitor cell differentiation. Alternatively, Six1 may indirectly inhibit differentiation through the regulation of other factors. In supporting this, Six1 enrichment was detected at the Ezh1 and Ezh2 promoter regions that are associated with H3K27ac-deposition and co-occupied by Six2 (Figure 7). Since previous studies have found that inactivation of both Ezh1/Ezh2 leads to unscheduled activation of Wnt4 (Liu et al., 2020), Six1 may regulate the expression of these two genes to prevent ectopic Wnt4 activation. These two possibilities could in conjunction explain the requirement of Six1 for preventing rapid epithelialization of the progenitors in Eya1Six1+/−;Six2−/−. This also suggests that Six1 is able to form transcriptional complexes capable of associating with the same regulatory DNA elements co-occupied by Six2 at the Ezh1 and Ezh2. Thus, our finding provides insights into why nephron progenitors undergo rapid RV formation in the absence of Six2.

In summary, our results indicate that the Six1 and Six2 genes of the Drosophila so subfamily have diverged in function by the acquisition of different regulatory patterns and special biochemical properties. Previous studies have focused on the conserved SD and HD in recognizing subgroup specific sites and common targets. In human fetal kidneys, SIX1 and
SIX2 cross-regulate each other (O’Brien et al., 2016), which is thought to be critical to maintain the absolute levels of SIX proteins that are necessary for generating approximately seventy times more nephrons in a human kidney than a mouse kidney. However, SIX2 apparently has a more prominent roles than SIX1 and ~79.2% of SIX2 target sites are not targeted by SIX1. As SIX1 and SIX2 physically interact when coexpressed in HEK293 cells (O’Brien et al., 2016), it is possible that small percentage of SIX1 and SIX2 may form complexes that may be below the detection threshold by co-immunoprecipitation. Nonetheless, our evidence for the functional difference between the mouse Six1 and Six2 suggests that the evolutionary division of these two genes has occurred at the levels of both protein function and their gene-expression programs, with particular cell types manifesting greater dependence on the expression of one or the other or both. Future research to explore whether the divergent C-terminal domains influence the molecular and biochemical rules governing the specificity of these proteins may help us understand how the mechanisms that control spatial aspects of nephrogenesis have been modified during evolution. Lastly, it is worth mentioning that a recent study explored the potential of SIX1 with EYA1 and SNAI2 in directly reprogramming HK2 cells into nephron progenitor-like cells (Vanslambrouck et al., 2019). Our finding that Six1 and Six2 have non-equivalent biochemical properties provides valuable information for the development of nephron reprogramming strategies.

DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found below: NCBI with GSE ID is GSE189131 (https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE189131).

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ETHICS STATEMENT

The animal study was reviewed and approved by All animal protocols were approved by Animal Care and Use Committee of the Icahn School of Medicine at Mount Sinai (protocol #06-0822).

AUTHOR CONTRIBUTIONS

JX and JL contributed to formal analysis and data validation. AR contributed to sequencing data processing and analysis. HY contributed to analysis and figure making. P-XX contributed to the conception and design of the study, data analysis, and funding acquisition. JX and P-XX wrote the first draft of the manuscript. All authors contributed to manuscript revision, read and approved the submitted version.

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SUPPLEMENTARY MATERIAL

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