

Published in final edited form as:

Dev Biol. 2010 August 15; 344(2): 1071–1087. doi:10.1016/j.ydbio.2010.05.495.

Comparative gene expression analysis of genital tubercle development reveals a putative appendicular Wnt7 network for the epidermal differentiation

Han Sheng Chiu¹, John C. Szucsik², Kylie M. Georgas¹, Julia L. Jones³, Bree A. Rumballe¹, Dave Tang¹, Sean M. Grimmond¹, Alfor G. Lewis², Bruce J. Aronow^{2,3}, James L. Lessard², and Melissa H. Little^{1,4}

¹ Institute for Molecular Bioscience, The University of Queensland, St. Lucia, 4072, Australia

² Division of Developmental Biology, Cincinnati Children's Hospital Research Foundation, Cincinnati, Ohio 45229, USA

³ Division of Biomedical Informatics, Cincinnati Children's Hospital Research Foundation, Cincinnati, Ohio 45229, USA

Abstract

Here we describe the first detailed catalogue of gene expression in the developing lower urinary tract (LUT), including epithelial and mesenchymal portions of the developing bladder, urogenital sinus, urethra and genital tubercle (GT) at E13 and E14. Top compartment-specific genes implicated by the microarray data were validated using wholemount *in situ* hybridization (ISH) over the entire LUT. To demonstrate the potential of this resource to implicate developmentally critical features, we focused on gene expression patterns and pathways in the sexually indeterminate, androgen-independent GT. GT expression patterns reinforced the proposed similarities between development of GT, limb and craniofacial prominences. Comparison of spatial expression patterns predicted a network of *Wnt7a*-associated GT-enriched epithelial genes, including *Gjb2*, *Dsc3*, *Krt5* and *Sostdc1*. Known from other contexts, these genes are associated with normal epidermal differentiation, with disruptions in *Dsc3* and *Gjb2* showing palmo-plantar keratoderma in the limb. We propose that this gene network contributes to normal foreskin, scrotum and labial development. As several of these are known regulated by, or contain cis elements responsive to retinoic acid, estrogen, or androgen, this implicates this pathway in the later androgen-dependent development of the GT.

Keywords

genital tubercle development; lower urinary tract development; gene expression; Wnt; appendage development

⁴Corresponding author: Professor Melissa H Little, NHMRC Principal Research Fellow, Institute for Molecular Bioscience, The University of Queensland, St. Lucia, 4072, Australia, Ph: +61 7 3346 2054, FAX: +61 7 3346 2101, M.Little@uq.edu.au.

Publisher's Disclaimer: This is a PDF file of an unedited manuscript that has been accepted for publication. As a service to our customers we are providing this early version of the manuscript. The manuscript will undergo copyediting, typesetting, and review of the resulting proof before it is published in its final citable form. Please note that during the production process errors may be discovered which could affect the content, and all legal disclaimers that apply to the journal pertain.

Introduction

Congenital anomalies of the urogenital tract are the third most common birth defects, with abnormalities of the external genitalia being the most prevalent manifestation. Cryptorchidism (undescended testes) and hypospadias (in which the urethral orifice is located on the underside of the penis or at the perineum) each occur in as many as 1 in 125 males born (Kurzrock et al, 1999; Cox et al, 2008) and the prevalence of these two conditions appears to be rising, potentially due to environmental exposure to anti-androgenic compounds (Miyagawa et al, 2002; Brouwers et al, 2007; Wang and Baskin, 2008). These and other less common anomalies of the external genitalia (diphallia, micropenis) create social as well as reproductive challenges for the affected individuals. The lower urinary tract (LUT) is defined as extending from the bladder to the meatal opening of the urethra. Congenital anomalies of the kidney and urinary tract (CAKUT) include such defects as vesicoureteric reflux (VUR) (improper insertion of the ureters into the bladder) and posterior urethral valves (fine outgrowths of urethral tissue that result in bladder outlet obstruction) (Kerecuk et al, 2008). Such conditions are quite common (VUR presents in 1–2% of all children (Kerecuk et al, 2008)), with these sequelae of most LUT problems presenting later in life (urge incontinence, stress incontinence, cystitis, neurogenic bladder, overactive bladder, erectile dysfunction) and being aetiologically poorly understood.

Despite a physiological relationship with the mesodermally-derived kidney and ureters, the LUT and GT have distinct embryological origins. The LUT develops from the endodermal cloaca, which becomes separated into the urogenital sinus and rectum by the downgrowth of the urorectal septum around E10.5 in mouse. The urogenital sinus gives rise to the internal urethra and bladder. While both become lined with a water-tight urothelial layer, these two regions differentiate into distinct structures with respect to musculature, stroma, epithelial layers and innervation. The role of the LUT is storage and voiding. Regulation of appropriate micturition requires complex neural connections and, unlike most internal organs, control of storage and voiding by the LUT involves voluntary regulation. Locally, this involves sympathetic and parasympathetic innervation of the bladder and somatic motor nerves to regulate contraction of the urethral sphincter (Fowler et al, 2008). As well as providing a barrier, the urothelial cells of the bladder also act as mechanosensors and are actively involved in triggering neural responses in response to bladder expansion. The importance of the urothelium in this process was recently highlighted by the presence of urothelial hyperplasia and non-voiding contractions in bladders from mice defective for either uroplakin II or IIIa (Aboushwareb et al, 2009). The genital tubercle (GT) gives rise to the male and female external genitalia (penis, clitoris, scrotum, labia, foreskin) and represents the intersection between the reproductive and urinary tracts. The primary swellings of the genital tubercle form the penis and clitoris, while the preputial swellings give rise to the foreskin in males and the labia minora in females. A second set of swellings, the labioscrotal swellings, arise later in development and give rise to the scrotum and labia majora in males and females, respectively. The LUT and GT physically intersect as the urethra passes through the GT. The GT is proposed to be derived from all three germ layers. The urethral plate (UP) is derived from urethral endoderm (Siefert et al, 2008). The GT surface other than the urethral plate is derived from surface ectoderm and the core of the extending appendage is derived from lateral plate mesoderm (Perriton et al, 2002). Murine GT development commences at approximately E10.75, with the appearance of paired swellings ventral to the cloacal membrane (Perriton et al, 2002). Without a urethral plate, the GT will fail to form. The urethral epithelium of the GT does not grow out from the urethra but forms *in situ* along the GT itself (Pennington and Hutson, 2002). GT development involves an early androgen-independent phase and a later androgen-dependent phase, after which sexual dimorphism is evident. In mouse, this dimorphism is not evident until

approximately E16, before which there has already been significant outgrowth and patterning during an androgen-independent phase.

The genital tubercle develops as an out-budding appendage, hence it has been proposed that there is congruence in the genetic basis of GT patterning with other out-budding appendages, including the limb and craniofacial prominences. Of these three appendages, limb patterning has been most highly studied. The limb has three key axes of development, the anterior-posterior axis (thumb to little finger), proximal-distal axis (arm to finger tips) and dorsal-ventral or plantar-palmar (back of hand to palm) axis. Major growth factor pathways involved in limb development include the hedgehog, fibroblast growth factor (FGF), bone morphogenetic protein (BMP) pathways and Wnt pathways (Goodman et al, 2000; Ahn et al, 2001; Hill et al, 2006; Geetha-Loganathan et al, 2008; Witte et al, 2009). Sonic hedgehog (*Shh*) is expressed in a posterior zone of polarizing activity (ZPA) and is critical for limb anterior-posterior (A-P) patterning. Duplication of this region on the anterior side of the limb leads to a mirror image duplication of digits (polydactyly). Extension in the distal direction involves signals from the apical ectodermal ridge (AER), notably *Fgf8*, but also *Fgf4*, *Fgf9* and *Fgf17* (Mariani et al, 2008). These same signaling pathways are involved in GT development. Here, *Shh* is expressed in the endodermal UP along the ventral surface of the GT (Haraguchi et al, 2001; Perriton et al, 2002). The UP has been proposed to act as an organizing centre similar to the ZPA in the limb. Transplantation of this region into the anterior limb of the chick induces digit duplication, however it also results in the dimpling of the skin and formation of a furrow similar in structure to the UP (Haraguchi et al, 2001; Perriton et al, 2002). The UP expression of *Shh* influences the development of the adjacent mesenchyme, eliciting extension to form the GT. *Shh* was proposed to regulate the expression of *Fgf8* and *Bmp7*, which are also expressed in the UP and distal urethral epithelium (DUE), and *Wnt5a*, *Bmp2*, *Bmp4* and *Fgf10*, which are expressed in the mesenchyme surrounding the urethral plate (Yamaguchi et al, 1999; Haraguchi et al, 1999; Perriton et al, 2002). Expression of *Shh* is required throughout the period of androgen-independent GT extension. By removing *smoothened* from different layers of the developing GT, it was also demonstrated that the primary target of *Shh* is the GT mesenchyme (Lin et al, 2009). *Shh* knockout mice show a loss of GT accompanied by a loss of expression of *Bmp4* and *Fgf10*, suggesting that *Shh* is upstream of both of these growth factor pathways. As for the limb, the GT undergoes considerable proliferative extension prior to structural differentiation. In the limb, this outgrowth is driven by the expression of *Fgf8* in the AER. The expression of *Fgf8* in the DUE was similarly presumed to be required for GT extension and outgrowth (Morgan et al, 2003; Yamada et al, 2006). Certainly the presumptive region of *Fgf8* expression is lost when *Shh* is deleted. Surprisingly, the conditional removal of *Fgf8* from the cloacal and urethral epithelium did not affect GT outgrowth, even though there was a loss of expression of putative downstream targets of *Fgf8* (*Bmp4*, *Wnt5a*, *Hoxd13*) and no evidence of a redundant role for other members of the *Fgf* family (Siefert et al, 2009). *Wnt5a* has been shown to be involved in outgrowth of GT and limb (Yamaguchi et al, 1999). The loss of *Shh* also reduces canonical Wnt signaling demonstrating that anterioposterior extension due to mesenchymal proliferation is likely require Wnt signaling from the DUE (Miyagawa et al, 2009). Indeed, concomitant loss of *Shh*, *Wnt5a* and *axin2* expression is seen in two models of disrupted GT and anorectal development, Danforth's short tail (Sd) and all-trans RA treated mice (Nakata et al, 2009).

GT expression of *Shh* also regulates *Hoxd13* and *Hoxa13* expression, although independently to Wnt and *Fgf* signalling. (Lin et al, 2009). The homeodomain genes *Hoxa13* and *Hoxd13* are also expressed in the UP, whereas *Msx1* and the *Shh* receptor, *Ptch1*, are expressed in the surrounding mesenchyme. Here again, there is a similarity to limb where *Hoxa13* and *Hoxd13* are required for the development and A-P patterning of the autopods

(digits) (Cobb and Duboule, 2005). Homozygote knockouts of these genes result in loss of the limbs and phallus, whereas heterozygotes display disrupted patterning in both locations (reviewed in Perriton et al, 2002). In humans, mutations in these genes result in hand-foot-genital syndrome (Goodman et al, 2000). In the posterior embryo, these homeodomain genes also control formation of the terminal vertebrae and the anus. The UP is therefore also thought to be critical for both dorsal-ventral patterning and proximal-distal extension of the GT.

As noted, congenital GT defects are very common. Based on observations in mice, hypospadias may result from disruptions in *Fgfr2*, *Fgf8*, *Fgf10* and *Bmp7*, all of which are involved in UP patterning (Beleza-Meireles et al, 2007). In humans, hypospadias has been observed in Opitz syndrome (*MID1* mutation), Pallister-Hall syndrome (*GLI3* mutation), Reiger syndrome, type 1 (*PITX2* mutation), Hand-foot-genital syndrome (*HOXA13*, *HOXD13* mutations) and Split hand/split-foot malformation type 1 and type 4 (mutations in *Dlx5*, *Dlx6* and *Dss1*) (Yamada et al, 2006; Suzuki et al, 2008). Micropenis occurs in Pallister-Hall syndrome, Robinow syndrome (*ROR2* mutation), Ulnar mammary syndrome (*TBX3* mutation) and X-linked lissencephaly (*ARX* mutation) (reviewed in Yamada et al, 2006). Mutations in *CXorf6* have also been identified in isolated cases of non-syndromic hypospadias (Kalfa et al, 2008).

Despite the clinical relevance and prevalence of defects, there have been almost no systems biology applied to the molecular basis of LUT/GT development. Li et al (2006) performed expression analysis of mouse GT (E14, E15, E16 and E17), but focused on those genes expressed during the later androgen-dependent phase of development. In this study, we have addressed this gap by creating a comprehensive dataset of microarray and *in situ* gene expression of the E13 and E14 LUT/GT. In order to test the utility of this dataset, we analysed in more depth gene expression in the GT. Ontological analysis of GT-enriched genes reaffirmed the importance of Wnt signaling and defined an epidermal, oestrogen-responsive, *Wnt7a*-associated gene network potentially involved in foreskin/labial development and possibly defective in hypospadias.

Materials and methods

At E13.5, a pregnant female FVB/N mouse was sacrificed by CO₂ asphyxiation and, following a midline abdominal incision, the gravid uteri were removed and placed in PBS on ice. Embryos were dissected under magnification. Bladder and urethra were dissected free of surrounding tissues and then divided at the level of ureteral insertion. Bladder segments were incubated in 20 mM EDTA in Tyrode's solution for 20 minutes at 37°C. Urethral segments were treated with Trypsin (1 mg/ml in PBS) for 20 minutes (37°C). In each instance, epithelium was separated from mesenchyme by rimming with a fine needle. Tissues were then placed in RLT Buffer containing β-ME, according to manufacturer's instructions (RNeasy Micro Kit, Qiagen Inc., Valencia, CA) and rapidly frozen in dry ice prior to storage at -80°C for subsequent RNA purification. This was achieved using a spin-column system (RNeasy Micro Kit, Qiagen Inc., Valencia, CA), according to manufacturer's directions, and stored at -80°C prior to amplification.

At E14, a pregnant female FVB/N mouse carrying the SMGA/EGFP transgene (Szucsik et al. 2004) was sacrificed by CO₂ asphyxiation and following a midline abdominal incision the gravid uteri were removed and placed in PBS on ice. Embryos were dissected under GFP illumination. The lower urinary tract was trisected into bladder (EGFP positive dome), UGS (EGFP negative) and urethra (EGFP positive). A separate, nontransgenic pregnant FVB/N female was used to obtain samples of the genital tubercle at E14. Tissues were rapidly frozen in dry ice and stored at -80°C for subsequent RNA purification using a spin-column

system (RNeasy Micro Kit, Qiagen Inc., Valencia, CA), according to manufacturer's directions, and stored at -80°C prior to amplification.

Expression profiling, data normalisation and analysis

Purified RNA was analyzed for concentration and integrity using the Agilent RNA 6000 Pico Kit (Agilent Technologies Inc., Santa Clara, CA) and then 0.5–1.5 ng (in a volume of 1 μl) underwent two-round amplification (Epicenter Biotechnologies, Madison, WI) according to manufacturer's directions except for the addition of Full SpectrumTM multistart primers (SBI System Biosciences, Mountain View, CA) per the manufacturer's directions. The resultant product was biotinylated and hybridized to Affymetrix Mouse MOE 430 2.0 microarray chips (Affymetrix Inc., Santa Clara, CA). CEL files were generated using Expression Console v1.1.1 (Affymetrix) and subjected to RMA normalization using Affymetrix CDF probeset definitions and GeneSpringGX v7.3.1 (Agilent). The relative expression of each probeset in each sample was calculated as the ratio relative to the median of its values as measured across the entire GUDMAP MOE 430 2.0 dataset that contains developing and postnatal tissues from kidney, lower urinary tract, and gonadal tissues. Probesets to be analyzed for differential expression across the developing lower urinary tract samples were then selected by filtering for those with RMA expression intensity greater than 6.2 in at least two replicates from any sample type. A general ANOVA was used to identify probesets that are differentially expressed between sample types with less than 5% false discovery using Benjamini Hochberg p-value correction. From the resulting 18K probesets, those with highest relative expression in the Genital Tubercle were identified based on normalized expression as referenced to the diverse panel of other developing and postnatal genitourinary samples. GeneSet enrichment analysis was performed using ToppGene (Chen et al, 2009). Genes that exhibited shared properties with respect to geneset enrichment features were transformed into network representations using CytoScape as previously described (Brunskill et al. 2008) using the ToppCluster server (<http://toppcluster.cchmc.org/>; Kaimal et al., submitted).

Collection of material for in situ hybridisation

For *in situ* hybridisation, outbred CD1 embryos were collected from two time points, TS17 (10.5 dpc) and TS21 (13.5 dpc), where 0.5dpc was defined as noon of the day on which mating plug was observed. Embryos were dissected in ice-cold PBS under a dissecting microscope and fixed in fresh 4% paraformaldehyde (PFA) at 4°C overnight. Tissues were then washed in PBTX and dehydrated through a PBTX/methanol series (25%, 50%, 75%) and stored in 100% methanol at -20°C until usage. TS17 whole embryos were collected with a puncture in the back of the head to prevent subsequent stain trapping in the brain. Tail somites were counted for TS17 litters to confirm embryos are within the range of 35–39 somites, as described by Kaufman (1994). The lower urogenital system plus genital tubercle were collected from TS21 and sex was determined by gonadal inspection.

Riboprobe Synthesis

The complete protocol for digoxigenin (Dig)-labelled riboprobe synthesis is available and described in detail on the GUDMAP gene expression database, (<http://www.gudmap.org/Research/Protocols/Little.html>). A brief overview, with minor changes to the protocol, is described here. Primers were ordered from Invitrogen according to the microarray output and were designed to amplify a 3' UTR region of a gene between 500–800 bp. Riboprobes were amplified from either the Fantom II plasmid pool or the combination of both 12.5 and 15.5 dpc whole embryonic mouse cDNA. The 3' primers were tagged with a T7 polymerase sequence which allows *in vitro* transcription of Dig-labelled riboprobes using T7 RNA polymerase. (Roche) Riboprobes were then purified with lithium chloride precipitation. A master mix was made with each riboprobe consisting of: 4M LiCl

(10 μ l), 0.2M EDTA, pH 7 (8 μ l), 100% Ethanol (300 μ l), and water (100 μ l) and stored at -20° C overnight. Samples were then spun for 20 minutes at 13,000 rpm at 4° C with supernatant discarded after the spin. Samples were gently washed with 1ml of chilled 70% ethanol, then spun for 5 minutes at 13000rpm at 4° C. Supernatant were discarded and samples dried for 10 minutes at room temperature. Pellets from each sample were resuspended with 25 μ l of water and stored at -70° C.

Whole mount *in situ* hybridisation and imaging of WISH

The complete protocol for whole mount *in situ* hybridisation is available and described in detail on the GUDMAP gene expression database. (<http://www.gudmap.org/Research/Protocols/Little.html>). In brief, whole mount *in situ* hybridisation was performed by using the BioLane HTI robot with digoxigenin-labelled antisense riboprobes. Tissues were rehydrated by a series of methanol/PBTX washes, followed by a short digestion of Proteinase K, and re-fixed by 0.2% gluteraldehyde/4% PFA before incubation in pre-hybridisation solution at 65° C for 2 hours. Tissues were later transferred into pre-hybridisation solution containing 0.2 μ g/ml of riboprobe, incubated at 65° C overnight. Following post-hybridisation washes, a 2 hours incubation in pre-block solution followed by incubation in pre-adsorbed anti-DIG/pre-block solution. After the post-antibody washes, to detect the hybridised alkaline phosphatase activity, chromogenic substrates NBT/BCIP were used. Time for colour development varies between 15 minutes up to 300 minutes depending on the specificity of the riboprobe. Once the tissue has reached its optimal intensity, the tissues were washed with 1% Triton X -100/PBS at 4° C for a duration up to 2 days, later fixed with 4% PFA and stored in PBS at 4° C. All whole mount tissues were photographed manually on a 1.5% agarose gel dish, using Nikon SMZ1500 stereomicroscope system with a Nikon DXM1200f, 12 megapixel digital camera, ACT-2U Image Application Software and Adobe Photoshop CS2. For imaging of bisected genital tubercles, this region was dissected away from the rest of the lower urinary tissues in PBS on a Petri dish, and each sample was bisected either longitudinally or transversely with the aid of a 30G needle and number 11 surgical blade.

Pseudo colored gene expression overlays

To compare multiple gene expression domains Adobe Photoshop CS2 software was used to produce pseudo colored gene expression images of overlaid, stage-matched tubercles at 13.5dpc. Experimental image data was first outlined to obtain the structural outer edge of the tubercle and inverted to produce a black and white image. These were then aligned with up to two other genes using the RGB image format.

Results

Tissue was collected for RNA extraction from E13 urethra and bladder in which the epithelial and mesenchymal layers were separated from each other using trypsin digestion or EDTA treatment, respectively (Figure 1A). Tissue was also collected from E14 lower urinary tract (LUT) micro-dissected into bladder, urogenital sinus (UGS; includes the junction between the urethra, bladder and ureters), urethra and GT as illustrated in Figure 1B. All data has been submitted to GEO and is available from the GUDMAP database (www.gudmap.org), from which the expression level of all genes present on the microarray can be queried individually.

Creation of an overall view of differential gene expression across the developing lower urinary tract

In order to identify sets of genes that are either highly expressed or comparatively over-expressed in one or more of the sampled individual LUT/GT compartments during

development, probesets were first filtered to identify those genes expressed at least at the 40th percentile of raw RMA normalized intensity in at least two replicates of each sample type. From these, probes were identified that exhibited the highest ranked relative expression as compared to the globally referenced dataset, i.e. their having normalized expression in the top 2% per each of the LUT sample types (~ 250–400 probesets per each of the 8 compartments). The combination of all these probesets were hierarchically clustered and Figure 1C shows a heat map of these 1420 probesets across all compartments. These are presented after hierarchical clustering to display the most highly expressed genes across the different LUT sample types. A full gene list and heatmap of the expression profiles manifested by these transcripts is provided in Supplementary Table 1. The expression of previously known compartment markers, including *Tbx20* and *Tbx18* in the bladder and the strong expression of *Hoxd13* in all compartments, supports the accuracy of the data (Figure 1C). To determine potential functional roles for these genes, the 1420 probeset list was enriched with public database annotations (e.g. GO, Jax Mouse, OMIM, TFBS, Drug Bank, TargetScan, MSigDB, etc.) using Toppgene (Chen et al, 2009). The results of these analyses are shown in Supplementary Table 2. This revealed expression of a large number of DNA binding transcription factors, particularly homeobox (*Hoxd13*, *Msx1*, *Six1*, *Irx3*, *Irx5*, *Msx2*, *Isl1*, *Pitx1*, *Pitx2*, *Hoxa10*, *Zfmx4*, *Dlx5*, *Zfmx3*, *Mkx*, *Satb2*, *Prrx1*, *Prrx2*, *Alx3*, *Alx1*, *Cux1*) and forkhead genes (*Foxf2*, *Foxd1*, *Foxa1*, *Foxn3*, *Foxq1*); fibronectin III (FnIII) domain encoding genes (*Ptprj*, *Ptpro*, *Flrt3*, *Flrt2*, *Cntn3*, *Dcc*, *Crlf1*, *Epha5*, *Epha4*, *Epha7*, *Robo1*) and genes encoding cell-cell adhesion molecules. The most active biological processes included cell-cell adhesion with 21/104 cadherin domain containing proteins being expressed in the developing LUT (*Pcdh10*, *Pcdh18*, *Cdh11*, *Pcdha10*, *Pcdha11*, *Pcdha13*, *Dsg2*, *Frem1*, *Dsc2*, *Pcdha6*, *Pcdha7*, *Pcdha9*, *Pcdha2*, *Pcdha3*, *Pcdha4*, *Pcdha5*, *Pcdha1*, *Pcdhb7*, *Pcdhc2*, *Pcdhac1*, *Pcdhgb2*). The most active pathways were the Wnt signaling (*Wnt5A*, *Gng4*, *Myh3*, *Pcdh10*, *Pcsh18*, *Nkd2*, *Wnt7A*, *Cdh11*, *Pcdha10*, *Pcdha11*, *Pcdha13*, *FrzB*, *Pcdha6*, *Pcdha7*, *Pcdha9*, *Pcdha2*, *Pcdha3*, *Pcdha4*, *Pcdha5*, *Pcdha1*, *Pcdhb7*, *Bcl9*, *Fzd6*, *Ankrd6*, *Wnt6*, *Myst4*, *Pcdhac2*, *Pcdhac1*, *Axin2*, *Lef1*, *Pcdhgb2*, *Dkk2*, *Dkk1*, *Dkk3*, *Axin2*, *Wif1*, *Nfat5*), hedgehog signaling (*Shh*, *Bmp5*, *Gli3*, *Wnt6*, *Wnt5A*, *Wnt7A*) and cadherin signaling pathways. Toppgene analysis identified many genes involved in limb/face/appendage and skeletal development. In addition, there were many genes previously implicated in neural development and ectodermal/epidermal/epithelial development genes networks, notably cadherins, collagens and keratins. Examining the data for genes linked to human or mouse phenotypes involving the urogenital tract revealed genes known to be critical for reproductive, genital, bladder/ureter and kidney development. This set of genes overlapped genes involved in anal development, presumably reflecting the common origin of the bladder and urethra from the cloaca, and body wall development. The strongest associations were seen between genes affecting external genitalia/reproductive tract, anal canal and body wall development (*TP63*, *Sall1*, *Sall4*, *Gli3*, *Pitx2*, *Tbx3*) (Figure 1E, Supplementary Table 3). *Sall1* and *p63* are involved in the development of all of these organs/tissues, *Sall4* is involved in all but body wall and *Six1* is involved in body wall, kidney and ureter development (Supplementary Table 3).

Defining the most active networks in each compartment

In Figure 1C there is both considerable overlap of gene expression between all compartments of the LUT as well as clusters of gene expression more active in specific compartments. Heat maps for each compartment are presented in Supplementary Figure 1 and complete gene lists for each compartment in Supplementary Table 1. To understand the major processes occurring in each compartment, Toppgene analysis of each compartment was performed to identify the most represented molecular functions, biological processes, cellular components, protein domains, pathways and phenotypes. A summary of this analysis is presented in Table 2. A complete analysis of gene ontology for each

compartment-enriched gene set is presented in Supplementary Table 2. This analysis reinforced the prevalence of cadherin and Wnt signaling, the predominance of transcription factor activity and the association of LUT compartments with genes involved in appendage, skeletal, neural, epithelial and urogenital development. Cluster analysis suggested strong alignment between E13 bladder epithelium and E13 urethra epithelium, but less of a relationship between the bladder and urethral mesenchyme of the same timepoint (Figure 1D). E13 bladder mesenchyme was most similar to E14 total bladder due to the expression of muscle genes (*Tnnt2*, *Kcnq5*, *Actg2*, *Myh11*). These two bladder samples were also outliers in their lack of association with cadherin or Wnt signalling. E14 UGS and urethra clustered together, whereas GT was distinct aside from a cluster of genes co-expressed in E13 urethra mesenchyme (*Fgf9*, *Alx3*, *Foxd1*, *Msx1*, *Wnt5a*). These genes have all previously been reported in GT development, reflecting the fact that the E13 urethra sample would have included GT. E13 urethral mesenchyme and UGS showed significant overlap (*Myrip*, *Mapk10*, *Prph*, *Phox2b*, *Sthm2*, *Gdap1*) with enrichment for genes involved in neurogenesis and neural crest, including the GDNF/Ret pathway (*Gfra1*, *Gfra2*).

LUT compartment-enriched gene expression highlights innervation and vascularisation

The expression profiling was then reanalyzed so as to specifically identify differences in gene expression between the compartments being assessed. The intention of identifying compartment-enriched genes was to identify unique markers of each compartment. Supplementary Table 4 lists all genes displaying compartment-enriched expression for each compartment. Figure 2 shows a heat map highlighting gene clusters with urethra-enriched or GT-enriched gene expression. Within the heatmap it is possible to identify E14 urethra-enriched genes that are likely to mark epithelium versus mesenchyme by examining their co-expression in E13 urethra and bladder epithelium. Wholemound *in situ* hybridisation (WISH) was used to validate enrichment of gene expression in each compartment. Riboprobes were generated for a total of 94 genes (Supplementary Table 5) and all WISH results were fully annotated for gene expression and submitted to the Genitourinary Development Molecular Anatomy Project website (www.gudmap.org; McMahon et al, 2008) and riboprobe sequences are shown in Supplementary Table 5). WISH was performed on E13.5 embryos for 11 urethra-specific genes (Figure 3C), including *Irs4* and *2310076G05Rik*, seventeen E14 UGS-enriched genes, including *Elavl4*, *Gap43* and *Ascl1* (Figure 3D), and eight E14 bladder-enriched genes, including *Maf* and *Klhl4* (Figure 3E). Quite a number of UGS-enriched genes were expressed in the pelvic ganglia. Bladder-enriched genes highlighted bladder innervation and vasculature as well as markers specific to the dome of the bladder (Figure 3E). Schematic representations of these regions of the LUT are shown in Figure 3A and B.

Analysis of androgen-independent genital tubercle development

To validate the utility of the dataset, we focused on the gene expression observed in the E14 GT given that this was the most unique set of compartment-enriched sets. As previous studies have reported expression of genes from within the BMP, FGF, Wnt and hedgehog pathways in GT development, we examined the expression of all members of these pathways present on the Affymetrix chips (Supplementary Table 6). This confirmed the expression of many pathway members known to be expressed in the GT although, as expected, few of these genes showed GT-enriched or -specific gene expression. Genes from the FGF/Hedgehog/Wnt pathways showing expression in E13 LUT and E14 GT included *Fgf9* (mesenchyme), *Pik3ca*, *Npc1*, *Fgfr3*, *Fgfbp1*, *Wnt4*, *Wnt5a*, *Wnt7a*, *Wnt7b* and *Wnt10a*. A smaller number of genes from the BMP pathway showed expression in the E13 LUT and E14 GT, including *Bmp2*, *Bmp4*, *Bmp7*, *Fstl1* and *Gsk3b*. *Bmp5* showed GT-restricted gene expression at E14, but expression levels were low. The genes selected as being GT-enriched were analysed for pathway, disease, or GO term associations

(Supplementary Table 7). As for the analysis of the most active pathways in this compartment, this revealed a strong link to Wnt pathway signaling. It also reinforced the previously proposed association between GT-enriched genes and genes involved in limb and craniofacial development (Supplementary Tables 2 and 7, Table 1).

WISH of E13.5 GT was performed to validate the microarray data and investigate the spatial relationship of the GT epithelial-enriched (“on” in GT and E13 urethral epithelium), GT mesenchyme-enriched (“on” in GT and E13 urethral mesenchyme) and GT-specific genes (only observed in E14 GT). As GT was not separated from LUT at E13, genes expressed in E13 urethral epithelium or mesenchyme, but not expressed in E14 urethra, were likely to represent early GT genes, possibly enriched for those expressed along the urethral plate. Those restricted to the epithelium at E13 were most likely to be expressed in the epithelium of urethral plate of the E14 GT, while those expressed in the mesenchyme at E13 were likely to be in the mesenchyme along the urethral plate in the E14 GT. WISH validated these assumptions. In order to fully annotate the gene expression patterns observed, we modified the previous genitourinary tract ontology (Little et al, 2006) to increase the level of anatomical resolution for the TS21 genital tubercle (Supplementary Figure 2). Figure 4 demonstrates examples of GT-enriched and GT-restricted gene expression patterns. A schematic representation of the features of the developing GT can be seen at http://www.gudmap.org/Internal/Consortium/Work_In_Progress/Organ_Summaries/index.html (currently requires username: Gudmap_pr, password: 888gudmap). In total, validation of GT-enriched or restricted expression was observed for 43/58 genes examined by WISH (74% validation of array data). Table 2 summarises the annotated expression patterns these genes in the developing GT. For those where validation was not observed, signal was either absent/very weak or expression was seen in additional sites. The former is likely to result from the reduced sensitivity of the ISH in comparison to microarray (negatives). The latter may result from the riboprobe detecting an isoform of the gene not identified by the Affymetrix oligonucleotide (additional sites of expression).

Novel gene expression domains on the dorsal surface of the presumptive glans

The dorsal side of the extending genital tubercle is longer than the ventral surface such that there is an overhanging dorsal lip and the orifice of the DUE is on the ventral side. At E13.5, there is a discernable glans that is most prominent from the dorsal side. Analysis of the gene expression patterns seen in E13.5 GT revealed some distinct patterns of expression as yet unreported. Of particular interest were very tightly delineated domains of expression on the dorsal surface of the genital tubercle. These genes did not show dorsally-restricted expression, but did reveal specific dorsal patterns of expression around the tip of the extending glans that allowed the subdivision of distal genital tubercle into a number of expression patterns not previously reported. *Dlx6os1*, *Cpm*, *Mup1*, *Sty13* and *Hs3st3b1* each showed distinct but non-overlapping expression in specific regions of the distal GT (some epithelial, some mesenchymal) other than the DUE (Figure 4, 6 and www.gudmap.org). Distinct gene expression patterns were also seen on the preputial swellings (e.g. *Sostdc1*, *Dsc3*, Figure 4) and preputial glands (e.g. *Gja1*, Figure 4).

Validation of Wnt5a signaling in GT mesenchyme and identification of an associated gene network

The WISH-confirmed GT-enriched genes contained many Wnt signaling pathways members (see Supplementary Figure 3). As previously reported, *Wnt5a* was expressed in the mesenchyme of the distal genital tubercle (Yamaguchi et al, 1999; Suzuki et al, 2003) (Figure 5A). In order to predict genes interacting with Wnt5a within this compartment, we generated pseudo-colored overlays of WISH results from all GT-enriched genes. This allowed the direct alignment of up to three genes to examine in detail their spatial

relationships (Figure 5B). *Wnt5a* expression overlapped with the distal GT mesenchymal expression of Frizzled b (*Frzb*), a secreted frizzled protein that acts as an inhibitor of the pathway (Figure 5Bc, c'). The proximal edges of the *Wnt5a* expression domain were also marked by a unique and distinct expression pattern of *Dkk1*, an inhibitor of canonical Wnt signaling (Figure 5Bb–e, b'–e'). This combination of ligand and inhibitor appears to delineate the GT equivalent of a progress zone in the glans/distal GT. Comparison of this domain of expression with other genes suggests a possible interaction with several other genes from within the GT mesenchyme-enriched cluster. *Kcnab1* shows overlapping expression with *Wnt5a* in the glans, but is expressed in a wider region of the GT mesenchyme, notably not overlapping the expression of *Dkk1* (Figure 5Bd, d'). The Wnt downstream signaling molecule *Lef1* also shows partial synexpression with *Wnt5a* (Figure 5Bb, b'). In contrast, the short stature homeobox 2 gene (*Shox2*) shows a complementary pattern of expression (Figure 5Be, e'). These visual comparisons of spatial expression patterns suggest an interaction between these genes. To both validate these predictions and define potentially *Wnt5a*-interacting genes, the initial GT-enriched dataset was re-analyzed to seek genes displaying synexpression across the LUT/GT with *Wnt5a* using K-means and hierarchical clustering. This identified a cluster of genes with moderate to high expression in the E13 urethral mesenchyme (*Wnt5a*, *Dkk2*, *Frzb*, *Wif1*, *Lef1*, *Phex*, *Runx2*, *Dlx2*, *Msx1*, *Dlx1*, *Shox1*, *Satb2*, *Osr2*, *Runx5*, *Cacna2d3*, *Prdm1*, *Kcnab1*, *Tfap2b*, *Nkx1.2*, *Kcna4*). Topgene analysis of this cluster identified enrichment for genes involved in skeletal and cartilage development/morphology (Figure 5C). For example, *Runx2* has previously been shown to be regulated by *Shox2* in limb chondrogenesis prior to long bone formation (Cobb et al, 2006). These genes often displayed differential anterior-posterior limb and dorsal-ventral GT expression, reinforcing congruence in the development of these two appendicular structures. Many also showed differential proximal-distal patterning in both structures, as is the case for *Wnt5a*. This suggests possible interactions between these genes in proximal-distal extension of the GT.

Evidence for a *Wnt7a*-associated gene network in the epidermis of the GT

In contrast to *Wnt5a*, *Wnt7a* was expressed in the surface epithelium of the GT with stronger expression on the dorsal surface (Figure 6A). Little attention has been paid to expression in the dorsal GT. *Wnt7a* was also expressed on the surface epithelium of the preputial swellings (Figure 6A). Pseudo-coloured overlays suggested co-expression of *Wnt7a*, the BMP antagonist sclerostin domain containing-1 (*Sostdc1*) and keratin 2–5 (*Krt5*) in tightly defined regions of the dorsal epithelial layer of the GT. Expression of these genes was also seen in the preputial swellings, but these genes were excluded from the dorsal distal glans other than at the DUE (Figure 6Bcd). Also epithelial in expression were several genes with complementary expression patterns on the dorsal distal glans. These included desmocollin 3 (*Dsc3*) (Figure 6Bb) and *Gjb2* (Figure 4), both genes previously associated with skin development. K-means and hierarchical clustering supported the association of these and other genes as representing a synexpression group in E13 urethral epithelium, therefore potentially restricted to surface epithelium (*Wnt7a*, *Dlx5*, *Msx2*, *Tfap2c*, *Krt2–5*, *Gja1*, *Irx3*, *Irx5*, *Tfap2a*, *Gabrp*, *Krt6a*, *Krt14*, *Col17a*, *Dsc3* and *Pkp1*). Topgene analysis of this *Wnt7a*-associated cluster showed a strong link with epithelial and epidermis development/morphology, including clear links to skin abnormalities (including hyperkeratosis) in humans (Figure 6C). Genes within this cluster are involved in cell-cell junction and intermediate filament biology and some (*Gjb2*, *Dsc3* and *Krt5*) have previously been reported to be involved in plantar/palmar expression in the developing limb, as has *Wnt7a*. Of particular note, many genes within this cluster are regulated by diethylstilbestrol (*Wnt7a*, *Msx2*, *Gja1*, *Gjb2*, *Tfap2c*, *Gabrp*, *Krt6a*) and/or retinoic acid (*Krt14*, *Krt6a*, *Wnt7a*, *Tfap2a*, *Krt5*, *Gja1*, *Gjb2*, *Tfap2c*). A final cluster of genes was identified using K means and hierarchical clustering. This represented GT-specific genes, many of which showed distinct

proximal-distal restriction of expression in the GT (Supplementary Figure 4). Heatmaps of each of these clusters (GT mesenchyme (Wnt5a) associated, GT epithelium (Wnt7a) associated and GT specific) are shown in Supplementary Figure 5.

A comparison of gene expression during appendage formation

Congruence of molecular processes in appendicular outgrowth was first observed between limb and branchial arch development with the observation of a key role for Fgf8 in the patterning of the first branchial arch (Trumpp et al, 1999). As for the limb, Fgf8 expression is seen in the overlying ectoderm of the first branchial arch is required for the survival and migration of the underlying mesenchyme. A similar congruence was then observed for genital tubercle development (Haraguchi et al, 2000; Yamada et al, 2006) with hedgehog and Fgf signaling showing considerable similarities in all three tissues. This began to explain the coincidence of defects within these tissues in many birth defects in humans (Schneider et al, 1999). Our analysis of the sites of expression of genes within the Wnt5a associated network in the genital tubercle would support a role for this cluster in appendage outgrowth. WISH was also performed for these genes in whole embryo to examine limb and craniofacial prominence gene expression. This showed very tight concordance in spatial expression patterns between limb and GT, as previously hypothesized (Trumpp et al, 1999; Schneider et al, 1999). Supplementary Figure 6 schematically represents this congruence in gene expression between two of these appendicular structures; limb and GT. To further examine the hypothesis that there are similarities in GT patterning with the other appendages in the embryo, we performed WISH on E10.5 and E11.5 embryos examining expression in limb, GT and branchial arches. In total, 74% (23/31) of GT-confirmed genes showed expression in limb and 71% (22/31) showed expression in the branchial arches/craniofacial prominences (Supplementary Table 8). In almost all cases, no other strong sites of gene expression were observed, supporting the notion of congruence in regulation of appendage development. As previously described, there was also a congruence the axis of expression between dorsal-ventral GT and anterior-posterior limb (Figure 7; *Shox2*, *Tfap2b*, *Msx1*, *Tfap2c*). However, a careful analysis of all GT gene expression patterns in comparison with limb expression patterns indicated that this was not always the case (Figure 7).

Discussion

This study describes the creation and analysis of the first comprehensive catalogue of gene expression profiles describing the development of the early murine lower urinary tract and genital tubercle. Li et al (2006) previously performed microarray analysis of GT development, however the data presented only listed genes that increased in E16 and E17 GT versus E14. In addition, this data was collected from cDNA arrays and not Affymetrix. Hence a comparison cannot be made with this data. A comparison of these analyses with previous literature, together with our extensive wholemount ISH analysis supports the validity of the data. Hence, this will be a valuable resource for investigating the molecular basis of morphogenesis in this organ system and to investigate correlations between development and disease. By way of example, mutations in the renin-angiotensin system, Robo2, HNF1 β (TCF2), PAX2 (renal-coloboma syndrome), EYA1 and SIX1 (branchio-otorenal syndrome (BORS)) and SALL1 (Townes-Brocks syndrome (TBS)) have been reported in CAKUT patients (Weber et al, 2008). Although not previously associated with bladder, urethra or genital defects, strong and compartment-enriched expression in the developing LUT/GT was identified for a number of these CAKUT-associated genes showed. SALL1 mutations in TBS result in renal dysplasia and VUR. Our data shows that Sall1 is clearly expressed in the urethra mesenchyme and GT, but not in the bladder. Only a single case report exists of a TBS patient with hypospadias (Salerno et al, 2000), but our data suggest that Sall1 may be more commonly involved in GT defects. Sall1 showed strong

synexpression with *Sox11*, suggesting that mutations in this gene may also cause similar defects. Expression of *SIX1* and *EYA1*, both mutated in BOR, showed highly similar gene expression patterns in the LUT. Neither was expressed in the bladder, but both were expressed in the E13 urethral mesenchyme, E14 GT, urethra and UGS. There have been no links between these genes and genital defects, but both are associated with defects in the development of other appendages (limbs, craniofacial structures) suggesting a possible involvement in GT as well.

As a further example of the validity and potential of this dataset, Toppgene analysis of the this dataset also annotated six genes as being involved in anal, body wall, reproductive and genital development (*TP63*, *Sall1*, *Gli3*, *Sall4*, *Pitx2*, *Tbx3*). Patients with anorectal malformations show high levels of genital (13.63%) or urologic (11.36%) anomalies (Saha et al, 2005), suggesting the need to carefully examine the urogenital tracts of newborns with anorectal anomalies. Nakata et al (2009) have recently also reported an association between *Wnt5a* expression in mesenchyme close to the cloacal membrane and normal anorectal development. The link between the body wall has been less noted, although it is known that prune belly syndrome, a male-associated anomaly in the musculature of the abdominal body wall, is also associated with cryptorchidism and urinary tract anomalies (Bogart et al, 2006). Prune belly syndrome has also been linked to posterior urethral valves (Weber et al, 2005) and loci on human chromosomes 1q and 11p, as well as occurring in patients with *Six1* mutations and renal defects (BOR syndrome).

To test the utility of this catalogue in investigating the development of an individual compartment, we focused on the genital tubercle and have validated and fully described the spatial expression pattern of 31 genes likely to be critical for GT extension and differentiation. GT development has been proposed to be similar to the development of other appendages, including the limb and face. Toppgene analysis confirmed this suggestion although the most enriched pathway in the GT was Wnt signaling, a growth factor family known to be involved in the limb (Witte et al, 2009). In the limb, *Wnt5a* is involved in proximal-distal limb extension as loss of *Wnt5a* reduces proliferation within the mesenchymal progress or undifferentiated zone below the AER, resulting in limb shortening and loss of more distal structures (digits) (Yamaguchi et al, 1999). This can be mimicked by inactivation of mesenchymal β -catenin (Hill et al, 2006), suggesting that this process involves canonical Wnt signaling (Geetha-Loganathan et al, 2008). The canonical Wnt inhibitor, *Dkk1*, is expressed in the AER and loss of this gene widens the AER but does not disturb plantar-palmar patterning (Adamska et al, 2003), linking *Dkk1* with *Wnt5a* rather than *Wnt7a* activity. Similarly in the GT, loss of *Wnt5a* is also known to disrupt proximal-distal extension (Yamaguchi et al, 1999). *Wnt5a* expression has been reported to be expressed in the GT mesenchyme and appears to be regulated, along with *Msx2*, *Lef1* and *Bmp4*, by signals from the distal urethral epithelium (Lin et al, 2008). Our data on *Wnt5a* agrees with the literature in that expression was seen in the GT mesenchyme. Our clustering and expression analyses implied interaction between *Wnt5a* and other known Wnt pathway members, including *Frzb* (secreted frizzled related protein 3, *Sfrp3*), *Dkk1*, *Dkk2*, *Msx1*, *Tcfap2b* and *Lef1*. An association between *Wnt5a* and *Frzb* in digit formation has already been described (Chimal-Monray et al, 2002), supporting the validity of this cluster. Alignment of the expression patterns of these genes showed a distinct band of *Dkk1* expression delineating the boundary of the *Wnt5a* expression in the emerging glans region, presumably delimiting the boundary of *Wnt5a*-driven canonical Wnt signaling.

Differentiation between the plantar and palmar (dorsal and ventral) surface of the limb requires dorsal *Wnt7a* signaling accompanied by repression of that signal on the ventral surface by the transcription factor, engaged-1 (*En-1*) (Geetha-Loganathan et al, 2008). Dorsal *Lmx1b* expression is also thought to be regulated by *Wnt7a* (Chen and Johnson,

2002; Geetha-Loganathan et al, 2008). Rare occasions have been reported where there has been a duplication of the palmar surface due to mutation in *Wnt7a* (Al-Qattan et al, 2009). The role of Wnt signaling in the developing ectoderm of the GT has not previously been investigated. Unlike the limb, *Wnt7a* expression was not restricted to the dorsal surface of the GT although in this region it did show a specific pattern of expression. Alignment of ISH and gene cluster analysis identified other genes expressed in the GT surface epithelium (*Sostdc1*, *Gjb2*, *Dsc3* and *Krt5*), most of which have known roles in epidermal differentiation in other tissues. *Gjb2* encodes a gap junction protein also called connexin 26. Mutations in connexin 26 result in deafness as a result of cornification of the epithelium within the cochlea of the ear (Richard, 2000; Houang et al, 2002). Connexin 26 and connexin 31 (*Gjb3*) are crucial for normal epidermal development and, as well as deafness, connexin 26 mutations can result in palmo-plantar keratoderma (Richard, 2000; de Zwart-Storm et al, 2008). *Dsc3* is a cadherin-like protein present in desmosomes and previously reported to be expressed in the epidermis of the foreskin (King et al, 1995). *Dsc3* is upregulated along with β -catenin in squamous cell carcinoma (Wang et al, 2007) and disruptions of squamous and stratified epithelium. This includes palmo-plantar keratoderma and immunobullous disease in a patient with circulating antibodies against *Dsc3* (Bolling et al, 2007). This patient presented hyperkeratotic ridges on their palms as well as intra-epidermal pustules. Based on these prior associations, we can predict that this cluster of genes is involved in differentiation of the epidermis of the GT, but not necessarily the epithelium of the UP.

Genes within the predicted *Wnt7a*-associated network are known to be modulated by retinoic acid (RA) and diethylstilbestrol (DES). Sexual dimorphism of the external genitalia is dependent on steroid hormones and there have been many reports linking imbalance between estrogen and androgen signaling to hypospadias, including the use of DES during pregnancy (Brouwers et al, 2007). This is thought to affect development during the androgen-sensitive sexually dimorphic stage of GT development (>E16) rather than during early development. An association between these epidermal GT genes and DES implies either a previously unreported early oestrogen-sensitive period or involvement in both the androgen-independent and subsequent androgen-sensitive phase of GT development. DES exposure during development also results in metaplasia of the normally simple columnar epithelium of the uterus, thereby preventing embryo implantation (Huang et al, 2005). Here *Msx2* is required for normal *Wnt7a* expression and DES can repress the expression of both *Msx2* and *Wnt7a* in turn (Huang et al, 2005). This effect is mediated via the estrogen receptor, since ER-KO mice treated with DES do not show a reduction in *Wnt7a* (Couse et al, 2004). Hence, *Wnt7a* and other DES-responsive genes in this network may also be involved in female reproductive tract development. Sexual dimorphism of the external genitalia is also thought to involve RA signaling due to the dynamic expression of RARs, RXRs, RALDH2 and CYP26 (Ogino et al, 2001). Administration of RA to mouse embryos as young as E9 does not affect genital tubercle extension, but does disrupt the formation of the urethral plate. Our re-examination of these data revealed undescribed defects in the development of the preputial swellings and the shape of the distal glans (Ogino et al, 2001). Studies on the effect of an absence of RA on limb development in the quail embryo showed that while all axes were disrupted, this particularly affected limb dorsal-ventral patterning genes, including *Wnt7a* and *En-1* (Stratford et al, 1999). While this does not eliminate a role for *Wnt7a* in other axes of GT development, it is suggestive of limb/GT congruence and strengthens the association between the genes in this cluster.

Overall, our observations suggest that *Wnt7a* may play a role in epidermal differentiation of the GT in collaboration with genes known to play roles in plantar-palmar epidermal dimorphism. The expression of *Wnt7a*-synexpressed genes in the epithelium of the preputial swelling may implicate this network in the differentiation of the foreskin and labia rather

than in extension of the glans. This division of expression may serve to demarcate the development of the smooth skin of the penis and clitoris and the epidermis that will give rise to the foreskin, labia minora and scrotum (Figure 6A, B). The link to androgen regulation may also suggest that at least some forms of hypospadias or other congenital glans malformations may involve disruptions to epidermal development. Indeed, a review of 137 non-diphalliac non-hypospadiac glans defects noted dermatological lesions in 15% of cases (Papali et al, 2008).

In this study, gene ontology, pathway and network analysis of the microarray data has reinforced the concept of molecular congruence in appendage development. However, while initially arising as paired genital swellings, the glans in both sexes (penis and clitoris) becomes a single midline structure, unlike the limbs. Significant morphological differences exist between the dorsal and ventral surfaces of the developing limb, both externally with respect to skin, hair and nail formation, and internally with respect to extensor and flexor muscle positioning. The same is true for the anterior-posterior axis of the limbs. This is not as prominent in the external genitalia. While duplication of the *Shh*-expressing ZPA on the posterior surface of the limb (duplication of the A-P axis) gives rise to a duplication of digits, a dorsal duplication of the *Shh*-expressing urethral plate never occurs. Duplication of the urethral plate can occur if there is an underlying duplication of the urethral endoderm, but this duplication occurs ventrally and gives rise to a total GT duplication, referred to as diphallia in males (Kaufman et al, 1990; Gryftopoulos et al, 2002; Mughal et al, 2003). This is usually accompanied by urethral or bladder duplication and imperforate anus. Such phallic duplications almost always occur in the lateral plane such that the two appendages are positioned side by side. Rare cases of duplication apparently in the dorsal-ventral axis usually involve incomplete or glans-only duplication with no accompanying urethral or bladder duplication (Melekos et al, 1986; Gavali et al, 2002). If the limb A-P axis is equivalent to GT D-V axis, lateral diphallia would be equivalent to a duplication of the hand in the D-V axis, which has never been reported. We therefore conclude that while early congruences in patterning exist between different appendages, particularly in relation to extension, there remain significant and anatomically obvious differences between these structures and hence the molecular regulation of these processes.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

We thank Emmanuelle Lesieur for assistance in material collection and Jane Brennan, Jane Armstrong, Chris Armit, Jamie Davies, Derek Houghton, Sue Lloyd-MacGilp, Mehran Sharghi, Xinguin Pi, Yogmatee Roochun, Ying Cheng, Simon Harding, Jamie Davies and Duncan Davidson for GUDMAP editorial and database support. We also thank Carol Wicking for scientific opinion on limb and facial development. ML is a Principal Research Fellow of the National Health and Medical Research Council. This work was supported by the National Institutes of Health via awards DK070136 (MHL) and DK070219 (JLL).

Grant information: Grant sponsor National Institute of Health; Grant numbers DK070136 (MHL) and DK070219 (JLL).

References

- Aboushwareb T, Zhou G, Deng FM, Turner C, Andersson KE, Tar M, Zhao W, Melman A, D'Agostino R Jr, Sun TT, Christ GJ. Alterations in bladder function associated with urothelial defects in uroplakin II and IIIa knockout mice. *NeuroUrol Urodyn*. 2009 Epub ahead of print.

- Adamska M, MacDonald BT, Meisler MH. Doubleridge, a mouse mutant with defective compaction of the apical ectodermal ridge and normal dorsal-ventral patterning of the limb. *Dev Biol.* 2003; 255:350–362. [PubMed: 12648495]
- Ahn K, Mishina Y, Hanks MC, Behringer RR, Crenshaw EB III. BMPR-1A signaling is required for the formation of the apical ectodermal ridge and dorsal-ventral patterning of the limb. *Development.* 2001; 128:4449–4451. [PubMed: 11714671]
- Al-Qattan MM, Al-Balwi M, Eyaid W, Al-Abdulkarim I, Al-Turki S. Congenital duplication of the palm syndrome: gene analysis and the molecular basis of its clinical features. *J Hand Surgery.* 2009; 34E:247–251.
- Beleza-Meireles A, Lundberg F, Lagerstedt K, Zhou X, Omrani D, Frisen L, Norddenschjold A. FGFR2, FGF8, FGF10 and BMP7 as candidate genes for hypospadias. *Eur J Hum Genetics.* 2007; 15:405–410. [PubMed: 17264867]
- Bogart MM, Arnold HE, Greer KE. Prune-belly syndrome in two children and review of the literature. *Pediatr Dermatol.* 2006; 23:342–345. [PubMed: 16918629]
- Bolling MC, Mekkes JR, Goldschmidt WF, van Noesel CJ, Jonkman MF, Pas HH. Acquired palmoplantar keratoderma and immunobullous disease associated with antibodies to desmocollin 3. *Br J Dermatol.* 2007; 157:168–173. [PubMed: 17578440]
- Brouwers MM, Feitz WF, Roelofs LA, Kiemeney LA, de Gier RP, Roeleveld N. Risk factor for hypospadias. *Eur J Pediatr.* 2007; 166:671–678. [PubMed: 17103190]
- Brunskill EW, Aronow BJ, Georgas K, Valerius MT, Aronow JE, Kaimal V, Jegga AG, Grimmond S, McMahon AP, Patterson L, Little MH, Potter SS. Atlas of Gene Expression in Developing Kidney at Microanatomic Resolution. *Developmental Cell.* 2008; 15:781–791. [PubMed: 19000842]
- Chen H, Johnson RL. Interactions between dorsal-ventral patterning genes *Imx1b*, *engrailed-1* and *wnt-7a* in the vertebrate limb. *Int J Dev Biol.* 2002; 46:937–941. [PubMed: 12455631]
- Chen J, Bardes EE, Aronow BJ, Jegga AG. ToppGene Suite for gene list enrichment analysis and candidate gene prioritization. *Nucleic Acids Res.* 2009; 37(Web Server issue):W305–311. [PubMed: 19465376]
- Chimal-Monroy J, Montero JA, Gañan Y, Macias D, Garcia-Porrero JA, Hurlle JM. Comparative analysis of the expression and regulation of *Wnt5a*, *Fz4*, and *Frzb1* during digit formation and in micromass cultures. *Dev Dyn.* 2002; 224(3):314–320. [PubMed: 12112461]
- Cobb J, Duboule D. Comparative analysis of genes downstream of the *Hoxd* cluster in developing digits and external genitalia. *Development.* 2005; 132:3055–3067. [PubMed: 15944189]
- Cobb J, Dierich A, Huss-Garcia Y, Duboule D. A mouse model for human short-stature syndromes identifies *Shox2* as an upstream regulator of *Runx2* during long-bone development. *Proc Natl Acad Sci U S A.* 2006; 103:4511–4515. [PubMed: 16537395]
- Couse JF, Korach KS. Estrogen receptor-alpha mediates the detrimental effects of neonatal diethylstilbestrol (DES) exposure in the murine reproductive tract. *Toxicology.* 2004; 205:55–63. [PubMed: 15458790]
- Cox MJ, Coplen DE, Austin PF. The incidence of disorders of sexual differentiation and chromosomal abnormalities of cryptorchidism and hypospadias stratified by meatal location. *J Urol.* 2008; 180:2649–2652. [PubMed: 18951572]
- De Zwart-Storm EA, van Geel M, van Neer PA, Steijlen PM, Martin PE, van Steensel MA. A novel missense mutations in the second extracellular domain of GFB2, p. Ser183Phe, causes a syndrome of focal palmoplantar keratoderma with deafness. *Am J Pathol.* 2008; 173(4):1113–1119. [PubMed: 18787097]
- Fowler CJ, Griffiths D, de Groat WC. The neural control of micturition. *Nature Reviews.* 2008; 9:453–466.
- Gavali S, Deshpande AV, Sanghani HH, Hirugade ST, Talpallikar MC, Borwankar SS. Glanular diphallus with urethral stricture. *Pediatr Surg Int.* 2002; 18:70–71. [PubMed: 11793071]
- Geetha-Loganathan P, Nimmagadda S, Scaal M. Wnt signaling in limb organogenesis. *Organogenesis.* 2008; 4(2):109–115. [PubMed: 19279722]
- Goodman FR, Bacchelli C, Brady AF, Brueton LA, Fryns JP, Mortlock DP, Innis JW, Holmes LB, Donnemfeld AE, Feingold M, Beemer FA, Hennekam RC, Scambler PJ. Novel HOXA13

- mutations and the phenotypic syndrome of hand-foot-genital syndrome. *Am J Hum Genet.* 2000; 67:197–202. [PubMed: 10839976]
- Gyftopoulos K, Wolffenbuttel KP, Nijman RJ. Clinical and embryologic aspects of penile duplication and associated anomalies. *Urology.* 2002; 60(4):675–679. [PubMed: 12385932]
- Haraguchi R, Suzuki K, Murakami R, Sakai M, Kamikawa M, Kengaku M, Sekine K, Kawano H, Kato S, Ueno N, Yamada G. Molecular analysis of external genitalia formation: the role of fibroblast growth factor (Fgf) genes during genital tubercle formation. *Development.* 2000; 127(11):2471–2479. [PubMed: 10804187]
- Haraguchi R, Mo R, Hui C, Motoyama J, Makino S, Shiroishi T, Gaffield W, Yamada G. Unique functions of Sonic hedgehog signaling during external genitalia development. *Development.* 2001; 128(21):4241–4250. [PubMed: 11684660]
- Haraguchi R, Kotoyama J, Sasaki H, Satoh Y, Miyagawa S, Nakagata N, Moon A, Yamada G. Molecular analysis of coordinated bladder and urogenital organ formation by hedgehog signaling. *Development.* 2007; 134:525–533. [PubMed: 17202190]
- Houang M, Gourmelen M, Moatti L, Le BY, Garabédian EN, Denoyelle F. Hypogonadotropic hypogonadism associated with prelingual deafness due to a connexin 26 gene mutation. *J Pediatr Endocrinol Metab.* 2002; 15(2):219–223. [PubMed: 11874189]
- Hill TP, Taketo MM, Birchmeier W, Hartmann C. Multiple roles of mesenchymal beta-catenin during limb patterning. *Development.* 2006; 133:1219–1229. [PubMed: 16495310]
- Huang WW, Yin Y, Bi Q, Chiang TC, Vuoristo J, McLachlan JA, Ma L. Developmental diethylstilbestrol exposure alters genetic pathways of uterine cytodifferentiation. *Mol Endocrinol.* 2005; 19:669–682. [PubMed: 15591538]
- Kalfa N, Liu B, Ophir K, Audran F, Wang MH, Mei C, Sulton C, Baskin LS. Mutations of CXorf6 are associated with a range of severities of hypospadias. *Eur J Endocrinol.* 2008; 159:452–458.
- Kaufman A, Guia R, Davila H, Kaufman A. Diphallus with third urethra. *Urology.* 1990; 35(3):257–260. [PubMed: 2316091]
- Kerecuk L, Schreuder MF, Woolf AS. Renal tract malformations: perspectives for nephrologists. *Nature Clin Prac.* 2008; 4:312–325.
- King IA, Sullivan KH, Bennett R Jr, Buxton RS. The desmocollins of human foreskin epidermis: identification and chromosomal assignment of a third gene and expression patterns of the three isoforms. *J Invest Dermatol.* 1995; 105(3):314–321. [PubMed: 7665906]
- Kurzrock EA, Baskin LS, Cunha GR. Ontogeny of the male urethra: theory of endodermal differentiation. *Differentiation.* 1999; 64:115–122. [PubMed: 10234808]
- Li J, Willingham E, Baskin LS. Gene expression profiles in mouse urethral development. *BJU International.* 2006; 98:880–885. [PubMed: 16978288]
- Lin C, Yin Y, Long F, Ma L. Tissue-specific requirements of β -catenin in external genitalia development. *Development.* 2008; 135:2815–2825. [PubMed: 18635608]
- Lin C, Yin Y, Veith GM, Fisher AV, Long F, Ma L. Temporal and spatial dissection of Shh signaling in genital tubercle development. *Development.* 2009; 136:3959–3967. [PubMed: 19906863]
- Little MH, Brennan J, Georgas K, Davies J, Davidson D, et al. A high-resolution anatomical ontology of the developing murine genitourinary tract. *Gene Expr Patterns.* 2007; 7(6):680–699. [PubMed: 17452023]
- Mariani FV, Ahn CP, Martin GR. Genetic evidence that FGFs have an instructive role in limb proximal-distal patterning. *Nature.* 2008; 453(7193):401–405. [PubMed: 18449196]
- McMahon AP, Aronow BJ, Davidson DR, Davies JA, Gaido KW, Grimmond S, Lessard JL, Little MH, Potter SS, Wilder EL, Zhang P. GUDMAP: the genitourinary developmental molecular anatomy project. *J Amer Society Nephrol.* 2008; 19(4):667–671.
- Melekos MD, Barbalias GA, Asbach HW. Penile duplication. *Urology.* 1986; 27(3):258–259. [PubMed: 3952917]
- Miyagawa S, Buchanan DL, Sato T, Ohta Y, Nishina Y, Iguchi T. Characterization of diethylstilbestrol-induced hypospadias in female mice. *Anat Rec.* 2002; 266:43–50. [PubMed: 11748570]
- Miyagawa S, Moon A, Haraguchi R, Inoue C, Harada M, Nakahara C, Suzuki K, Matsumaru D, Kaneko T, Matsuo I, Yang L, Taketo MM, Iguchi T, Evans SM, Yamada G. Dosage-dependent

- hedgehog signals integrated with Wnt/b-catenin signaling regulate external genitalia formation as an appendicular program. *Development*. 2009; 136:3969–3978. [PubMed: 19906864]
- Morgan EA, Nguyen SB, Scott V, Stadler HS. Loss of Bmp7 and Fgf8 signaling in Hoxd13-mutant mice causes hypospadias. *Development*. 2003; 130:3095–3109. [PubMed: 12783783]
- Mughal SA, Soomro S, Shaikh JM. Double phallus. *J Coll Physicians Surg Pak*. 2003; 13(9):534–535. [PubMed: 12971880]
- Nakata M, Takada Y, Hishiki T, Saito T, Terui K, Sato Y, Koseki H, Yoshida H. Induction of Wnt5a-expressing mesenchymal cells adjacent to the cloacal plate is an essential process for its proximodistal elongation and subsequent anorectal development. *Pediatr Res*. 2009; 66(2):149–154. [PubMed: 19390486]
- Ogino Y, Suzuki K, Haraguchi R, Satoh Y, Dolle P, Yamada G. External genitalia formation: role of fibroblast growth factor, retinoic acid and distal urethral epithelium. *Ann NY Acad Sci*. 2001; 948:13–31. [PubMed: 11795391]
- Papali AC, Alpert SA, Edmondson JD, Maizels M, Yerkes E, Hagerty J, Chaviano A, Kaplan WE. A review of pediatric glans malformations: a handy clinical reference. *J Urol*. 2008; 180:1737–1742. [PubMed: 18721953]
- Pennington EC, Hutson JM. The urethral plate – does it grow into the genital tubercle or within it? *BJU International*. 2002; 89:733–739. [PubMed: 11966634]
- Perriton CL, Powles N, Chiang C, Machonochie MK, Cohn MJ. Sonic hedgehog signaling from the urethral epithelium controls external genital development. *Dev Biol*. 2002; 247(1):26–46. [PubMed: 12074550]
- Richard G. Connexins: a connection with the skin. *Exp Dermatol*. 2000; 9(2):77–96. [PubMed: 10772382]
- Saha SR, Roy AK, Saha S. Incidence of associated congenital anomalies in anorectal malformations. *J Indian Med Assoc*. 2005; 103:690–691. [PubMed: 16821668]
- Salerno A, Kohlhase J, Kaplan BS. Townes-Brocks syndrome and renal dysplasia: a novel mutation in the SALL1 gene. *Pediatr Nephrol*. 2000; 14:25–28. [PubMed: 10654325]
- Schneider RA, Hu D, Helms JA. From head to toe: conservation of molecular signals regulating limb and craniofacial morphogenesis. *Cell Tissue Res*. 1999; 296(1):103–109. [PubMed: 10199970]
- Siefert AW, Harfe BD, Cohn MJ. Cell lineage analysis demonstrated an endodermal origin of the distal urethra and perineum. *Dev Biol*. 2008; 318:143–152. [PubMed: 18439576]
- Siefert AW, Yamaguchi T, Cohn MJ. Functional and phylogenetic analysis shows that Fgf8 is a marker of genital induction in mammals but is not required for external genital development. *Development*. 2009; 136:2643–2651. [PubMed: 19592577]
- Stratford T, Logan C, Zile M, Maden M. Abnormal anteroposterior and dorsoventral patterning of the limb bud in the absence of retinoids. *Mech Dev*. 1999; 81:115–125. [PubMed: 10330489]
- Suzuki K, Bachiller D, Chen YP, Kamikawa M, Ogi H, Haraguchi R, Ogino Y, Minami Y, Mishina Y, Ahn K, Crenshaw EB 3rd, Yamada G. Regulation of outgrowth and apoptosis for the terminal appendage: external genitalia development by concerted actions of BMP signaling. *Development*. 2003; 130(25):6209–6220. [PubMed: 14602679]
- Suzuki K, Haraguchi R, Ogata T, Barbieri O, Alegria O, Vieux-Rochas M, Nakagata N, Ito M, Mills AA, Kurita T, Levi G, Yamada G. Abnormal urethra formation in mouse models of split-hand/split-foot malformation type 1 and type 4. *Eur J Human Genet*. 2008; 16:36–44. [PubMed: 17878916]
- Szucsik JC, Lewis AG, Marmer DJ, Lessard JL. Urogenital tract expression of enhanced green fluorescent protein in transgenic mice driven by a smooth muscle gamma-actin promoter. *J Urol*. 2004; 171(2 Pt 1):944–949. [PubMed: 14713859]
- Trumpp A, Depew MJ, Rubenstein JLR, Bishap JM, Martin GR. Cre-mediated gene inactivation demonstrates that FGF8 is required for cell survival and patterning of the first branchial arch. *Genes & Dev*. 1999; 13:3136–3148. [PubMed: 10601039]
- Wang MH, Baskin LS. Endocrine disruptors, genital development and hypospadias. *J Androl*. 2008; 29:499–505. [PubMed: 18497336]
- Wang L, Liu T, Wang Y, Cao L, Nishioka M, Aguirre RL, Ishikawa A, Geng L, Okada N. Altered expression of desmocollin 3, desmoglein 3 and beta-catenin in oral squamous cell carcinoma:

- correlation with lymph node metastasis and cell proliferation. *Virchows Arch.* 2007; 451(5):959–966. [PubMed: 17846785]
- Weber S, Mir S, Schlingmann KP, Nurnberg G, Becker C, Kara PE, Ozkayin N, Konrad M, Nurnberg P, Schaefer F. Gene locus ambiguity in posterior urethral valves/prune belly syndrome. *Pediatr Nephrol.* 2005; 20:1036–1042. [PubMed: 15912376]
- Weber S, Moriniere V, Knüppel T, Charbit M, Dusek J, Ghiggeri GM, Jankauskienė A, Mir S, Montini G, Peco-Antic A, Wühl E, Zurowska AM, Mehls O, Antignac C, Schaefer F, Salomon R. Prevalence of mutations in renal developmental genes in children with renal hypodysplasia: results of the ESCAPE study. *J Am Soc Nephrol.* 2006; 17:2864–2870. [PubMed: 16971658]
- Witte F, Dokas J, Neuendorf F, Mundlos S, Stricker S. Comprehensive expression analysis of all Wnt genes and their major secreted antagonists during mouse limb development and cartilage differentiation. *Gene Exp Patterns.* 2009; 2:215–223.
- Yamaguchi TP, Bradley A, McMahon AP, Jones SA. Wnt5a pathway underlies outgrowth of multiple structures in the vertebrate embryo. *Development.* 1999; 126:1211–1223. [PubMed: 10021340]
- Yamada G, Suzuki K, Haraguchi R, Miyagawa S, Satoh Y, Kamimura M, Nakagata N, Kataoka H, Kuroiwa A, Chen Y. Molecular genetic cascades for external genitalia formation: an emerging organogenesis program. *Dev Dyn.* 2006; 235:1738–1752. [PubMed: 16598715]

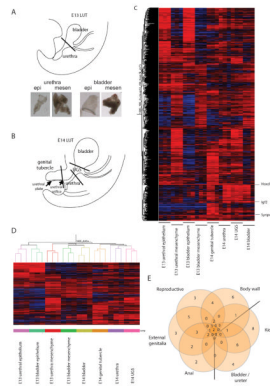


Figure 1. Defining the gene expression profile of the developing lower urinary tract

A) Diagram of the E13 LUT indicating the point of dissection to separate E13 urethra from E13 bladder and the resultant separation of epithelium from mesenchyme achieved using trypsin digestion and EDTA dissociation, respectively. B) Diagram of the E14 LUT indicating the point of dissection to delineate four spatial regions; genital tubercle (GT), urethra, UGS and bladder. C) Heat map of the 1420 most highly expressed genes across all eight compartments of the developing LUT. D) Cluster diagram highlighting the hierarchical relationships between all samples analysed (eight compartments in triplicate). Note that all replicates correctly clustered with their compartment of origin. E) Venn diagram indicating the degree of gene overlap between the processes of reproductive, external genital, anal, body wall, kidney and bladder/ureter development identified in the GT dataset.

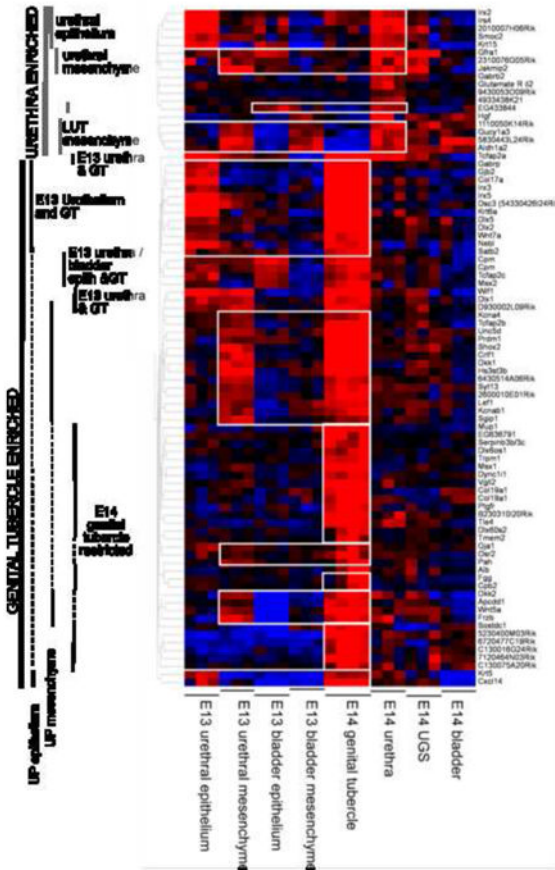


Figure 2. Heat map of gene expression across E14 urethral-enriched and genital tubercle-enriched genes

Urethral-enriched genes can be subdivided into epithelial versus mesenchymal based on the expression in E13 urethra. Genital tubercle-enriched genes were subdivided into urethral plate (UP) epithelium (expressed in genital tubercle and E13 urethral plate epithelium but not E14 urethra), UP mesenchyme (expressed in genital tubercle and E13 urethral mesenchyme but not E14 urethra) and genital tubercle-specific genes (expressed only in GT).

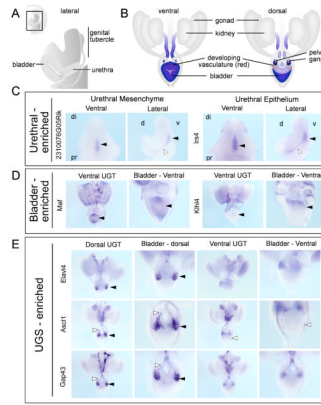


Figure 3. Validation of E14 urethral, bladder and UGS-enriched genes

A) Schematic diagram of a bisected LUT/GT, lateral view, illustrating the bladder, UGS and urethra. up, urethral plate; due, distal urethral epithelium B) Schematic diagram of whole urogenital tracts, ventral and dorsal views. C) Urethral-enriched genes with epithelial (*Irs4*) and mesenchymal (*2310076G05Rik*) expression displayed as ventral (left) and lateral (right) views. Expression can be seen along the UP (solid arrowheads) and in the urethra itself (open arrowheads). D) E13 bladder-enriched genes *Maf* and *Khlh4* showing expression in vasculature and differential expression in the dome of the bladder (solid arrowheads, rostral end). E) UGS-enriched genes *Elav4*, *Ascl1* and *Gap43* revealing specific expression in migrating neural crest cells (open arrowheads) and developing ganglia (solid arrowheads) at the base of the dorsal surface of the bladder. di, distal; pr, proximal; d, dorsal; v, ventral.

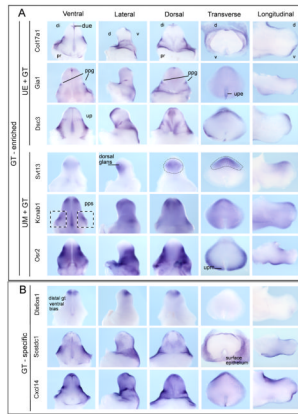


Figure 4. Validation of E14 genital tubercle-enriched gene expression

A) Genital tubercle-enriched genes predicted to be expressed in UP epithelium and GT (*Coll17a*, *Gjal*, *Dsc3*) or UP mesenchyme and GT (*Syt13*, *Kcnab1*, *Osr2*). Expression can also be seen in preputial gland (*Gjal*), distal urethral epithelium (*Coll17a*, *Gjal*, *Dsc3*), surface epithelium of glans and preputial swelling (*Coll17a*, *Gjal*, *Dsc3*) and preputial swelling mesenchyme (*Kcnab1*, *Osr2*). B) Genital tubercle-restricted genes (*Dlx6os1*, *Sostdc1*, *Cxcl14*) showed expression in preputial swelling epithelium (*Sostdc1*, *Cxcl14*), distinct domains across the dorsal GT (*Dlx6os1*, *Sostdc1*, *Cxcl14*) and distal genital tubercle (glans) (*Dlx6os1*). Material is presented from left to right as wholemounts of ventral GT, lateral GT (dorsal to the left side), dorsal GT, and transverse and longitudinal sections through wholemount specimens. di, distal; pr, proximal; d, dorsal; v, ventral; due, distal urethral epithelium; up, urethral plate; upe, up epithelium; upm, up mesenchyme; ppg, preputial gland; pps, preputial swelling.

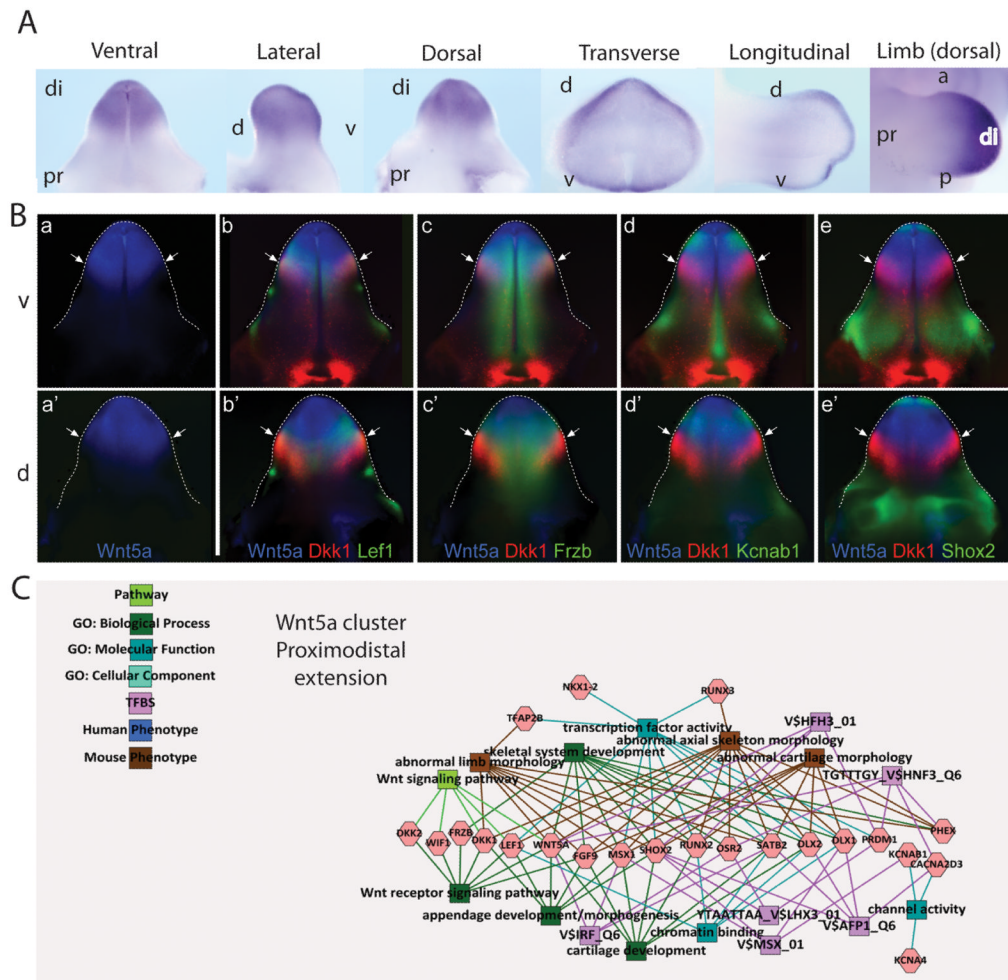


Figure 5. Identification of *Wnt5a* associated gene networks in GT development

A. Wholemount ISH of *Wnt5a* in the developing E13.5 GT showing ventral, lateral, dorsal, transverse and longitudinal views, as well as *Wnt5a* expression in the limb (dorsal view). di, distal; pr, proximal; d, dorsal; v, ventral; a, anterior; p, posterior. B. Pseudo-colored expression pattern overlays of known and candidate genes of a *Wnt5a* network. Experimental WISH images of stage-matched genital tubercles at 13.5dpc were pseudo-colored and overlaid with either *Wnt5a* (a–e, a'–e'). Ventral (v; a–e) and dorsal (d; a'–e') views are shown and the edge of the GT is outlined with a dotted line. *Wnt5a* network genes include *Wnt5a* alone (a, a'), *Wnt5a/Dkk1/Lef1* (b, b'), *Wnt5a/Dkk1/Frzb* (c, c'), *Wnt5a/Dkk1/Kcnab1* (d, d') and *Wnt5a/Dkk1/Shox2* (e, e'). Arrows indicate the border of the *Wnt5a* distal GT expression domain. All genes showed some degree of overlap with the *Wnt5a* domain. *Dkk1* showed overlapping expression with *Lef1* (b, b') and *Frzb* (c, c'), in contrast to *Kcnab1* (d, d') and *Shox2* (e, e'), which did not overlap with the distal *Dkk1* domain. C. GT mesenchyme-enriched gene cluster associated via hierarchical clustering decorated with GO terms and known transcription factor binding sites and known human and mouse phenotype associations. This cluster shows strong links to limb morphogenesis, particularly of mesodermal elements including cartilage and bone.

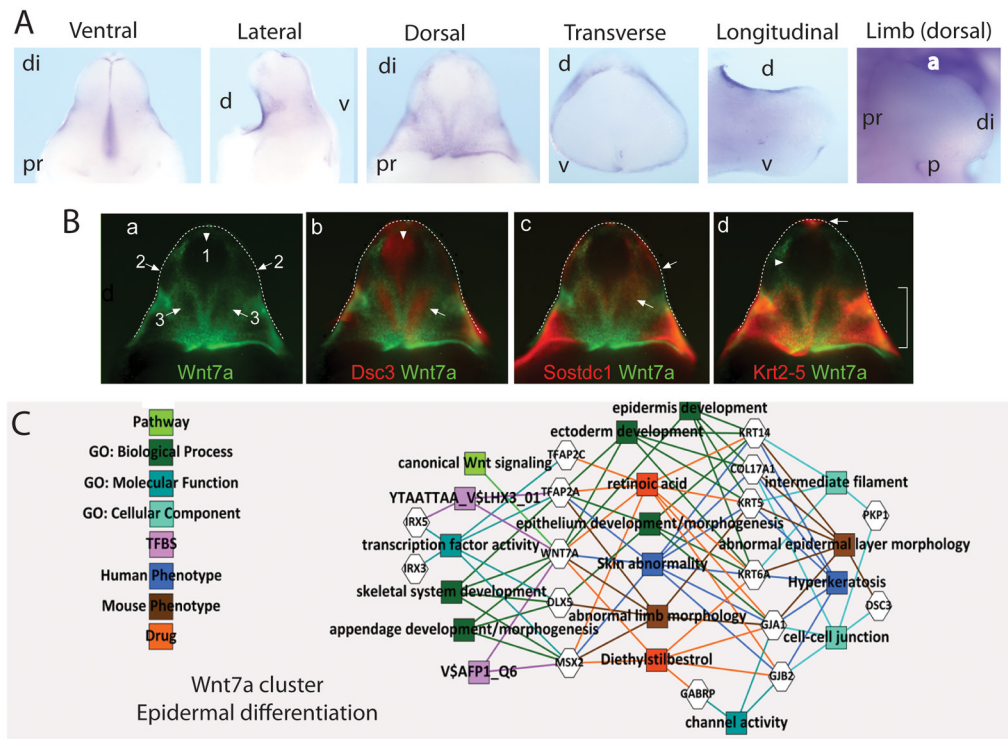


Figure 6. Identification of *Wnt7a* associated gene networks in GT development

A. Wholemount ISH of *Wnt7a* in the developing E13.5 GT showing ventral, lateral, dorsal, transverse and longitudinal views, as well as *Wnt5a* expression in the limb (dorsal view). **B.** Pseudo-colored expression pattern overlays of known and candidate genes of a *Wnt7a* network. Experimental WISH images of stage-matched genital tubercles at 13.5dpc were pseudo-colored and overlaid with *Wnt7a* (a–d). Dorsal (d; a–d) views are shown and the edge of the GT is outlined with a dotted line. *Wnt7a* network genes include *Wnt7a* alone (a), *Dsc3/Wnt7a* (b), *Sostdc1/Wnt7a* (c) and *Krt5/Wnt7a* (d). (a) Arrows indicate the main regions (1–3) on the dorsal side of the GT where *Wnt7a* was not expressed. (b) *Dsc3* showed a complementary pattern to *Wnt7a* with strong, tightly restricted expression in regions 1 and 3 (arrows), but was expressed in the epithelium. (c,d) *Sostdc1* and *Krt5* showed overlapping expression with *Wnt7a*. *Sostdc1* was expressed more broadly than *Wnt7a* and extended into regions 2 and 3 (arrows), however like *Wnt7a*, expression was absent from the central region 1. (d) *Krt5* expression closely resembled *Wnt7a* with strong expression in the proximal *Wnt7a* domains. In the distal *Wnt7a* domains, only weak *Krt5* expression was seen (arrowhead), except for a small region of strong expression at the distal GT tip (arrow). **C.** GT epithelium-enriched gene cluster associated via hierarchical clustering decorated with GO terms and known transcription factor binding sites and known human and mouse phenotype associations. This cluster shows strong links to retinoic acid and diethylstilbestrol regulation as well as epithelium and epidermal differentiation, notably in the limbs.

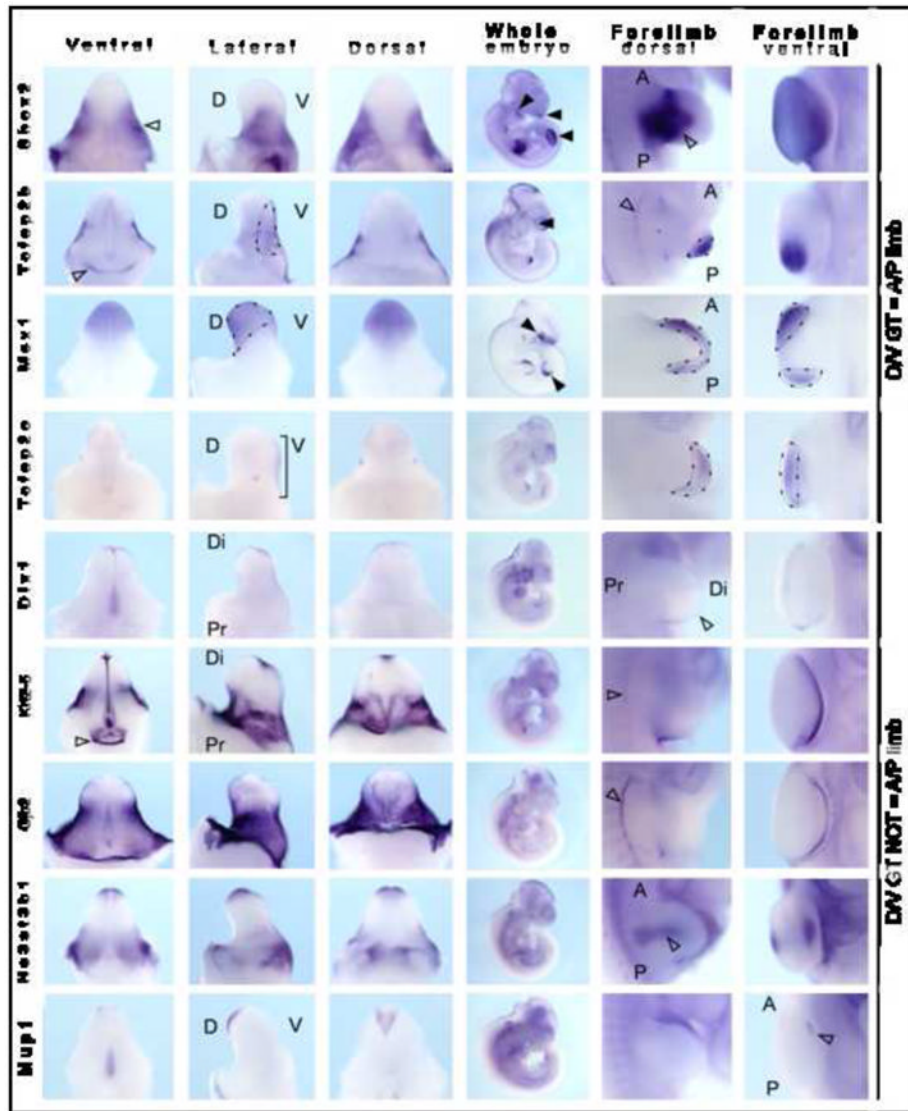


Figure 7. Examination of the congruence between gene expression polarity in genital tubercle versus limb

Wholemout ISH was performed on E13.5 genital tubercle, E10.5 whole embryo and E10.5 forelimb. Specimens are presented from left to right as ventral GT, lateral GT, dorsal GT, E10.5 whole embryo, E11.5 dorsal limb and apical/ventral forelimb limb views. Expression of *Shox2*, *Tcfap2b* and *Msx1* showed congruence of dorsal/ventral GT and anterior/posterior limb (lateral GT = DV limb). *Shox2* was expressed in the mesenchyme of the paired lateral preputial swellings (open arrowhead) and was also expressed in central mesenchyme of the developing limb. *Tcfap2b* showed preputial swelling expression. D, dorsal; V, ventral; A, anterior; P, posterior; Di, distal; Pr, proximal.

Table 1
Overview of Topgene ontological analysis of the most active genes in each compartment of the developing LUT examined

For each category, features that are common between different compartments are highlighted in distinct colours to assist in visual assessment of similarities between compartments. No pathways were strongly evident in E13 urethral mesenchyme, E13 bladder mesenchyme or E14 bladder. No human phenotypes were linked to genes expressed in the E13 bladder mesenchyme.

	E13 Ureth Epith	E13 Ureth Mesen	E13 Bladder Epith	E13 Bladder Mesen	E14GT	E14 Urethra	E14 UGS	E14 Bladder
Molecular Function	TF activity structural molecules calcium ion binding	TF activity calcium ion binding ephrin R activity	TF activity cytoskeleton Cl channel activity	VEGFR binding DNA binding calmodulin binding	TF activity calcium ion binding chromatin binding	TF activity calcium ion binding chemorepellant	calcium ion binding TF activity receptor protein kinase	TF activity TGF beta binding transferase activity
Biological Process	epithelial/epidermal devel cell adhesion regulation of transcription	neural development cell adhesion axonogenesis	epithelial/epidermal dev regulation of transcription autonomic NS/neural crest	axon guidance cell motion/localisation cell adhesion	limb/appendage devel neural development cell adhesion	neural development cell-cell adhesion tube development	neural development cell adhesion axonogenesis	lung development epithelial development regulation of transcription
Cellular Component	apicolateral junction cell junction intercalated disc	neuron projection synapse plasma membrane	apicolateral junction cell-cell junction basolateral membrane	extracellular matrix chromosome passenger complex extracellular space	plasma membrane cell-cell junction intercalated disc	extracellular space plasma membrane extracellular matrix	neuron projection intrinsic to plasma memb nucleosome	extracellular matrix extracellular space plasma membrane
Domain	cadherin bZIP Plectin	cadherin nuclear orphan receptor stathmin	plectin marvel LDLa	bZIP TSP1 HELP	cadherin homeobox OAR	cadherin forkhead thyroglobulin	cadherin stathmin Fnl1l	ETS domain 3 ZnF GATA Fnl1l
Pathway	cadherin signaling wnt signaling breast cancer E2 signaling		breast cancer E2 signaling cell adhesion tight junction		wnt signaling cadherin signaling basal cell carcinoma	cadherin signaling wnt signaling axon guidance	cadherin signaling wnt signaling SLE	
Human Phenotype	abnormal keratinisation abnormalities of skin abnormal mouth/teeth	Hirschsprung disease Anal stenosis Abnormal limb muse	abnormality of the uterus renal dysplasia cutaneous syndactyly	onset in 1 st or 2nd decade	abnormal phalanges/limb hyperkeratosis abnormality of the head/face	abnormality of face external genital abnorm abnormality of phalanges	renal dysplasia/VUR genital abnormality Polydactyly	cardiovascular dilatation muscular hypotonia molluscoid pseudotumors
Mouse Phenotype	abnormal skin morphology abnormal vagus ganglion abnormal esophagus morph	abnormal neurons abnormal musculature perinatal lethality	abnormal mesonephros morphole abnormal epidermis morph thickened epidermis	impaired smooth muscle abnormal umbilical artery abnormal skeleton	abnormal limb abnormal face/mouth/head abnormal skeleton	perinatal lethality abnormal oral/facial abnormal digestive sys	abnormal neuron neonatal lethality conductive hearing loss	absent tracheal rings abnormal ischium abnormal vertebrae
Disease	Epidermolysis Bullosa Keratoderma, Palmoplantar	Hirschsprung Disease Autistic Disorder	renal adysplasia nonsyndromic cleft lip +/-palate	autistic disorder hypertrophic neuropathy	autistic disorder Branchio-Oto-Renal syn	salivary gland neoplasms Branchio-Oto-Renal syn	salivary gland neoplasms Branchio-Oto-Renal syn	Autistic Disorder Ehlers-Danlos Type 2

E13 Ureth Epith	E13 Ureth Mesen	E13 Bladder Epith	E13 Bladder Mesen	E14GT	E14 Urethra	E14 UGS	E14 Bladder
neoplasia	Wardenburg Shah syn	prostate cancer	oral submucous fibrosis	Epidemolysis Bullosa	autonomic control loss	Williams Syndrome	oral submucous fibrosis
insulin resistance	protein QT loci	infertility, male	intervertebral disc disease	female UG disease	head/neck Neoplasms	Autistic Disorder	reperfusion Injury
infertility, male	Pheochromocytoma	prostatic neoplasms	pulmonary function traits	Keratoderma, Palmoplantar	neoplasm metastasis	Hirschprung disease	Obesity

VEGFR: vascular endothelial growth factor receptor; SLE: systemic lupus erythematosus; VUR: vesicoureteric reflux; QT: quantitative trait; TGF: transforming growth factor; CI: chloride

Table 2
Observed gene expression patterns for genital tubercle-enriched genes

Affy ID, Affymetrix ID; DGTE, distal genital tubercle epithelium; DGTm, distal genital tubercle mesenchyme; DUE, distal urethral epithelium; PUE, proximal urethral epithelium; UPM, mesenchyme underlying the urethral plate; PPSM, preputial swelling mesenchyme; PPSE, preputial swelling epithelium; PPG, preputial gland.

Symbol	Cluster	Affy ID	Distal GT		Urethral epithelium			Preputial swelling			
			DGTm	DGTE	DUE	PUE	UPM	PPSM	PPSE	PPG	
Col17a1	GT epithelium - enriched	1418799_a_at	-	+	+	-	-	-	-	+	-
Gjb2	GT epithelium - enriched	1423271_at	-	+	-	-	-	-	-	+	-
Krt2-6a	GT epithelium - enriched	1422784_at	-	-	+	+	-	-	-	-	-
Krt2-5	GT epithelium - enriched	1424096_at	-	+	+	+	-	-	-	+	-
Dlx5	GT epithelium - enriched	1449863_a_at	+	-	-	-	-	+	-	-	-
Dsc3	GT epithelium - enriched	1434534_at	+	+	+	-	-	-	-	+	-
Wnt7a	GT epithelium - enriched	1423367_at	-	+	+	+	-	-	-	+	-
Serpinc3b	GT epithelium - enriched	1422940_x_at	+	-	-	+	+	+	-	-	-
Cxcl14	GT epithelium - enriched	1418457_at	-	+	+	-	-	-	-	+	-
Shox2	GT mesenchyme - enriched	1438042_at	-	-	-	-	-	-	+	-	-
Lef1	GT mesenchyme - enriched	1454734_at	+	-	-	-	-	+	+	-	+
Syt13	GT mesenchyme - enriched	1434470_at	+	-	-	-	-	-	-	-	-
Prdm1	GT mesenchyme - enriched	1420425_at	+	-	-	-	-	+	+	-	-
Dkk1	GT mesenchyme - enriched	1420360_at	+	-	+	-	-	+	-	-	-
Tefap2b	GT mesenchyme - enriched	1435670_at	+	-	-	-	-	+	-	+	-
Hs3st3b1	GT mesenchyme - enriched	1433977_at	+	-	-	-	-	+	+	-	-
Wnt5a	GT mesenchyme - enriched	1436791_at	+	-	+	-	-	+	-	-	-
Kcnab1	GT mesenchyme - enriched	1448468_a_at	+	-	-	-	-	+	+	-	-
Apcdd1	GT mesenchyme - enriched	1449070_x_at	+	-	-	-	-	+	+	-	-
	GT mesenchyme - enriched	1440630_at	+	-	-	-	+	+	-	-	-
Unc5d	GT mesenchyme - enriched	1440484_at	+	-	-	-	+	+	-	-	-
Kcna4	GT mesenchyme - enriched	1438613_at	+	-	-	-	+	-	+	-	-
D930002L09Rik	GT mesenchyme - enriched	1444452_at	+	-	-	-	+	-	-	-	-

Symbol	Cluster	Afy ID	Distal GT		Urethral epithelium		UPM		Preputial swelling		
			DGTM	DGTE	DUE	PUE	UPM	PPSM	PPSE	PPG	
Satb2	GT mesenchyme - enriched	1427017_at	+	-	-	+	+	-	-	-	-
Frzb	GT mesenchyme - enriched	1416658_at	+	-	-	-	+	-	-	-	-
Mup1	E14 GT restricted	1420465_s_at	+	-	-	-	-	-	-	-	-
Dlx6os1	E14 GT restricted	1438799_at	+	-	-	-	-	-	-	-	-
Trpm1	E14 GT restricted	1437445_at	+	-	-	-	-	-	-	-	-
Msx1	E14 GT restricted	1448601_s_at	+	-	-	-	-	+	-	-	-
Coll19a1	E14 GT restricted	1456953_at	+	-	-	+	+	-	-	-	-
Coll19a1	E14 GT restricted	1421698_a_at	+	-	-	+	+	-	-	-	-
Pigfr	E14 GT restricted	1420349_at	+	-	-	+	+	-	-	-	-
	E14 GT restricted	1445804_at	+	-	-	+	+	-	-	-	-
Tle4	E14 GT restricted	1430384_at	-	-	-	+	+	-	-	-	-
Dlx6os2	E14 GT restricted	1440797_at	+	-	-	+	+	-	-	-	-
Sostdc1	E14 GT restricted	1449340_at	-	+	+	-	-	-	+	+	-
Wif1	E13 urethra and GT	1425425_a_at	+	-	-	+	+	-	+	-	-
Dlx1	E13 urethra and GT	1449470_at	-	+	+	-	-	-	-	-	-
Gjal	E13 urethra and GT	1415801_at	+	-	+	-	-	-	+	-	+
Osr2	E13 urethra and GT	1426155_a_at	+	-	-	-	-	-	+	-	-
Msx2	E13 bladder and GT	1449559_at	+	+	+	-	-	-	-	-	-
Tcfap2c	E13 bladder and GT	1448977_at	+	-	-	-	-	-	-	-	+
Cpm	E13 bladder and GT	1429413_at	+	-	+	-	-	-	-	-	-