Basic Helix-Loop-Helix Transcription Factor TCF21 Is a Downstream Target of the Male Sex Determining Gene SRY

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Abstract

The cascade of molecular events involved in mammalian sex determination has been shown to involve the SRY gene, but specific downstream events have eluded researchers for decades. The current study identifies one of the first direct downstream targets of the male sex determining factor SRY as the basic-helix-loop-helix (bHLH) transcription factor TCF21. SRY was found to bind to the Tcf21 promoter and activate gene expression. Mutagenesis of SRY/Sox9 response elements in the Tcf21 promoter eliminated the actions of SRY. SRY was found to directly associate with the Tcf21 promoter SRY/Sox9 response elements in vivo during fetal rat testis development. TCF21 was found to promote an in vitro sex reversal of embryonic ovarian cells to induce precursor Sertoli cell differentiation. TCF21 and SRY had similar effects on the in vitro sex reversal gonadal cell transcriptomes. Therefore, SRY acts directly on the Tcf21 promoter to in part initiate a cascade of events associated with Sertoli cell differentiation and embryonic testis development.

Introduction

Mammalian embryos have bipotential gonads which can develop into either a testis or an ovary depending on paternal and maternal chromosomal contributions. Paternal transmission of a Y chromosome triggers testicular differentiation, whereas the contribution of paternal X chromosome results in ovarian differentiation. The bipotential gonad contains a pool of undifferentiated somatic cell precursors that during development differentiate into either testicular (Sertoli, Leydig, and peritubular myoid) or ovarian (theca and granulosa) somatic cells in response to extrinsic and intrinsic factors. The gene for the transcription factor Sry is located on the Y chromosome and is sufficient to induce testis fate in the bipotential gonad [1,2,3]. The nuclear orphan receptor steroidogenic factor 1 SF1 and SRY cooperatively upregulate an immediate downstream gene Sox9 through an interaction on its testis-specific enhancer [4]. Subsequently, Sox9 regulates the differentiation of precursor cells into the Sertoli cells. Sox9 expressed by Sertoli cells cooperates with Fgf9 and prostaglandins [5,6] to promote testis development. Somatic cells undergo dramatic molecular interactions during gonadal sex differentiation to ultimately influence the differentiation of germ cells. Several sex-specific genes have been shown to interact during the differentiation of gonads. For example, SRY, Sox9, Fgf9, and prostaglandins have an important role in the differentiation of Sertoli cells and testis. In contrast, Rspo1, Wnt4, beta-catenin, and Foxl2 antagonize Sox9 and Fgf9 to promote ovarian differentiation [4,7,8]. Despite extensive efforts made to uncover the molecular mechanisms underlying sex determination in mammals, the downstream target genes of SRY remain poorly understood.

The process of sex determination starts with differentiation of somatic cells. Since SRY binds to HMG box sequences on the DNA, the cell fate associated genes containing HMG box binding sites in their promoter are considered potential downstream target candidates. Numerous cell type-restricted basic helix-loop-helix (bHLH) transcription factors have been identified and served as target candidates. Numerous cell type-restricted basic helix-loop-helix (bHLH) transcription factors have been identified and shown to control cell fate specification, differentiation and morphogenesis during development [9,10,11]. Some bHLH genes function early in embryonic development and influence a wide variety of tissues, whereas others act later in development and are required in the adult cells to maintain cellular differentiation. In our previous microarray analysis, a bHLH gene Tcf21 (also called, bHHLH23, Pod1, capsulin, epicardin) was found to be highly expressed in the gonads coinciding with the expression of SRY and Sox9 [12]. A genome wide phylogenetic classification of the entire bHLH gene family has recently developed a uniform nomenclature with Tcf21 named as bHHLH23 and put it in perspective to other species [11,13]. Tcf21 null mutants exhibit male-to-female sex reversal with some germ cells entering meiosis [14]. An analysis of human, mice and rat Tcf21 promoters showed at least three global SRY-binding sites within a 2 kb promoter region. Therefore we hypothesize that SRY interacts with Tcf21 to promote somatic cell differentiation during male gonadal sex determination. Observations demonstrate Tcf21 is a downstream target gene for SRY action and promotes Sertoli cell development.
Results

Expression of Tcf21 gene and localization of the TCF21 protein in the rat embryonic testis

Tcf21 expression has been detected quite early in development in the embryonic mesoderm, showing a similar pattern to Wnt1 and Gata4 [15]. In a microarray analysis with rat gonadal RNA [12], relatively higher levels of Tcf21 transcripts were found at the embryonic day 13 (E13) stage of testis development, after which levels are maintained during this stage of development (Fig. 1). At embryonic day 13, which is equivalent to the 13 to 18 tail somites stage, TCF21 was localized to somatic cells adjacent to germ cells (Fig. 2A). At E14, TCF21 immunopositive cells were found in the Sertoli cells and at E16 in the interstitium including Leydig cells (Fig. 2, B&D). As a positive control, AMH was used and localized to Sertoli cells (Figure 2C). The differential localization of TCF21 protein at different times of sex determination suggests differential functions at different periods of testis development. Expression patterns of TCF21 in the E13 and E14 testes indicate a transient role in Sertoli cell differentiation, whereas expression at E16 in the interstitium suggests a role during later Leydig cell differentiation [14,16].

Characterization of the rat Tcf21 promoter

Analysis of the Tgf21 promoter revealed three putative SRY binding sites [17] in the 2 kb upstream region from the transcription start site. A comparison was made of the genomic region of the rat Tgf21 promoter with that of mouse and human (Fig. 3A). SRY binding sites are conserved among human, mouse and rat. There are multiple overlapping putative SRY binding sites in mouse and rat promoters. Most of the SRY binding sites are paired and separated by one or two nucleotides. In addition, the rat Tgf21 promoter contained a binding site for SRY/SOX9 at a −1.3 k upstream region and GATA 4 in multiple locations. Multiple E-box binding sites are present within the −2 kb promoter regions.

SRY stimulates Tcf21 promoter activity in vitro

Primary cell cultures were derived from E13 testes, equivalent to 12–18 tail somite stage of male rat embryos, and used for promoter reporter assays. Binding sites for SRY and SOX9 on the Tcf21 promoter were investigated with site specific mutagenesis for their ability to regulate the Tcf21 promoter. E13 cultures derived from testicular primary cells or postnatal 20-day-old (P20) primary Sertoli cells were co-transfected with an Sry expression plasmid together with different Tcf21 promoter fragments in a luciferase reporter system. Similar results were obtained from both E13 testis cultures or purified Sertoli cell cultures. The longer promoter constructs had lower luciferase activity (Figure 3B). Interestingly, over-expression of rat SRY (Figure 3D), mouse SRY, mouse SOX9 and human SRY (data not shown) resulted in an increased activation of the Tcf21 promoter (Figure 3D). Point mutations of SRY motifs significantly perturbed promoter activity (Fig. 3C and D). Mutation of one of the three SRY binding sites did not result in significant change in promoter activity, while mutation of two binding sites significantly decreased the activity of the Tcf21 promoter. Observations suggest the requirement for having at least two SRY binding sites for activating the promoter in vitro. Overexpression of SRY increased the activity of the Tcf21 promoter having paired SRY binding sites (Fig. 3D).

SRY recombinant protein binds Tcf21 promoter in vitro

The possibility that SRY binds to the defined SRY binding sites on the fragment of the Tgf21 promoter was investigated. Reactions were set up to bind HA tagged SRY-HA recombinant protein to the Tgf21 promoter oligonucleotides with or without mutated SRY binding sites followed by immunoprecipitation (IP) with anti-HA antibody. Enriched fragments were analyzed by PCR with primers flanking the binding sites (Fig. 4A). Affinity purified non-immune mouse IgG was used as a negative control in the experiment. Anti-HA antibody pulled down the oligonucleotide with intact SRY binding sites, but not the one with mutated SRY binding sites (Fig. 4A).

The specific SRY antibody developed by Dr. Peter Koopman’s laboratory, Queensland University, Brisbane, Australia was shown to bind SRY, but not SOX9 [6]. In order to validate a commercially available anti-SRY (Santa Cruz, Santa Cruz CA) antibody for ChIP assay, in vitro immunoprecipitation was performed using recombinant SRY and SOX9 proteins. Enrichment of target protein was detected by western blot using antibodies against the HA tags for the recombinant proteins. The commercial SRY antibody did pull down SRY, but did not pull down SOX9 recombinant protein (Figure 4B). The SOX9 antibody recognized both SRY and SOX9 recombinant proteins (Figure 4B). Therefore, the SRY antibody was specific for SRY and thus used in the ChIP assay.

SRY binds in vivo to the Tcf21 promoter during male sex determination

There has been little progress toward identifying in vivo downstream targets of SRY. This is in part due to technical limitations of procedures such as in vivo ChIP assay which traditionally requires a large amount of starting material from the embryonic tests. For example, a conventional ChIP assay requires a large amount of chromatin which is equivalent to pooled gonads of five to six hundred embryos [6,18]. The techniques used in the present study overcome these technical difficulties. The present study used a modified in vivo ChIP assay with isolated testis samples from 20 to 25 testes at the 12 to 18 tail somite stage of rat embryo development. The promoters of any
genes that SRY directly regulates at the time of gonadal sex determination should be pulled down from the chromatin mixture by anti-SRY antibody. To test whether SRY associates with a fragment of Tcf21 promoter, PCR primers were designed to amplify the Tcf21 promoter regions flanking an SRY binding site. As a negative control, primers were designed to a promoter region which did not contain any HMG box sites or putative SRY binding sites. The SRY antibody obtained from Dr. Peter Koopman, Brisbane, Australia, pulled down the fragment of Tcf21 promoter containing the predicted SRY binding site, but not the negative control region without its binding site (Fig. 4C). As a positive control in the ChIP assay, anti GATA4 antibody was

Figure 2. Immunolocalization of TCF21 protein in embryonic testis. (A) TCF21 protein in somatic cells of the E13 testis (arrows). (B) Localization of TCF21 protein in E14 testis. Insert shows magnified section of E14 testis. (C) AMH localization in E14 testis. AMH was stained with DAB, so brown color represents Sertoli cells. (D) TCF21 protein in E16 testis (green). (E) AMH localization in E16 testis. Abbreviations: GC = Germ Cells. SC = Sertoli Cells. LC = Interstitial cells. Dotted circles indicate testis cord structures. Data are representative of a minimum of three different experiments.
doi:10.1371/journal.pone.0019935.g002
Figure 3. Tcf21 promoter mutagenesis analysis. (A) Analysis of mouse, rat and human Tcf21 promoter and SRY binding sites. Gonadal (testis) cell culture and transfection assays with relative luciferase activity compared to wild type (WT) promoter presented. Bars with asterisks are significantly different from −0.5 kb control promoter. (B) Promoter activity decreased with the length of the promoter. (C) Deletion of SRY binding motif and relative promoter activity. (D) Over-expression of SRY in E13 male primary cells increase overall luciferase activity of the mutant promoter, except for the mutant which lacks first uncoupled TAACAAA site (gray response element). Data are presented as the mean ± SEM from a minimum of three different experiments performed in replicate. The asterisks indicate (*) p<0.05, (**) p<0.01, (***) p<0.001 with a Student’s t-test.
doi:10.1371/journal.pone.0019935.g003
Figure 4. SRY in vitro and in vivo binding to Tcf21. (A) Immunoprecipitation-based in vitro pulldown assay was designed to test the physical binding of SRY protein with Tcf21 promoter SRY/SOX9 response element oligonucleotide. Recombinant SRY-HA tagged protein was bound in vitro to 250 bp Tcf21 promoter fragment containing three SRY binding sites. An anti-HA antibody immunoprecipitated a fragment of Tcf21 promoter containing all three SRY-binding sites, whereas mutation of the bind site eliminated SRY binding. (B) Antibody specificity for validating in vivo ChIP assay. Recombinant SRY and SOX9 immunoprecipitation with anti-SRY (Santa Cruz, CA) and anti SOX9 (Koopman) antibodies followed by western blot with anti-HA antibody which was attached as a tag to those recombinant proteins. The SRY antibody specifically pulled down SRY, whereas SOX9 antibody recognized both SRY and SOX9. (C and D) In vivo carrier ChIP assay was performed using chromatin from the embryonic testis from E13 male embryos (14–18 tail somite stage). The anti-SRY antibody (Koopman) (C) and commercial anti-SRY antibody (Santa Cruz, CA) (D) immunoprecipitated a fragment of Tcf21 promoter containing SRY binding sites, in vivo. Enriched DNA was amplified by primers flanking SRY binding site and for non-specific enrichment PCR primers were designed between 0.7 and 0.9 kb promoter which did not contain any consensus binding site for HMG box proteins (no Sry site). A control non-immune IgG (IgG) had no precipitation, while a positive control input DNA (Input) did have Tcf21 PCR product. A positive control GATA4 antibody (GATA4) precipitation also pulled down a GATA4 site in Tcf21 promoter. (E) Over-expression of TCF21 induced AMH expression in the E13 primary cells derived from embryonic ovaries. The TCF21 or SRY over-expression (TCF21) or (Sry) indicated, along with controls (No Primer, No DNA, Water and Empty Plasmid), with positive PCR control of L19 gene (Positive). Data is representative of a minimum of three different experiments.

doi:10.1371/journal.pone.0019935.g004
used. Three GATA4 response element sites are present at different regions of the Tcf21 promoter and were used as a positive control. A ChIP assay for GATA4 was found to immunoprecipitate and identify the in vitro Tcf21 promoter (Fig. 4C). The anti SRY from Dr. Peter Koopman was previously shown to be specific for mouse SRY and was found to pull down the Tcf21 promoter from developing rat E13 gonads (Fig. 4C). A commercially available anti SRY antibody (Santa Cruz, Santa Cruz, CA) was found to be specific for SRY (Fig. 4B) and also pulled down the Tcf21 promoter in the ChIP assay from developing E13 testis (Fig. 4D).

Therefore, two different anti SRY antibodies in the ChIP assay demonstrated that SRY directly binds the Tcf21 promoter during male gonadal sex determination. Positive controls such as TESCO which is considered a downstream target for SRY [4], has not been characterized in the rat, so current study did not focus on TESCO as a positive control. Instead we have initiated genome-wide ChIP-Chip analysis of SRY targets in embryonic testis during sex determination.

**TCF21 promotes in vitro sex reversal of embryonic ovarian cells to induce Sertoli differentiation**

The above in vitro and in vivo results demonstrate that SRY interacts with the Tcf21 promoter during male sex determination and testis differentiation. Tcf21 expression level was similar to that of Wnt and Gata4 at E9.5 and E10.5 [15], whereas during male sex determination transcripts for this gene remained elevated in the rat. The homozygous knockout mutant for Tcf21 showed abnormal gonadogenesis, lost testis cord structures, abnormal vascular innervations into the testis, and reduced Sox9 expression in the testis [14]. These results suggest that Tcf21 is involved in male sex determination and testis differentiation. An embryonic E13 ovary culture system was used to test whether Tcf21 can promote in vitro sex reversal and the induction of the expression of a male marker gene of testis differentiation. Expression plasmids for Tcf21 or Sry were over-expressed in subcultures (less than 12) of primary cell cultures derived from E13 rat ovaries. The objective was to promote an in vitro sex reversal from an ovarian somatic cell population to a testicular somatic cell population. Tcf21 over-expression induced the expression of anti Mullerian hormone (AMH) which is a specific marker of Sertoli cell differentiation in the embryonic testis, but not expressed by the embryonic ovary or ovarian culture, Figure 4E. As expected, SRY over-expression in the embryonic ovarian cells also induced AMH expression (Figure 4E). This observation suggests that Tcf21 promotes embryonic Sertoli cell differentiation in a similar manner as SRY.

The cell culture transcriptomes were examined to assess the in vitro sex reversal of the embryonic ovarian cell culture to a testis cell differentiated state on a genome wide level. Microarray analysis of the embryonic cell cultures were investigated in the absence or presence of SRY, Tcf21 or Tcf21 with its potential binding partner Tcf12/Reb-alpha. All bHLH transcription factors dimerize and due to the high level of Tcf12 expression in the embryonic testis (Figure 1), the combined actions of Tcf21 and Tcf12 were also assessed. Tcf12 is also known as Reb-alpha which is somewhat ubiquitously expressed and a binding partner of numerous bHLH factors [19]. Over-expression of SRY in the embryonic ovary cell culture promoted a transcriptome more similar to the testis with approximately 800 significantly increased transcripts, Figure 5A. Interestingly, Tcf21 also promoted a transcriptome with approximately 20% overlap with that induced by Sry, Figure 5A. The combined actions of Tcf21 and Tcf12 induced a more dramatic increase in AMH expression, data not shown, and a transcriptome with overlap with Tcf21 and Sry, Figure 5. Observations support a potential role for Tcf12 as a potential partner for Tcf21. A list of the genes with the most highly increased expression induced by SRY and Tcf21 are presented in Table S1. An extended list of the genes altered after Tcf21, SRY or Tcf21 and Tcf12 over-expression are presented in Table S2. Observations demonstrate Tcf21 and Tcf12 promote some similarities in the in vitro sex reversal transition in the transcriptome as SRY, supporting a role of Tcf21 as a downstream target of Sry during male gonadal sex determination.

**Discussion**

Since the discovery of SRY as a master male sex determining factor several efforts have been made to identify its downstream target genes. Whether Sox9 is the only downstream gene that carries out all functions necessary for initiating male sex development or whether there are multiple SRY target genes that cooperatively act to induce testis-specific somatic cell differentiation remains to be determined. Recent developments suggest that SRY and SF1 cooperatively activate Sox9 expression through interactions with its enhancer region, confirming Sox9 as one of the SRY target genes [4] (Figure 6). The SRY protein has been shown to interact in vivo with the promoter enhancer region of the Cerebellin 4 precursor gene (Cbln4), which is present in Sertoli cells during early embryonic development [18] (Figure 6). Although, the exact function of Cbln4 gene in testis differentiation is not clearly understood yet, this gene has been suggested as one of the downstream target genes of SRY. Using a similar approach to that taken in the current study we have found SRY directly interacts and regulates the promoter for neurotrophin 3 (Ntf3) [20], that has previously been suggested to have a critical role in testis cord formation [21,22] (Figure 6). A major hindrance in finding SRY target genes is the technical limitation to in vivo chromatin immunoprecipitation (ChIP) methods. Using an in vivo carrier ChIP method the current study demonstrates that SRY interacts with the bHLH family transcription factor Tcf21 during male sex determination. Observations provide insights into the mechanism of SRY regulation of Sertoli cell differentiation and the factors regulating (e.g. Tcf21) cell differentiation during male gonadal sex determination.

Numerous cell type-restricted basic helix-loop-helix (bHLH) transcription factors have been identified and shown to control cell fate specification, differentiation and morphogenesis during embryonic development, while others act later in development and are required in the adult cells to maintain cellular differentiation. Examples include Mesp in heart cell differentiation, MYOD in muscle differentiation, and Neurogenin in neuron differentiation [23,24,25]. In an embryonic rat and mouse testis microarray Tcf21 was found as one of the most highly expressed transcription factor genes [12], Figure 1. The presence of Tcf21 protein in the somatic cells of the embryonic testis was confirmed immunohistochemically. Nuclear localization of Tcf21 was observed in the Sertoli cell in testis of the E13 and E14 testis, whereas it appeared in the interstitial cells in E16 testis suggesting it has multiple functions during testis differentiation. Prior to male sex determination Tcf21 mRNA is localized in the gonadal ridges and its expression pattern was found to be similar to Wnt1 and Gata4 [15]. Gene knockout studies suggested that Tcf21 null mutant male mice were partially sex-reversed [14]. Combined observations suggest Tcf21 has multiple roles in male sex determination and testis morphogenesis.

Most of the genes that are highly expressed during the period of male sex determination have SRY or SOX9 binding sites in their promoters [17]. Since the Tcf21 promoter harbors three SRY
binding sites in the −2 Kb promoter region, it was hypothesized that SRY may bind these response elements and regulate the activity of the core promoter. Over-expression of SRY in the primary embryonic testis cell culture increased the Tcf21 promoter activity. This Tcf21 promoter activity was significantly reduced when all three SRY binding sites were mutated. SRY seemed to be able to activate the Tcf21 promoter if any two of the three binding sites were intact. An oligonucleotide pull-down assay was used to confirm SRY directly binds to the Tcf21 promoter SRY binding sites (Figure 4A). Although these in vitro promoter actions and binding studies support a potential role for SRY regulation of Tcf21, SOX9 can have similar activities and a direct role of SRY is uncertain in vivo. An example of a gene that has similar in vitro observations, but was found in vivo to primarily respond to SOX9 is prostaglandin synthase (PGDS) [6] (Figure 6). To investigate the potential direct SRY binding to the Tcf21 promoter in vivo a ChIP assay with chromatin samples from embryonic testis undergoing sex determination and cellular differentiation was performed. The major limitation in running ChIP assays with embryonic gonad samples is the amount of chromatin required. This limitation has hindered efforts made to find downstream target genes of SRY. The current study adopted the ChIP protocol developed by O’Neill et al (2006) [26] and modified to meet conditions with rat SRY protein. Observations demonstrate a ChIP with SRY antibody immunoprecipitated from embryonic testes the Tcf21 gene promoter. Tcf21 appears to be one of the downstream target genes of SRY. To identify other SRY target genes an SRY ChIP is currently being subjected to ChIP-Chip whole genome promoter tiling arrays.

**Figure 5. Transcriptome analysis of SRY and TCF21 actions.** Venn diagram (A) with overlap of total gene sets (p<0.05) altered in response to various expression constraints (Sry, Tcf21, Tcf21 plus Tcf12, Tcf12). Total genes in the gene sets and overlap numbers are presented. These gene sets are statistically significant increase genes, but with no further cut-off parameters. (B) Restricted gene sets overlapped in female E13 cultures involving gene sets with altered expression with >1.2 fold change, p<0.05, and mean difference >10.

doi:10.1371/journal.pone.0019935.g005
(Figure 4E). *Amh* is not expressed in the ovary at this time of development, but Sertoli cells in the embryonic testis do express *Amh*. In contrast, *Tcf21* null mutants did not completely lack SOX9 and AMH expression because the loss of TCF21 was likely compensated for by yet undefined tissue specific binding partners. This phenomenon has been seen for heart muscle differentiation in *Mesp1* mutants [23,27] and prostate differentiation in *Ncoa1* mutants [28]. The over-expression of *Tcf21* in the fetal ovarian cell culture promoted an SRY-like transcriptome and cellular differentiation similar to normal testis development. SRY and TCF21 both induced similarities in the transcriptomes and *in vitro* gonadal sex reversal in the ovarian cell culture. The group of specific genes identified will be useful to consider in further elucidation of the cascade of gene expression in testis differentiation. In addition to the identification of the *in vitro* and *in vivo* association of SRY with the *Tcf21* promoter, a functional role of TCF21 in promoting *in vitro* sex reversal and a SRY-like gene expression profile was observed.

A distinct function of TCF21 in testis differentiation is associated with fetal Leydig cell differentiation. Excessive proliferation of P450SCC and SF1 expressing cells were observed in the gonads of *Tcf21* null mice [29]. TCF21 expression allows fetal Leydig cells to maintain stem cell like characteristics. These results can be explained by observations that TCF21 suppresses SF1, NOTCH1 and hedgehog signaling which are required for differentiation of fetal Leydig cells [16,30]. In the rat we observed differential immunolocalization of TCF21 during male sex determination and testis differentiation. At the onset of sex determination when SRY is critical for the induction of Sertoli cell fate determination, TCF21 is primarily localized to Sertoli cells during testis cord formation (E13–E14). Later in testis differentiation as the interstitial cells and Leydig cell differentiation is initiated the TCF21 is primarily localized in Leydig cells (E16). Therefore, TCF21 has a transient role in the initial differentiation of Sertoli cells in response to SRY, and then a later role in the initiation of Leydig cell differentiation. The hypothesis is that TCF21 likely promotes a cascade of other factors, such as bHLH proteins, to continue the subsequent differentiation.

Combined *in vitro* and *in vivo* observations have demonstrated that TCF21 is a downstream target of SRY and plays an important role in testis differentiation (Figure 6). Our results also provide insights into the possibility for bHLH protein involvement in Sertoli and Leydig cell differentiation. Since bHLH transcription factors have a critical role in the differentiation of many tissues and cell types, a role in mammalian sex determination and gonadal cell differentiation will be an interesting future area of research. Experiments are in progress to determine the cascade of genes following TCF21 and SRY expression and their roles in testis differentiation and male sex determination.

**Materials and Methods**

**Tissue preparation and Histology**

Harlan Sprague-Dawley rats were kept in a temperature controlled environment and given food and water *ad libitum*. Estrous cycles of female rats were monitored by cellular morphology from vaginal smears [31]. Rats in early estrus were paired with males overnight and mating confirmed by sperm positive smears, denoted day 0 of pregnancy. Pregnant rats were euthanized at E13, E14, and E16 of pregnancy, and embryonic gonads were collected for histological analysis. Sex was determined by PCR using primers specific for *Sry* on genomic DNA isolated from embryo tails as previously described [32]. All procedures were approved by the Washington State University Animal Care and Use Committee (IACUC approval # 02568-018). Tissue specimens were fixed in Bouin’s solution for 1 h and embedded in paraffin using standard procedures. Serial sections of 4 μm were stained with hematoxylin and eosin (H&E) using standard procedures [33].

**Immunohistochemistry**

Embryonic testis sections [34] were deparaffinized, rehydrated through a graded ethanol series, boiled 10 minutes in 10 mM sodium citrate buffer to expose the antigens, washed with 0.1% Triton-X solution, and then blocked with 10% serum of the species secondary antibody was raised in for 30 min prior to incubation with 0.5 μg/ml primary rabbit anti-TCF21 antibody for 18 h (ABcam). The sections were then washed with PBS and incubated with 1:3000 diluted Alexa Fluor 488 labeled secondary antibody for 45 min (goat anti-rabbit IgG; Invitrogen, Eugene, OR). Slides were mounted with Vecta-Shield plus DAPI (Vector Laboratories Inc.), sealed with coverslips, and analyzed using fluorescence confocal microscopy (Zeiss). Negative control experiments were performed using a non-immune IgG at 0.5 μg/ml (rabbit IgG; Sigma, *St. Louis, Mo*). AMH localization was
performed using 5 μg/ml primary anti-MIS antibody for 18 h (R&D Systems, Minneapolis, MN) and 1:3000 diluted Alexa Fluor 488 labeled secondary antibody (donkey anti-goat IgG; Invitrogen, Eugene, OR) using the protocol above. Non-fluorescent staining was performed using DAB staining method.

**Reporter Plasmid Preparation**

All the primers used in plasmid preparation are listed in the Table S3. In order to prepare Tf21 promoter/reporter vectors, different size fragments (≈2.7 k, ≈1.9 k, ≈1.7 k, ≈1.6 k, and ≈0.5 k) of the rat genomic DNA upstream of the coding region was amplified (for PCR primers, see Table S3) and cloned into the pGL3-Basic luciferase reporter vector (Promega, Madison, WI) using the SmaI and KpnI sites of the multiple cloning region [35]. To generate mutant promoter/reporter constructs, the SRY binding consensus sites in the promoter were mutated using complementary oligos (Table S3). Mutagenesis was performed using i-proof Taq polymerase (BioRad) according to the manufacturer’s directions. All constructed promoter vectors were sequence verified. Mouse Sry and Sox9 expression vectors were generously gifted by Dr. Peter Koopman (University of Queensland, Australia) to M.K.S. Rat Sry expression construct was prepared as described [20]. Briefly, a full length rat Sry expression plasmid with a MYC tag was produced by amplifying the single exon from rat genomic DNA and cloned into pcMV-MYC expression vectors (Clontech, Mountain View, CA) using the BglII and NotI restriction sites. In order to prepare recombinant rat SRY protein, Myc-tagged rat SRY coding region was inserted into pcDNA 3.1 vector and in vitro transcribed & translated using a TnT T7 quick in vitro Transcription and Translation kit (Promega).

**Cell Preparation and Culture**

**Sertoli cell culture.** All cell preparation and transfections were performed according to the protocol developed in the laboratory as previously described [36]. All animal procedures and protocols were approved by the Washington State University Animal Care and Use Committee. Decapsulated testes were minced with razor blades. Fragments were then digested with trypsin (1.5 mg/ml, Life Technologies, Gaithersburg, MD) to remove the interstitial cells followed by collagenase (1 mg/ml type I, Sigma) for removal of peritubular cells and then hyaluronidase (1 mg/ml, Sigma) for removal of germ cells. Sertoli cells were plated under serum free conditions in 24-well Tissue Culture Plates (Falcon Plastics, Oxnard, CA) at 1 x 10^6 cells/well. Cells were maintained in 5% CO2 atmosphere in Ham’s F-12 medium (HyClone) at 32°C [37].

**E13 testis cell culture.** E13 embryos (12 to 18 tail somite-stage) were collected from timed-pregnant females as described above [34,38]. Gonads from E13 animals were dissected and each pair of gonads from individual animals was placed into one well of a 24 well plate with 500 μl Ham F-12 medium until embryos could be sexed as described above. The male gonads were then pooled and digested with trypsin (2.5%) and collagenase (1 mg/ml type I) plus DNase (3 mg/ml) to dissociate the cells. All the cells from the digested testes were then cultured on 100 mm plates in Ham’s F-12 with 10% bovine calf serum (Sigma). Cells initially multiplied well in culture and were split two times (1:2) as they reached confluence, at which point cell division slowed considerably. Cells were maintained in culture, changing medium every three days, until growth plaques were observed at approximately one month. These growth plaques were then collected for further propagation and frozen stocks were prepared for subsequent cell splitting such that cells could be maintained at fewer than 12 subcultures. E13 female primary cells were also prepared accordingly using female gonads from E13 (14 to 18 tail somite-stage female embryos).

**Transfection Procedures**

**Sertoli cell transfection.** Sertoli cells cultured for 48 hours were transfected by the calcium phosphate method coupled with hyper osmotic shock (10% glycerol) as previously described [35,36]. Briefly, 2 μg promoter reporter plasmid with or without 1 μg expression plasmid in 150 μl transfection buffer (250 mM CaCl2 mixed 1:1 (v/v) with 2× HEBES (28 mM NaCl, 50 mM HEPES, and 1.47 mM Na2HPO4 at pH 7.05)) was added to each well of a 24 well plate containing 1 x 10^6 Sertoli cells in 1 ml Ham’s F-12 medium and incubated at 32°C for 3.5 hours. Following incubation, the cells were subjected to hyper-osmotic shock. The medium was aspirated and 1 ml of 10% glycerol in Hanks’ Balanced Salt Solution (HBSS, Invitrogen) was added for 3 minutes. Wells were washed twice in HBSS before fresh Ham’s F-12 with 0.01% BSA was added. Following a 72 hour incubation cells were harvested for luciferase assays. Medium was aspirated and 100 μl of 1 x cell lysis solution (Promega) was added per well. Plates were frozen and thawed before cell lysate was collected. For detection of luciferase reporter activity 20 μl of Sertoli cell lysate was mixed with 100 μl of luciferase substrate (Promega) and luciferase activity detected on a Wallac Victor II 1420 instrument.

**E13 cultured testis cell transfection.** Cells between sub 8 and 12 were transfected using Lipofectamine 2000 (Invitrogen, Carlsbad, CA) [39]. Two μg promoter reporter plasmid with or without 0.5 μg expression plasmid was mixed with 2 μl Lipofectamine 2000 in 100 μl Opti-MEM medium (Invitrogen) for each well of a 24 well plate. This 100 μl mix was added to each well containing ~90% confluent cultured E13 testis cells in 1 ml Ham’s F-12 medium without antibiotics and incubated at 32°C for 24 hours. After 24 hours medium was aspirated from cells and replaced with 1 ml Ham’s F-12 with 10% serum. Cells were incubated 72 hours and collected for luciferase assays as described above for Sertoli cells. For over-expression, cells were transfected in 6 well plates with 4 μg expression construct for TCF21-pCMV-myc, rat SRY-pCMV-myc and rat TCF12-pCMV-HA. Cells were collected after 6 days and RNA was extracted using Trizol. RNA was reverse transcribed and cDNA was used for PCR. At least three over-expression experiments were made with both E13 male and female cell cultures.

**Immunoprecipitation Pull-Down Assay**

SRY recombinant protein with HA tag was synthesized in vitro by using Sry expression construct and an in vitro transcription and translation kit according to the manufacturer’s instruction (Promega) [40]. Oligonucleotides corresponding to Tf21 promoter with or without SRY binding site were generated by PCR amplification with primers listed in the Table S3. DNA-protein binding was performed in 30 nL reaction mixture containing 100 mM KC1, 1 mM MgCl2, 10 μM ZnSO4, 10 mM Tris, pH 7.5, 4% glycerol, 0.1% Triton X-100, 1 mg/ml BSA, 1 μg of poly(dIdC)/poly(dAdT), 0.5 mM DTT and protease inhibitor cocktail mini tabs (Sigma). Binding reaction continued for an hour at room temperature. Protein-A Sepharose beads were pres swollen in the cChIP incubation buffer (50 mM NaCl, 20 mM Tris HCl pH 7.5, 20 mM Na-butylate, and 5 mM EDTA) overnight. Before use, beads were incubated with anti-HA antibody or with non-immune IgG as a negative control. Following gentle centrifugation (600 x g) for 5 minutes, beads were treated with protein-oligo mixture and incubated with gentle rotation overnight at 4 degree Celsius. In this way the antibodies, while bound to the beads, would attach to their matching proteins and capture the
oligo fragments specifically bound to the proteins. Beads were washed three times with wash buffer containing 50 mM, 100 mM and 150 mM NaCl. Pulled-down oligos were eluted in a buffer containing 0.5% SDS and purified in PCR purification columns according to manufacturer’s instruction (Promega). Purified oligo DNA was subjected to PCR with primers designed to flank that particular sequence of Tcf21 promoter.

Chromatin Immunoprecipitation (ChIP) Assay

In vivo. Carrier ChIP (cChIP) assay was adopted from O’Neill et al., (2006) [26] and conditions were modified to meet the condition for immunoprecipitating SRY. To run ChIP assay, male and female gonads (gonad+mesonephros) were dissected from approximately twenty to twenty five 13 dpc (12–18 tail somite stage) rat embryos per array and snap-frozen in liquid nitrogen. Drosophila SL2 cells were used as a carrier. Densely-grown cells (approximately 5×10^7 cells) were pelleted and washed three times in ice-cold PBS, 5 mM sodium butyrate and resuspended in 0.5 ml NB buffer (15 mM Tris-HCl, pH 7.4, 60 mM KCL, 15 mM NaCl, 5 mM MgCl₂, 0.1 mM EDTA, 0.5 mM 2-mercaptoethanol, 0.1 mM PMSF). Frozen testis samples were thawed, mixed with SL2 cells and homogenized to make single cell suspension. Nuclei were pelleted, resuspended in 20 ml NB buffer, 5% (vol/vol) sucrose, pelleted and resuspended again in 5 ml digestion buffer (50 mM Tris-HCl pH 7.4, 0.32 M sucrose, 4 mM MgCl₂, 1 mM CaCl₂, 0.1 M PMSF). Following micrococcal nuclease digestion (NEB, USA) for 10 minutes at 37 degree, the digested samples were gently spun (800xg) for 10 minutes and supernatant set aside on ice. The pellet was resuspended in 250 ul incubation buffer (50 mM NaCl, 20 mM Tris HCl pH 7.5, 20 mM Na-butyrate, 5 mM Na2EDTA, and 0.1 mM PMSF) and again centrifuged gently for 10 minutes. Both the supernatants were pooled and a fraction (50 ul) out of it was kept aside to use as input. The remaining supernatant was incubated with either non-immune IgG or anti-SRY or anti-Gata4 antibodies at 4°C overnight. After incubation with 100 μl of preswollen protein A-Sepharose beads (Sigma) for 2 h at 4°C, the bead-bound immunoprecipitates were centrifuged gently and washed five times with wash buffer (50 mM Tris HCl pH 7.5, 10 mM EDTA, 5 mM Na butyrate and 50–150 mM NaCl). The protein-DNA complexes were incubated at room temperature with elution buffer (1% SDS in TE) and centrifuged at 11500 xg at 4°C for 10 minutes. Elution was repeated two times and eluted DNA was pooled. Co-immunoprecipitated DNA was purified by proteinase K digestion, phenol/chloroform extraction, and ethanol precipitation. Final concentration of immunoprecipitated DNA varied from 100 to 300 ng per assay. Two rounds of PCR reactions were run for 25 cycles before evaluating samples by gel electrophoresis. Identity of the PCR-amplified fragments was verified by sequencing. Primers used to investigate the enrichment of Tcf21 promoter fragments by PCR are listed in Table S3.

Microarray Analysis

mRNA processing and hybridization were performed at Genomics Core Laboratory, Center for Reproductive Biology, Washington State University, Pullman, WA using standard Affymetrix reagents and protocol [41]. Briefly, mRNA was transcribed into cDNA with random primers, from the later, cRNA was transcribed, and from that, single-stranded sense DNA was synthesized which was fragmented and labeled with biotin. Biotin-labeled fragmented ssDNA was then hybridized to the Rat Gene 1.0 ST microarrays containing more than 27,000 transcripts (Affymetrix, Santa Clara, CA, USA). Hybridized chips were scanned on Affymetrix Scanner 3000. CEL files containing raw data were then pre-processed and analyzed with Partek Genomic Suite 6.5 beta software (Partek Inc, St. Louis, MO) using RMA, GC-content adjusted algorithm. Lists of differentially expressed genes treatment were generated using following cut off criteria: signal ratio Control/Treatment >1.20 or <0.83, mean difference for un-logged signals between control and treatment >10, t-test p-values<0.05. CEL files from this study have been deposited with the NCBI gene expression and hybridization array data repository (GEO, http://www.ncbi.nlm.nih.gov/geo, GSE, GSE24666 and can also be accessed through www.skinner.wsu.edu, and is MIAME compliant. For genes annotation, Affymetrix annotation file RaGene1_0stv1.na30.mt4.transcript.csv was used unless otherwise specified. Generation of affected KEGG pathways (Kyoto Encyclopedia for Genes and Genome, Kyoto University, Japan) used Pathway-Express, a web-based tool freely available as part of the Onto-Tools (http://vortex.cs.wayne.edu) [42]. Previous studies have demonstrated that microarray data are validated with quantitative PCR data [38,43]. Due to the presence of 11 different oligonucleotide sets for each specific gene being used on the microarray versus only a single primer set for a gene in a quantitative PCR, the microarray is more effective at eliminating false positive or negative data and provides a more robust quantification of changes in gene expression.

Supporting Information

Table S1 Genes differentially expressed in F-E13 with at least 2 Treatments.

Table S2 (S2A) Differential regulation of female E13 cell culture transcriptome by rat Sry expression construct. (S2B) Differential regulation of female E13 cell culture transcriptome by Tcf21 expression construct. (S2C) Differential regulation of female E13 cell culture transcriptome by Tcf21 plus Tcf2 expression construct.

Table S3 List of primers used in the present study.

Acknowledgments

We acknowledge the technical assistance of Dr. Marina Savenkova and Ms. Ellyn Schinke and thank Ms. Heather Johnson for assistance in the preparation of the manuscript. Gene expression data has been deposited in the NCBI GEO (www.ncbi.nlm.nih.gov/geo) and are accessible with NCBI GEO (www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE24666).

Author Contributions

Conceived and designed the experiments: MKS RKB. Performed the experiments: RKB ISR TMC. Analyzed the data: MKS RKB ISR TMC. Wrote the paper: MKS RKB ISR TMC. Reviewed the manuscript and approved the final version: RKB ISR TMC MKS.

References


