TSHZ3 and SOX9 Regulate the Timing of Smooth Muscle Cell Differentiation in the Ureter by Reducing Myocardin Activity

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Abstract
Smooth muscle cells are of key importance for the proper functioning of different visceral organs including those of the urogenital system. In the mouse ureter, the two transcriptional regulators TSHZ3 and SOX9 are independently required for initiation of smooth muscle differentiation from uncommitted mesenchymal precursor cells. However, it has remained unclear whether TSHZ3 and SOX9 act independently or as part of a larger regulatory network. Here, we set out to characterize the molecular function of TSHZ3 in the differentiation of the ureteric mesenchyme. Using a yeast-two-hybrid screen, we identified SOX9 as an interacting protein. We show that TSHZ3 also binds to the master regulator of the smooth muscle program, MYOC, and displaces it from the coregulator SRF, thereby disrupting the activation of smooth muscle specific genes. We found that the initiation of expression of smooth muscle specific genes in MYOC-positive ureteric mesenchyme coincides with the down regulation of Sox9 expression, identifying SOX9 as a possible negative regulator of smooth muscle cell differentiation. To test this hypothesis, we prolonged the expression of Sox9 in the ureteric mesenchyme in vivo. We found that Sox9 does not affect Myocd expression but significantly reduces the expression of MYOC/SMF-dependent smooth muscle genes, suggesting that down-regulation of Sox9 is a prerequisite for MYOC activity. We propose that the dynamic expression of Sox9 and the interaction between TSHZ3, SOX9 and MYOC provide a mechanism that regulates the pace of progression of the myogenic program in the ureter.

Citation: Martin E, Caubit X, Airik R, Vola C, Fatmi A, et al. (2013) TSHZ3 and SOX9 Regulate the Timing of Smooth Muscle Cell Differentiation in the Ureter by Reducing Myocardin Activity. PLoS ONE 8(5): e63721. doi:10.1371/journal.pone.0063721

Editor: Qingzhong Xiao, William Harvey Research Institute, Barts and The London School of Medicine and Dentistry, Queen Mary University of London, United Kingdom

Received March 26, 2012; Accepted April 11, 2013; Published May 6, 2013

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Funding: Research in the laboratory of LF was supported by the Centre National de la Recherche Scientifique (CNRS) with additional funding from the Association Francaise contre les Myopathies (AFM, grants 12545, 13013) and the Agence Nationale de la Recherche (ANR, grant ANR-09-GENO-027-01). Work in the laboratory of AK was supported by a grant (DFG Ki728/7-1) from the Deutsche Forschungsgemeinschaft (German Research Council). E. Martin was a fellow of the AFM. Part of this work was performed using France-BioImaging infrastructure supported by the Agence Nationale de la Recherche (ANR-10-INSB-04-01, call “Investissements d’Avenir”). The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: The authors have declared that no competing interests exist.

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Introduction
Smooth muscle (SM) is a frequent investment of visceral organs, and is required to assure their rigidity and contractility. In the urinary tract, SM is present in the renal pelvis, the ureter, the bladder and the urethra and plays a crucial role in the functional and structural integrity of these organs.

In vivo characterization of visceral smooth muscle cell (SMC) development in the ureter has provided evidence for a role for Sonic hedgehog (SHH) and Bone morphogenetic protein 4 (BMP4) signaling in the differentiation of the mesenchymal cells that surround the urothelium [1,2,3,4,5,6,7,8]. Targeted removal of Shh signal production in the ureteric epithelium resulted in a reduction of ureteral mesenchymal cell proliferation and delayed SM differentiation, which is likely to play a causal role in the observed hydroureter [8]. Moreover, removal of Shh resulted in a deficit of Bmp4-expressing peri-ureteral mesenchyme, while conditional deletion of Bmp4 dramatically reduced expression of SM specific genes [1,8]. Four transcription factors, the T-box protein TBX18, the zinc-finger protein Teashirt 3 (TSHZ3), the homeodomain-containing protein SIX1 and the high-mobility-group (HMG) domain transcription factor SOX9 are expressed in SMC precursors and are important for the cytodifferentiation of ureteric mesenchyme [9,10,11,12]. Phenotypic and molecular analyses of Tbx18 mutant ureters have indicated that Tbx18 is required for specifying mesenchymal cell precursors to a ureteric fate [9]. Six1 acts synergistically with Tbx18 in regulating the proliferation and differentiation of SMC precursors [12]. Teashirt 3 (Tshz3) is expressed in precursors, and in differentiating and differentiated SMCs; its expression is maintained in adult ureteric SM. In Tshz3 mutant ureters the myogenic program is not activated in the proximal region due to the absence of expression of myocardin (Myocd), a key regulator of SM differentiation [11,13]. It was suggested that SM differentiation depends on a signaling axis starting with secreted SHH and ending with ureteric SMC differentiation, with Tshz3 downstream of Bmp4 but upstream of Myocd and SMG contractile protein synthesis [11,14]. This work also provided some indication concerning the temporal control of SM differentiation. In the wild type ureter, MYOC and SMC markers are detected at E15.5, shortly before onset of urine
production at E16.0 to E16.5. \textit{Tshz3}^{−/−} mice as well as other mutants with defects in ureteric SM differentiation consequently develop hydroureter at around E16.5 [11]. Recently, it was shown that the high mobility group domain transcription factor gene \textit{Sox9} plays an important role both in kidney and ureter development in the mouse [10,15]. \textit{Sox9} is transiently expressed in the undifferentiated ureteric mesenchyme. Conditional inactivation of \textit{Sox9} in this domain led to a significant down-regulation of \textit{Myocd} expression and perturbed differentiation into SMC, which resulted in strong proximal hydroureter formation due to functional obstruction. The involvement of visceral SMCs in pelvi-ureteric junction obstruction (PUJO), a frequent cause for persisting dilatation of the upper urinary tract (hydronephrosis), underlies the significant interest in understanding the mechanisms that control the differentiation of visceral SMCs [16,17,18,19].

Previous data suggested that both TSHZ3 and SOX9 critically regulate the onset of the myogenic program in the developing ureter. However it is not clear whether TSHZ3 and SOX9 act independently or are part of a regulatory network that controls cytodifferentiation of the ureteric mesenchyme. Here, we set out to characterize the molecular function of TSHZ3 in the development of the ureteric mesenchyme. Using a yeast-two-hybrid screen, we identified SOX9 as an interacting protein. We verify this interaction in vitro and in vivo, and define its temporal nature in the ureter. We show that TSHZ3 also binds to MYOCD, and that the ternary complex between SOX9, TSHZ3 and MYOCD disrupts the activation of SM specific genes. Therefore, despite controlling the onset of the myogenic program by inducing the expression of \textit{Myocd}, TSHZ3 and SOX9 subsequently delay the differentiation of visceral SMCs by inhibiting Myocardin activity. We therefore propose that the dynamic expression of \textit{Sox9} and the interaction between TSHZ3, SOX9 and MYOCD provide a mechanism that regulates the pace of progression of the myogenic program in the ureter.

\section*{Materials and Methods}

\subsection*{Ethics Statement}

All experiments were performed in accordance with the European legislation (86/609/CE), the guidelines established by the French Ministry of Agriculture (Animal Rights Division) and approved by H. Hedrich, state head of the animal facility at Medizinische Hochschule Hannover and performed according to German legislation. The architecture and functioning rules of our animal house, as well as our experimental procedures have been approved by the “Direction Départementale des Services Vétérinaires” (ID number E-13-055-21). Experiments were not performed on living animals.

\subsection*{Mouse strains}

Wild-type and \textit{Tshz3}^{−/−} mouse lines used in this study were maintained on a CD1 outbred background. The \textit{Tshz3}^{−/−} allele carries an in-frame insertion of \textit{lacZ} coding sequence within the second exon of \textit{Tshz3} [11]. For conditional prolonged \textit{Sox9} expression, females homozygous for an \textit{Hprt}^{−/−} allele were crossed with \textit{Tbx18Cre}/\textit{Tshz3lacZ} males as previously described [10].

\subsection*{Cell lines}

The murine C5H1/1T1/2 cell line was purchased from American Type Culture Collection (ATCC number CCL-226); the human HEK293T cell line was a gift from Dr. J. Iovanna (CRCM, Marseille, France). Cell lines were cultivated following the ATCC culture protocol and maintained in Dulbecco's Modified Eagle's Medium with supplementation of 10% fetal bovine serum and 50 units/ml penicillin, 50 mg/ml streptomycin, 1% sodium pyruvate.

\subsection*{Yeast Two-hybrid Screen (Y2H)}

The screen has been previously reported in [20].

\subsection*{Immunological and \textit{in situ} hybridization analysis}

Tissues were fixed in 4% paraformaldehyde/PBS. Immunostaining was performed either on 10 μm cryosections of tissues or on paraffin embedded sections after quenching endogenous peroxidase activity and antigen retrieval followed by reaction with secondary antibodies. Primary and secondary antibodies used were mouse anti-smooth muscle α actin (SMaA, 1A4, Sigma; /1000), guinea-pig anti-TSHZ3 antibody (1/50000) produced by A. Garratt's laboratory (Max-Delbrueck-Center, Berlin, Germany). Rabbit anti-SOX9 antibody (1/1000) was kindly provided by Dr Pascal de Santa Barbara (Inserm, ER125, Montpellier, France). Other primary antibodies were rabbit anti-β-galactosidase (Cappel, 1/500), mouse anti-β-galactosidase (Sigma Aldrich, 1/1000), goat anti-MYOCD (sc-21559, Santa-Cruz, 1/200), rat anti-HA (3F10 Roche, 1/1000), mouse anti-Flag M2 (F3165, Sigma Aldrich, 1/1000). Secondary antibodies used were Alexa-Fluor-488 or 488 donkey anti-rabbit, Alexa-Fluor-546 or 488-IgG, goat anti-mouse, Alexa-Fluor-488 goat anti-guinea pig and Alexa-Fluor-555 goat anti-rat (Molecular Probes). Secondary antibodies were diluted at [1/1000]. Cultured cells were fixed in 1% PFA for 10 min. Cryosections and cultured cells were washed with 0.1% Tween/PBS for 15 min and then blocked in 10% fetal calf serum, 0.1% Tween/PBS for 1 hour. Fixed samples were incubated with primary antibodies overnight at 4°C, followed by incubation with secondary antibodies for 1 hour (RT) and then counterstained with DAPI.

\textit{In situ} hybridisation using digoxigenin-labeled probes was performed on sections as previously described [21]. The \textit{Myocd} probe was kindly provided by Dr Eric Olson (University of Texas Southwestern Medical Center, Dallas, USA).

\subsection*{Construction of plasmids}

The mouse \textit{Myocd} 856-Flag pcDNA3.1 (smooth muscle \textit{Myocardin} isofrom, [22]) was kindly provided by Dr Joseph Miano (Aab Cardiovascular Research Institute, University of Rochester, USA). The human \textit{Sox9}-Flag expression vector was provided by Dr Andreas Schell (U638, Inserm, Nice, France). The Flag-\textit{Sox9AC} pcDNA3.1 was given by Dr Benoit de Crombrugghe (University of Texas M. D. Anderson Cancer Center, Houston, USA). The mouse mamalian expression plasmid for myocardial response factor (Sfy) was kindly provided by Dr Zhenlin Li (University Pierre and Marie Curie, Paris VII, France). The mouse \textit{Myod} 856-HA, Sfy-HA, \textit{Sox9-HA}, and Flag-\textit{Sox9HM} were generated by PCR amplification and cloned into pcDNA3.1. The pcDNA3.1-HA-\textit{Tshz3} and pCX-\textit{Tshz3} clones were produced by subcloning a 3.4 kbp PCR-generated fragment covering the complete HA-\textit{Tshz3} open reading frame into the pcDNA3.1(+)(Invitrogen) and the pCX-MCS2 (gift of Dr Xavier Morin, ENS, Paris, France) plasmids, respectively. An N-terminal 6XHis tag was ligated to \textit{Tshz3} by PCR. N-\textit{Tshz3} and G-\textit{Tshz3} were cloned by PCR and ligated to pcDNA3.1. Human TSHZ3-trunc pCMV was a gift from Dr Joseph D. Busbaum (Laboratory of Molecular Neuropsychiatry, Mount Sinai School of Medicine, New York, USA). Tshz3-trunc was cloned into the pcDNA3.1 vector in frame with an amino-terminal HA epitope. HA-Tshz3 dZNF was initially generated in pBS SK vector by a QuikChange multi-site mutagenesis kit (Stratagene) and then transferred to the pcDNA3.1
vector. All the resulting plasmids were sequenced to check the insert integrity.

mRNAs extraction and qRT-PCR

Before qRT-PCR analysis on cells, C3H10T1/2 cells were nucleofected using Amaxa™ nucleofector Kit V (Lonza) following the manufacturer’s protocols with designed program T-020. For qRT-PCR on ureters, embryonic ureters were dissected from E16.5 Tbx18Cre/+ HprtSox9/+ and wild type embryos. Total RNA was isolated using TRIsure Reagent (Bioline) according to the manufacturer’s instructions. RNA was treated with RQ1 RNase-free DNase1 (Promega). 750 ng of total RNA was reverse-transcribed using Superscript II reverse transcriptase (Invitrogen) with oligo dT(12–18mers) and random primers (15mers). cDNA was submitted to RNAseH treatment during 20°C at 37°C to remove the complementary RNA strand. Real-time PCR was performed on an IQ5 System (Bio-Rad Laboratories) using SYBR Green Supermix (Invitrogen). qRT-PCR was performed for 40 cycles of 95°C for 30 s, 60°C for 30 s, and 72°C for 30 s. Analyses were performed in triplicate on RNA samples from several independent experiments. Results were normalised to Gadph housekeeping gene [23] and then normalised to their respective control group. RQ = 2^−ΔΔCt and ΔΔCt = (Ct experimental−Ct Gadph)−(Ct control−Ct Gadph).

Primer sequences used for Sybr qRT-PCR are as follows:

Mouse Tshz3 S-GCGCGCAGCAGCCTATGTTTC and AS-TCAGCCATCCGGTCACTCGTC;

Mouse Myocd S-CGGATTCGAAGCTGTTGTCTT and AS-AAACCAGGCCCCCTCCC;

Human Sox9 S-AACGCCTTCATGGTGTGG and AS-TCGCTCTCGTTCAGAAGTCTC;

Mouse Sox9 S- CGAACGCACATCAAGACGG and

Figure 1. TSHZ3 and SOX9 physically interact in vitro and in vivo. (A–I) Mapping of the TSHZ3 interaction domain with SOX9. (A) Sequence analysis of the two Sox9 clones clA47 (amino acids 1–163) and clA45 (amino acids 1–168) showed that the selected interaction domain corresponds to amino acids 1 to 163 of SOX9 that contains part of the HMG domain. The SOX9DC construct contains the HMG domain but not the transactivation (TA) domain (B) Coimmunoprecipitation experiment shows SOX9DC interacting with TSHZ3 protein. (C) Schematic structure of the TSHZ3 full length (TSHZ3 fl) and TSHZ3 truncated proteins used in this study. N-TSHZ3 harbours the N-terminal half of TSHZ3 (amino acid: 1–483), C-TSHZ3 harbours the C-terminal half of TSHZ3 (amino acid: 484–1081), TSHZ3 d2NF arbours N-terminal half of TSHZ3 (amino acid: 1–483) and mutated zinc finger motifs and, TSHZ3-trunc lacks the amino acids 1–182. AD = acidic domain; Znf = zinc finger domain; HD = homeodomain. (D) GST pulldown assays show that TSHZ3 interacts with in vitro translated SOX9. (E) TSHZ3-HA, N-TSHZ3-HA and SOX9-Flag localize to the nucleus in HEK293T transfected cells. Cells were counterstained with DAPI to detect nuclei. (F–I) HEK293T cells were transfected with HA-tagged TSHZ3 constructs and Flag-tagged SOX9 or control empty plasmids. Proteins were immunoprecipitated with a Flag antibody, followed by immunoblotting as indicated; in: input. doi:10.1371/journal.pone.0063721.g001

Figure 2. Comparative expression analysis of SOX9 and TSHZ3 in the developing ureter. (A–I) Immunostainings for SOX9 (red), TSHZ3 (green) on sagittal sections at E12.5, and on transverse sections of the ureter at E14.5 and E15.5. Nuclei were stained with DAPI (blue). (A, D, G) TSHZ3 antibody staining reveals mesenchymal expression in the ureter. (B, E, H) SOX9 antibody staining reveals a highly dynamic expression pattern in the developing ureter. (C, F, I) SOX9 expression overlaps with that of TSHZ3 in undifferentiated mesenchymal precursors from early stages whereas SOX9 signal decreases from E15.5 in the cm of the ureter. Broken line in A, B, C, D, E, G, H delineates metanephric mesenchyme (mm), ureter (ur), ureteric epithelium (ue), condensed ureteric mesenchyme (cm), loose ureteric mesenchyme (lm).

doi:10.1371/journal.pone.0063721.g002
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Figure 3. Expression of SOX9 in SM progenitors. (A–G) Transverse tissue sections of E14.5 (A–C) and E15.5 (E–G) proximal ureters showing co-immunostaining of SOX9 (red) and MYOCD (green) in the condensed mesenchyme at E14.5. (D, H) Smooth muscle co-immunostaining of SOX9 (red) and MYOCD (green) in the condensed tissue sections of E14.5 (A–C) and E15.5 (E–G) proximal ureters showing GST protein-binding assays

Coimmunoprecipitation assays and Western blot

HEK293T cells were transiently transfected with plasmids encoding the Flag or HA-tagged MYOCD, HA-tagged TSHZ3, Flag-tagged SOX9, HA-SRF with Promofectin (Promocell). pCDNA3.1 parental plasmid was transfected to equalize the amount of DNA in each condition. For protein isolation, after 24 hours of culture, 100 mm plates of cells were scraped into 1 ml of PBS and centrifuged at 2000 rpm for 5 min. The cell pellet was immediately resuspended in 500 μl of lysis buffer containing 50 mM Tris-HCl pH 7.6, 200 mM NaCl, 1 mM EDTA, 0.25% Na-deoxycholate, 1 mM PMSF, 1 mM Na3VO4 and protease inhibitors (Complete; Roche). Cell extracts were incubated with anti-Flag agarose M2 beads (Sigma) or anti-HA (3F10, Roche) and protein G beads prewashed in lysis buffer according to the manufacturer’s protocol. Samples were incubated overnight at 4°C on a rotator. For Western blot, total cellular protein was subjected to SDS/8% PAGE and then transferred to nylon membranes. 4% total extract was loaded as input. Immunoblotting was carried out using rat anti-HA (3F10, Roche) or mouse anti-Flag M2 (F1804, Sigma) antibodies, followed by secondary anti-rat HRP (Molecular Probes) or anti Mouse TrueBlot ULTRA HRP (eBioscience) antibodies and detection using the Western Lightning plusTM ECL (Perkin Elmer).

Mammalian expression and reporter gene assays

The mouse telokin promoter − luciferase reporter gene including nucleotides from −190 to +181 (T370, telokin-Luc) of the telokin gene was a gift from Dr Paul Herrling (Indiana University School of Medicine, Indianapolis, USA). The rat SM α-actin promoter fragment extends from nucleotide 2,555 to 2,813 and the rabbit SMMHC promoter from 4,200 to 11,600, the SM22-luciferase reporter gene includes nucleotides 475 to 61 of mouse SM22 were generously provided by Pr Gary Owens (Molecular Physiology and Biological Physics, University of Virginia, USA). Plasmids were transformed into mouse C3H10T1/2 fibroblasts cells using FuGENE6 (Roche). 10T1/2 cells were seeded at 25000 cells/well in 24-well plates. Half an hour post-seeding, each dish was incubated with a total of 1 μg of plasmid DNA (0.125 μg of promoter-luciferase plasmid, 0.25 μg of expression plasmid, and 0.025 μg of phRLTK plasmid as an internal control) and 2 μl of FuGENE in 45 ml of Dulbecco’s modified Eagle’s medium. 24 h later, extracts (100 μl/well) were prepared for measurement of luciferase activity with Tristar LB941 multiplate reader (Berthold Technologies). The level of promoter activity was evaluated by determining the level of firefly luciferase activity relative to the control Renilla using the dual luciferase assay system essentially as described by the manufacturer (Promega). Results are reported as the mean and SEM.
Figure 4. TSHZ3 and SOX9 modulate the transcriptional and myogenic activity of Myocardin. (A) Overexpression of Sox9 and Tshz3 suppresses the induction of endogenous SM gene expression by MYOCD. 10T1/2 cells were cotransfected with indicated combination of expression constructs. Cells were harvested 24 hours post nucleofection for qRT-PCR analysis of transcripts of endogenous SmA, telokin, SMMHC and SM22α (Gadph was used to normalize (n = 6, mean ± S.E.M). (B) SOX9 and TSHZ3 suppress the MYOCD and SRF-induced activation of SM specific promoters. 10T1/2 cells were cotransfected with indicated combination of expression constructs, luciferase reporters controlled by SmA, telokin, SMMHC and SM22α The amount of each plasmid was 250 ng for Myocd, Sox9 and Tshz3, 10 ng for Srf, 125 ng for promoter reporter plasmid and 25 ng for PhRLTK renilla luciferase (n = 8, mean ± S.E.M). Asterisk indicates statistical significance as determined by a Wilcoxon test (p<0.05).

doi:10.1371/journal.pone.0063721.g004
Statistical analyses
Statistical analyses were carried out using StatEL software for Mac OSX (www.adscience.eu). The non parametric Wilcoxon test was used to evaluate the significance of differences between control and tested samples. In all these analyses, P<0.05 or P<0.01 were regarded as significant.

Results

TSHZ3 and SOX9 form a molecular complex
We recently reported on the important role of the transcription factor TSHZ3 for ureteric SM differentiation [11], but the molecular function of TSHZ3 in this developmental context remained unclear. Data obtained in invertebrate and vertebrate models showed that TSHZ-mediated gene regulation depends on interaction with different protein binding partners [24,25,26,27]. To get insight into the molecular function of TSHZ3, we searched for protein interaction partners using a yeast-2-hybrid screen [20]. We identified 84 independent clones among which 47 (56%) clones encode for high mobility group (HMG) containing proteins, corresponding to SMARCE1/BAF57 (40 clones) [20] and five SOX proteins (7 clones). Sequence analysis of the 47 clones revealed that the selected interaction region contained an HMG domain. Two of the cDNAs (cl A47 and cl A45; Figure 1A) coded for SOX9 (Genbank accession number NM_011448.3), which has been implicated in SMC differentiation in the proximal ureter [10]. Consistent with the yeast two-hybrid data, co-immunoprecipitation assays performed in HEK293T cells using HA-epitope tagged TSHZ3 and Flag-epitope tagged N-terminal SOX9 (Sox9 DC; Figure 1A) showed that a fragment of SOX9 containing the HMG domain interacts with TSHZ3 (Figure 1B).

To delineate the region of TSHZ3 that accounts for SOX9 binding, we generated various Tshz3 deletion constructs (Figure 1C) and performed a series of in vitro and in vivo binding assays (Figure 1D, F–I). We found that in vitro translated SOX9 binds to a fusion protein of full-length TSHZ3 (TSHZ3fl) with GST (GST-TSHZ3fl) but not to GST alone (Figure 1D). In a reciprocal manner, TSHZ3fl bound to GST-SOX9 fusion protein (Figure S1A). In these assays, SOX9 strongly interacted with GST-N-TSHZ3 harbouring the N-terminal half of TSHZ3 (amino acid: 1–483; N-TSHZ3) and weakly with the C-terminal half of TSHZ3 (amino acid: 484–1081; C-TSHZ3) (Figure 1D); N-
TSHZ3 interacted with GST-SOX9 (Figure S1A). Next we performed coimmunoprecipitation (co-IP) assays in HEK293T cells transfected with a HA-tagged TSHZ3 construct alone, or in the presence of Flag-tagged SOX9. Tagged versions of both SOX9 and TSHZ3 localized to the nucleus of HEK cells (Figure 1E). An enrichment of HA-tagged TSHZ3 protein was detected only upon co-transfection of the Sox9 expression construct (Figure 1F). The specificity of the association was confirmed by performing reciprocal co-IP assay with anti-HA antibody (Figure S1B). Co-IP assays confirmed that N-TSHZ3 interacts with SOX9 in vivo (Figure 1G) and showed that the zinc finger motif was dispensable for interaction (Figure 1H). The N-terminal deletion mutant of TSHZ3 (amino acid: 183–1081; TSHZ3-trunc) did not interact with SOX9 (Figure 1I). These data demonstrate that the interaction with SOX9 takes place within the first 102 amino acids of the TSHZ3 protein.

SOX9 and TSHZ3 show overlapping expression in SM progenitors of the developing ureter

Genetic and molecular studies suggested that Tshz3 and Sox9 are required for the cytodifferentiation of the proximal ureteric mesenchyme into SM [10,11]. These data together with our in vivo binding studies suggested that TSHZ3 and SOX9 interact molecularly in ureteric mesenchymal cells in vivo. To determine when this interaction takes place in the developing ureter, we performed a detailed analysis of the expression of TSHZ3 and SOX9 from E12.5 to E16.5 in wild-type (WT) ureters. From E12.5 to E14.5, TSHZ3 (Figure 2A, D) and SOX9 (Figure 2B, E) were co-expressed (Figure 2C, F) in the condensed mesenchyme (cm) around the epithelium (ue) as well in the outer ring of loose mesenchymal cells (lm). SOX9 was also detected in the epithelium, until E14.5 (Figure 2E). From E15.5 onwards, SOX9 expression was down-regulated in the condensed mesenchyme (Figure 2H, I) that undergoes SM differentiation [8,28]. Downregulation of SOX9 coincides with the activation of the expression of MYOCD target genes

The decrease of SOX9 protein at the onset of SMC differentiation prompted us to compare the expression of SOX9 with that of SM markers and MYOCD, a key transcriptional activator of several SM specific genes [13,29,30,31]. In wild-type ureters at E14.5, we found that SOX9 and MYOCD proteins were co-detected in the SM progenitors of WT ureters (Figure 3A–C), as supported by the absence of SMαA (Figure 3D). One day later at E15.5, the SOX9 level decreased and the protein was excluded from the MYOCD expression domain (Figure 3E–G); SMαA was now detected (Figure 3H). At E16.5, SOX9 was detected in the loose mesenchyme that will mature into connective tissue but SOX9 was absent from the SM layer (Figure S2H–J). In contrast to SOX9, expression of Tshz3 was maintained from E15.5 onwards in the SM layer ([11]; Figure S2 C, D, F, G). As the expression of SM genes depends on the presence of the serum response factor (SRF), we confirmed that SRF was expressed at E14.5 and E15.5 in ureteric mesenchymal cells (Figure S2 K–P). In the urogenital tract, the initiation of SM development is first detected in the presumptive bladder and 48 h later, it occurs in the developing ureter [28]. Accordingly, immunostaining of mouse embryonic urogenital tracts at E13.5 showed Myocardin staining in the bladder but not in the ureter (Figure S2 Q, R). This in vivo analysis temporally delineates the multistep process of the SM progenitor cell differentiation.

TSHZ3 and SOX9 repress the myogenic activity of MYOCD

A large number of studies established that MYOCD activates SM genes directly through an obligate interaction with SRF [32,33]. Thus, we sought to analyse the consequence of the expression of TSHZ3 and SOX9 on the MYOCD/SRF-mediated transcriptional activation of SM-specific genes in more detail. This analysis was performed in the C3H10T1/2 fibroblastic cell line,
which represents a relevant cellular model for studying SM differentiation [34,35]. Sox9 or Tshz3 expression plasmids were transfected individually or in combination with or without Myocd expression plasmid in 10T1/2 cells. Evaluation of mRNA levels of SM genes by quantitative real-time PCR (qRT-PCR) showed that forced expression of TSHZ3 and SOX9 significantly diminished the MYOCD/SRF-mediated transcriptional activation of SM genes (Figure 4A). qRT-PCR analysis showed that the relative levels of exogenous Myocd mRNA were comparable in the different conditions, indicating that the observed repression was not related to variation of Myocd expression (Figure S3). Based on the above findings, we examined the effect of TSHZ3 and SOX9 on the luciferase activity of reporter plasmids containing the minimal promoters of a series of SM specific genes. We found that TSHZ3 and SOX9 potently repressed MYOCD/SRF-mediated luciferase activity of reporter plasmids (Figure 4B). We conclude that TSHZ3 and SOX9 interfere with the transcriptional activity of MYOCD/SRF.

TSHZ3 interacts with MYOCD

The regulation of SM gene expression depends on the interaction of SRF with cofactors and the displacement of MYOCD from SRF can lead to the repression of SM genes [13,36,37]. We hypothesised that TSHZ3 and SOX9 modulate the transcriptional activity of MYOCD/SRF by interacting with either MYOCD or SRF. To test this hypothesis, we performed GST pull-down and coimmunoprecipitation assays. As shown in Figure 5A, TSHZ3 interacted efficiently with MYOCD. In contrast, SRF neither bound to SOX9 nor to TSHZ3 (not shown); and SOX9 did not bind to MYOCDS (Figure S4A).

We mapped the MYOCD-binding domain of TSHZ3 in coimmunoprecipitation and GST pull-down assays using truncated constructs of TSHZ3. GST pull-down experiments showed that MYOCD interacts with the N-terminal half of TSHZ3 (Figure 5A and Figure S4B). To investigate the interaction of MYOCD and TSHZ3 further, we performed coimmunoprecipitation assays. HEK cells were transfected with expression plasmids for the SM isoform of MYOCD (Flag-tagged MYOCD-S56, [22]), and TSHZ3-HA. We verified that MYOCD and TSHZ3 transfected proteins predominantly localized to the nucleus of HEK cells (Figure S4C). Specific interaction of MYOCD and TSHZ3 was observed by co-IP with anti-Flag antibody followed by immunoblotting with anti-HA (Figure 5B and Figure S4D). Co-IP assays showed that N-TSHZ3-HA and a TSHZ3 truncated protein lacking the three zinc finger domains (TSHZ3 dZNF-HA) bound to MYOCD-Flag (Figure 5C, D). Co-IP experiments revealed, in contrast, that a TSHZ3 N-terminal deletion mutant lacking residues 1–182 (TSHZ3-trunc-HA) lost both the ability to interact with MYOCDS and to repress the transcriptional activity of MYOCD/SRF (Figure 5, E, F). Thus, the ability of TSHZ3 to inhibit SM gene promoter correlated with the ability to interact with MYOCDS and was probably specific for MYOCDS activity, as TSHZ3 did not affect SRF trans-activation properties (Figure S4E).

TSHZ3 and SOX9 interfere with MYOCD/SRF complex formation

As the displacement of MYOCD from SRF can lead to the repression of SM genes, we analysed whether TSHZ3 and SOX9 could affect the interaction of MYOCD with SRF [37]. We tested this hypothesis by performing co-IP experiments. We first determined the effect of TSHZ3 and found that the amount of immunoprecipitated SRF decreased with increasing amounts of TSHZ3 (Figure 6A). In addition, we found that the TSHZ3-trunc protein lacking the MYOCD interaction domain did not affect SRF binding to MYOCD (Figure 6B), suggesting that the direct interaction between TSHZ3 and MYOCD prevents the association of SRF with MYOCD. We next analyzed the effect of both SOX9 and TSHZ3 on MYOCD/SRF complex formation. We found that MYOCD was less efficient to immunoprecipitate SRF. In SOX9 transfected cells, SOX9-HA was detected in the immunoprecipitate, probably via its interaction with TSHZ3 (Figure 6C). Together, these results suggest that the presence SOX9 and TSHZ3 can displace MYOCD from SRF, and as a consequence, prevents the activation of SM genes.

Prolonged expression of SOX9 in the ureteric mesenchyme does not prevent initiation of Myocd expression but reduces expression levels of SM genes

Our in vitro data suggest that MYOCD is displaced from SRF in the presence of SOX9 and TSHZ3 (Figure 6C), and that expression of SM genes correlates with the downregulation of Sox9 in vivo (Figure 3 and 7A). If this was of functional importance, prolonged expression of Sox9 should affect the transcriptional activation of MYOCD/SRF target genes without affecting the expression of Myocd (Figure 7B). To test this, Sox9 expression was maintained in the ureteric mesenchyme beyond E14.5 using a conditional Cre/loxP-based Sox9 misexpression approach [10], and expression of MYOCD/SRF target genes was measured. E16.5 ureters from wild-type and Tbx18Cre/+;HprtSox9/+ mice were collected and the levels of Myocd and SM gene transcripts measured by qRT-PCR. We found that prolonged expression of Sox9 significantly affects SM gene but not Myocd expression (Figure 7C), suggesting that down-regulation of Sox9 is important to trigger the expression of SM genes and the progression of myogenic program.

Discussion

Myocd encodes a transcriptional coactivator that physically associates with the MADS-box transcription factor, SRF, to promote myogenic differentiation. Expression and activity of Myocd/MYOCD is tightly controlled in space and time by a multitude of signaling pathways and transcriptional moieties in different SMC programs. Here, we have shown that in the ureteric mesenchyme the two transcription factors TSHZ3 and SOX9 control MYOCD transcriptional activity, thus, SMC differentiation, by ternary complex formation. Together with the previous...
findings that TSHZ3 and SOX9 are required for Myocd expression, this points to a crucial and dynamic function of these two transcription factors in ureteric SM development.

**A TSHZ/SOX9/MYOCD pathway controls SM differentiation of progenitor cells**

The differentiation of ureteric mesenchyme into SMCs is sensitive to extracellular cues. In response to SHH from the ureteric epithelium, surrounding mesenchymal cells activate Bmp4 [8]. TSHZ3 and SOX9 are likely to act downstream of BMP4 to trigger the initiation of the myogenic program through the transcriptional activation of Myocd [10,11]. To date, a functional requirement for Myocd or Srf has not been established, but the absence of Myocd expression in Tshz3- and Sox9-mutant ureters, that show perturbed SMC differentiation, argue for a key role for Myocd in the initiation of the myogenic program [10,11]. However, the precise nature of the molecular pathways regulating Myocd expression in the ureter remain poorly described and the regulatory regions responsible for transcriptional activation of Myocd have not been identified [22,39]. In Sox9-deficient ureters, the expression of Tshz3 is not affected [10], and in Tshz3-mutant ureters Sox9 expression is unchanged (Figure S5), indicating that Tshz3 and Sox9 do not regulate each other. Physical interaction between TSHZ3 and SOX9, as identified in this study, may provide a clue for an upstream module for transcriptional activation of Myocd. This heterodimeric complex may directly activate Myocd transcription, alternatively it may provide a permissive environment for the expression of Myocd. Further characterization of Tshz3- and Sox9-mutant ureters may contribute to our understanding of the transcriptional regulation of Myocd in visceral SMC progenitors and the initiation of their differentiation program.

The results of this study indicate that TSHZ3 and SOX9 regulate the progression of the SM differentiation program by controlling the activity of MYOCD. The following observations support this conclusion. (1) TSHZ3 and SOX9 reduce the ability of MYOCD to activate expression of endogenous SM genes, as well as luciferase reporter gene controlled by SM gene promoter. (2) TSHZ3 and SOX9 form a ternary complex with MYOCD that strongly reduces the ability of MYOCD to interact with SRF. (3) Prolonged expression of Sox9 prevents MYOCD to activate expression of endogenous SM genes. (4) TSHZ3 interacts directly with MYOCD and there is a positive correlation between the domain of TSHZ3 required for interaction with MYOCD and repression of SM gene expression. These data add to the notion that regulation of MYOCD activity provides another layer in the control mechanisms of SMC differentiation. They also explain why both ablation and overexpression of Sox9 in ureteric mesenchymal cells affect SMC differentiation.

Based on these results, we propose a mechanistic model on the initiation and progression of ureteric SM differentiation by TSHZ3 and SOX9 (Figure 7C). TSHZ3 and SOX9 regulate the expression of Myocd, which is a critical step for the initiation of the myogenic program. Then TSHZ3 and SOX9 form a ternary complex with MYOCD, which prevents MYOCD from activating SM specific genes. The subsequent down-regulation of Sox9 expression allows the interaction between MYOCD and SRF to activate the expression of SM genes. Interestingly, analysis of Shh-mutant ureters has led to the hypothesis that SM progenitors require a signal for differentiation, suggesting that in absence of this signal SM progenitors do not differentiate [8]. One possibility could be that the differentiation signal triggers the downregulation of Sox9 expression. In absence of the signal, inhibition of MYOCD activity by TSHZ3 and SOX9 may maintain a progenitor-like state. Clearly, identification of this signal through which down-regulation of Sox9 occurs is critical and would represent a major advance in understanding the mechanisms underpinning ureteric SM differentiation.

Cellular and biochemical experiments suggest that despite the presence of the potent activator MYOCD, TSHZ3 and SOX9 impair the activation of SM specific genes, favoring the maintenance of a progenitor state. In vivo, there is a close correlation between the downregulation of Sox9 and the initiation of the expression of SRF/MYOCD target genes. Prolonged expression of Sox9 clearly provided in vivo evidence for the inhibitory role of Sox9 in visceral myogenesis. Of note, the correlation between the downregulation of Sox9 and the mesenchymal cell differentiation has previously been described in osteoblast differentiation [39]. In skeletal muscles, it has been proposed that Tshz3 and Sox9 might be responsible for maintaining myoblasts in an undifferentiated stage and prevent differentiation into myotubes. Notably, in the presence of TSHZ3 or SOX9, MYOD is unable to activate the myogenin promoter [40,41].

These data suggest that Tshz3 and Sox9 direct ureteric mesenchymal cells to the visceral SM lineage but Sox9 inhibits differentiation of SM progenitor cells into mature SM.

**Similar strategies in vascular and visceral myogenic programs**

We identified a dynamic transcriptional network, in which transcription factors are required in a first step to initiate the expression of Myocd and in a second step to trigger the progression of the myogenic program by controlling the activity of MYOCD. This two-step mechanism bears similarity with a mechanism previously described for vascular smooth and cardiac muscles, where the FOXO4 factor participates in the transcriptional activation of Myocd and FOXO4 can interact with and repress the transcriptional activity of MYOCD [22,42]. It has been proposed that the interaction between FOXO4 and MYOCD might help to fine tune the potent transcriptional activity of MYOCD [22]. The capacity of TSHZ3 to modulate the activity of MYOCD/SRF suggests that in visceral SMC TSHZ3 might help to fine-tune the transcriptional activity of MYOCD at late stages of ureter development.

Based on current knowledge this model might not occur in other context of visceral SM development. In the future, it would be important to consider whether the combinatorial action of TSHZ3 and SOX9 could apply to other TSHZ and SOX factors.

**Supporting Information**

**Figure S1** TSHZ3 and SOX9 form a complex in vivo and in vitro. (A) GST-SOX9 or GST alone were incubated with N-TSHZ3, TSHZ3II, or C-TSHZ3 in vitro expressed in reticuloocyte. (B) HEK cells were transfected with Flag-tagged SOX9 and HA-tagged TSHZ3. TSHZ3 proteins were immunoprecipitated using anti-HA antibody, as indicated. Immunoprecipitated proteins were identified by Western blotting using antibodies against the HA and Flag epitopes. (TIF)

**Figure S2** Maintenance of TSHZ3 protein in ureteric SM progenitors and SM cells, and loss of SOX9 detection in SM cells at E16.5. (A-D) Transverse sections of proximal ureters showing β-galactosidase immunostaining (brown), which recapitulates TSHZ3 expression, and expression of myocardin by in situ hybridization (purple signal) at E14.5 and E15.5. (E-J) Detection of TSHZ3 protein (green, F, G) contrasting with
absence of SOX9 (red, I, J) in SM cells stained with the SM marker: SMA (magenta, E, H) on transverse sections of ureters at E16.5. (K–P) Detection of SRF (green, K, M, N, P) and SMA (magenta, E, H) on transverse sections of ureters at E14.5 (K–M) and E15.5 (N–P). Q, R Expression of Myocardin in the ureter and the bladder at E13.5. Q and R, Immunostaining for Myocardin (red) and nuclei (DAPI, Blue). Myocd was not detected in the ureteral mesenchyme (A), in contrast to the bladder (B). Dotted line in A separates the epithelial layer and the mesenchymal layer. Asterisk indicates the lumen of the ureter. (TIF)

**Figure S3** Myocd expression is stable. 10T1/2 cells were cotransfected with the indicated constructs. 24 hours post nucleofection, total RNA was harvested from cells and expression levels of exogenous Myocd were measured by qRT-PCR. (TIF)

**Figure S4** TSHZ3 interacts with MYOCD. (A) HEK cells were transfected with expression vectors encoding Flag-tagged SOX9 and HA-tagged MYOCD. Cell lysates were incubated with anti-Flag (SOX9) antibody, and the immunoprecipitates were probed with anti-HA (MYOCD) or anti-Flag (SOX9) antibodies by Western analysis. (B) Aliquots of in vitro translated 35S-labelled full-length (TSHZ3 fl) and deletion mutants of TSHZ3 (N-TSHZ3, C-TSHZ3) were incubated with glutathione-Sepharose beads loaded with bacterially expressed GST alone or GST-MYOCD. The retained 35S-labeled TSHZ3 proteins were separated on SDS-PAGE followed by autoradiography. 10% of input (in) is shown on the left. In control experiments, no binding of TSHZ3 proteins to the GST beads was observed. (C) TSHZ3-HA (red, anti-HA epitope tag), N-TSHZ3-HA (red) and MYOCD-Flag (green, anti-Flag epitope tag) localize to the nucleus in HEK293T transfected cells. Cells were counterstained with DAPI to detect nuclei. (D) HEK cells were transfected with expression vectors encoding Flag-tagged MYOCD and HA-tagged TSHZ3. Cell lysates were incubated with anti-HA (TSHZ3) antibody, and the immunoprecipitates were probed with anti-HA (TSHZ3) or anti-Flag (MYOCD) antibodies by Western analysis. (E) 10T1/2 with indicated combination of expression constructs and a luciferase reporter controlled by SMaA. Luciferase activity was measured. SRF significantly activated SMA luciferase promoter around 1.5 fold relative to the basal activity of SMA promoter alone. This activation was unaffected by co-transfection with TSHZ3. An arbitrary value of 1 was assigned to the basal activity of SMA promoter alone. (n = 12, mean ± S.E.M). (TIF)

**Figure S5** Expression of Sox9 is not affected in Tshz3 mutant ureters. (A, B) Sox9 expression in wild type ureter. (C, D) Sox9 expression in Tshz3 mutant ureter. (TIF)

**Acknowledgments**

We are thankful to Helen Skaer, Teddy Fauquier, Andreas Schiedl and Adrian Woolf for helpful discussions.

**Author Contributions**

Conceived and designed the experiments: AK LF. Performed the experiments: EM XC RA CV AF AK LF. Analyzed the data: EM XC RA CV AF AK LF. Contributed reagents/materials/analysis tools: EM XC RA CV AF AK LF. Wrote the paper: EM XC RA CV AF AK LF.
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